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MICROBIOME TARGETED THERAPIES IN GULF WAR ILLNESS

by

Diana Agnes Kimono

Bachelor of Science Nelson Mandela University, 2010

Master of Science Nelson Mandela University, 2013

Submitted in Partial Fulfillment of the Requirements

For the Degree of Doctor of Philosophy in

Environmental Health Sciences

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Accepted by:

Saurabh Chatterjee, Major Professor

Geoff Scott, Committee Member

Dwayne Porter, Committee Member

Stephen Lasley, Committee Member

Cheryl L. Addy, Vice Provost and Dean of the Graduate School



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DEDICATION

To my beloved parents, Mrs. Rachel Wamburu and Mr. Perez Wamburu. And to my amazing husband William Akoto. Without you, I do not think I could have achieved this.



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I would like to offer my sincere gratitude to my supervisor, Dr. Saurabh Chatterjee for his steadfast support and advisement. I would also like to thank my PhD committee members: Dr. Geoff I. Scott, Dr. Dwayne Porter and Dr. Stephen Lasley, for their valuable insights and support in my work.

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ABSTRACT

Gulf war illness (GWI) is a chronic multisymptomatic disorder affecting about 30% of veterans of the 1990-1991 Persian Gulf war. Affected veterans complain of chronic symptoms which begun during or shortly after the war and persist 30 years later. This dissertation is a report of three studies which use a murine model to investigate the microbiome as a therapeutic target in GWI. Mice were exposed to pesticides and the prophylactic drug pyridostigmine bromide (PB) and studied these chemical's impact on the microbiome in both an acute and persistence model of GWI.

The *first* study looks at the effect of altered microbiome on metabolism and proposes short chain fatty acids as a therapy for GWI. Results show that mice exposed to GWI showed toll like receptor activation, inflammation and metabolic reprogramming in the liver. These symptoms were alleviated with sodium butyrate, a short chain fatty acid. The *second* study looked at the effect of altered microbiome on the enteric nervous system and proposes the use of SsnB a TLR4 antagonist in combination with sodium butyrate as a possible therapy. Results show that mice which were treated with GW chemicals had reactive enteric glia which produced reactive oxygen species and proinflammatory cytokines, thereby modulating the expression of tight junction proteins in the intestine. Further, administration of SsnB and butyrate led improved EGC states and therefore improving tight junction protein integrity. The *third* study looks at the altered microbiome



V

in the persistence of GWI neurological symptoms. Results show that mice exposed to GW chemicals presented with decreased relative abundance of *Akkermansia muciniphila*, a probiotic bacterium associated with good health, and this correlated with HMGB1 levels, neuroinflammation and neurotrophins level such as BDNF which are key players in maintaining neurological health



TABLE OF CONTENTS

DEDICATIONiii
ACKNOWLEDGEMENTS iv
ABSTRACTv
LIST OF TABLES
LIST OF FIGURES ix
CHAPTER 1: INTRODUCTION
CHAPTER 2: INCREASED BUTYRATE PRIMING IN THE GUT STALLS MICROBIOME ASSOCIATED GASTRO-INFLAMMATION AND HEPATIC METABOLIC REPROGRAMMING IN A MOUSE MODEL OF GULF WAR ILLNESS
CHAPTER 3: DYSBIOSIS-ASSOCIATED ENTERIC GLIAL CELL IMMUNE ACTIVATION AND REDOX IMBALANCE MODULATE TIGHT JUNCTION PROTEIN EXPRESSION IN GULF WAR ILLNESS PATHOLOGY41
CHAPTER 4: HOST <i>AKKERMANSIA MUCINIPHILA</i> ABUNDANCE CORRELATES WITH GULF WAR ILLNESS SYMPTOM PERSISTENCE VIA NLRP3 MEDIATED NEUROINFLAMMATION AND DECREASED BRAIN DERIVED NEUROTROPHIC FACTOR
CHAPTER 5: CONCLUSION
REFERENCES
APPENDIX A: PERMISSION TO REPRINT CHAPTERS



LIST OF TABLES

Table 2.1 Real time PCR primer sequences.	17
Table 3.1. Rat Primer sequences	53
Table 4.1 Primer sequences	98



LIST OF FIGURES

Figure 1.1 Summary of all three projects discussed
Figure 2.1. Gut microbiome alteration in mice model of Gulf War Illness (GWI)33
Figure 2.2. Change in gut microbiome in GWI alter niacin receptor (GPR109A) and tight junction proteins in the intestine
Figure 2.3. Sodium butyrate priming in a rodent model of GWI attenuates TLR4 activation in the small intestine
Figure 2.4. Sodium butyrate priming in a rodent model of GWI improves proinflammatory phenotype in small intestine mediated by the TLR4 pathway
Figure 2.5. Sodium butyrate treatment in GWI improves circulatory DAMPs37
Figure 2.6. Sodium butyrate treatment in a rodent model of GWI attenuates TLR4 activation in Liver
Figure 2.7. TLR4 activation is associated with metabolic changes and inflammatory response in the liver but the phenotypic liver injury is predominantly absent
Figure 2.8. TLR4 drives the metabolic alterations in GW-chemical exposed liver40
Figure 3.1. Altered microbiome associated increase PAMPS and DAMPS71
Figure 3.2. Altered microbiome induced change in EGC phenotype to a reactive phenotype
Figure 3.3. Expression of Toll-like receptors in small intestine and EGC
Figure 3.4. Expression of S100B and RAGE in the small intestine74
Figure 3.5. Formation of S100B/RAGE complex in small intestine
Figure 3.6. Activation of NOS-2 in small intestine
Figure 3.7: Exposure to DAMPS and PAMPS cause activation of TLR4 in EGC77
Figure 3.8. EGC exposed to LPS or HMGB-1 change to a reactive phenotype
Figure 3.9 . Activation of inducible nitric oxide synthase by LPS or HMGB-1



Figure 3.10. NADPH oxidase 2 and peroxynitrite mediated oxidative stress <i>in vivo</i> 80
Figure 3.11. NADPH oxidase 2 activation in rat EGC
Figure 3.12. Peroxynitrite formation in EGC 82
Figure 3.13. ROS mediated activation of NLRP-3 inflammasome and inflammation in EGC
Figure 3.14. DNA fragmentation of rat EGC
Figure 3.15. Expression of claudin 2, occludin and aquaporin 3 in mouse small intestine
Figure 3.16. Protein expression of aquaporin 3, claudin 2 and occludin in intestinal epithelial cells
Figure 4.1 . Exposure to GW chemicals results in decreased relative abundance of <i>Akkermensia muciniphila</i> and chronic high levels of circulatory HMGB1112
Figure 4.2 . Exposure to GW chemicals is associated with altered claudin 5 levels in the frontal cortex
Figure 4.3. Activation of Macrophages and associated HMGB1/RAGE complex formation
Figure 4.4. Increased ROS is associated with NLRP3 inflammasome activation115
Figure 4.5 . GW chemical exposure is associated with chronic neuroinflammation and decreased Brain derived neurotrophic factor (BDNF) levels in Frontal cortex116
Figure 4.6. Decreased relative abundance of <i>Akkermansia muciniphila</i> correlates with IL-1β and BDNF levels in the frontal cortex
Figure 4.7 . Deletion of NLRP3 is associated with decreased neuroinflammation and lower BDNF levels
Figure 4.8. Serum Endotoxin levels in GW chemical treated mice (GWP) compared to the vehicle control (CONT) only treated mice



CHAPTER 1

INTRODUCTION

Gulf war illness is a chronic multisymptomatic disorder affecting about 30% of veterans of the 1990-1991 Persian Gulf war (1)(2) (3). Affected veterans complain of acute and chronic symptoms which begun during or shortly after the war and still persist 30 years later (1)(3)(2)(4). These symptoms include fatigue, muscle pain, cognitive problems, insomnia, rushes, gastrointestinal problems etc. Gulf war illness has particularly been a challenge because although veterans complain of the above symptom, there is no significant clinical pathology that can be medically detected (4). For many years, this challenge resulted in the disorder being dismissed as post-traumatic stress disorder (PTSD) or other psychological problems that the veterans may have developed due to the war (1)(2). However, in the last 15 years, epidemiological studies have established a compelling link between GWI with environmental and chemical exposures that occurred shortly before (in preparation for the war) or during the war. These exposures include vaccines, desert storm dust, depleted uranium, combustion products from oil wells, pesticides and prophylactic drugs(5).

Currently, research is focused on understanding the mechanisms that drive this disorder by studying veteran cohorts, using animal models and invitro work. Other studies



are also aiming to determine possible therapies to relieve affected veterans of these discomforts (6).

To date, our lab research on GWI focusses on the role of the microbiome in GWI. The microbiome consists of all the microorganisms in the gastrointestinal tract (7). The gut is populated by billions of microorganisms which exist in symbiotic and commensal relations with their host. These can be bacteria, viruses, fungi, protozoans etc. These microorganism populations affect the health and influence disease states of their host (8)(9)(10)(11). Although the mechanisms by which they influence health are not fully understood, it is now known that these microorganisms act through. Enabling digestion of complex foods eg xyloglucans to release nutrients eg short chain fatty acids (12) antiinflammatory substances, manufacture of vitamins (13) etc. They also produce and consume signaling molecules which can be detected by the brain eg neurotransmitters such as gama amino butyric acid (GABA), dopamine, serotonin etc which are important in neurological health (14). For example, the study by Vogt et al found elevated levels of the gut microbiota metabolite trimethylamine N oxide in cerebral spinal fluid of AD patients (15), which is a known risk factor for the disease. Another study found that key metabolites had been altered (amino acids and fatty acids) significantly in IBS patients stools and this correlated with the observed dysbiosis (16). With this critical role of the microbiome in health and chronic disease, we investigated its role in driving and influencing GWI pathology.

This dissertation is a report of three studies which use a murine model to investigate the microbiome as a therapeutic target in GWI. We exposed mice to pesticides and the prophylactic drug pyridostigmine bromide (PB) and studied these chemical's



impact on the microbiome. Although animal models of GWI still remain imperfect, this model is generally accepted because it is known that GW veterans were exposed to a wide range of pesticides eg permethrin, deet etc. These chemicals were applied to the veteran's uniforms and tents and skin to protect them from insects while in the war.

Permethrin, which we use in our study is a pyrethroid which acts on the central nervous system as a neurotoxin by disrupting sodium channels in neurons of insects resulting in paralysis. In small quantities, it is not significantly toxic to humans and larger animals, however, to insects it is lethal since they are unable to metabolize this chemical quickly. We also used a pyridostigmine bromide (PB) an investigational product (at that time) which was supplied as prophylactic drug to GW warriors to protect them from the harmful effects of nerve agents eg sarin gas. About 250, 000 personnel reported having used the drug in the GW at a dosage of 30 mg per 8 hours. However, there was some variation because the drug was self-administered, and some people could have consumed more or less than required. It is a reversible inhibitor of acetylcholinesterase (AchE), an enzyme which is targeted by nerve agents which irreversibly bind AchE (17).

Generally, we hypothesized that exposure to GW chemicals led to altered microbiome alterations which fuels inflammation in the gut and other organs. This is through the release of Damage associated molecular patterns (DAMPs) and pathogen associated molecular patterns (PAMPs) which escape the gastrointestinal tract and leak out into the circulation through compromised gut barrier integrity.

The first study looks at the effect of an altered microbiome on metabolism and also proposes short chain fatty acids as a therapy for GWI. We found that mice exposed to GWI



showed toll like receptor activation, inflammation and metabolic reprogramming in the liver. These symptoms were alleviated with sodium butyrate which acted through its receptor (18).

The second study looked at the effect of the altered microbiome on the enteric nervous system (ENS) and proposes the use of sparstolonin B (SsnB) a TLR4 antagonist in combination with sodium butyrate as a possible therapy. We found that mice which were treated with GW chemicals had reactive enteric glia which produced reactive oxygen species (ROS) and proinflammatory cytokines, thereby modulating the expression of tight junction proteins and aquaporins in the small intestine. In our previous studies we had found that poor gut integrity was an important symptom in GWI and is the likely portal for immunostimulatory particles to reach other distant organ beyond the gut. We also found that administration of SsnB and butyrate led improved EGC states and therefore improving tight junction protein integrity (19).

The third study looks at the altered microbiome in the persistence of GWI symptoms. Using a persistence model of GWI, we found that mice exposed to GW chemicals presented with decreased relative abundance of *Akkermansia muciniphila*, a probiotic bacterium associated with good health, and this correlated with HMGB1 levels, neuroinflammation and neurotrophins level such as BDNF which are key players in maintaining neurological health. Figure 1 is a graphical summary of these three studies.





Figure 1.1. Summary of all three projects discussed. Exposure to GW chemicals results in microbiome alterations (dysbiosis) with increased DAMPs and PAMPS, inflammation and a leaky gut (leaky gut is shown by dotted red line). DAMPs eg HMGB1 and PAMPs eg LPS cross into the blood stream reaching distant organs such as the liver and the brain. This drives inflammation and associated symptoms such as carbohydrate and fat metabolic reprograming in liver, decrease in neurotrophin levels e,g brain derived neurotrophic factor (BDNF) in the brain. Brown boxes represent organs/systems on which research was focused, blue boxes list pathological markers and findings of studies white boxes represent conclusions and our thoughts for future studies. *Abbreviations used GW (Gulf war), PB (pyridostigmine bromide), DAMPs (Damage associated molecular patterns) PAMPs (Pathogen associated molecular patterns), LPS (lipopolysaccharides), HMGB1 (High mobility group box 1 protein), ROS (reactive oxygen species), GI (gastrointestinal), CNS (central nervous system), BBB (blood brain barrier), BDNF (Brain derived neurotrophic factor)*



CHAPTER 2

INCREASED BUTYRATE PRIMING IN THE GUT STALLS MICROBIOME ASSOCIATED-GASTROINTESTINAL INFLAMMATION AND HEPATIC METABOLIC REPROGRAMMING IN A MOUSE MODEL OF GULF WAR ILLNESS¹

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¹ Seth, R. K[#]., Kimono, D[#]., Alhasson, F., Sarkar, S., Albadrani, M., Lasley, S. K., Horner, R., Janulewicz, P., Nagarkatti, M., Nagarkatti, P., Sullivan, K., & Chatterjee, S. (2018). Increased butyrate priming in the gut stalls microbiome associated gastrointestinal inflammation and hepatic metabolic reprogramming in a mouse model of Gulf War Illness. *Toxicology and applied pharmacology*, *350*, 64–77. https://doi.org/10.1016/j.taap.2018.05.006 # authors contributed equally

Running Title: Butyrate priming and metabolic reprogramming in GWI

Key words: Permethrin, Pyridostigmine bromide, Gut dysbiosis, TLR4, Claudin-2, Cytokines

Author for correspondence:

*Dr. Saurabh Chatterjee, Ph.D. Environmental Health and Disease Laboratory, NIEHS Center for Oceans and Human Health on Climate Change Interactions, Department of Environmental Health Sciences, University of South Carolina, Columbia 29208 USA. Email: <u>schatt@mailbox.sc.edu</u>; Tel: 803-777-8120; Fax: 803-777-3391

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ABSTRACT

Most of the associated pathologies in Gulf War Illness (GWI) have been ascribed to chemical and pharmaceutical exposures during the war. Since an increased number of veterans complain of gastrointestinal (GI), neuroinflammatory and metabolic complications as they age and there are limited options for a cure, the present study was focused to assess the role of butyrate, a short chain fatty acid for attenuating GWIassociated GI and metabolic complications. Results in a GWI-mouse model of permethrin and pyridostigmine bromide (PB) exposure showed that oral butyrate restored gut homeostasis and increased GPR109A receptor copies in the small intestine (SI). Claudin-2, a protein shown to be upregulated in conditions of leaky gut was significantly decreased following butyrate administration. Butyrate decreased TLR4 and TLR5 expressions in the liver concomitant to a decrease in TLR4 activation. GW-chemical exposure showed no clinical signs of liver disease but a significant alteration of metabolic markers such as SREBP1c, PPAR-α, and PFK was evident. Liver markers for lipogenesis and carbohydrate metabolism that were significantly upregulated following GW chemical exposure were attenuated by butyrate priming in vivo and in human primary hepatocytes. Further, Glucose transporter Glut-4 that was shown to be elevated following liver complications were significantly decreased in these mice after butyrate administration. Finally, use of TLR4 KO mice completely attenuated the liver metabolic changes suggesting the central role of these receptors in the GWI pathology. In conclusion, we report a butyrate specific mechanistic approach to identify and treat increased metabolic abnormalities in GWI veterans with systemic inflammation, chronic fatigue, GI disturbances, metabolic complications and weight gain.



2.1. INTRODUCTION

Gulf War Illness (GWI) has been characterized as a chronic multi-symptom illness with pathology that includes neuronal inflammation leading to cognitive deficiencies, chronic fatigue, joint and muscle pain and gastrointestinal complications (3)(20). GWI research has identified toxicant chemical exposures in the war theater including sarin nerve gas, pyridostigmine bromide (PB) anti-nerve gas pills, insecticides and insect repellents, to be prime reasons for most symptoms reported by the veterans (3). A study of military pesticide applicators from the GW recently reported increased cognitive decrements in attention, memory and information processing speed in veterans with combined exposures to PB, pesticides and insect repellents (21). Animal models of GWI have also shown chronic impairment in learning and memory, fatigue and gastrointestinal dysfunction when exposed to GW-relevant chemicals including PB, pesticides and the insect repellent permethrin (22)(23). GWI has emerged as a primarily neuroimmune disorder with greater inflammatory effects noted when GW-relevant toxicants were combined with corticosterone in animal models to mimic the physical and mental stressors of war deployment (24). However the basis of gastrointestinal complications and parallel networks for neuronal complications have remained largely elusive (25) (26)(25). Though the larger epidemiological studies list GI complications as a major health problem in only a subset of GW veterans, there are hardly any reports that show hepatic metabolic disturbances as a widely reported symptom (3);(27). This may be due to the silent nature of the presentation of these symptoms and unknown links to a wider systemic complication in GW veterans following sedentary lifestyles/physical disabilities/diet/aging over a longer period of time (28). However, with increased incidences of obesity and persistence of



symptoms in GW-veterans and the US population in general, the causes, outcomes, and extent of metabolic disturbances can no longer be ignored. The long-ranging implications of such silent changes in the GI tract and the liver following chemical exposure form the basis of the present study.

We have shown recently the unambiguous role of the gut microbiome in causing neuronal inflammation largely due to gut leaching and systemic endotoxemia (29). The altered gut bacterial signature obtained following Gulf War chemical exposure caused a TLR4-linked inflammatory surge in the GI tract and could be traced in the frontal cortex (29). The observations reported paved the way for newer investigations into wider systemic inflammatory complications in extraneuronal organs that might not have a clear phenotype yet may be a basis for multisymptomatic illness as described in GWI.

GWI is also characterized by the presence of chronic fatigue. Most classifications in the past have listed chronic fatigue as one of the most widely reported symptom burdens in GW veterans (20)(30) . Interestingly, changes in the microbiome of affected patients with chronic fatigue syndrome/fibromyalgia have been strongly associated with the causes of this illness (31)(32). Chronic fatigue is also strongly linked to widescale changes both in the gut bacteria and the systemic metabolism with the latter believed to have roots in the liver though skeletal muscles also play a major role (33). Persistent alteration of liver metabolism following changes in gut microbiome and its subsequent effects on systemic metabolism may affect the development of chronic fatigue via the altered availability of NADPH, ATP, and cofactors for various biochemical pathways (33).



Most chronic liver diseases like fatty liver disease and biliary fibrosis are silent in a presentation at the clinic and remain asymptomatic until it reaches an irreversible stage. Nonalcoholic fatty liver disease or cholestatic liver disease have chronic fatigue as one of the symptoms (34). The fatigue associated with these silent liver complications have been assigned to lipotoxicity, insulin and leptin resistance, endocrinopathies and metabolic syndrome (35). Interestingly, we have shown that environmental chemicals alter liver metabolism by increasing glucose transporters in the liver fibroblasts, elevating expressions of PPAR- α , PFK and decreasing PPAR- γ levels with a concomitant rise in leptin (36). These changes in the liver have been associated with an altered microbiome following consumption of a diet rich in high fat and low fiber for a long period of time at least in the murine models (37)(38).

With strong emphasis on the altered microbiome being associated with pathologies of inflammatory bowel disease, chronic liver disease, chronic fatigue syndrome and neuronal complications (all have an inflammatory component) via the gut-brain axis, it is important that studies be focused on bacterial metabolites within the gut that might be linked to some or all the pathways that link these systemic complications. Interestingly, GWI patients present significant symptoms that resemble some or all these in the clinics. Thus bacterial metabolites like butyric acid, propionic acid and acetate need to be considered as molecules of interest in treating multiple symptoms of GWI since these molecules have shown promising results in the clinic to cure IBD and dysbiosis related complications (39) (40)(41). Further, Butyrate-producing bacteria such as Roseburia species supplementation rescued patients from IBD (41). Butyrate, in particular, is a short chain fatty acid (SCFA) that primarily interacts with the GPR109A and is an



immunosuppressant widely known to increase T-regulatory cells in the intestine and a prominent HDAC1 inhibitor (42)(43).

The present study tests the hypothesis that GW chemical exposure causes a decrease in butyrate-producing bacteria and concomitant butyrate priming in the gut through oral supplementation attenuates GI inflammation, gut leaching and metabolic abnormalities in the liver and higher systemic leptin levels. The study uses state of the art genomic approaches and an oral priming by butyrate for elucidating genus-specific changes in gut bacteria, and human hepatocytes treated with an insect repellent permethrin, (used in Gulf War theater) for mechanistic investigations.

2.2. MATERIALS AND METHODS

2.2.1. Materials

Pyridostigmine bromide (PB), Permethrin (Per), Lipopolysaccharides (LPS), Corticosterone and Sodium butyrate (NaBT) were purchased from Sigma-Aldrich (St. Louis, MO). Anti-claudin-2, anti-occludin, anti-TLR4, anti-flotillin, anti-HMGBl, anti-Leptin and anti-IL1β primary antibodies were purchased from Abeam (Cambridge, MA). Anti-TLR5 primary antibody was purchased from Santacruz Biotechnology (Dallas, TX). Species-specific biotinylated conjugated secondary antibodies and Streptavidin-HRP (Vectastain Elite ABC kit) were purchased from Vector Laboratories (Burlingame, CA). Fluorescence-conjugated (Alexa Fluor) secondary antibodies, ProLong Gold antifade mounting media with DAPI were purchased from Thermofisher Scientific (Grand Island, NY) and all other chemicals which were used in this project purchased from sigma only if otherwise specified. Paraffin-embedding of tissue sections on slides were done by AML



laboratories (Baltimore, MD). Microbiome analysis was done by Second Genome, the microbiome company (San Francisco, CA).

2.2.2. Animals

Adult wild-type male (C57BL/6J mice) and adult mice that contained the disrupted TLR4 gene (TLR4 KO) (B6·B10ScN-Tlr4^{lps-del} / JthJ) were purchased from the Jackson Laboratories (Bar Harbor, ME). Mice were implemented in accordance with NIH guideline for human care and use of laboratory animals and local IACUC standards. All procedures were approved by The University of South Carolina at Columbia, SC. Mice were housed individually and fed with chow diet at 22–24 °C with a 12-h light/ 12-h dark cycle. All mice were sacrificed after animal experiments had been completed. Right after anesthesia, blood from the mice was drawn using cardiac puncture, in order to preserve serum for the experiments. The mice liver was collected for further experiments immediately after terminal euthanasia. Fecal pellets and luminal contents were collected from the animals, followed by dissection of the small intestine. The tissues were fixed using 10% neutral buffered formalin. Distal segments of small intestines were used for the staining and visualizations.

2.2.3. Rodent model of Gulf War Illness (GWI)

Mice were exposed to Gulf War chemicals based on established rodent models of Gulf War Illness with some modifications (25)(44). The treated wild-type mice group (GWI) and treated TLR4 KO mice group (TLR4 KO) were dosed triweekly for one week with PB (2 mg/kg) and Permethrin (200 mg/kg) via the oral route. After completion of PB/Permethrin dosages, mice were administered corticosterone intraperitoneally (i.p.) with



a dose of 100 µg/mice/day for 5 days of the week for one week. The dose of corticosterone was selected from the study which exposed mice to 200mg/L of corticosterone through drinking water. The i.p. dose of corticosterone had similar immunosuppression as examined by low splenic T cell proliferation (data not shown). The vehicle control group (Veh) of mice received saline injections and vehicle for oral gavage in the same paradigm. Another group of wild-type mice was exposed with PB, Permethrin and corticosterone similar to GWI group of mice and co-exposed with sodium butyrate (GWI + NaBT) 10 mg/kg via the oral route.

2.2.4. Microbiome analysis

Fecal pellets and luminal contents were collected from the animals of each group after sacrifice and then sent to Second Genome and School of Medicine, the University of South Carolina for microbiome analysis. The second Genome performed nucleic acid isolation with the MoBio PowerMag Microbiome kit (Carlsbad, CA) according to manufacturer's guidelines and optimized for high-throughput processing V3-V4 sequencing and bioinformatic analysis.

2.2.5. Cell culture

Freshly isolated primary human hepatocytes were obtained from Liver Tissue Cell Distribution System, University of Minnesota, Minneapolis, MN. Plated hepatocytes were maintained in DMEM media supplemented with 10% FBS until treated. Cells were then serum starved in DMEM supplemented with 1.5% FBS for 8h and exposed to vehicle control and chemicals. Cells were then treated with vehicle (Veh Cont), LPS (1 μM), LPS



+ NaBT (LPS 1 μ M and Sodium Butyrate 0.2mg/mL) for 24 h. After experiment cells were harvested for mRNA extraction and gene expression analysis.

2.2.6. Laboratory methods

Immunohistochemistry

The distal part of small intestine was collected from mice and fixed in 10% neutral buffered formalin. The fixed tissues swiss rolled, paraffin embedded and cut in 5 μ M thick section. These sections were subjected to deparaffinization using a standard protocol. Epitope retrieval solution and steamer (IHC-Word, Woodstock, MD) were used for epitope retrieval for deparaffinized sections. 3% H₂O₂ was used for the recommended time to block the endogenous peroxidase. After serum blocking, the tissue was incubated overnight at 4.0 °C with primary antibody IL1B. Species-specific biotinylated conjugated secondary antibodies and streptavidin conjugated with HRP were used to implement antigen-specific immunohistochemistry. 3,3'-Diaminobenzidine (DAB) (Sigma Aldrich, St Louis, MD) was used as a chromogenic substrate. Mayer's Hematoxylin solution (Sigma Aldrich) was used as a counterstain. Sections were washed between the steps using phosphate buffered saline 1×. Finally, stained sections were mounted with Simpo-mount (GBI laboratories, Mukilteo, WA). Tissue sections were observed using Olympus BX51 microscope (Olympus, America). Cellsens software from Olympus America (Center Valley, PA) was used for morphometric analysis of images.

Immunofluorescence staining

Paraffin-embedded distal part of the small intestine or liver sections were deparaffinized using a standard protocol. Epitope retrieval solution and steamer were used



for epitope retrieval of sections. Primary antibodies such as anti-Claudin-2, anti-Occludin, anti-GPR109A, anti-TLR4, anti-Flotillin, and anti-TLR5 were used at the recommended dilution. Species-specific secondary antibodies conjugated with Alexa Fluor (633-red and 488-green) were used at advised dilution. In the end, the stained sections were mounted using Prolong gold antifade reagent with DAPI. Sections were observed under-Olympus fluorescence microscope using $20\times$, $40\times$ or $60\times$ objective lenses.

Real-time quantitative PCR

mRNA expression in small intestine, liver, and human primary hepatocytes was examined by quantitative real-time PCR analysis. Total RNA was isolated from each 25 mg liver tissue or 15 mg small intestine tissue or 1×10^6 primary human hepatocytes cell by homogenization in TRIzol reagent (Invitrogen, Carlsbad, CA) according to the manufacturer's instructions and purified with the use of RNeasy mini kit columns (Qiagen, Valencia, CA). cDNA was synthesized from purified RNA (1µg) using iScript cDNA synthesis kit (Bio-rad, Hercules, CA) following the manufacturer's standard protocol. Real-time qPCR (qRTPCR) was performed with the gene-specific primers using SsoAdvanced SYBR Green Supermix and CFX96 thermal cycler (Bio-rad, Hercules, CA). Threshold Cycle (Ct) values for the selected genes were normalized against respective samples internal control (18S). Each reaction was carried out in triplicates for each gene and for each sample. The relative fold change was calculated by the 2- $\Delta\Delta$ Ct method. The sequences for the primers used for Real-time PCR are provided in Table 1



Gene ^a	Primer sequence (5' to 3' orientation)
MM_IL-1β	Sense: CCTCGGCCAAGACAGGTCGC Antisense: TGCCCATCAGAGGCAAGGAGGA
MM_MCP-1	Sense: CACAGTTGCCGGCTGGAGCAT Antisense: GTAGCAGCAGGTGAGTGGGGGC
MM_TNF-α	Sense: CAACGCCCTCCTGGCCAACG Antisense: TCGGGGCAGCCTTGTCCCTT
MM_SREBP1c	Sense: GGAACAGACACTGGCCGA Antisense: AAGTCACTGTCTTGGTTGAT
MM_PPAR-a	Sense: AGACCTTCGGCAGCTGGTCAC Antisense: GTGGCAACGGCCTGCCATCT
MM_PPAR-γ	Sense: TTCGCTGATGCACTGCCTAT Antisense: GGAATGCGAGTGGTCTTCCA
MM_GLUT-1	Sense: CCTGTCTCTTCCTACCCAACC Antisense: GCAGGAGTGTCCGTGTCTTC
MM_GLUT-4	Sense: CACCGGCAGCCTCTTATCAT Antisense: CACCGAGACCAACGTGAAGA
MM_PFK	Sense: GCCGTGAAACTCCGAGGAA Antisense: GTTGCTCTTGACAATCTTCTCATCAG
Hs_SREBP1c	Sense: CATGGATTGCACTTTCGAA Antisense: GGCCAGGGAAGTCACTGTCTT
Hs_PPAR-γ	Sense: GGCTTCATGACAAGGGAGTTTC Antisense: AACTCAAACTTGGGCTCCATAAAG

 Table 2.1. Real-time PCR primer sequences.

^aMM: Mouse specific primers, Hs: Human specific primers.

Elisa

Serum Leptin and serum HMGB1 was estimated using ELISA kits from Abclonal Biotechnologies (Woburn, MA) following manufacturer protocol. Serum IL1 β was estimated using an ELISA kit from ProteinTech (Rosemont, IL) following manufacturer protocol.



Serum biochemistry tests

Biochemical analysis of mouse serum was done for ALT, urea Nitrogen, creatinine, cholesterol, triglycerides and glucose from the University of Georgia college of veterinary medicine.

2.2.7. Statistical analysis

Prior to initiation of the study, we conducted calculations for each experimental condition with appropriate preliminary data to confirm that the sample number is sufficient to achieve a minimum statistical power of 0.80 at an alpha of 0.05. All in vivo and in vitro experiments were repeated three times with 3 mice per group (N = 3; data from each group of three mice were pooled). Student's *t*-test was used to compare means between two groups at the termination of treatment. A one-way ANOVA was applied as needed, to evaluate differences among treatment groups followed by the Bonferroni post-hoc correction for intergroup comparisons.

2.3. RESULTS

2.3.1. Butyrate production is key to gut health in GW-chemical exposure and microbial dysbiosis

We have shown previously that GW chemical exposure caused a significant alteration in microbial population when compared to untreated controls with significant increases in Firmicutes-Bacteriodetes ratio, a trend that is uniformly observed in IBD, neuroinflammation and metabolic syndrome. The changes were consistent with the neuroinflammatory phenotype in the mouse model of GWI. On in-depth analysis of the microbial data, we found that GW-chemical exposed group showed a marked decrease in



Lactobacillus, and Bifidobacterium sp., the genus being responsible for producing the short chain fatty acid butyrate. Interestingly butyrate has been shown to attenuate IBD and resists proinflammatory changes in the small intestine (Fig. 1A) (p < 0.05). The two genus showed a > 5 fold (log scale) decrease in abundance (Fig. 1A). Butyrate priming through oral gavage and its presence during exposure significantly elevated the levels of Bifidobacterium, butyrogenic bacteria Roseburia sp. and Lactobacillus (i, ii and iii) (Fig. 1B) (p < 0.05) when compared to GWI alone with the first two genera showing an increase up to > 60% when compared to GWI. The percentages noted in the figure are compared to the overall abundance of all genus detected in the metagenomic analysis. The comparisons between GWI and GWI + NaBT groups were done using GWI as the base line. Such a comparison showed a > 60% increase of these genus in GWI + NaBT group when compared to GWI alone (Fig. 1B). The observations in Fig. 1A led to the rationale for using Butyrate as a viable molecule for attenuating microbiome-associated inflammatory phenotype and the subsequent changes observed in the GWI model. Butyrate exerts its actions via binding to the niacin receptor. GPR109A has been recently discovered to bind butyrate and stimulate the activation of Treg cells thus suppressing TH17 mediated proinflammatory events (43). Our results showed that there is a significant decrease in the protein levels of GPR109A in GW-chemical exposed group when compared to untreated controls (Fig. 2A and D) (p < 0.05). Butyrate presence in the intestine via feeding GWexposed mice through an oral gavage significantly increased GPR109A protein levels in the villi regions when compared to GWI-group (Fig. 2A and D) (p < 0.05) suggesting that butyrate presence resisted the decrease in GPR109A protein levels thus helping butyrate to



exert its actions in the dysbiosis-affected small intestine and restore gut-epithelial cell integrity and metabolic homeostasis.

2.3.2. Butyrate priming through oral route restores tight junction protein levels

The epithelial tight junction determines the paracellular water and ion movement in the intestine and also prevents uptake of larger molecules, including antigens, in an uncontrolled manner where Claudin-2 and Occludin play a major role and are perceived as a marker for leaky gut (45). Our results from immunofluorescence microscopy for the immunoreactivity of Claudin-2 showed a significant increase in GW-chemical exposed group when compared to untreated controls (Fig. 2B and E) (p < 0.05), thus confirming our previously reported data. Butyrate presence in GW-chemical exposed group showed a significant decrease in that group when compared to GW-chemical exposed group alone suggesting a parallel role of butyrate in Claudin-2 protein levels in the small intestine (Fig. 2B and E) (p < 0.05). The results also showed that butyrate priming nearly restored the Claudin-2 levels to untreated controls (Fig. 2B). Similarly, the protein level of another tight junction protein Occludin was significantly decreased in GW-chemical exposed groups (Fig. 2C and F) and sodium butyrate treatment significantly restored the levels of Occludin in the intestine. The results suggested that Butyrate may have a previously unconfirmed role in modulating Claudin-2 and Occludin proteins in the small intestine though ILIOAdependent repression of Claudin-2 has been shown (46)



2.3.3. Butyrate priming in the intestine attenuates proinflammatory phenotype in the intestine via a decrease in TLR4 activation

Since gut leaching was predominant in GWI mouse model and resulted in endotoxemia, we studied whether butyrate priming helped in attenuating the proinflammatory microenvironment in the small intestine (29). Results showed that butyrate administration through an oral route decreased TLR4 colocalization (as shown by white circles), a hallmark of its activation in GW-chemical exposed group when compared to GW-group alone (Fig. 3A and B) (p < 0.05). Notably, the results also confirmed our earlier observations of an increased TLR4 trafficking to lipid rafts in GW chemical exposed group when compared to untreated controls (Fig. 3A and B) (p < 0.05). TLR4 activation was followed by increased IL-1 β protein levels in the villi regions but not in crypts of GW chemical exposed group when compared to untreated controls (Fig. 4A and B) (p < 0.05). Also, butyrate priming significantly decreased the IL-1 β levels in the same regions when compared to GW-Chemical exposed group (Fig. 4A and B). Gene expressions of IL-1B, monocyte chemoattractant MCP-1 and TNF- α were significantly decreased in Butyrate administered group when compared to GW-chemical exposed group (Fig. 4C) (p < 0.05). Interestingly, serum IL-1 β significantly increased in GW-chemical exposed groups (66.71) \pm 1.98pg/mL) as compared to vehicle control group (39.95 \pm 1.8pg/mL) (Fig. 4D). However, Butyrate exposure to the GW-Chemical exposed mice showed a significant decrease in the serum IL-1 β (34.56 ± 1.26pg/mL) (Fig.4D). The results suggested that butyrate presence helped attenuate intestinal inflammation primarily from a TLR4 pathway however it could not rule out other parallel inflammatory pathways in the gut such as histone deacetylases.



Similar to pathogen-associated molecular patterns (PAMPs), that can trigger a proinflammatory response, sterile inflammation can be triggered by endogenous molecules from a necrotic or damaged cell that can activate several proinflammatory pathways including TLR4 (47). Such endogenous molecules are collectively called Damageassociated molecular patterns or DAMPs. We have shown previously that HMGB1 and leptin can be released from several organ systems and can trigger a proinflammatory cascade (48)(49)(50). We quantified the released HMGB1 and leptin in mouse serum using competitive ELISA techniques. Result showed that GW-chemical exposed groups had significantly higher levels of HMGB1 and leptin in the serum when compared to untreated controls (Fig. 5A and B) (p < 0.05). Butyrate priming significantly decreased the level of HMGB1 while a decrease in leptin levels was not significant when compared to GWchemical exposed group (Fig. 5A and B) (p < 0.05). The results suggested that circulatory DAMPs can be soluble mediators of ectopic inflammatory events distant to the small intestine while butyrate priming may attenuate such effects and help identify therapeutic targets in the systemic inflammatory phenotype seen in GWI.

2.3.4. Increased TLR activation in the liver is attenuated by butyrate priming in the intestine

TLR activation is observed in organ systems following gut dysbiosis (51)(52). TLRs especially TLR4, TLR2, and TLR5 have been shown to increase tissue inflammation (52). Interestingly, TLR-induced metabolic deregulation is increasingly seen as an important event in metabolic syndrome (53). Our results showed that GW chemical exposed group showed a significantly increased TLR4 activation (trafficking to lipid rafts) in the liver especially in the sinusoidal cells (white circles) (Fig. 6A and B). TLR4



trafficking significantly decreased in butyrate administered group when compared to GWchemical exposed group (Fig. 6A and B). Interestingly, butyrate administration markedly increased TLR4 protein (red) levels in the liver but could not be observed in the rafts of the membrane (yellow), a sign that TLR4 protein was increased but the activation was attenuated by butyrate administration (Fig. 6A). TLR5 a protein that is activated the following binding with flagellin also increased in the liver of GW exposed mice but was significantly decreased in the butyrate administered group (Fig. 6C and D). The results suggested that increased circulatory levels of DAMPs or a leaky gut-associated flagellin might have resulted in activation of TLR4 and TLR5 in the liver of GW chemical exposed group but was blocked by the presence of butyrate.

2.3.5. TLR4 activation is associated with metabolic changes and inflammatory response in the liver but the phenotypic liver injury is predominantly absent

The liver is the principal organ for gluconeogenesis, lipogenesis and cholesterol metabolism (54). Recent studies have put a great deal of emphasis on liver metabolic reprogramming in conditions of metabolic syndrome that include hepatic expression of lipid and glucose metabolism markers, hepatic insulin and leptin resistance (36)(55). Interestingly, the fatty liver disease is associated with a long-term metabolic alteration (often years to manifest) and inflammatory response in the liver till the disease phenotype surfaces and is rightly called a silent disease (56). We studied both the changes in hepatic metabolic markers and the inflammatory response arising from a TLR4 activation following GW-chemical exposure and microbial dysbiosis to ensure whether we can detect early responses in the liver that can manifest into liver disease years later. Results showed that hepatic SREBP1c, a molecule predominantly responsible for lipogenesis was



upregulated following GW-chemical exposure when compared to untreated controls (Fig. 7A) (p < 0.05) (57). Butyrate administration significantly decreased SREBP1c gene expression in the hepatic lobule when compared to GW-chemical exposed group (Fig. 7A) (p < 0.05). PPAR- α is a transcription factor and a major regulator of lipid metabolism in the liver (58). PPAR- α is activated under conditions of energy deprivation and is necessary for the process of ketogenesis (58). PPAR- α was significantly upregulated in the GWchemical exposed group when compared to untreated controls while butyrate administration significantly decreased and restored the PPAR- α levels when compared to GW-chemical exposed group (Fig. 7A) (p < 0.05). PPAR- γ is an important player in liver fat metabolism and is known to be increased in benign steatosis but is significantly downregulated in models of liver injury (36). Our results showed that PPAR- γ was significantly decreased in GW chemical exposed group when compared to untreated controls and butyrate priming reversed this downregulation when compared to GW-chemical exposed group (Fig. 7A) (p < 0.05). The results were in agreement with our previous studies in a liver metabolic disease that had similar decreases in PPAR- γ (36).

We have shown previously that liver metabolic disorders triggered by environmental contaminants can increase the expression of Phosphofructokinase (PFK) (36). PFK, a rate-limiting enzyme in the glycolytic pathway was significantly upregulated in GW chemical exposed group when compared to untreated controls while butyrate administration significantly restored the PFK levels (Fig. 7A) (p < 0.05). Hepatic class I glucose transporters (GLUT) have limited role in the liver but recent studies show their importance in hepatic disease states (59). We and others have shown that GLUT-1 and GLUT-4 are regulated by leptin and purinergic signaling and they are upregulated in fatty



liver disease primarily in hepatic stellate cells (50)(60). Our results show that both GLUT-1 and GLUT-4 were upregulated in the GW chemical exposed groups when compared to untreated controls however butyrate administration significantly decreased the GLUT-1 and GLUT-4 levels when compared to GW-chemical exposed groups (Fig. 7A) (p < 0.05). Further, to investigate the role of GW-Chemical exposure in exacerbating the inflammatory response in liver, hepatic mRNA expression profiles of interleukin (IL)-lp, monocyte chemotactic protein 1 (MCP-1), tumor necrosis factor (TNF)- α and Kupffer cell activation marker CD68 were analyzed. Results indicated that there was a significant increase in the mRNA expression profiles of IL-1 β , MCP-1, TNF- α and CD68 in GW Chemical exposed mice livers compared with vehicle treated mice livers (Fig. 7B) (p < 0.05). Interestingly, mice groups co-exposed with GW chemicals and sodium butyrate showed significantly decreased level of IL-1 β , MCP-1, and CD68 but not TNF- α . The results suggested a similar role of higher leptin and/or heightened inflammation in causing the increase but remained to be seen whether it was cell or organ specific.

The liver has multiple cell types and includes cells of epithelial, endothelial, fibroblast and macrophage lineages. They perform multiple functions including metabolic, cellular defense and wound healing. The liver lobule comprises of 90% hepatocytes which are epithelial in origin and is a center for most of the metabolic functions. We used human primary hepatocytes, primed with lipopolysaccharide (LPS) (concentrations found in our previous study (29) to study the effects of metabolic dysregulation if any due to GW-chemical exposure. Results showed that LPS primed hepatocytes showed a significant increase in lipogenesis mediator SREBP1c while butyrate co-exposure decreased these levels (Fig. 7C) (p < 0.05). Similar to our in vivo data, LPS primed hepatocytes showed a


significant increase in PPAR- γ gene expression when co-exposed to butyrate while LPS only or untreated controls showed no change in the PPAR- γ levels (Fig. 7C) (p < 0.05).

Hematoxylin and Eosin stains of liver tissue sections obtained from GW-chemical exposed group showed no signs of lipid accumulation or macrophage infiltration or Mallory body formation signifying the absence of advanced stage inflammatory foci or liver disease (Fig. 7D). Though histopathology of the liver section from each mice group clearly showed that there was no sign of liver damage or development of Nonalcoholic steatohepatitis, we resorted to clinical chemistry analysis for more detailed outcomes. To confirm such observations, we performed clinical chemistry analysis of mouse serum samples for ALT, BUN, creatinine, total cholesterol, triglyceride and serum glucose. The clinical chemistry data showed (Table 2) that there was no significant difference in serum ALT (showed a marked increase in GWI group but was not significant between groups), BUN, creatinine and total cholesterol upon GW Chemical exposure as compared to vehicle control group. However, co-exposure with sodium butyrate decreased the levels of ALT, BUN but not the total cholesterol (Table 2). The triglyceride levels were significantly increased in GWI chemical exposed mice groups as compared to untreated control groups, while co-exposure with sodium butyrate caused the triglyceride level to be decreased significantly when compared to GWI group (p < 0.05) suggesting a slow but incremental risk of fatty liver in the future (Table 2). The glucose levels showed no significant difference between groups (data not shown).



3.6. TLR4 drives the metabolic alterations in GW-chemical exposed liver

TLR4 induced downstream proinflammatory signaling has been found to aid in insulin resistance (61). Prolonged insulin resistance has been shown to cause metabolic disturbances in the liver, skeletal muscle and adipose tissue (53). Since microbiome associated gut leaching and systemic endotoxemia were reported in the mouse model of GWI and the present study found metabolic changes in the liver, we studied the direct role of TLR4 in causing the metabolic changes. The results showed that TLR4 knockout (TLR4 KO) mice had decreased TLR5 expression in the liver when compared to GW-chemical exposed group (Fig. 8A and B) (p < 0.05). TLR4 KO mice had significantly decreased expression of Class I glucose transporter GLUT-4, PFK, PPAR-α, and lipogenesis mediator SREBP1c when compared to GW-chemical exposed group while GLUT-1 showed no change in the expression suggesting GLUT-1 might not be regulated by TLR4 (Fig. 8C) (p < 0.05). The results suggested that TLR4 activation following systemic endotoxemia might be responsible for the ectopic metabolic alterations in the liver but is unable to present any significant changes in liver disease phenotype. The results also are in agreement with epidemiological studies where veterans deployed in GW don't report liver abnormalities in the clinics based on the typical symptoms.

2.4 DISCUSSION

Epidemiological studies have shown a strong correlation between GW toxicant exposures and cognitive/neurological complications but there are also reports of chronic fatigue, gastrointestinal disturbances and occasional cases of metabolic syndrome (3). Our study shows that microbial dysbiosis owing to GW-chemical exposure causes a significant



decrease in healthy gut bacteria like Bifidobacterium and Lactobacillus (62). Interestingly, they are a class of bacteria that generate butyrate in the gut (62). Recent studies have shown a beneficial effect of butyrate in preclinical studies involving colitis and IBD (39) The above results prompted us to use sodium butyrate administration through an oral route as a priming agent throughout the chemical exposure time so that a restored butyrate in the gut could prevent and prime the gut against the dysbiosis, inflammatory leaching, and generation of systemic mediators in the small intestine. Results also showed the role of butyrate receptor GPR109A, decreasing Claudin- 2 and decreasing TLR4 activation. We have shown recently that GW chemical exposure causes gut dysbiosis, the disintegration of gut membrane causing leaching and systemic endotoxemia (29) that eventually led to TLR4 activation.

We also showed a causal role of dysbiosis to the neuroinflammation in frontal cortex thus raising a possibility of the existence of a "Gut-Brain-Axis" in GWI similar to other pathological conditions (29). This axis may act in parallel to some of the direct toxic effects of GW chemical exposures on the brain tissue (63)(44). Sodium Butyrate priming, as shown in our data might reverse the pathology associated with GW chemical exposure since it restored gut health, reversed gut barrier integrity and decreased SI inflammation (decreased IL-1 β) while increasing the possibility of increased butyrate binding to GPR109A due to higher availability of this receptor in SI. Notably, butyrate priming also decreased the release of HMGB1 and leptin though slightly in circulation albeit from the intestinal epithelial cells but other sources like liver cannot be ruled out. The source might be the damaged epithelial cells in the small intestine since the potential generation of free



radical species has been shown before and oxidative stress in the intestinal epithelial cells and macrophages could release HMGB1 (64).

On the other hand, HMGB1 release due to gut integrity changes also causes oxidative stress and cell necrosis as have been reported in other studies (64). Though leptin is primarily released from adipocytes and liver, chemical/food-induced leptin release have been shown in the gut and has been traced in duodenal juice (65)(66). Thus, our finding of increased leptin in circulation following GW chemical exposure might be a result of the leaky gut or liver though the exact source remains to be determined at this time. The release of both leptin and HMGB1 and its modulation by butyrate priming in the gut points to the intestine as a source of these inflammatory mediators along with endotoxin and has tremendous implication determining ectopic/endocrine pathology of GWI.

In spite of well-coordinated symptom reporting in GWI about chronic fatigue in most of the studies, the causes of such chronic fatigue have been limited to abnormalities in neurological pathways or mitochondrial dysfunction without an organ-specific definition (67). Presence of symptoms related to metabolic syndrome or liver diseases is rare (68). Interestingly, fatigue is also associated with metabolic syndrome and various liver diseases (69)(70)(71). Though we hardly see literature reporting liver abnormalities in GWI, asymptomatic metabolic abnormalities in the liver (as evident in silent liver diseases like NAFLD), can contribute to chronic fatigue. These facts mentioned above led us to examine the liver pathology likely affected by higher circulatory mediators like endotoxin, leptin, and HMGB1. Owing to the tremendous role of the liver in carbohydrate metabolism, we focused on the role of circulatory HMGB1 and leptin on (a) hepatic TLR4 activation and (b) alterations in both lipid and carbohydrate metabolism.



Our results showed a significant increase in TLR4 trafficking to the lipid rafts, a hallmark of activation of the TLR4 pathway in the liver following exposure to GWchemicals. Also, there was a subsequent increase in TLR5 levels in the liver with both TLR4 activation and TLR5 levels showing decreases after butyrate priming. These results assume huge significance since TLR4 activation has been found to cause insulin resistance, uptake of free fatty acids for triglyceride production in macrophages and sterol biosynthesis (72)(53). On the other hand, stearic acid has been shown to promote TLR4 mediated inflammation (73) (74). Our results of increased expression of genes such as SREBP1c and PPAR- α which play a major role in liver lipogenesis (cholesterol biosynthesis and import) might be the result of the increased TLR4 activation and the subsequent cascade of events that alter liver metabolism following GW-chemical exposure. Interestingly, both SREBP1c and LXRs control lipid metabolism and it remains to be seen whether an increase in SREBP1c in the GW-chemical exposed liver was an adaptive way to suppress a chronic TLR4 activation thus mounting an anti-inflammatory response as is seen in some studies (75)(76). There are numerous reports which find increased glucose metabolism following NF-kB activity which is downstream of TLR4 activation (77).

Our results of increased expression of phosphofructokinase, a rate-limiting enzyme for glycolysis show that a TLR4 mediated mechanism might play a role in driving a glycolytic pathway in the liver. Notably, isolated hepatocytes when stimulated with TLR4 ligand LPS or GW-chemical Permethrin did not show an increase in PFK but exhibited a 3-fold increase in SREBP1c over vehicle control suggesting that hepatocytes along with macrophages may be targets of TLR4 activation thus playing a vital role in the reprogramming of lipid metabolism. Further, Class I glucose transporters GLUT-1 and



GLUT-4 was elevated in the GW-chemical exposed liver. The result assumes significance since inflammation in the liver has been shown to increase glucose uptake in hepatic stellate cells in a mouse model of fatty liver disease (50). Our results of decreased inflammation and subsequent metabolic disturbances in the liver following butyrate priming may shed some light on the inhibitory role of butyrate on histone deacetylase activity (78). Butyrate is a known HDAC inhibitor (78).

Studies show that butyrate can act as an HDAC inhibitor and decrease NFkß activity. We found a decrease in NFx β activity following butyrate administration but was found to be insignificant (data not shown). Also, butyrate can act independently of TL4 activation by inhibiting HDACs (79). Future studies should target the extensive role of butyrate in HDAC inhibiton in Gulf War Illness that may be independent of TLR4 activation. Importantly, the results of altered expressions of the metabolic genes failed to induce any histological changes that support inflammatory or metabolic liver disease following GW-chemical exposure. This is of high significance since liver diseases take years to manifest and most remain asymptomatic (silent) thus evading most clinical observations. It remains to be seen whether the unavailability of reports related to liver complications in GWI is due to the silent nature of the manifestations that are only limited to changes in the expressions of metabolic genes and would perhaps take years to show any phenotypic disease. Finally, our studies with TLR4 gene-deficient mice exposed to GW chemicals reversed the levels of TLR5 and expressions for SREBP1c, PPAR- α , PFK and GLUT-4 emphasizing the fact that TLR4 activation was indeed responsible for the metabolic reprogramming in the liver. Further, the reversal of TLR4-inducible systemic



release of DAMPS and metabolic changes in the liver bodes well for a potential use of this compound for a gut-targeted therapy in GWI veterans.

In summary, we show that GW-chemical exposure in mice and subsequent systemic inflammation following a dysbiosis in the gut could cause significant changes in the way the liver metabolizes lipid and carbohydrate with no detectable pathology while butyrate resists those changes. The study will help us advance our efforts to scrutinize clinical symptom reporting in the liver and re-evaluate the way we approach the therapeutic aspect of GWI by targeting multiple physiological pathways. Uses of short-chain fatty acids or probiotics can help in such pursuits.

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Figure 2.1. Gut microbiome alteration in mice model of Gulf War Illness (GWI). A. The proportional abundance of microbial genera: Graphical representation of the most abundant taxa of bacteria at the genus level. Groups compared are gulf war illness group (wild-type mice exposed to gulf war chemicals) (GWI, n = 3) and control group fed with vehicle (Veh, n = 3) (p-value: < 0.05). B. Percentage abundance of gut bacteria Bifidobacterium (i), Roseburia (ii), and Lactobacillus (iii) in a group of mice co-exposed with Gulf war chemicals and Sodium butyrate (GWI + NaBT, n = 3) as compared with GWI mice (n = 3) (p-value: < 0.05).





Figure 2.2. Change in gut microbiome in GWI alter niacin receptor (GPR109A) and tight junction proteins in the intestine. A. The expression pattern of butyrate and niacin receptor GPR109A was assessed by immunofluorescence microscopy. The representative images showed immunoreactivity of GPR109A in the distal part of the small intestine of veh control group of mice (veh, n = 3), gulf war illness group of mice (GWI, n = 3) and a group of mice co-exposed with GWI and sodium butyrate (GWI + NaBT, n = 3). B and C. The expression pattern of Claudin-2 and Occludin (tight junction proteins) was assessed by immunofluorescence microscopy. Tissue levels of Claudin-2 (B) and Occludin (C) in Vehicle control group of mice co-exposed with GWI and sodium butyrate (GWI + NaBT, n = 3) and a group of mice co-exposed with GWI and sodium butyrate (GWI + NaBT, n = 3) was assessed by immunofluorescent microscopy after labeling the protein with the red fluorescent secondary antibody and counterstained with DAPI (blue). D–F. The bar diagram shows the quantitative morphometric analysis of fluorescence intensities of GPR109A (D), Claudin-2 (E), and Occludin (F) immunoreactivity in the region of interest (ROI) in the small intestine. *(p < 0.05).





Figure 2.3. Sodium butyrate priming in a rodent model of GWI attenuates TLR4 activation in the small intestine. A. Immunofluorescence microscopy of small intestine showing TLR4 (red) trafficking to the lipid rafts (green) of the small intestine tissue, an essential process for TLR4 activation causing a co-localization of TLR4 in flotillin-rich rafts (yellow). Representative images of TLR4-flotillin co-localization in the small intestine of vehicle control group of mice (Veh, n = 3), gulf war chemical treated group of mice (GWI, n = 3) and a group of mice co-exposed with GWI and sodium butyrate (GWI + NaBT, n = 3) shown by white circles covering the yellow spots created by an overlay of red (TLR4) and green (Flotillin). Images were taken at higher magnification ($40 \times$ oil). B. Graphical representation of the quantitative morphometric analysis of colocalization events in the region of interest (ROI) in the small intestine. Images for analysis were randomly chosen in different microscopic fields. Data is represented as Mean \pm SEM and *(p < 0.05). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.





Figure 2.4. Sodium butyrate priming in a rodent model of GWI improves proinflammatory phenotype in small intestine mediated by the TLR4 pathway. A. Small intestine tissue slices were probed for IL-1 β immunoreactivity in vehicle control group of mice (Veh, n = 3), gulf war chemical treated group of mice (GWI, n = 3) and a group of mice co-exposed with GWI and sodium butyrate (GWI + NaBT, n = 3) using immunohistochemistry. Specific immunoreactivity to IL-1 β is evident by dark brown spots. B. Graphical representation of morphometric analysis of the IL-1β immunoreactivity in tissue slices. Data normalized against vehicle control (veh) *(p < 0.05). C. Quantitative real-time PCR (gRTPCR) analysis of inflammatory markers in the small intestine. mRNA expression of IL-1 β , MCP-1, and TNF- α was analyzed in the samples of vehicle control group of mice (Veh, n = 3), gulf war chemical treated group of mice (GWI, n = 3) and a group of mice co-exposed with GWI and sodium butyrate (GWI + NaBT, n = 3). Normalized mRNA expression is represented as a fold change of vehicle control (veh) on Y-axis. Data points represented with Mean \pm SEM *(p < 0.05). D. Graphical representation of serum IL-1 β in pg/mL of the samples of vehicle control group of mice (Veh, n = 3), gulf war chemical treated group of mice (GWI, n = 3) and a group of mice co-exposed with GWI and sodium butyrate (GWI + NaBT, n = 3)





Figure 2.5. Sodium butyrate treatment in GWI improves circulatory DAMPs. Similar to the pathogen-associated molecular pattern (PAMPs), the endogenous molecules called damage-associated molecular patterns or DAMPs (such as HMGB1) are linked with proinflammatory responses in distal organs. A. Western blot analysis of serum high mobility group box 1 protein (HMGB1) and serum adipokine leptin from samples of vehicle control group of mice (Veh, n = 3), gulf war chemical treated group of mice (GWI, n = 3) and a group of mice co-exposed with GWI and sodium butyrate (GWI + NaBT, n =3). Ponceau S staining was done to see the equal loading of serum proteins and used for normalization of protein expression. B-C. Graphical representation of morphometric analysis of HMGB1 and leptin western blot bands. The data was normalized to a total serum protein (Ponceau S). Y-axis depicts the HMGBl/Ponceau S ratio (B) and leptin/Ponceau S ratio (C) from Veh, GWI and GWI + NaBT groups. *p < 0.05 is considered statistically significant. D-E. Graphical representation of serum HMGB1 (D) and serum leptin (E) in ng/mL of the samples of vehicle control group of mice (Veh, n =3), gulf war chemical treated group of mice (GWI, n = 3) and a group of mice co-exposed with GWI and sodium butyrate (GWI + NaBT, n = 3).





Figure 2.6. Sodium butyrate treatment in a rodent model of GWI attenuates TLR4 activation in Liver. A. Immunofluorescence microscopy of liver slices showing TLR4 (red) trafficking to the lipid rafts (green), an essential process for TLR4 activation causing a co-localization of TLR4 in flotillin-rich rafts (yellow). Representative images of TLR4flotillin co-localization in the liver of vehicle control group of mice (Veh, n = 3), gulf war chemical treated group of mice (GWI, n = 3) and a group of mice co-exposed with GWI and sodium butyrate (GWI + NaBT, n = 3) shown by white circles covering the yellow spots created by an overlay of red (TLR4) and green (Flotillin). Images were taken at higher magnification ($60 \times$ oil). B. Graphical representation of the quantitative morphometric analysis of colocalization events in the liver. Images for morphometric analysis were randomly chosen in different microscopic fields. Data is represented as Mean \pm SEM *(p < 0.05). C. Tissue levels of TLR5 immunoreactivity in vehicle control group of mice (Veh. n = 3), gulf war chemical treated group of mice (GWI, n = 3) and a group of mice coexposed with GWI and sodium butyrate (GWI + NaBT, n = 3) mouse liver samples as observed by immunofluorescent microscopy after labeling the TLR5 protein with the green fluorescent secondary antibody and counterstained by DAPI (blue). D. The bar diagram shows the quantitative morphometric analysis of fluorescence intensities of TLR5 immunoreactivity in the liver tissue. Images for morphometric analysis of TLR5 were randomly chosen in different microscopic fields. Data is represented as Mean ± SEM *(p < 0.05)





Figure 2.7. TLR4 activation is associated with metabolic changes and inflammatory response in the liver but the phenotypic liver injury is predominantly absent. A. Quantitative real-time PCR (qRTPCR) analysis principle carbohydrate metabolic markers (PFK, GLUT-1, and GLUT-4) and fat metabolic markers (SREBP1c, PPAR- α , and PPAR- γ) in the liver tissue. mRNA expression of SREBP1c, PPAR- α , PPAR- γ and PFK, GLUT-1, GLUT-4 and B. mRNA expression analysis of inflammatory marker IL-1B, MCP-1, TNF- α , and Kupffer cell activation marker CD68 were analyzed in the liver sample of vehicle control group of mice (Veh, n = 3), gulf war chemical treated group of mice (GWI, n = 3) and a group of mice co-exposed with GWI and sodium butyrate (GWI + NaBT, n =3). Normalized mRNA expression is represented as a fold change of Vehicle control (veh) on Y-axis. Data points represented with Mean \pm SEM *(p < 0.05). C. mRNA expression of SREBP1c, PPAR- γ were analyzed in the primary human hepatocytes cells treated with lipopolysaccharide (LPS) and Co-treated with LPS and sodium butyrate (LPS + NaBT). Normalized mRNA expression is represented as a fold change of Vehicle control (Veh Cont) on Y-axis. Data points represented with Mean \pm SEM *(p < 0.05). D. Representative Hematoxylin and Eosin stained (H&E) images of liver sections showed liver pathophysiology of vehicle control group of mice (Veh, n = 3), gulf war chemical treated a group of mice (GWI, n = 3). Images were taken at $10 \times$ magnification.





Figure 2.8. TLR4 drives the metabolic alterations in GW-chemical exposed liver. A. Tissue levels of TLR5 in Gulf War chemical treated a group of wild-type mice (GWI, n = 3) and a group of TLR4 knockout mice (TLR4 KO, n = 3). Mouse liver samples as observed by immunofluorescence microscopy after labeling the TLR5 protein with green fluorescent secondary antibody and nuclear counterstaining by DAPI (blue). Images were taken at $60 \times$ (oil) magnification. B. The bar diagram showed the quantitative morphometric analysis of fluorescence intensities of TLR5 immunoreactivity in the liver tissue. *(p < 0.05). C. mRNA expression analysis of principle carbohydrate metabolic markers (PFK, GLUT-1, and GLUT-4) and fat metabolic markers (SREBP1c, PPAR- α) in the liver tissue of vehicle, GWI and TLR4 KO mice. Normalized mRNA expression is represented as a fold change of vehicle control (veh) on Y-axis. Data points represented with Mean ± SEM *(p < 0.05).



CHAPTER 3

DYSBIOSIS-ASSOCIATED ENTERIC GLIAL CELL IMMUNE-ACTIVATION AND REDOX IMBALANCE MODULATE TIGHT JUNCTION PROTEIN EXPRESSION IN GULF WAR ILLNESS PATHOLOGY²

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Running Title: Reactive enteric glial cells contribute to GWI pathology

Key words: dysbiosis, peroxynitrite, Rage, S100B, nitric oxide, p47 phox, SsnB, Sodium butyrate.

Author for correspondence:

*Dr. Saurabh Chatterjee, Ph.D. Environmental Health and Disease Laboratory, NIEHS Center for Oceans and Human Health on Climate Change Interactions, Department of Environmental Health Sciences, University of South Carolina, Columbia 29208 USA. Email: <u>schatt@mailbox.sc.edu</u>; Tel: 803-777-8120; Fax: 803-777-3391

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ABSTRACT

About 14% of veterans who suffer from Gulf war illness (GWI) complain of some form of gastrointestinal disorder but with no significant markers of clinical pathology. Our previous studies have shown that exposure to GW chemicals resulted in altered microbiome which was associated with damage associated molecular pattern (DAMP) release followed by neuro and gastrointestinal inflammation with loss of gut barrier integrity. Enteric glial cells (EGC) are emerging as important regulators of the gastrointestinal tract and have been observed to change to a reactive phenotype in several functional gastrointestinal disorders such as IBS and IBD. This study is aimed at investigating the role of dysbiosis associated EGC immune-activation and redox instability in contributing to observed gastrointestinal barrier integrity loss in GWI via altered tight junction protein expression. Using a mouse model of GWI and in vitro studies with cultured EGC and use of antibiotics to ensure gut decontamination we show that exposure to GW chemicals caused dysbiosis associated change in EGCs. EGCs changed to a reactive phenotype characterized by activation of TLR4-S100B/RAGE-iNOS pathway causing release of nitric oxide and activation of NOX2 since gut sterility with antibiotics prevented this change. The resulting peroxynitrite generation led to increased oxidative stress that triggered inflammation as shown by increased NLRP-3 inflammasome activation and increased cell death. Activated EGCs in vivo and in vitro were associated with decrease in tight junction protein occludin and selective water channel aquaporin-3 with a concomitant increase in Claudin-2. The tight junction protein levels were restored following a parallel treatment of GWI mice with a TLR4 inhibitor SsnB and butyric acid that are known to decrease the immunoactivation of enteric glial cells. Our study demonstrates that immune-



redox mechanisms in EGC are important players in the pathology in GWI and may be possible therapeutic targets for improving outcomes in GWI symptom persistence.

3.1. INTRODUCTION

Gastrointestinal disturbances are one of the most commonly reported chronic symptoms among veterans who returned from the Persian Gulf war of 1990–1991(3,80,81). About 14% of veterans who suffer from Gulf War illness (GWI) complain of some form of gastrointestinal (GI) problems such diarrhea, pain and gas etc. (80,81). According to (30), the most commonly reported gastrointestinal issues reported among United States and British Gulf war (GW) veterans were diarrhea, vomiting, and stomach problems. A study by (80) showed veterans of the Persian GW presented with diarrhea and had rectal hypersensitivity as did (82) who reported increased somatic hypersensitivity and pain among some GW veterans with GI issues.

Although the veterans report these symptoms, the prospective study by(81) did not find any significant clinical markers of disease pathology in blood or intestine tissue of deployed participants. Similarly, one of our own studies which reported metabolic reprogramming in liver as a result of leaky gut and endotoxemia did not find any biochemical markers of liver damage or altered metabolism in a mouse model of GWI. This was surprising because we had had previously shown that exposure to GW theater chemicals resulted in an alteration of gut microbiome and concomitant TLR4 mediated gastrointestinal and neuroinflammation with endotoxemia (29)(18). This elusive nature of



GWI is a strong reason for further studying underlying mechanisms of this condition in order to obtain effective therapies.

Of emerging interest in inflammatory gastroenterology are enteric glial cells (EGC) which reside in close proximity with the neurons of the enteric nervous system. These cells are similar in structure and physiology to astrocytes of the brain but are not excitable(83)(84). Initially, the principal function of EGC was thought to be providing mechanical support to enteric neurons. However, recent studies have shown that these cells play an important role in regulating the gastrointestinal microenvironment through several mechanisms, which have been extensively reviewed (83)(85)(86). EGC were found to significantly modulate gut homeostasis through release of growth factors (87) cytokines and prostaglandins (87)(86) but may also play a pathogenic role by contributing to nitrosative stress and proinflammatory cytokines when exposed to stressful or toxic stimuli in the gut. Moreover, studies have found that EGC have the ability to "sense" a change in microbiome from probiotic to pathogenic, possibly through toll like receptors (TLRs). A study by (88), found that adhesive E. Coli seem to activate a TLR-S100β/RAGE-iNOS signaling pathway in human EGC, while probiotic lactobacillus did not. Another study found that when EGC were treated with lipopolysacharrides (LPS), there was activation of TLRs with a release of S100B and nitric oxide (NO) (89). In this reactive state, EGC produce proinflammatory cytokines and chemokines e.g., (IL-1 β , TNF- α , MCP-1) and release of inducible NO which may contribute to oxidative stress in the gut (86,87,90)(84).

In irritable bowel syndrome (IBS) and inflammatory bowel disease (IBD), it is well known that an altered microbiome plays a significant role in the pathogenesis of the disease (90). In IBS for example, patients were found to have a decrease in abundance of



Bifidobacteria and Lactobacillus but an increased prevalence of pathogenic species like *Escherichia* spp., Shigellas, and several Clostridia(91). Furthermore, it has been observed that metabolic diseases e.g., diabetes and obesity also present with increased ratio of Firmicutes to Bacteriodetes (92,93). Studies concerning the mechanisms of these gastrointestinal diseases have found that change of EGC phenotype from homeostatic to pathogenic is a characteristic of these diseases (94–96). A study by Wang et al. reported a significantly increased expression of glial fibrillary acidic protein (GFAP), Tyrosine receptor kinase B and Substance P in the colon of IBS patients with a correlated increase in intestinal inflammation(97). Other studies show that a loss in EGC resulted in poor gastrointestinal health characterized by loss of gut barrier integrity (98).

Our previous research reported an altered microbiome in a murine model of GWI with increase in Firmicutes over Bacteriodetes and a decrease in several butyrogenic bacteria. This dysbiosis was accompanied by activation of TLR4, increased inflammation, a leaky gut, endotoxemia with release of damage associated molecular patterns (DAMPS) such as HMGB1 in gulf war chemical treated mice compared to controls (18,29,63). Interestingly, a recent study by Hernandez et al., showed that exposure to pyridostigmine bromide a known gulf war chemical exposure resulted in enteric neuronal and glial reactivity and inflammation (99).

This current study investigates the contribution of EGC in observed inflammatory phenotype which we and others have observed in GWI. We test the hypothesis that, the altered microbiome which results in increased pathogen associated molecular patterns (PAMPS) (e.g., LPS, flagellin and other immunostimulatory bacterial parts), leaky gut and increase in circulatory DAMPS (e.g., HMGB-1) in GW-chemical (Permethrin and



pyridostigmine bromide) treated mice results in a reactive EGC phenotype compared to mice treated with vehicle control treated mice and mice co-exposed with GW chemicals and antibiotics. Through this reactive EGC phenotype intestinal cells such as enteric neurons and epithelial cells might be further affected leading to a vicious cycle of consistent proinflammatory state. This constant proinflammatory state of intestinal cells might answer the persistence of gastrointestinal, systemic and neuro inflammation in gulf war illness. The study uses a murine model of GWI and *in vitro* studies with EGCs and intestinal cells to elucidate possible mechanisms to explain this observed inflammation observed in GWI.

3.2. MATERIALS AND METHODS

Pyridostigmine bromide (PB), Permethrin (Per), Sodium Butyrate, Sparstolonin B (SsnB), Corticosterone, Neomycin trisulfate hydrate, Enrofloxacin, Apocynin (APO), Phenylboronic acid (FBA) were purchased from Sigma-Aldrich (St. Louis, MO, United States). Lipopolysaccharides (LPS), LPS-RS (TLR4 inhibibitor) were purchased from Cayman chemical company (Ann Arbor, MI, United States), Rat High mobility group box 1 protein (HMGB-1) Rat (rec) (His) was purchased from Chimirigen, Mediatech, Inc. (Manassas, VA, United States), Anti-TLR4, anti-flotillin-1, anti-S100B, anti-GFAP, anti-ASCII and anti-Caspase 1, anti-TLR5, anti-3NT, anti-GP91, anti-P47phox, anti-NOS 2, anti-HMGB-1 and anti-aquaporin-3 primary antibodies were purchased from Santa Cruz Biotechnology (Dallas, TX, United States), anti-claudin 2, anti-TLR2 and anti-occludin were purchased from Abclonal Technology (Woburn, MA, United States) was while anti-NLRP-3, anti-RAGE were purchased from Abcam (Cambridge, MA, United States). Fluorescence-conjugated (Alexa Fluor) secondary antibodies, ProLong Gold antifade



mounting media with DAPI were purchased from Thermofisher Scientific (Grand Island, NY, United States), Apoptosis ApopTag[®] Fluorecein *in situ* detection kit from Millipore (Billerica, MA, United States), The Pierce LAL Chromogenic Endotoxin Quantification Kit from Thermo Scientific (Waltham, MA, United States) and Griess reagent system from Promega corporation (Madison, WI). All other chemicals which were used in this study were purchased from Sigma unless otherwise specified. Paraffin-embedding of tissue sections on slides were done by AML laboratories (Baltimore, MD, United States).

3.2.1. Animals

Adult wild-type male (C57BL/6J mice) were purchased from the Jackson Laboratories (Bar Harbor, ME, United States). Mice were implemented in accordance with NIH guideline for human care and use of laboratory animals and local IACUC standards. All procedures were approved by The University of South Carolina at Columbia, SC, United States. Mice were housed individually on 7090 Sani-Chip bedding from Teklad (Madison, WI, United States) and fed with 8904 irradiated chow diet from Teklad (Madison, WI, United States) at 22–24°C with a 12-h light/12-h dark cycle. All mice were sacrificed after animal experiments had been completed. Immediately after terminal anesthesia, mice's small intestine was collected and dissected for further experiments, while fecal pellets were collected from the colon and immediately stored in sterile Eppendorf tubes for microbiome analysis. The tissues were fixed using 10% neutral buffered formalin. Distal segments of small intestines were used for the staining and visualizations.



3.2.2. Rodent Model of Gulf War Illness (GWI)

Mice were exposed to Gulf War chemicals based on established rodent models of Gulf War Illness with some modifications (44,63). The treated wild-type mice group (GW) were dosed tri-weekly for 1 week with PB (2mg/Kg) and Permethrin (200 mg/kg) by oral gavage. After completion of PB and Permethrin dosages, mice were administered corticosterone intraperitoneally (i.p.) with a dose of 100μ g/mice/day for 5 days of the week for 1 week to represent war stress. The dose of corticosterone was selected from the study which exposed mice to 200 mg/L of corticosterone through drinking water. The i.p. dose of corticosterone had similar immunosuppression as examined by low splenic T cell proliferation (data not shown). The vehicle control group (CONT) of mice received saline injections and vehicle for oral gavage (6% DMSO) in the same paradigm. Similarly, another group of mice (GW + AB) were exposed to PB/Permethrin and corticosterone as in above mentioned dosages along with antibiotics (Neomycin 45 mg/kg and Enrofloxacin 1mg/Kg) thrice per week for 2 weeks for intestinal decontamination and obtaining gut sterility, while another group (AB) were exposed to antibiotics (Neomycin, 45 mg/kg and Enrofloxacin 1 mg/Kg) for 2 weeks. A fifth group of mice was treated with PB/Permethrin and corticosterone, but with Sodium butyrate (10 mg/Kg) and Sparstolonin B (SsnB) 3 mg/Kg (GW + SsnB + BT). We have shown before that SSnB is a potent TLR4 antagonist and Butyrate decreases intestinal inflammation in GWI.



Enteric Glial Cell Culture

Immortalized rat EGC were obtained from ATCC[®] (ATCCCRL-2690). Plated EGC were maintained in DMEM media supplemented with 10% FBS until treated. Cells were serum starved in DMEM supplemented with 1% FBS for 12 h and then exposed to vehicle control and chemicals. Cells were then treated with vehicle control-PBS (VEH), LPS (1 µg/mL), HMGB-1 (100 ng/mL), SsnB (10 µg/mL), Sodium butyrate (5 mM)and inhibitors FBA (100 µM) and Apocynin (100 µM) with either HMGB-1, LPS or antibiotics (neomycin and enrofloxacin cocktail) at different dilutions ranging from (1X to 1000X) (see Supplementary Figures S2–S7) for 24 h. After which the experiment was terminated and cells were harvested for mRNA extraction, gene expression analysis and protein expression studies. Nitric oxide production was estimated from culture fluids by measuring nitrite formation using the Griess assay.

Intestinal Epithelial Cell Culture

Immortalized rat intestinal epithelial cells (IEC-6) ATCC© CRL-1592, were obtained from ATCC. The cells were maintained in DMEM media supplemented with 10% FBS and 1x ITS until treated. Cells were serum starved in DMEM supplemented with 1% FBS for 12 h and then primed with LPS (100 ng/mL) for 12 h. Cells were then treated with culture fluids from EGC (above) for 24 h, then harvested for further analyses.



3.3.4. Microbiome Analysis

Microbiome was analyzed from fecal pellets and luminal contents collected from animals after sacrifice and sent to Second Genome for 16S rRNA sequencing. Microbial analyses were performed from isolated nucleic acids using the MoBio PowerMag Microbiome kit (Carlsbad, CA, United States), according to manufacturer's instructions. The microbiome data is in NCBI EBI under the accession number <u>PRJEB19474</u>.

3.3.5. Laboratory Methods

Immunofluorescence Staining

Paraffin-embedded distal part of the small intestine sections were deparaffinized using a standard protocol. Epitope retrieval solution and steamer were used for epitope retrieval of sections. Primary antibodies such as anti- GFAP, anti-S100β, anti-NOS2, anti-NLRP-3, anti-ASCII, anti-GP91, anti-P47phox anti-TLR4, anti-Flotillin, anti-aquaporin3 were used at the recommended dilution (1:200). Species-specific secondary antibodies conjugated with Alexa Fluor (633-red and 488-green) were used at advised dilution (1:300). Finally, the stained sections were mounted using Prolong gold anti-fade reagent with DAPI. Sections were observed under–Olympus fluorescence microscope using 20X, 40X or 60X objective lenses, or under confocal microscopy using Leica SP-8 with LasX-10 software at magnification of 63X.

Western Blot

Serum HMGB-1 levels were estimated from 35 μ g of denatured mouse serum protein, while TLR2, 4 and 5 were estimated from 25 μ g of denatured small intestine tissue



by a Western Blot analysis following standard protocols. Briefly, serum was concentrated and then diluted 1:5. Tissue protein or serum protein was then separated on a Novex 4– 12% bis-tris gradient gel and subjected to standard SDS-PAGE. The separated proteins were then transferred to a nitrocellulose membrane by a Trans-Blot Turbo transfer system. The membrane was then stained with Ponceau S, and then blocked with 5% BSA solution for 1 h, then incubated with primary antibody overnight at 4°C. Species-specific anti-IgG secondary antibody conjugated with HRP was used to tag primary antibody. ECL western blotting substrate was used to develop the blot The blot was then imaged using G:Box Chemi XX6 and subjected to densitometry analysis using Image J software.

Real-Time Quantitative PCR

Messenger RNA expression in small intestine and rat EGC was examined by quantitative real-time PCR analysis. Total RNA was isolated from each 15 mg small intestine tissue or 1×10^6 EGC by homogenization in Trizol reagent (Invitrogen, Carlsbad, CA, United States) according to the manufacturer's instructions and purified with the use of RNeasy mini kit columns (Qiagen, Valencia, CA, United States). cDNA was synthesized from purified RNA (1 µg) using iScript cDNA synthesis kit (Bio-Rad, Hercules, CA, United States) following the manufacturer's standard protocol. Real-time qPCR (qRTPCR) was performed with the gene-specific primers using SsoAdvanced SYBR Green Supermix and CFX96 thermal cycler (Bio-Rad, Hercules, CA, United States). Threshold Cycle (Ct) values for the selected genes were normalized against respective samples internal control (18S). Each reaction was carried out in triplicates for each gene and for each sample. The relative fold change was calculated by the 2- $\Delta\Delta$ Ct method. The sequences for the primers used for Real-time PCR are provided below in Table 1



TABLE 3.1. Rat primer sequences.

Rat_IL-1β	Sense: CCTCGGCCAAGACAGGTCGC Antisense: TGCCCATCAGAGGCAAGGAGGA
Rat_NLRP-3	Sense: TGCATGCCGTATCTGGTTGT Antisense: ATGTCCTGAGCCATGGAAGC
Rat_TNF-α	Sense: CAACGCCCTCCTGGCCAACG Antisense: TCGGGGCAGCCTTGTCCCTT
Rat_ASCII	Sense: GGACAGTACCAGGCAGTTCG Antisense: GTCACCAAGTAGGGCTGTGT
Rat_Caspase 1	Sense: GACAGGTCCTGAGGCCAAAG Antisense: AAAAGTTCATCCAGCAATCCATTT
Rat_MCP 1	Sense: TAGCATCCACGTGCTGTCTC Antisense: CAGCCGACTCATTGGGATC
Rat_18S	Sense: GGATCCATTGGAGGGCAAGT Antisense: ACGAGCTTTTTAACTGCAGCAA
Rat_NOS2	Sense: AGCAGAGTTGGTGCAGAAGC Antisense: GGGAATAGCACCTGGGTTTT
Rat_Claudin1	Sense: AGGTCTGGCGACATTAGTGG Antisense: CGTGGTGTTGGGTAAGAGGT
Rat-ZO-1	Sense: GGAAATGTGTAAATCACCTGGAAGA Antisense: CCAAAGAACAGAAGACCACCAAC
mm_18S	Sense: TTCGAACGAACGTCTGCCCTATCAA Antisense: ATGGTAGGCACGGCGATA
mm_Claudin1	Sense: TTTCGCAAA GCACCGGGCATACA Antisense: GCCACTAATGTCGCCAGACCTGAAA
mm_ZO-1	Sense: CCACCTCTGTCCAGCTCTTC Antisense: CACCGGAGTGATGGTTTTCT
mm_TLR2	Sense: ACCAAGATCCAGAAGAGCCA Antisense: CATCACCCGGTCAGAAAACAA
mm-TLR4	Sense: GGAGTGCCCGCTTTCACCTC Antisense: ACCTTCCGGCTCTTGTGGAAGC
mm-TLR5	Sense: TGTAAAGTACTGGTGCCCGTGTGT Antisense: ACTGCGCAAACATTCTGCTGGC
mm-NOS-2	Sense: CGCTGGCTACCAGATGCCCG Antisense: GCCATAGCGGGCTTCCAGC

The primer sequences are given as 5'-3' orientation; Sense: Forward primer. Antisense:

Reverse primer.



Endotoxin Level Detection by Litmus Amebocyte Lysate Assay

Bacterial endotoxins (EU/mL) were detected in mouse stool samples for mice which were treated with vehicle control, gulf war chemicals, and mice co-exposed with gulf war chemicals and antibiotics using the Pierce LAL Chromogenic Endotoxin Quantification Kit according to the manufacturer's instructions. Briefly, stool samples were obtained from mice and equal weights were homogenized in endotoxin free water. The supernatant was then collected, and heat inactivated at 70°C. This was then diluted 1:300 and endotoxins quantified.

Nitrite Estimation by Griess Assay

Nitric oxide release was estimated from the cell culture fluids by measuring nitrite formation immediately after the experiment was terminated. Nitrite was measured using the Griess reagent system from Promega corporation (Madison, WI, United States) and experiments were performed according to manufacturer's protocols.

Tunel Assay

DNA fragmentation was detected using the ApopTag[®] Fluorescein *in situ* detection kit from Millipore (Billerica, MA, United States) by following the manufacturer's standard protocol.

3.2.4. Statistical Analysis

All *in vivo* experiments were repeated three times (N = 3) with at least 3 mice per group (n = 9; data from each group of three mice were pooled). All *in vitro* and laboratory analysis experiments were repeated at least three or four times. The statistical analysis was



carried out by analysis of variance (ANOVA) (see Supplementary Table S1 for F-statistics) and a Turkey's HSD test to determine specific group differences. Further we performed an unpaired student *t*-test, using Graph pad prism software (GraphPad Software Inc., La Jolla, CA, United States). For all analyses *P < 0.05 was considered statistically significant and are marked as (*).

3.3 RESULTS

3.3.1 Altered Microbiome Is Associated with Increase in PAMPs and DAMPs in Gulf War Chemical Exposed Mice

Studies have shown an association between altered microbiome and increase in endotoxin levels in serum or feces (18). In this study, using the LAL assay, we estimated the endotoxin levels (PAMPS e.g., LPS) in the stools of mice which were treated with GW chemicals in comparison with the controls and found that there was a significant increase in endotoxin levels of mice treated with GW chemicals compared to the controls (Figure 1A; P < 0.05). We further assessed the amount of HMGB-1 which was released in the small intestine (Figures 1B,C) using immunofluorescence microscopy and in the blood circulation (Figures 1D,E) using a western blot analysis for serum HMGB-1 levels in the circulation. These high amounts of DAMPS and PAMPS in the body will reach the EGC and cause persistent glial reactivity.



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3.3.2. Altered Microbiome (Dysbiosis) Correlates with an increased expression of GFAP while gut decontamination with antibiotics decreases GFAP in Intestinal enteric glial cells

Enteric glial cells which are found in close proximity with enteric neurons are very abundant in the lamina propria, mucosa and sub mucosal regions of the small intestine (83). Using immunofluorescence microscopy, we found that there was a significant increase (P < 0.05) in GFAP expression in the small intestine of mice treated with GW chemicals (PB + BER) compared to the control group, and mice co-exposed to GW chemicals and antibiotics (Figures 2A,B). The increased expression of this protein has been associated with a reactive EGC phenotype in IBS and IBD (100); (87).

3.3.3. Altered Microbiome Correlates with a Reactive EGC Phenotype Through Activation of Toll-Like Receptors While Gut Decontamination via Antibiotic Usage Reversed Activation

Our previous studies showed that the altered microbiome was associated with an activation of Toll like receptors such as TLR4 and TLR5 in GW chemical treated mice (18). In this study we show that there was a significant increase mRNA (Figure 3A) and protein expression (Figures 3B,E) levels of TLR 2, 4, and 5 in mice which were exposed to gulf war chemicals (Permethrin and pyridostigmine bromide) compared to mice treated with only vehicle control and mice co exposed with GW chemicals and antibiotics (P < 0.05). We further detected a significant increased expression of TLR4 on EGC (TLR4/GFAP colocalizations) in GW chemical treated (GW) mice compared to Vehicle (CONT) and mice co-exposed with GW chemicals and antibiotics (P < 0.05) (Figures 3C,D) by immunofluorescence microscopy.



3.3.4 Altered Microbiome Associated Increased Expression of S100B in Reactive EGC Resulting in NOS-2 Expression While Antibiotic Usage for Depletion of Bacteria Reversed Such Activation

Using immunofluorescence microscopy, we found that there was a significant increase in the expression of S100B in GW chemical treated mice compared to mice treated with vehicle control and mice which were co exposed with GW chemicals and antibiotics (P < 0.05) (Figures 4A,C). We also found that there was a significant increase in RAGE expression in GW chemical treated mice in EGC by co-staining RAGE and GFAP (P < 0.05) compared to vehicle control treated mice. However, this increase was not significant for mice treated with both GW chemicals and antibiotics (Figures 4B,D).

We then studied the interaction between RAGE and S100B using immunofluorescence microscopy assuming that a co-localization of these two proteins would suggest complex formation and aid interaction. We showed that there was significant increase (P < 0.05) in S100 β /RAGE complex formation in GW chemical exposed mice (GW) and mice treated with vehicle control (CONT) or mice co-exposed with GW chemicals and antibiotics (GW + AB) (Figures 5A,B). In Figure 6A using RTqPCR we found that there was a significant increase in mRNA expression of inducible nitric oxide synthase in the small intestine of GW chemical treated mice and mice treated with vehicle control and mice co exposed with antibiotics and gulf war chemicals (n =9, p < 0.05). Further, we showed that there was a marked increase in inducible nitric oxide synthase (NOS-2) expression in the intestine tissues of mice treated with GW chemicals (GW) compared to mice treated with vehicle control (CONT) and mice co-exposed with GW chemicals and antibiotics (GW + AB), although this increase was not significant (P =0.075, and 0.11 respectively) (Figures 6B,C).



These results are evidence of activation of a TLR-S100 β /RAGE-iNOS pathway in association to an altered microbiome *in vivo* as suggested by a decrease of activation following the use of antibiotics to ensure gut decontamination.

3.3.5. Exposure to PAMPS (e.g., Lipopolysaccharides) and DAMPS (e.g., HMGB-1) Causes the Activation of TLR4-s100β/RAGE-NO Pathway in EGC

EGC can respond to an over balance in gut microorganisms by detecting PAMPS on/from the pathogen these bacteria such as cell wall, nucleic acid, flagella etc and mount an effective immune response through toll like receptors or NOD-like receptors (88).

Using immunofluorescence microscopy, we found that there was significant increase in TLR4 expression when we treated rat EGC with LPS or HMGB-1 (Figures 7A,B, P < 0.05). We also found an increase in S100 β /RAGE complex formation in LPS and HMGB1 treated cells compared to cells treated with vehicle control (P < 0.05) (Figures 8A–C). However, the difference between expression of these receptors was not significantly different between the cells treated with HMGB1 alone compared to those treated with HMGB1 + LPS-RS to block the TLR4 receptor. This indicates that possibly, DAMPS like HMGB1 can trigger inflammatory pathways in EGCs via several other receptors apart from TLR4.

We further evaluated the activation of inducible nitric oxide synthase and release of nitric oxide in the rat EGC treated with LPS or HMGB1 (Figures 9A–D). We used RT q PCR to evaluate the expression of nitric oxide synthase in rat EGC (Figure 9A). Our results showed a significant increase in the expression of iNOS in cells treated with LPS or HMGB1 compared to vehicle control (P < 0.01). We also found that there was a



significant increase in the protein expression of NOS-2 in LPS and HMGB-1 treated cells compared to cells treated with vehicle control only as evaluated by immunofluorescence microscopy (Figures 9B,C) (n = 3, P < 0.05). Finally, we investigated whether there was a release of nitric oxide by the cells (Figure 9D). We found that NO release was significantly increased LPS (2.6 fold) (*P < 0.05), but only a marked increase in cells treated with HMGB1 (P = 0.07) compared to vehicle control treated cells. Together, these results indicate the activation of a TLR-S100B/RAGE pathway that subsequently led to the increased production of nitric oxide, especially in response to microbial PAMPS.

3.3.6. Activation of NOX-2 and Increased peroxynitrite formation in Rat EGC

Studies have showed that NADPH oxidases are activated in response to pathogenic stimuli in human EGC (101). We found that treatment of EGC with LPS or HMGB1 significantly increased their expression of NOX-2 (Figures 11A–D) (P < 0.05). This activation was observed by immunofluorescence microscopy by detecting co-localization events (per 100 cells) between two key subunits of the NOX-2 enzyme complex. One in the lipid membrane GP91phox (labeled with green secondary antibody) and P47phox (labeled with red antibody). We found a significant increase in these co-localizations in LPS or HMGB1 treated cells compared to vehicle control treated cells (VEH) and cells treated with LPS/HMGB1 and Apocynin (LPS + APO) a NADPH inhibitor (Apocynin blocks the transport of p47 phox to the membrane) (*P < 0.05).

NOX-2 induced superoxide and nitric oxide react rapidly to form peroxynitrite, an indicator of redox related formation of tyrosyl radical and subsequent formation of tyrosine nitration. We also observed that there was a significant increase in formation of



peroxynitrite (shown by increased 3- nitrotyrosine formation) in LPS or HMGB1 treated EGC compared to Vehicle control (VEH) treated and LPS or HMGB1 and apocynin (LPS + APO) or (HMGB1 + APO) (Figures 12A–D) treated EGC (P < 0.05, n = 6).

3.3.7. Oxidative Stress Triggers Activation of NLRP-3 Inflammasome Which Results in Increased Inflammation

Reactive oxygen species (ROS) can trigger activation of inflammasomes resulting in caspase 1 mediated cleavage of Il-1 β and IL-18 proinflamatory cytokines (102). Our results (Figures 13A,B) showed significant increase in mRNA expression of NLRP-3, Caspase-1, IL-1 β and TNF- α in LPS treated EGC but not HMGB1 treated cells which only showed an increase in TNF- α expression compared to the vehicle control (P > 0.05). Treatment of EGC with LPS and apocynin (LPS + APO) and FBA (LPS + FBA) significantly decreased the observed mRNA expression (P < 0.05).

We then investigated the protein expression of NLRP-3 and ASCII and adaptor protein of NLRP-3 in rat EGC using immunofluorescence microscopy. We found that cells treated with LPS but not HMGB1 treated cells showed a significant increase in NLRP-3 and ASCII complex formation compared to Vehicle control treated cells (VEH) indicating activation of the NLRP-3 inflammasome, when EGC encounter PAMPS. We further found that treatment of cells with LPS and FBA (LPS + FBA) showed a significant decrease in NLRP-3 protein activation (P < 0.05), (Figures 13C–E) suggesting the role of NOX-2 derived peroxynitrite as a candidate for the inflammasome formation.



3.3.7. Increased DNA Fragmentation in Reactive Rat EGC Following Stimulation With LPS and Its Dependence on NOX-2-Induced Oxidative Stress

Increased pathogenic stimuli were found to initiate apoptosis in EGC (101). We also investigated the fate of these reactive EGCs when continually exposed to PAMPs or DAMPs though a tunel assay to detect fragmented DNA.

We found that LPS or HMGB1 treated cells showed a significant increase in colocalization events per 100 cells compared to cells treated with only vehicle control (P = 0.043) (Figures 14A–D). There was a significant decrease in tunel events when cells were treated with LPS and Apocynin but not FBA. And when cells were treated with HMGB1 and apocynin or FBA, there where was no significant decrease in tunel events.

3.3.8. Reactive EGC Contribute to Inflammation and Intestinal Barrier Integrity in Small Intestine: Gut Decontamination by Antibiotics and Blocking EGC Immune Activation Restores Gut Barrier Protein Levels in GWI Mice

Cytokines, ROS and growth factors etc affect tight junction proteins, water channels and processes such as differentiation, apoptosis etc. (86). In this study, we showed that when LPS primed intestinal epithelial cells were treated with culture fluid from EGC, there was a significant increase in mRNA expression of proinflammatory cytokines in IEC-6 cells (Figure 14E) (P < 0.05). LPS primed IEC-6 cells which were treated with culture fluids from EGC treated with LPS (LPS-SN) showed a significant increase in mRNA expression of IL-1 β , MCP1 and TNF- α when compared to the vehicle control (VEH) (P <0.05). LPS primed cells treated with culture fluids from Vehicle control treated EGC showed a significant decrease in MCP-1 expression (P < 0.05) and a marked decrease in


TNF- α but no significant decrease in IL-1 β expression compared to the LPS primed IEC-6 cells treated with culture fluids from LPS treated EGC.

To ensure that EGC immune activation via an altered microbiome plays a significant role in gut barrier protein expression in the intestine, we studied the GW chemical treated mice for their protein levels of aquaporin, a selective water channel, occludin and claudin-2. Results showed that administration of antibiotics was associated with significantly restored the levels of aquaporin 3 in the intestine of GWI-treated mice when compared to GW-treatment (Figures 15A–C). Levels of occludin were also restored when compared to controls but were significantly elevated when compared to GW-mice only (Figures 15D–F). Claudin-2 levels have been found to be increased in association with gut integrity loss. Our results showed that use of antibiotics significantly decreased the levels of claudin-2 in antibiotic treated mice when compared to GWI-mice alone (Figures 15G–I).

To show that EGC immune activation as a result of TLR4 and specific inflammation was responsible in part in causing differential expression of tight junction proteins that may play a significant role in gut barrier protein integrity loss, we chose to use two significant compounds that has been studied for their TLR-antagonism (SSnB) and anti-inflammatory properties (Sodium Butyrate-BT) specifically in the intestine. Results showed that a combined use of TLR4 antagonist and butyrate markedly increased aquaporin levels (Figure 15A) in the small intestine when compared to GW-mice while levels of Claudin-2 were significantly decreased in the small intestine following SSnB + BT administration when compared to the same group (Figure 15D). Occludin which is decreased in GW mice and plays a significant role in maintaining gut barrier integrity was



also restored to normal levels in the diseased mice following administration of SSnB + BT (Figure 15G). The results suggested that blocking TLR4 and subsequent immunoactivation that resulted in a reactive EGC phenotype in mice due to dysbiosis can be reversed by the use of these antagonists. Also, the results show that reactive EGCs might have a significant role in causing gut barrier dysfunction following activation via release of PAMPs and DAMPs and can be a cause of symptom persistence in GWI.

3.3.9 Reactive EGC Modulate Tight Junction Proteins and Aquaporins in Intestinal *Epithelial Cells*

We investigated the hypothesis that EGC which are exposed to DAMPS (e.g., HMGB-1) and PAMPS (e.g., LPS) modulate intestinal tight junction proteins and selective water channels by treating LPS primed IEC-6 cells with culture fluids freshly collected from EGC which have been treated with LPS (LPS-SN), HMGB1 (HMGB1-SN), vehicle (VEH-SN), LPS + SsnB + Butyrate (LPS + SsnB + BT) or HMGB1 + SsnB + Butryate (HMGB1 + SsnB + BT). Protein expression was studied by immunofluorescence microscopy observed at a total magnification of 400X; scale 10 µm. We found that expression of aquaporin-3 was significantly increased when IEC-6 cells were treated with LPS-SN, while when they were treated withHMGB1-SN the expression was significantly decreased compared to IEC 6 cells only treated with vehicle control (VEH) (n = 3, p < 10.05). IEC-6 cells treated which were treated with culture fluids from vehicle control treated EGC (VEH-SN) showed only a slight increase in aquaporin-3 protein expression, while IEC-6 cells treated with inhibitors SsnB and Butyrate together with LPS or HMGB1 restored expression of aquaporin 3 almost back to similar levels as IEC-6 cells treated with vehicle control (VEH) (Figures 16A,B). Claudin 2 expression increased significantly when



IEC-6 cells were treated with culture fluids from EGC treated with LPS-SN (n = 3; p < 10.05), but only markedly, when with HMGB1 treated culture fluids (HMGB1-SN). And when IEC-6 cells were treated with culture fluids for EGC treated with SsnB and butyrate together with HMGB1 or LPS, claudin 2 levels were decreased similar to vehicle control treated cells (VEH) (Figures 16C,D). Higher magnification (630X and scale bar 20 µm) images taken under confocal microscopy are included to show the localization of this protein in the membrane (Figure 17A). IEC-6 cells that were treated with culture fluids from EGC treated with LPS (LPS-SN) or HMGB1(HMGB1-SN) showed a significant decrease in occludin protein expression compared to vehicle control treated IEC-6 cells (VEH) (Figures 16E,F). Treatment with culture fluids from EGC which had been treated with LPS or HMGB1 together with inhibitors (SsnB and butyrate) showed a restoration in occludin levels compared to the control (VEH), although the effect was stronger in IEC 6 cells treated with LPS + SsnB + BT compared to HMGB1 + SsnB + BT treated EGC culture fluids. Higher magnification (630X and scale;20 µm) images taken under confocal microscopy were used to show the localization of this protein in the membrane (Figures 17A,B). The tubulin staining used in occludin images to show the extent of occludin traversing the tubulin outline thus signifying their apical membrane localization as widely perceived (Shown by white arrows).

These results indicate that reactive EGC are strong players in modulating tight junction protein expression through production of factors that may influence gut barrier integrity.



3.4 DISCUSSION

Our results propose a possible molecular mechanism to explain the altered microbiome associated inflammation in a cellular level and poor gastrointestinal health which we observed in our studies on GWI (18,29). The results reported in this study are an advancement to our previous reported work in Gulf War illness pathology. Since gut sterility by antibiotics reversed immunopathology in GWI we used the same approach to correlate the observed gut dysbiosis with EGC immunoactivity. We found a novel role of altered microbiome in causing a reactive EGC phenotype characterized by activation of toll-like receptors, RAGE/S100B and increased expression of nitric oxide synthase. This pathway contributes to NADPH oxidase mediated generation of ROS which trigger inflammation. The mechanism of NOX2 mediated inflammation is well accepted as we and others have shown the role of peroxynitrite mediated inflammation in liver, kidney and gastrointestinal disturbances (29). Further, we propose that this increased inflammation and ROS may result in enteric gliopathy and a later episode of enteric neuropathy although this needs to be investigated further and is a speculation at this point. With continued production of proinflammatory cytokines and other destructive factors (e.g., ROS) by reactive EGC, the entire or part of the epithelial barrier in the gut might lose its integrity. This hypothesis is further strengthened by our results from the supposed blockage of TLR4 and inflammation by using SSnB and Butyrate (thus blocking EGC immune activation), further exacerbating the observed gastrointestinal pathology in gulf war illness. This explanation not only helps us understand the acute phase of gulf war associated gastrointestinal inflammatory disorders, but also could explain why these symptoms may



persist for long since a vicious cycle might exist following a continuous assault on the intestinal epithelial cells.

Enteric glial cells are important regulators of the gastrointestinal tract health. They can influence the gut microenvironment both positively or negatively depending on surrounding conditions (84,89). Remarkably they can respond to the presence of bacteria pathogens through this TLR-RAGE/S100β-iNOS pathway. This is through the recognition of bacterial parts such as cell wall, high bacterial populations, DNA etc via toll like receptors. With an altered microbiome, there is proliferation of certain bacterial species at the expense of others and this change upsets the natural healthy balance in microbiome. This disruption happens due to several stressful stimuli e.g., infection, diet or exposure to chemicals such as in the case of gulf war illness.

Our previous studies have clearly shown that exposure to GW chemicals indeed results in altered microbiome (18, 29).We found increase the an in Firmicutes/Bacteriodetes ratio with significant increases in several Firmicutes genera in gulf war chemical treated mice compared to the vehicle controls. Further we found an associated loss in bacteria populations such as Bacteroides, Oscillibacter and Ruminiclostridia. Increased abundance of Bacteroides for example are associated with healthy gastrointestinal states (103), while Oscillibacter and Ruminiclostridia have also been shown to be abundant in healthy controls in studies of Crohn's disease. The decline in beneficial microbiota may have allowed for the proliferation of several bacteria populations at genus level, which usually exist in low percentages. There was a rise in several Coriobacteria, Bacilli and Verrucomicrobia bacteria. These have all been associated to increase in IBS and IBD (91). This upset balance of bacteria population



dynamic results in normally benign bacterial populations to become pathogenic and could cause EGC to change to a reactive phenotype through toll-like receptor signaling (104).

Both the altered microbiome and reactive EGC phenotypes have been linked to several diseased states of the gut such as IBS, IBD, gut hypersensitivity etc. (92). However, there is scanty information concerning the true mechanism of how they contribute to these diseases. Our current study showed a correlation between altered microbiome and a reactive EGC phenotype in small intestine. Mice treated with GW chemicals (GW) had a higher expression of GFAP a protein whose increased expression has been associated with IBS, S100β/RAGE complex formation and finally an increase in nitric oxide synthase activity in EGC. These proteins were not increased in vehicle control treated mice (CONT) and their expression was significantly less in mice treated with GW chemicals and antibiotics (GW + AB). This emphasizes the role of microbiome in contributing to EGC reactive phenotype. Though we have used antibiotics to ensure gut contamination or sterility, the use of such approach may not be ensuring complete gut sterility in mice. Often the use of such antibiotics can selectively lead to bacteriostatic effects in healthy fauna while elevating the abundance of harmful bacteria in the gut. The use of germ free mice is the best approach for conducting studies where the endpoint is to assess the role gut bacteria in the pathology. Though it has to be admitted that antibiotic use for ensuring gut sterility is a standard approach where use of germ-free mice is a constraint. The use of antibiotics in our study is thus a limitation and needs further corroborative studies in future using the germ-free model.

Mechanistically, we showed that the increased activation of NOX-2 following an altered microbiome and associated activation of the EGCs in GW chemical exposed mice



plays a significant role in contributing to the observed intestinal inflammatory phenotype. NOX-2 in EGC and in adjacent intestinal cells, participates in oxidative stress which results from the increase in nitric oxide production. We found that the generated ROS triggered activation of the NLRP-3 inflammasome which further caused increase in inflammation and programmed cell death as showed by increased DNA fragmentation (tunel assay) in rat EGCs stimulated with LPS and/or HMGB1.

This increased inflammation and direct loss in enteric glia has been reported as in Chron's disease (84,94). The reactive inflammatory glial phenotype is detrimental to the health of the gastrointestinal tract because it produces destructive factors which interact with surrounding cells in the intestine e.g., intestinal epithelial cells, enteric neurons etc. In our study we showed that when EGC conditioned media was applied to primed epithelial cells, there was an increase in proinflammatory cytokine expression such as IL1B, MCP-1 and TNF- α which can be conducive to a leaky gut microenvironment. Furthermore, we observed that a reactive EGC phenotype can also have detrimental effects on the EGCs as shown by increased DNA fragmentation and cell death through the increased inflammation and ROS generated. This ultimately may result in programmed cell death in glia by pyroptosis or apoptosis as shown elsewhere (101). The general loss in enteric glia could lead to suboptimal functioning of enteric neurons and even enteric neuropathy (105). This mechanism could be a possible explanation for the symptoms of GWI which continue to persist for 25 years though the present report does not study the role of an activated EGC on intestinal neurons. However, the effect of the reactive EGCs on intestinal epithelial cell barrier integrity can be profound as a blockade of the EGC activation mechanisms by SSnB and butyrate prevented protein alterations in the tight junctions. The results are also



interesting since we observe a cyclical pattern of epithelial cell damage-activation of EGCs and a link to altered expression of tight junction proteins such as claudin-1,2, occludin or ZO-1 that may contribute to gut-leakiness, that eventually might fuel a continuous persistence of inflammation in the local intestinal microenvironment.

3.5. CONCLUSION

We report that EGC are important players in GWI gastrointestinal disease pathology and respond to the altered microbiome in the host gut by converting to a reactive phenotype which greatly affects the healthy functioning of the gastrointestinal tract. This reactive phenotype significantly contributes to oxidative stress which further triggers inflammation, loss of gut barrier integrity and possibly death of enteric glia and enteric neurons, although further investigations need to be carried out to confirm these neuronal effects. Further, these findings provide insights into how a possible altered microbiome may be contributing to the observed GWI intestinal epithelial cell inflammatory phenotype by destabilizing the redox status of glial cells and adjacent epithelial cells via NOX-2 mediated peroxynitrite generation, inflammasome activation and release of pro inflammatory cytokines. It has to be further realized that more concrete evidence will be needed that involves germ free mice to conclude with certainty that microbiome alterations definitely dictate the observed EGC effects and therefore remains a limitation in this study.

Antibiotic use for gut sterility should be overcome with more stringent experimental designs such as germ-free mouse models and gnotobiotic mice. Nevertheless, the present evidence will therefore be valuable to consider EGC nitric oxide production,



formation of peroxynitrite, a redox signaling intermediate and inflammation pathways as therapeutic targets in gulf war illness.





Figure 3.1. Altered microbiome associated increase PAMPS and DAMPS

A and B. Stool endotoxin levels. Endotoxin levels in stool samples were determined by the LAL assay. Graph A show the levels of endotoxin in Endotoxin Units (Eu) in vehicle control (CONT, n=9) treated mice, gulf war chemical exposed mice (GW, n=9) and mice co-exposed with antibiotics (GW+AB, n=9) (*P<0.05). C. Expression of HMGB-1 in small intestine tissues. Expression of HMGB1 was assessed by immunofluorescence microscopy at (total Magnification 200X; scale bar 100 µm). Images show immunoreactivity in the distal part of the small intestine for vehicle control treated mice (CONT, n=9), GW chemical treated mice (GW, n=9) and mice co-exposed with gulf war chemicals and antibiotics (GW, n=9). C. Quantitative morphometric analysis of HMGB-1 immunoreactivity represented as arbitrary light units in the region of interest (%ROI) *P<0.05. **D**. Serum High mobility group box 1 (HMGB1) levels. Serum HMGB-1 levels were estimated by western blot analysis for mice treated with control (CONT, n=3), Gulf war chemical exposed mice (GW, n=5) and mice co-exposed to antibiotics and GW chemicals (GW+AB, n=3). Ponceau red staining was used for normalization of protein. E. Quantitative morphometric analysis of western blot bands normalized against total Ponceau. The Y axis shows HMGB-1/Ponceau S ratio. (*P<0.05).





Figure 3.2. Altered microbiome induced change in EGC phenotype to a reactive phenotype.

A. Expression GFAP. Expression of GFAP was assessed by immunofluorescence microscopy at (top panel magnification 200X; scale 100 μ m and bottom panel magnification 600X; scale 20 μ m). Images show immunoreactivity of the distal part of the small intestine for vehicle control treated (CONT, n=9), gulf war chemical treated mice (GW n=9), mice treated with only antibiotics (AB, n=4) and mice co-exposed with gulf war chemicals and antibiotics (GW+AB n=9). **B.** Quantitative morphometric analysis of GFAP immunoreactivity represented as arbitrary light units as observed in the region of interest (%ROI). (*P<0.05).





Figure 3.3. Expression of Toll-like receptors in small intestine and EGC

A and B General mRNA and protein expression levels of toll-like receptors TLR2, TLR4 and TLR5 in small intestine of mice treated with vehicle control (CONT, n=9), gulf war chemical treated mice (GW, n=9) and mice treated with antibiotics only (AB, n=4) and mice co-exposed with GW chemicals and antibiotics (GW+AB, n=9). mRNA expression was determined by RTqPCR, while protein expression was determined by western blot analysis. E. Quantitative morphometric analysis of western blot bands normalized against β -actin The Y axis shows protein/ β -actin ratioResults are expressed as mean +/- SEM for n=9. (* P<0.05). C. Tissue level expression of TLR4 in EGC in small intestine. Expression of TLR4 in EGC was observed by in dual labelling of TLR4 and EGC cells marker GFAP via immunofluorescent microscopy visualized at (top panel magnification 200X; scale 100 μ m and bottom panel magnification 600X; scale 20 μ m). in small intestine tissues obtained from mice treated with vehicle control (CONT, n=9); mice treated with GW chemicals (GW n=9) mice, mice treated with antibiotics only (n=4) and co-exposed with GW chemicals and antibiotics (GW+AB, n=3). D Quantitative morphometric analysis of immunoreactivity of GFAP/TLR4 (yellow) is represented as colocalizations events per field from randomly chosen microscopic fields. (*P<0.05)





Figure 3.4. Expression of S100B and RAGE in the small intestine

A. Expression levels of S100B in EGC. Protein expression levels of S100B in EGC was determined by co-staining S100B with GFAP and assessed by immunofluorescence microscopy at (top panel magnification 200X; scale 100 µm and bottom panel magnification 600X; scale 20 µm). Images show immunoreactivity in distal part of the small intestine for vehicle control treated mice (CONT, n=9), gulf war chemical treated mice (GW, n=9) and gulf war chemical treated mice, mice treated with antibiotics only (AB, n=4) and mice co-exposed with antibiotics (GW+AB, n=9). C. Quantitative morphometric analysis of immunoreactivity of GFAP/S100B (vellow) is represented as colocalizations events per field from randomly chosen microscopic fields (% ROI). *(P<0.05). **B.** Expression of RAGE in EGC Protein expression levels of RAGE in EGC was determined by co-staining RAGE with GFAP and assessed by immunofluorescence microscopy at (top panel magnification 200X; scale 50 um and bottom panel magnification 600X; scale 20 μm). . Images show immunoreactivity in distal part of the small intestine for vehicle control treated mice (CONT, n=9), gulf war chemical treated mice (GW, n=9) mice treated with antibiotics only (AB, n=4) and gulf war chemical treated mice coexposed with antibiotics (GW+AB, n=3). **D.** Corresponding quantitative morphometric analysis of immunoreactivity of GFAP/RAGE (yellow) is represented as colocalizations events per field from randomly chosen microscopic fields (% ROI). *(P<0.05)





Figure 3.5. Formation of S100B/RAGE complex in small intestine

A. S100B(Green)/RAGE(Red) complex formation expression in EGC in small intestine tissues. Protein expression levels were assessed by immunofluorescence microscopy of tissues at (top panel magnification 200X; scale 50µm and bottom panel magnification 600X; scale 20 µm). Images show immunoreactivity the distal part of the small intestine for gulf war chemical treated mice (GW, n=9), vehicle control (CONT, n=9), mice treated with antibiotics only (n=4) and mice co-exposed GW chemicals and antibiotics (GW+AB, n=9). **B.** Quantitative morphometric analysis of immunoreactivity for S100B/RAGE is represented as colocalization events per field for randomly chosen fields (% ROI). (*P<0.05).





Figure 3.6. Activation of NOS-2 in small intestine.

A. NOS-2 mRNA expression in the small intestine of intestine of mice treated with vehicle control (CONT, n=9), gulf war chemical treated mice (GW, n=9), mice treated with antibiotics only (AB, n=4) and mice co-exposed with GW chemicals and antibiotics (GW+AB, n=9. *P<0.05) was determined by RTqPCR. **B.** Protein expression levels of NOS-2 in enteric glial cells was determined by immunofluorescence microscopy of tissues and imaged at (top panel magnification 200X; scale 50µm and bottom panel magnification 600X; scale 20µm). **C.** Quantitative morphometric analysis of immunoreactivity for GFAP/NOS-2 represented as colocalization events per field for randomly chosen fields (% ROI)





Figure 3.7: Exposure to DAMPS and PAMPS cause activation of TLR4 in EGC

A. TLR4 activation in rat Enteric Glial Cells (EGC) (n=6). Colocalization of TLR4 and Flotillin in EGC treated with either vehicle control (VEH) or LPS, LPS+LPS-RS, HMGB-1 or HMGB1+LPS-RS determined by immunofluorescence microscopy at (magnification 400X; scale 50μ m).**B.** Quantitative morphometric analysis of TLR4/Flotillin colocalizations between red and green immunoreactivity were determined for every 100 cells per field. Fields were chosen randomly, and colocalizations represented as immunoreactivity in the region of interest (%ROI). (*P<0.05)





Figure 3.8. EGC exposed to LPS or HMGB-1 change to a reactive phenotype

A. S100B/RAGE complex formation in rat Enteric Glial Cells (EGC) (n=6). Colocalization of S100 β and RAGE in EGC treated with either vehicle control (VEH) or LPS, LPS+LPS-RS, HMGB1 or HMGB-1+HMGB-1+LPS-RS was determined by immunofluorescence microscopy at (total magnification 400X and scale 20 μ m) **B and C** Quantitative morphometric analysis of S100B/RAGE complex formation. Colocalizations between red and green immunoreactivity were determined for every 100 cells per field for randomly chosen fields and represented as immunoreactivity in the region of interest (% ROI). (*P<0.05)





Figure 3.9. Activation of inducible nitric oxide synthase by LPS or HMGB-1

A. mRNA expression of NOS-2 in EGC exposed to vehicle control (VEH), LPS, LPS+LPS-RS or HMGB-1 and HMGB-1+LPS-RS (n=6) expressed as x fold of the vehicle control. mRNA expression was determined by qRTPCR. **B.** NOS-2 protein expression in the cells was detected by staining with green fluorescent antibody and counterstained with DAPI (blue) and viewed at (total magnification 400 μ m and scale 20 μ m). Fluorescence expression of NOS-2 in rat EGC per 100 cells in different fields and represented as immunoreactivity in the region of interest (% ROI). **D**. Nitrite concentration in EGC (n=8). Nitric oxide production in EGC supernatants was estimated by griess assay from freshly harvested supernatants. Nitrite concentration is reported as X fold increase over the vehicle control (VEH) (*P<0.05)





Figure 3.10. NADPH oxidase 2 and peroxynitrite mediated oxidative stress in vivo.

A. NOX-2 activation in small intestine assessed by immunofluorescence microscopy (n=9) (at total magnification 200X and scale 50μ m). Activation of NOX 2 was studied through observing colocalizations between GP91phox (labelled with green fluorescent antibody) and P47 phox (labelled with red fluorescent antibody) subunits of the NADPH 2 oxidase complex resulting in a yellow region. Colocalizations were determined in small intestine tissues of CONT, GW, AB and GW+AB chemical exposed mice. **B** Graphical representation of morphometric analysis of colocalization events of GP91phox and P47phox in the region of interest. **C** Immunoreactivity of 3-nitrotyrosine (3NT) in EGC assessed through observing colocalizations between GFAP (labelled with green fluorescent antibody) and 3NT phox (labelled with red fluorescent antibody) at top panel magnification 200X, scale 50μ m and bottom panel magnification 600X and scale 20μ m) Colocalizations were determined in small intestine tissues of CONT, GW, AB and GW+AB chemical exposed mice. **D** Graphical representation of morphometric analysis of colox and scale 20µm) colocalizations were determined in small intestine tissues of CONT, GW, AB and GW+AB chemical exposed mice. **D** Graphical representation of morphometric analysis of colocalizations were determined in small intestine tissues of CONT, GW, AB and GW+AB chemical exposed mice. **D** Graphical representation of morphometric analysis of colocalizations were determined in small intestine tissues of CONT, GW, AB and GW+AB chemical exposed mice. **D** Graphical representation of morphometric analysis of colocalization events of GFAP (green) and 3NT (red).





Figure 3.11. NADPH oxidase 2 activation in rat EGC

A and B Activation of NOX-2 in rat EGC (n=6). Activation of NOX 2 was studied through observing colocalizations between GP91phox (labelled with green fluorescent antibody) and P47 phox (labelled with red fluorescent antibody) subunits of the NADPH 2 oxidase complex resulting in a yellow region. Colocalization (yellow) of GP91phox and P47phox was detected in vehicle VEH, LPS, LPS + apocynin (LPS+APO), HMGB-1, HMGB-1+APO treated cells at total magnification 400X; scale 20 μ m). C and D Morphometric analysis of GP91/p47phox colocalization events in rat EGC per 100 cells in different fields. (*P<0.05)





Figure 3.12. Peroxynitrite formation in EGC

A and B Expression of 3NT in rat EGC (n=6). Immunoreactivity of 3NT was detected in vehicle (VEH), LPS, LPS+APO, HMGB-1 and HMGB-1+APO treated cells by staining with red fluorescent antibody and counterstained with DAPI (blue) viewed at total magnification 400X; scale 20 μ m). C and D. Morphometric analysis of 3NT in rat EGC per 100 cells in different fields. (*P<0.05)





Figure 3.13. ROS mediated activation of NLRP-3 inflammasome and inflammation in EGC

A and B. Quantitative real time PCR (qRT-PCR) analysis of inflammasome and inflammation (n=6). mRNA expression of NLRP-3, caspase-1, IL-1 β and TNF- α in rat EGC which were treated with vehicle (VEH), LPS+ apocynin (LPS+APO), LPS+ phenylboronic acid (LPS+FBA), HMGB-1, HMGB-1+APO. mRNA expression is represented as a fold change of the vehicle control. Data points are represented as mean ±SEM (n=3). (*P>0.05) C. NLRP-3/ASC2 protein expression in rat EGC assessed by immunofluorescence microscopy and viewed at total magnification 400X; scale 20µm). Colocalization events were determined for every 100 cells per field in cells treated with LPS, LPS+APO, LPS+FBA, HMGB-1, HMGB-1+APO, HMGB1+FBA. **D and E**. Quantitative morphometric analysis of fluorescence intensity of NLRP-3/ASC2. Fields for morphometric analysis were randomly selected from different fields per slide and represented as % region of interest (% ROI). (*P<0.05)





Figure 3.14. DNA fragmentation of rat EGC.

A and B Tunel assay showing DNA fragmentation (n=6). DNA fragmentation was determined by the Tunel assay in Vehicle (VEH), LPS, LPS+APO, LPS+FBA, HMGB-1, HMGB-1+APO, HMGB-1+FBA treated cells viewed at total magnification 400X; scale 10 μ m. **C and D.** Quantitative morphometric analysis of fluorescence expression of Tunel positive cells represented as Tunel events per field. (*P<0.05). **E. Effect of EGC culture fluids on LPS primed intestinal epithelial cells.** mRNA expression of IL-1 β , MCP-1 and TNF- α in IEC-6 cells which have been primed with LPS and treated with culture fluids from EGC treated with LPS and vehicle control (n=6) represented as x Fold of the vehicle control.





Figure 3.15. Expression of claudin 2, occludin and aquaporin 3 in mouse small intestine.

A, C and F. Protein expression of occludin, claudin 2 and aquaporin 3 in mouse small intestine was determined by immunofluorescence microscopy and visualized at total 200X; scale 50 μ m in tissues obtained from mice treated with vehicle control (CONT, n=6); mice treated with GW chemicals (GW n=9) mice co-exposed with GW chemicals and antibiotics (GW+AB, n=9) and mice treated with GW chemicals, Sparstalonin B (SsnB) and Sodium butyrate (GW+SsnB+BT)(n=6) (A, D and G). Higher magnification images for GW chemical treated group (GW) focusing on the apical or apical-lateral membranes total magnification 600X; scale 20 μ m) (C, F, I). Quantitative morphometric analysis of immunoreactivity of occludin or aquaporin 3 is represented as % ROI. (*P<0.05). (B, E, H)





Figure 3. 16. Protein expression of aquaporin 3, claudin 2 and occludin in intestinal epithelial cells.

A, C and E: Protein expression of aquaporin-3, claudin-2 and occludin in IEC 6 cells treated with culture fluids from Vehicle (VEH-SN), LPS (LPS-SN), HMGB1 (HMGB1-SN) and inhibitors SsnB and butyrate (LPS+SsnB+BT and HMGB1+SsnB+BT) treated EGC. The expression of these proteins was studied by immunofluorescence microscopy and viewed at 400X total magnification; scale 10μ m. Yellow arrows indicate the localization of the proteins in the cell membrane. B, D and F: Quantitative morphometric analysis of immunoreactivity of aquaporin, claudin 2 and Occludin represented as % ROI (n=3, *P<0.05).



Data Availability Statement

The datasets generated for this study can be found in the Supplementary Material.

Ethics Statement

The animal study was reviewed and approved by the USC Institutional Research Board (USC IACUC).

Author Contributions

SC and DK designed and performed the experiments. DK, SS, DB, RS, YL, AM, MA, and AK performed the experiments. KS, PJ, and SL analyzed and interpreted the data. PN and MN analyzed the data and edited the manuscript. NK, KS, and RH edited the manuscript. SC and DK wrote the manuscript, edited and analyzed the data.

Conflict of Interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest. The reviewer GW and handling Editor declared their shared affiliation at the time of the review.

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CHAPTER 4

HOST AKKERMENSIA MUCINIPHILA ABUNDANCE CORRELATES WITH GULF WAR ILLNESS SYMPTOM PERSISTENCE VIA NLRP3 MEDIATED NEUROINFLAMMATION AND DECREASED BRAIN DERIVED NEUROTROPHIC FACTOR³

³ Diana Kimono, Dipro Bose, Ayan Mondal, Ratanesh Seth, Punnag Saha, Patricia Janulewicz, Kimberly Sulivan, Stephen Lasley, Ronnie Horner, Nancy Klimas, Saurabh Chatterjee. Submitted to *Brain Behavior and immunity* 02/23/2020



Running title: Akkermensia muciniphila and GWI persistence

Author for correspondence:

*Dr. Saurabh Chatterjee, Ph.D. Environmental Health and Disease Laboratory, NIEHS Center for Oceans and Human Health on Climate Change Interactions, Department of Environmental Health Sciences, University of South Carolina, Columbia 29208 USA. Email: schatt@mailbox.sc.edu; Tel: 803-777-8120; Fax: 803-777-3391 Key words: dysbiosis; *Akkermansia muciniphila*; BDNF; RAGE; 3-nitrotyrosine;

peroxynitrite; inflammasomes; symptom persistence.



ABSTRACT

Neurological disorders are commonly reported among veterans who returned from the Gulf war. Veterans who suffer from GWI, complain of continued symptom persistence that include neurological disorders, muscle weakness, headaches and memory loss among many others that developed during or shortly after the war. Our recent research showed that chemical exposure associated microbial dysbiosis accompanied by a leaky gut connected the pathologies in the intestine, liver and brain. However, the mechanisms that caused the symptoms to persist even thirty years after the war remained elusive to investigators. In this study, we used a persistence rodent model of GWI to investigate the persistence of microbiome alterations, resultant chronic inflammation and its effect on neurotrophic and synaptic plasticity marker BDNF. The results showed that exposure to GW-chemicals (pesticides such as permethrin and prophylactic drugs e.g. pyridostigmine bromide) resulted in persistent pathology characterized by low relative abundance of the probiotic bacteria Akkermansia muciniphila in the gut, which correlated with high circulatory HMGB1 levels, blood brain barrier dysfunction, neuro-inflammation and lowered neurotrophin BDNF levels. Mechanistically, use of mice lacking the NLRP3 gene had significantly decreased inflammation and a subsequent increase in BDNF in the frontal cortex. This suggests that a persistently low species abundance of Akkermensia muciniphila and associated chronic inflammation due to inflammasome activation might be playing a significant role in contributing to persistent neurological problems in GWI. A therapeutic approach with various small molecules that can target both the restoration of a healthy microbiome and decreasing inflammasome activation might have better outcomes in treating GWI symptom persistence.



4.1. INTRODUCTION

Neurological disorders are commonly reported among veterans who returned from the gulf war (GW) of 1990-1991 (3,106),(107). Afflicted veterans complain of problems including neuralgias, repeated seizures, migraine headaches, muscle weakness and coordination, memory problems etc (107,108)(109). These issues occur in combination with other gulf war illness (GWI) symptoms and their pathology is not very well understood. Most veterans who suffer from GWI developed their symptoms during or shortly after the war and these symptoms persist thirty years later. Although the causes of these symptoms are difficult to pinpoint, epidemiological studies have established a compelling link between these symptoms in different GW veteran cohorts and environmental exposures which occurred during the war, or chemicals that were applied to the warriors before or shortly after the war (110–112). Such exposures include desert storm dust, depleted uranium, combustion byproducts from oil wells, possible chemical weapons, pesticides, vaccines and prophylactic medicines such as pyridostigmine bromide etc (113,114) (4).

In recent years, research has focused on studying symptoms as well as elucidating mechanisms of these disorders, using human GW veteran cohorts as well as animal, and in vitro studies. For example, Van Riper et al, reported widespread disruption in white matter microstructure distribution across brain regions involved in processing and modulating chronic pain (13). James et al found that there was a significant positive correlation between C-reative protein (CRP), pain and neurocognitive mood in GW veterans. Another study by Abou-Donia et al reported elevated autoantibodies to neurons and other brain cells eg tau proteins, glial acidic fibilliary protein (GFAP) and myosin basic protein (MBP)



which indicate neuronal injury or gliosis in GW veterans (115). In other recent animal studies, Zakirova et al found cognitive deficits in mice several weeks after treatment with GW chemicals and these deficits were associated with increased astrogliosis and a reduction in synaptophysin in mouse hippocampi and cerebral cortex (44). Further Madhu et al., found that cognitive impairments which persisted 10 months post-exposure to GW chemicals were associated increased density of activated microglia and astrocytes in rats and inflammation with elevated levels of HMGB1 in cerebral cortex (116). These studies provide evidence that exposure to GW chemicals plays a significant role in persistence of neurological dysfunction. This may be generally through the disruption of neuronal networks, reactive glia which fuel inflammation or poor neuronal growth and neuroplasticity.

Many neurological disorders such as Parkinson's disease (PD), Alzheimer's disease (AD), bipolar disorders, neuropathic pain etc (117)(118,119) (120,121) are associated with decreased levels of neurotrophins or impairments in their signaling pathways (122)(22). These disorders also commonly present with chronic neuroinflammation (123–125). Brain derived neurotropic factor (BDNF) is the most prevalent neurotrophins in the brain and has been very widely studied (122,126). It is produced by neurons and it plays important functions in neuroplasticity, growth and survival of neurons (127). To date, our previous research on GWI has largely focused on the possible role of an altered microbiome (bacteriome and virome) in contributing to a persistent inflammatory phenotype as observed in our studies (18,19,29,128).

We proposed that exposure to GW chemicals alters the microbiome which then drives inflammation through production of immunostimulatory particles i.e. damage



associated molecular pattern (DAMPS) and pathogen associated molecular patterns (PAMPS), these continuously fuel inflammation observed in different organ systems. Although our results largely supported our speculations, we were still limited in knowing whether the observed changes and mechanisms in microbiome and associated chronic inflammation due to an altered microbiome indeed persisted. In this present study, we used a persistence rodent model of GWI in which mice were first exposed to GW chemicals for two weeks (war phase) and then left in their cages on a normal chow diet for 20 weeks (post war phase). We then investigated the persistence of microbiome alterations, chronic inflammation and its effect on neuronal health (BDNF levels). We further used a NLRP3 KO mouse to study its potential role in contributing to observed neuroinflammation.

4.2. MATERIALS AND METHODS

Materials

We purchased Pyridostigmine bromide (PB) and Permethrin from Sigma- Aldrich (St. Louis, MO). Anti- BDNF, anti-RAGE, anti-claudin 5, anti-HMGB1, anti-IL-1β, anti-ASC-2 were purchased from Sigma-Aldrich while anti-NLRP3, anti-3-nitrotyrosine, anti-TMEM 119 primary antibodies were purchased from Abcam (Cambridge, MA). Species specific biotinylated conjugated secondary antibodies and Streptavidin-HRP (Vectastain Elite ABC kit) was purchased from Vector Laboratories (Burlingame, CA). Fluorescence conjugated (Alexa fluor) secondary antibodies, ProLong Diamond antifade mounting media with DAPI and Pierce LAL chromogenic endotoxin quantitation kit was bought from Thermo Fisher Scientific (Waltham, MA). Unless otherwise specified, all other chemicals used were purchased from Sigma. Paraffin-embedding of tissue and sectioning



were done by AML laboratories (Baltimore, MD) and at the Instrument Resources Facility, University of South Carolina School of medicine (Columbia, SC). Microbiome analysis was done by Cosmos ID (Rockville, MD).

4.2.1. Animal experiments

Adult (10 weeks old) wild type male (C57BL/6J mice) and NLRP3 deficient adult (10 weeks) male (B6N.129-Nlrp3tm3Hhf/J) mice were purchased from the Jackson Laboratories (Bar Harbor, ME). Mice experiments were implemented in accordance with NIH guidelines for humane care and use of laboratory animals and local IACUC standards. All procedures were approved by The University of South Carolina at Columbia, SC. Mice were housed individually and fed on a chow diet at 22–24°C with a 12-h light/ 12-h dark cycle. All mice were sacrificed after animal experiments had been completed. Right after anesthesia, blood from the mice was drawn using cardiac puncture, to preserve serum for further experimentation. Their brains were removed immediately, and frontal cortex dissected out and were fixed using Bouin's fixative solution. We also collected the fecal pellets and luminal contents for microbiome analysis.

Treatments and rodent model of Gulf war illness

Mice were exposed to GW chemicals (pyridostigmine bromide and permethrin) based on established rodent models of GWI with some modifications (44,63,129). The treated mice group (GWP) and NLRP3KO (GWP-NLR3KO) mice group were dosed triweekly for two weeks with PB (2mg/kg) and permethrin (200 mg/kg) by oral gavage, then mice were left in cages to feed on a normal chow diet for 20 weeks. The control group



(CONT) of mice received vehicle (6% DMSO in PBS) by oral gavage as in other experiments above.

Microbiome analysis

Microbiome analysis was done by CosmosID (Rockville, MD, USA) from fecal pellets and luminal contents which were collected from the animals of each group after sacrifice. DNA isolation, sequencing and analysis of gut microbiome were performed according to vendor optimized protocol. Briefly, DNA was isolated from fecal samples using the ZymoBIOMICS Miniprep kit, following the manufacturer's instructions. 16S sequencing was carried out on the V3-V4 (341 nt-805 nt) region of the 16S rRNA gene with a twostep PCR strategy. First, PCR was performed using 16S-optimized primer set to amplify the V3–V4 regions of 16S rDNA within the metagenomic DNA. Then the PCR products from the previous steps were mixed at equal proportions and used as templates in the second step to produce Illumina dual-index libraries for sequencing, with both adapters containing an 8-bp index allowing for multiplexing. The dual-indexed library amplification products are purified using Ampure beads (Beckman Coulter). Library quantification was performed using Qubit dsDNA HS assay (ThermoFisher) and qualified on a 2100 Bioanalyzer instrument (Agilent) to show a distribution with a peak in the expected range. A final qPCR quantification was performed before loading onto a MiSeq (Illumina) sequencer for PE250 (v2 chemistry). The sequences for each sample were then run on the 16S pipeline of the CosmosID GENIUS software, and results were analyzed.



4.2.2. LABORATORY METHODS

Immunohistochemistry

The fixed brain tissues were embedded in paraffin and sliced into 5 µM thick sections. These sections were deparaffinized following optimized standard protocols. Epitope retrieval solution and steamer (IHC-Word, Woodstock, MD) were used for epitope retrieval for deparaffinized sections. 3% H2O2 was used for the recommended time to block the endogenous peroxidase. After serum blocking, the primary antibodies were applied at recommended and optimized concentrations. Species- specific biotinylated conjugated secondary antibodies and streptavidin conjugated with HRP were used to implement antigen specific immunohistochemistry. 3,3'-Diaminobenzidine (DAB) (Sigma Aldrich, St Louis, MD) was used as a chromogenic substrate. Mayer's Hematoxylin solution (Sigma Aldrich) was used as a counter stain. Sections were washed between steps using phosphate buffered saline 1X. Finally, stained sections were mounted in Simpomount (GBI laboratories, Mukilteo, WA). Tissue sections were observed using Olympus BX63 microscope (Olympus, America). Cellsens software from Olympus America (Center Valley, PA) was used for morphometric analysis of images. Immunofluorescence staining Paraffin embedded sections were deparaffinized using standard protocol. Epitope retrieval solution and steamer were used for epitope retrieval of sections. Primary antibodies were used at recommended dilutions. Species specific secondary antibodies conjugated with Alexa Fluor (633-red and 488-green) were used at advised dilution. In the end, the stained sections were mounted using prolong diamond antifade reagent with DAPI. Sections were observed under-Olympus florescence microscope BX63 using 20X, 40X, 60X objective

lens.


Real time Quantitative PCR

Total RNA was isolated from frontal cortex tissue homogenization in TRIzol reagent (Invitrogen, Carlsbad, CA, USA) according to the manufacturer's instructions and purified with the use of RNeasy mini kit columns (Qiagen, Valencia, CA, USA). cDNA was synthesized from purified RNA (1 μ g) using iScript cDNA synthesis kit (Bio-rad, Hercules, CA, USA) following the manufacturer's standard protocol. Real-time qPCR (qRTPCR) was performed with the gene-specific primers using Sso Advanced SYBR Green Supermix and CFX96 thermal cycler (Bio-rad, Hercules, CA, USA). Threshold cycle (Ct) values for the selected genes were normalized against respective samples internal control 18S. Each reaction was carried out in triplicates for each gene and for each sample. The relative fold-change was calculated by the 2– $\Delta\Delta$ Ct method. The sequences for the primers used for real-time PCR are provided in table 1.

Table 4.1: Primer sequences

mm Claudin 5	Sense: TTCGCCAACATTGTCGTCC
	Antisense: TCTTCTTGTCGTAGTCGCCG
mm18S	Sense: TTCGAACGAACGTCTGCCCTATCAA
	Antisense: ATGGTAGGCACGGCGATA

Endotoxin level detection by Litmus Amebocyte Lysate assay

Serum bacterial endotoxin levels (EU/mL) were detected using the Pierce LAL Chromogenic Endotoxin Quantification Kit (Waltham, MA) according to the



manufacturer's instructions. Briefly, serum samples were obtained from mice and diluted 1:80 with endotoxin free water. The endotoxins were then quantified.

Western Blot analysis

30 mg of tissue from each brain tissue sample was immediately homogenized in 300 µl of RIPA buffer with protease and phosphatase inhibitors cocktail (Pierce, Rockford, IL) using slow speed mechanical homogenizer. The homogenate was centrifuged, and the supernatant was collected and saved for experimental use. 30 µg of denatured protein from each sample was loaded per well of Novex 4–12% bis-tris gradient gel (Life technologies, Carlsbad, CA) and subjected for standard SDS-PAGE. Separated protein bands were transferred to nitrocellulose membrane using precut nitrocellulose/filter paper sandwiches (Bio-Rad, Hercules, CA) and Trans-Blot Turbo transfer system (Bio-Rad) using 30 minutes transfer protocol. Further, blots were blocked with 3% bovine serum albumin solution prepared in Tris-buffered saline with 0.05% tween-20 (TBS-T). Primary antibodies were used at recommended dilutions in 1.5% blocking buffer and incubated overnight at 4°C. Species-specific anti-IgG secondary antibody conjugated with HRP were used at recommended dilutions in 1% blocking buffer and incubated for 2h at room temperature. Pierce ECL Western Blotting substrate (Thermo Fisher Scientific Inc., Rockford, IL) was used in dark to develop the blot. Finally, the blot was imaged using G:Box Chemi XX6 (Syngene imaging systems) and subjected to densitometry analysis using Image J software.



4.2.3. Statistical analysis

We conducted calculations for each experimental condition prior to initiation of the study, using preliminary data to confirm that the sample size was enough to achieve a minimum statistical power of 0.80 at an alpha of 0.05. One-way analysis of variance (ANOVA) was used with post-hoc comparisons among different exposure conditions or treatments (e.g., least significant differences (LSD) and Bonferroni correction) to compare means among multiple groups. Student's t-tests was used to compare means between two groups at the termination of treatment. Correlative associations were tested using Pearson's Rank Product moment coefficient analysis using Graph pad prism software (GraphPad Software Inc., La Jolla, CA, United States)

4.3. RESULTS

4.3.1. Gulf war chemical exposure results in a decreased relative abundance of Akkermansia muciniphila which negatively correlates with increased circulatory HMGB1 levels.

Our previous studies have strongly suggested that exposure to GW chemicals alters the microbiome and these alterations may contribute to persistence of GWI symptoms through release of DAMPs and PAMPs (18,29,128). In this study we analyzed the microbiome for alterations in specific bacterial species that have a pronounced role in inflammation persistence in chronic diseases of the gut, metabolic reprogramming and neuronal deficiencies. We analyzed 10 distinct bacterial species that had a fold change difference in abundance and has been found to contribute to inflammation and metabolic responses (Fig. 1A) We found that mice treated with GW chemicals (GWP) had significantly lower abundance of A. muciniphila (significant), Bacteroides



Thetaiotomicron and *Dorea Sp* (not significant) when compared to mice treated with only vehicle control (Figure 1A) (*P<0.05; n=6). Notably, *A. muciniphila* has been associated with several health benefits (130,131) (38). Furthermore, we found that mice which were treated with GW chemicals (GWP) had significantly higher HMGB1 levels in their serum compared to mice treated with vehicle control only (CONT) *P<0.05, n=6 (Figure 1B and C). We then carried out statistical analyses to determine whether the increased levels of HMGB1 were related to the observed decreased relative abundance of *A. muciniphila*. In Figure 1D, we found that there was a negative correlation between *A. muciniphila* abundance and circulatory HMGB1 levels (Peason's r = -0.50; R square COD=0.255).

4.3.2. Exposure to GW chemicals is associated with blood brain barrier tight junction protein dysregulation and the changes persist 5 months after exposure

The study by Abou-Donia et al suggests the presence of a leaky BBB among veterans who returned from the GW and this may be a portal for immunostimulatory particles such as DAMPS and PAMPS to continuously fuel neuroinflammation (115). We studied the mRNA and protein expression levels of claudin 5, the major tight junction protein in the complex that makes up the BBB. We found that GWP mice also exhibited significantly lower claudin 5 mRNA and protein levels compared to vehicle control treated mice (CONT) P<0.05, n=6 (Figure 2 A, B and C). Further, we studied the levels of claudin 5 in the BBB by observing colocalizations between claudin 5 and CD31 a marker for endothelial cells that make up the lining of the blood vessels (Figure 2 D and E). We found that there was a significant decrease in the number of colocalizations (yellow spots) constituting claudin 5 and CD31 in GWP mice compared to controls (CONT). This result points to the fact that at least one major component of BBB integrity is repressed at the



protein level paving the way for possible dysfunctional BBB, and this may lead to the passage of DAMPS such as HMGB1 leaking into the brain and triggering several immune responses.

4.3.3. GW chemical exposure is associated with persistent activation of microglia via the HMGB1-RAGE pathway resulting in increased reactive oxygen species and triggering of the NLRP3 inflammasome.

Chronic neurological disorders such as AD and PD etc, are characterized by activation of immune cells such as microglia, the resident macrophages of the brain (132). These cells may be activated by the presence of pathogens or DAMPS such as HMGB1. We studied the protein expression levels of activated microglia marker TMEM119. The results showed that there was a significant increase in activated microglia in the frontal cortex of mice treated with GW chemicals (GWP) compared to controls (CONT) (Figure 3 A and B) even after 5 months of exposure. Further, we found that there was evidence of activated HMGB1-RAGE signaling as indicated by colocalization events. Figure 3 C, D shows a high expression of RAGE protein levels and subsequent increased RAGEHMGB1 colocalizations in GWP mice compared to controls (Figure 3 E, F). RAGE is a receptor, which binds several ligands including HMGB1. Interaction of HMGB1-RAGE can lead to increased reactive oxygen species (ROS) generation eg peroxynitrite which reacts with tyrosine in proteins to form the stable adduct 3 nitrotyrosine. In Figure 4A, we detected significantly higher levels 3-nitrotyrosine (3NT) in GW chemical treated mice (GWP) compared to vehicle control treated mice (CONT) Figure 4B and C; (*P<0.05; n=6). We also found that mice exposed to GW chemicals had significantly higher inflammasome activation compared to vehicle control treated mice (CONT) Figure 4D and E; (*P<0.05; n=6). This activation was detected as yellow dots indicating a colocalization between the



NLRP3 protein complex (labelled with red antibody) and the adapter protein ASC2 (marked with green), which facilitate processing of proinflammatory cytokines from their basal inactive to more active form via protein cleavage.

4.3.4. Exposure to GW chemicals is associated with a persistent increased neuroinflammation and low levels of BDNF

Inflammasomes are large immune complexes found in several cell types and are responsible for processing the inflammatory cytokines IL-1 β and IL-18 by cleaving them from their precursors (133). Uncontrolled activation of these complexes can lead to chronic inflammation as has been found in cancer, diabetes and neurodegenerative disease (51,134,135). In our study, we found that NLRP3 inflammasome activation were also significantly associated with increased IL-1 β protein levels in GWP mouse frontal cortex when compared to mice treated with only vehicle control (CONT) (Figure 5A, B); *P<0.05; n=6. Neurodegerative diseases are often characterized by neuroinflammation accompanied by decreased levels of neurotrophins (119)(136). Similarly, in this study we observed that mice which were treated with GW chemicals had significantly lower levels of the neurotrophin BDNF when compared to mice treated with only vehicle control (CONT) (Figure 5-F) *P<0.05; n=6.

4.4.5. Akkermensia muciniphila relative abundance correlates with BDNF levels and persistent neuroinflammation

High abundance of *A. muciniphila* has been linked to decreased inflammation in chronic diseases (137–139). To study whether the host bacteria's abundance played a role in affecting chronic inflammation and sustained BDNF levels in our model of GWI, we



carried out correlative analyses to determine whether there were statistically significant relationships between the bacterial abundance, inflammation and BDNF levels. The results showed that there was a significant positive correlation between BDNF levels and abundance of *A. muciniphila* (Pearson's r=0.83, R square (COD)=0.73; p=0.0024), and a significant negative correlation with IL-1 β protein levels (Pearsons r=-0.684; R square (COD)=0.46; p=0.02) (Figure 6A and B). Shaded area represents 95% confidence bands.

4.4.6. Deletion of NLRP3 is protective against persistent neuroinflammation and is associated with increase in BDNF levels

To study the role of NLRP3 in driving inflammation and lowering BDNF levels, we treated mice lacking NLRP3 with GW chemicals and then subjected them to our experimental conditions for 20 weeks or 5 months. We then studied the protein levels of IL-1 β and BDNF by western blot analysis and immunohistochemistry. Our results show that there were significantly lower levels of IL-1 β (Figure 7A, B), and increased BDNF (Figure 7 CF) levels in the frontal cortex of NLRP3 KO mice treated with GW chemicals (GWPNLR3KO), when compared to wild type mice treated with GW chemicals (GWP)

(*P<0.05; n=6).

4.5. DISCUSSION

In our previous studies, we reported that there was a general alteration in microbiome accompanied by endotoxemia, a leaky gut, and inflammation in different organs such as the small intestine, brain and liver (18,29,128). We found significant increases in phyla Firmicutes and Tenericutes over Bacteriodetes in GW chemical exposed mice when compared to controls (29). However, it was not clear whether these alterations



persisted over a longer period of lifespan or would eventually resolve over time through repopulation and reconstitution of the host microbiome. We also were eager to study the mechanisms that would connect the altered microbiome and the persistent inflammatory changes in the intestine and the neural immune network. In this study, mice were exposed to the GW chemicals and allowed to ad libitum diet and water for 5 months. This was to simulate the period of exposure (war phase) and the post war period in GW veterans. We found that exposure to GW-chemicals (pesticides such as permethrin and prophylactic drugs eg pyridostigmine bromide) resulted in persistent pathology characterized by low abundance of *A. muciniphila*, high circulatory HMGB1 levels, blood brain barrier dysfunction, neuroinflammation and lowered neurotrophin BDNF levels.

We report that exposure to GW chemicals caused a decrease in *A. muciniphila* or resulted in conditions that favor other bacteria populations repopulate over *A. muciniphila* (Figure1A). *A. muciniphila* is a mucin degrading bacterium which exists as part of the normal human gut flora and is abundant in healthy individuals (16,139–141). In recent years the herein reported bacterium is emerging as an important probiotic which can be consumed to improve health (142). This bacterium was found to improve ulcerative colitis in mice (143) and restored colonic mucus layer thickness with decreased inflammation in aging mice (144). In another study, the abundance of this bacterium inversely correlated with inflammation and altered lipid metabolism in obese mice (142). Although the mechanism by which *A. muciniphila* promotes these health benefits is not fully understood, studies report that the bacterium strengthens gut barrier integrity through its association with enterocytes and also produces high amounts of anti-inflammatory cytokine IL-8 (139,145). It is possible that the low levels of this bacterium in the gut compromised gut



105

barrier integrity, a condition that we have observed in our previous acute models of GWI (146)(29). This condition of compromised gut barrier integrity has also been reported among veterans who suffer from gastrointestinal problems in GWI (55). And yet another study found that *A. muciniphila* treatment normalized diet-induced metabolic endotoxemia, adiposity, and the adipose tissue marker CD11c in obese mice which otherwise had increased inflammatory indicators in the intestine and aided in the metabolic disease development (137).

Similarly, our studies of GWI mouse models have consistently found that altered levels of tight junction proteins in the gut was associated with increase in endotoxins and DAMPs such as HMGB1 and inflammation that was related to an alteration of gut microbiome abundance (146)(18). Notably, mice in the present persistent model study that were exposed to similar chemicals showed a slight increase in serum endotoxin levels (low level endotoxemia consistent with an obesity phenotype) in GW chemical treated mice when compared to vehicle control treated mice (Figure S1). In addition, a significant increase in circulatory HMGB1 levels were observed in GW-chemical exposed mice which negatively correlated with A. muciniphila relative abundance (Figure 1B, C and D) suggesting that a sustained and consistent low inflammatory trigger was closely associated with a persistent change in the microbiome and decreased A muciniphilia abundance in these mice. Further, the persistence of systemic inflammatory indicators such as endotoxin levels and serum HMGB1 for such a long period that failed to ease even 5 months after exposure (equivalent to >20 human years) and its connection with an altered microbiome triggered our interest to study their effects on neuronal structures and their networks.



Interestingly, before we could study the neuroinflammatory indicators for persistence, we needed to assess the integrity of the blood brain barrier, a vital interface of neuronal physiology and pathology. Results showed that expression of Claudin 5, a key tight junction protein of the BBB in the brain was decreased in the frontal cortex of mice treated with GW chemicals compared to controls (Figure 2A-E). This protein, together with other tight junction proteins such as claudin 1, zona occludens, occludins etc make up tight junctions in the BBB. The BBB is a selective barrier found at the interface of blood vessels in the brain and brain tissue. It is made of a single cell layer of endothelial cells, astrocyte and pericytes. Its unique properties allow it to tightly regulate the movement of particles between the circulation and brain tissue (56) (57). Claudin 5 and other tight junction proteins are found between adjacent endothelial cells of blood vessels and help to anchor these cells to create a tightly regulated selective barrier which allows the passage of particles between the blood and the brain(147). Low levels of this protein have been found in neurodegenerative and neuronflammatory diseases such as AD, PD and schizophrenia (148,149). We found decreased mRNA and protein levels of this this protein in GWP mice compared to controls (Figure 1A-F). This provides strong evidence that at least one key component of the BBB is dysregulated and this possibly compromised the barrier's integrity, likely caused by the serum mediators endotoxins and HMGB1 causing it to become leaky though we have no direct evidence of such an event in an in vitro experimental set up using BBB endothelial cells. We hypothesized that this leaky BBB allowed the passage of unwanted particles such as DAMPs and PAMPS, HMGB1 as one such example, which we found to be greatly increased in the serum. It is also worth noting that even though serum endotoxin levels are not significantly higher in GWP mice



compared to controls, even low levels of endotoxins over a long time can be a toxic stimulus to the body and may contribute to observed pathology(150), (62).

HMGB1 is a DAMP known to trigger proinflammatory pathways through toll like receptors (TLRs) eg TLR4 and through the receptor for advanced glycation end products (RAGE) (49,151). We found that there was increased activation of microglia in GWP mice compared to controls (Figure 3A and B) with an increased expression of RAGE receptors in the frontal cortex Figure 3 C,D. We also detected HMGB1/RAGE complex formation using immunofluorescence microscopy, which may indicate activation of RAGE signaling (Figure 3E and F). The activation of this pathway is known to result in transcription of proinflammatory cytokines and generation of reactive oxygen species (ROS) (152) such as nitric oxide. High levels of nitric oxide in the presence of superoxide could result in the formation of peroxynitrite a potent ROS which attacks tyrosine to form 3 nitrotyrosine denaturing them and rendering them dysfunctional. We found significantly higher levels of ROS in our GWP mice compared to controls (Figure 4A and B). High amounts of ROS are a known trigger of the NLRP3 inflammasome (102). Inflammasomes are large protein complexes which are assembled in response to infections etc and are involved in processing of proinflammatory cytokines such as IL-1 β and IL-18. We found that mice treated with GW chemicals had higher expression of NLRP3/ASC2 complex formation compared to controls (Figure 4C and D). Although inflammasomes are triggered as a defense mechanism, chronic activation of these complexes has been implicated in conditions characterized by chronic low-grade inflammation such as diabetes, PD, ALS etc (153,154). This activation of the NLRP3 inflammasomes was followed by increased IL-1 β levels in the brain (Figure 5A and B) which might contribute to the persistent neuroinflammation in



108

GWI. In neurological diseases, inflammation has been associated with poor neuronal health, with fewer neurons, decreased neuronal plasticity and growth (155). This in part is due to decreased levels of neurotrophins such as BDNF. BDNF is a neurotrophin produced by neurons and is involved in neuronal growth, survival and plasticity (122). Studies by Guan and Fang, and another by Lapchak suggests that increased IL-1 β levels interfered with BDNF synthesis (156), while Tong et al., showed that increased IL-1 β interfered with BDNF signaling through the PI3K/AKT pathway by preventing its activation of AKT. The above mechanisms resulted in decreased growth and survival of neurons (157). In this study, we report low levels of BDNF in the frontal cortex of mice which were treated with GW chemicals compared to controls even 5 months post exposure (Figure 5C-F). We also studied any correlations that BDNF, IL-1 β with A. muciniphila relative abundance may have to connect the intestinal, microbiome changes and neuronal levels of these mediators referenced above. In Figures 6A and B, we show that A. muciphila relative abundance correlated negatively with IL-1 β levels and positively with BDNF levels. The above result indicates that A. muciniphila relative abundance might have played a role in modulating neuroinflammation and neurotrophin levels in GWI as has been the case in obesity and other diseases described previously (46). This could be through the bacterium's production of anti-inflammatory compounds that counter inflammation or modulation of the intestinal barrier integrity, although more studies need to be done to determine the exact mechanism. Finally, to study the role of NLRP3 inflammasome in contributing to neuroinflammation that persist for a prolonged period of time and its association with increased abundance of A. muciniphila, we used a mouse model with the systemic knockout of NLRP3 gene (KO mouse). The results found that deletion of this gene was associated with increased BDNF



levels and protected the mice from neuroinflammation (Figure 7A-F). Deleting NLRP3 is protective through preventing inflammasome activation and subsequent processing of proinflammatory cytokines and distinctly proves that the persistent inflammation in GWI chemical exposed mice that had altered *A. muciniphila* abundance is due to NLRP3mediated inflammasome activation though there may be multiple molecular mediators for triggering such an activation. Our current study identifies some of these mediators such as gut derived endotoxins, HMGB1 or peroxynitrite to name a few but the molecular mechanism of such a trigger remains speculative at this time.

In conclusion, we report that persistence of GWI symptoms is characterized by low relative abundance of *A. muciniphila* and chronic high circulatory HMGB1 levels which trigger NLRP3 mediated neuroinflammation and decreased levels of neurotrophins through RAGE signaling. These findings not only provide insight into the mechanism of persistent neurological disturbances in GWI but also provide possible therapeutic targets. Firstly, the microbiome can be targeted through replacement or enhancing of *A. muciniphila* gut bacterial populations and other significantly lowered probiotic bacteria, and secondly through therapies that target RAGE signaling or NLRP3 inflammasomes to relieve persistent inflammation and improve quality of life.

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Conflict of interest

The authors declare no conflict of interest





Figure 4.1. Exposure to GW chemicals results in decreased relative abundance of *Akkermensia muciniphila* and chronic high levels of circulatory HMGB1

A. Percentage abundance of gut bacteria species. Percentage abundance of 10 most abundant species in the gut bacteriome are represented comparing GW chemical treated groups (GWP) to vehicle control treated group groups (CONT). Data is represented as the mean of 6 mice per group. **B.** Relative abundance of *A. muciniphila*. Relative abundance was determined from duplicate fecal samples of six mice per group treated with GW chemicals (GWP) compared to mice treated with vehicle control only (CONT). Data is represented as Mean ±SEM. (*p<0.05; n=6). **C.** Serum HMGB1 levels. Western blot of HMGB1 levels in serum for mice treated with GW chemicals (GWP) compared to mice treated with Vehicle control (CONT) only. Data is represented as Mean ±SEM. **D.** Densitometry of HMGB1 immunoblots, normalized against ponceau red (*p<0.05; n=6). **E.** Relationship between *A. muciniphila* and circulatory HMGB1 levels. A correlative analysis was carried out to determine how *A. muciniphila* is related to serum HMGB1 levels. We found a negative correlation between *A. muciniphila* and serum HMGB1 levels (Peason's r = -0.50; R square COD=0.255 shaded area represents 95% confidence bands).





Figure 4.2. Exposure to GW chemicals is associated with altered claudin 5 levels in the frontal cortex

A. Claudin 5 mRNA levels in the frontal cortex: mRNA levels of claudin 5 as studied by RTqPCR show that mice exposed to GW chemicals (GWP) had significantly decreased claudin 5 mRNA levels compared to mice treated with vehicle control only (CONT). **B.** Claudin 5 levels in protein levels in frontal cortex. Western blot analysis of Claudin 5 protein levels in CONT and GWP treated mice. **C.** Morphometry analysis of claudin 5 immunoblots, normalized against β-actin (*p<0.05; n=6). Data is represented as Mean ±SEM. **D.** Representative immunofluorescence micrographs of the blood brain barrier showing colocalization of tight junction protein Claudin 5 (labelled in red) and endothelial cell marker CD31 (labelled in green) as yellow spots around a BBB (Magnification 60X and scale bar 10µm). Inset (Magnification (40X and scale bar 20 µm) shows the whole micrograph field, from which the main image was obtained. **E.** Quantitative morphometry analysis of colocalizations for every 100 cells, represented as % ROI (*p<0.05; n=6). Data is represented as Mean ±SEM.





Figure 4.3. Activation of Macrophages and associated HMGB1/RAGE complex formation

A. TMEM119 immunoreactivity in frontal cortex. Representative immunohistochemistry micrographs of TMEM 119 reactivity in control (CONT) and GW chemical treated (GWP) mice (Magnification 40X and scale bar 50µm). **B.** Morphometric analysis (represented as % ROI) obtained from 10-15 images from different microscopy fields from each mouse sample. Data is represented as Mean ±SEM. (*p<0.05; n=6). **C**. Western blots of RAGE protein levels in the frontal cortex of GWP and CONT treated mice. **D.** Morphometry analysis of all immunoblots normalized against β -actin (n=5). (*p<0.05; n=6) Data is represented as Mean ±SEM. **E**. RAGE/HMGB1 complex formation. Immunofluorescence micrographs showing activation of NLRP3 inflammasome protein. HMGB1 (red) and RAGE (green) in GWP and CONT mice. Colocalizations are shown as yellow dots and marked with arrows in the micrographs (Magnification 40X and scale bar 20µm). **F.** Morphometric analysis (represented as % ROI) obtained from 6-8 images from different microscopy fields from each mouse sample. Data is represented as Mean ±SEM. (*p<0.05; n=6).





Figure 4.4. Increased ROS is associated with NLRP3 inflammasome activation

A. 3-nitrotyrosine immunoreactivity in frontal cortex. Representative immunohistochemistry micrographs of 3-nitrotyrosine reactivity in control (CONT) and GW chemical treated (GWP) mice (Magnification 40X and scale bar 50µm). B. Morphometric analysis (represented as % ROI) obtained from 10-15 images from different microscopy fields from each mouse sample. Data is represented as Mean \pm SEM. (*p<0.05; n=6). C. Immunofluorescence micrographs showing activation of NLRP3 inflammasome protein. NLRP3 (red) and its ASC2 (green) in GWP and CONT mice. Colocalizations are shown as yellow dots and marked with arrows in the micrographs. Magnification 60X and scale bar 20µm. D. Morphometric analysis (represented as % ROI) obtained from 6-8 images from different microscopy fields from each mouse sample. Data is represented as Mean ±SEM. (*p<0.05; n=6).





Figure 4.5. GW chemical exposure is associated with chronic neuroinflammation and decreased Brain derived neurotrophic factor (BDNF) levels in Frontal cortex.

A. Immunohistochemistry micrographs of frontal cortex tissues of GWP and CONT treated mice showing immunoreactivity of IL-1β (Magnification 20X and scale bar 50µm). **B.** Morphometric analysis (as % ROI) obtained from 10-15 images from different microscopic fields from each mouse sample. (*p<0.05; n=6). Data is represented as Mean ±SEM. **C.** BDNF immunoreactivity. Representative immunohistochemistry micrographs showing BDNF in frontal cortex tissues of GWP and CONT treated mice (Magnification 40X and scale bar 50µm). **D.** Morphometric analysis (represented as % ROI) obtained from 10-15 images from different microscopy fields from each mouse sample. **E.** Western blots of BDNF protein levels in the frontal cortex of GWP and CONT treated mice. **F.** Morphometry analysis of all immunoblots normalized against β-actin (n=5). (*p<0.05; n=6) Data is represented as Mean ±SEM.





Figure 4.6. Decreased relative abundance of *Akkermansia muciniphila* correlates with IL-1β and BDNF levels in the frontal cortex.

A. Correlation between *A. muciniphila* and IL-1 β levels in GW chemical (GWP) and vehicle control (CONT) treated mice We carried out a linear regression analysis to determine the relationship between IL-1 β and *A. muciniphila* in GWP and CONT mice. There was a negative correlation between *A. muciniphila* and IL-1 β in the FC (Pearson's r=-0.68; R square (COD)=0.46 and p=0.02). **B.** Correlation between *A. muciniphila* and BDNF levels in GW chemical (GWP) and vehicle control (CONT) treated mice. We carried out a linear regression analysis to determine the relationship between BDNF and *A. muciniphila* in GWP and CONT mice. There was a positive correlation between *A. muciniphila* and BDNF levels in GWP and CONT mice. There was a positive correlation between *A. muciniphila* in GWP and CONT mice. There was a positive correlation between *A. muciniphila* relative abundance and BDNF levels (Pearson's r= 0.83, R square (COD)=0.7 and p=0.0024)





Figure 4.7. Deletion of NLRP3 is associated with decreased neuroinflammation and lower BDNF levels.

IL-1 β immunoreactivity in frontal cortex. Representative immunohistochemistry micrographs showing IL-1 β in frontal cortex tissues of GW chemical treated (GWP), GW chemical treated NLRP3KO (GWP-NLR3KO) and vehicle control (CONT) treated mice (Magnification 40X and scale bar 50 μ m). **B.** Morphometric analysis (represented as % ROI) obtained from 10-15 images from different microscopy fields from each mouse sample. Data is represented as Mean ±SEM. (*p<0.05; n=6). **C**. BDNF immunoreactivity in frontal cortex tissues of GW chemical treated (GWP), GW chemical treated NLRP3KO (GWP-NLR3KO) and vehicle control (CONT) treated mice (Magnification 40X and scale bar 50 μ m). **D.** Morphometric analysis (represented as % ROI) obtained from 10-15 images from different microscopy fields from each mouse sample. Data is representative immunohistochemistry micrographs showing BDNF immunoreactivity in frontal cortex tissues of GW chemical treated (GWP), GW chemical treated NLRP3KO (GWP-NLR3KO) and vehicle control (CONT) treated mice (Magnification 40X and scale bar 50 μ m). **D.** Morphometric analysis (represented as % ROI) obtained from 10-15 images from different microscopy fields from each mouse sample. Data is represented as Mean ±SEM. (*p<0.05; n=6). **E.** Western blots of BDNF protein levels in the frontal cortex of GWP and CONT treated mice. **F.** Morphometry analysis of all immunoblots normalized against β -actin. Data is represented as Mean ±SEM. (*p<0.05; n=6).



Supplementary Figures



Figure 4.8. Serum endotoxin levels in GW chemical treated mice (GWP) compared to the vehicle control (CONT) only treated mice.

Data is represented as Mean ±SEM. (*p<0.05; n=6, ns=not significant)



CHAPTER 5

CONCLUSION

Veterans who suffer from GWI report that their symptoms begun during, or shortly after the war. Although it is difficult to trace back the exact causes, research agrees that environmental exposures did in fact contribute significantly to causing the condition. It is perplexing however that such a brief war (about 1 month of active combat) resulted in such far reaching persistent health consequences.

This dissertation reports on three studies which look at at gulf war chemical exposure-induced gut microbiome alterations as a significant driver of GWI pathology. We further investigated possible microbiome targeted therapies to alleviate the symptoms of GWI.

In these three studies, we hypothesized that exposure to GW-chemicals irreversibly altered the microbiome, an important "organ" in modulating health. These alterations in the microbiome resulted in an increase in immunostimulatory particles ie DAMPs and PAMPs and dysfunctional tight junctions in the gut. These particle cross into the blood circulation, affecting the enteric nervous system and distant organs such as the liver and the brain.



In the enteric nervous system, we found that enteric glial cells became reactive in response to an altered microbiome. These cells can detect an alteration in microbiome through toll like receptors such as TLR4. These reactive glia produced large amounts of reactive oxygen species and proinflammatory cytokines which modulate tight junction proteins and aquaporins in the gut. The second study found an association between GW chemical exposure and hepatic inflammation and lipid and glucose metabolic reprogramming but with no significant clinical pathology. The third study found that there were persistent alterations in microbiome at the species levels. Several important gut bacteria were altered in the GW chemical exposed mice. However, this study focused on *Akkermasia muciniphila* a known probiotic bacterium whose levels are usually high among healthy individuals. We found that there was a lowered relative abundance of *A. muciniphila* which correlated negatively with NLRP 3 mediated neuroinflammation and positively with neurotrophin BDNF levels in the brain.

In these studies, we proposed the use of different kinds of therapies.

1. The use of natural therapies such as SsnB, a TLR4 antagonist polyphenol isolated from a Chinese herb *Scirpus yagara*. Since we found that TLR4 plays a significant role in triggering inflammation in GWI. 2. The use of probiotics eg butyrate which support gut bacteria. We found that these short chain fatty acids play an important role in protecting mice against GW chemical induced inflammation and metabolic reprogramming. 3. Replacement of gut bacteria through consumption of depleted probiotic bacteria such as *A*. *muciniphila*, which we found to correlated with neuroinflammation and loss of neurotrophins such as BDNF which are important in neurological health. 4. Finally, the use of innate immune system receptor antagonists such as TLR4, TLR5, NLRP3 and RAGE



can be explored further, since we found that these receptors played a role in promoting GWI pathology. However, this method needs to be applied cautiously since the innate immune system receptors are also critical to health, and long-term inhibition could worsen pathology.

Although a lot of research still remains to be done, these three studies advance the knowledge on GWI pathology, and inform us on how to consider similar conditions, where the use of chemical combinations together with stress and other under lying conditions could result in unexplainable syndromes such as GWI.



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Dysbiosis-Associated Enteric Glial Cell Immune-Activation and Redox Imbalance Modulate Tight Junction Protein Expression in Gulf War Illness Pathology

Diana Kimono, ¹ Sutapa Sarkar, ¹ Muayad Albadrani, ¹ Ratanesh Seth, ¹ Dipro Bose, ¹ Ayan Mondal, ¹ Yuxi Li, ² Amar N. Kar, ³ Mitzi Nagarkatti, ⁴ Prakash Nagarkatti, ⁴ Kimberly Sullivan, ⁵ Patricia Janulewicz, ⁵ Stephen Lasley, ⁶ Ronnie Horner, ⁷ Nancy Klimas, ⁸ and Saurabh Chatterjee^{1,*}

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