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TITLE: Down-Regulation of Olfactory Receptors in Response to Traumatic Brain Injury Promotes Risk for Alzheimer's Disease

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# Down-Regulation of Olfactory Receptors in Response to Traumatic Brain Injury Promotes Risk for Alzheimer’s Disease

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## ABSTRACT
Traumatic Brain Injury (TBI) is a risk factor for subsequent development of Alzheimer’s disease (AD). Abnormal tau processing is a common pathological feature of TBI and AD and tau neuropathology plays a key role in both TBI complications and AD dementia. This study is based on our recent findings of aberrant down-regulation of specific olfactory receptors (OR) as biological indices for TBI and down-regulation of OR TBI biomarkers following TBI might contribute to TBI-related tau neuropathology. We propose that down-regulation of select OR TBI biomarkers in the brain may contribute to the elevation of tau neuropathological phenotypes, thereby promoting the development of AD dementia among Operation Enduring Freedom (OEF) and Operation Iraqi Freedom (OIF) veterans with exposure to TBI. In Year 1, we found that activation of OR4M1 by a low affinity ligand resulted in reduced tau phosphorylation via JNK signaling pathway and a manuscript was published based on this finding. We constructed a virtual 3D structure model for OR4M1 to screen high affinity ligands. Fifty Seven (57) compounds were identified and clustered into 32 clusters based on their structure similarities. We found significant decrease of the blood OR contents in a rat model of TBI. Outcomes from our study will provide a better understanding of TBI complications and how it is related to AD.

## SUBJECT TERMS
Olfactory receptor, tau phosphorylation, virtual screen

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INTRODUCTION
Traumatic Brain Injury (TBI) is an acquired injury caused by a sudden trauma to the head which disrupts normal brain functioning and leads to either transient or chronic impairments in physical, cognitive, emotional, and/or behavioral functions. In the civilian population, TBI is typically associated with direct, close impact mechanical trauma to the brain due to falls, motor vehicle accidents and sports. In contrast, TBI among military personnel, particularly among Veterans returning from the Persian Gulf region as part of service in Operation Enduring Freedom (OEF) or Operation Iraqi Freedom (OIF), is primarily due to exposure to blast pressure waves stemming from blast-producing weaponry, such as improvised explosive devices (IEDs). There is growing evidence of persistent long-term impacts of TBI on cognition. In particular, there is strong epidemiological evidence (Van Den et al., 2007) and growing pathophysiological (Johnson et al., 2012) and experimental (Sivananda and Thakur, 2012) evidence linking prior exposure to traumatic brain injury (TBI) to increased risk for development of Alzheimer’s disease (AD). However, the details of the biological interrelationship between TBI exposure and long-term complications such as AD, how to identify individuals with TBI who are at higher risk for these long-term complications, and how to alter the prognosis of those individuals are poorly understood. We recently found among a civilian TBI study cohort that exposure to TBI is associated with long-term aberrant down-regulations of specific olfactory receptors (ORs) (e.g., OR11H1, OR4M1 and OR52N5) in peripheral blood mononuclear cells (PBMCs) (Zhao et al., 2013). In addition, we found down-regulation of these ORs is significantly correlated with the severity of brain injury and TBI-specific symptoms, and a two-biomarker panel, comprised of OR11H1 and OR4M1, provides the best criterion for segregating the TBI and control cases, with 90% accuracy, 83.3% sensitivity, and 100% specificity (Zhao et al., 2013; Appendix 1). More importantly, we also found lower contents of these ORs in select brain regions among cases who have suffered from TBI, suggesting that down-regulation of these ORs in the brain following TBI might contribute to TBI-related neuropathological alterations leading to long-term neurological complications, such as AD. Collectively our novel observation implicates, for the first time, select ORs as surrogate biological indices (biomarkers) with the potential to reflect long-term physiological alterations in the brain following TBI. Based on this, our overall study is designed to explore whether down-regulation of ORs post-TBI may contribute to long-term TBI complications such as AD, and clarify the underlying mechanisms for this.

BODY
Both AD and TBI exhibit tau neuropathology, which plays a key role in TBI complication and AD dementia. Thus, the study was designed to test the relationship between long-term down regulation of ORs and tau-mediated neuropathogenesis. The overall study is separated into three parts: identification of high affinity, selective ligands for OR4M1, OR11H1 and OR52N5; in vitro study using this high-affinity OR ligands to investigate the impacts of OR downstream signaling on tau neuropathogenic mechanisms; and in vivo assessment of OR and tau signaling in a rodent model of TBI.

In vivo assessment of OR: OR expression in a blast-induced rat model of TBI
Adult male Long Evans hooded rats (250–350g; 10–12 weeks of age) were exposed to overpressure injury using the Walter Reed Army Institute of Research (WRAIR) shock tube, which simulates the effects of air blast exposure under experimental conditions (Elder et al., 2012). Consistent with our observations in human TBI subjects, we found significantly lower contents of select ORs in the blood from the blast-injured group compared to control rats, over 22 weeks following TBI exposure (Fig 1). Our observation demonstrated, for the first time, a direct cause-and-effect relationship between TBI exposure and long-term down regulation of select ORs. More importantly, this observation validates our hypothesis that down regulation of select OR in PBMCs and in the brains of our human TBI study cohort may be caused by prior TBI exposure. Ongoing studies are examining the impacts of blast exposure in modulating the regulation of these ORs in the brains from the blast-induced rat model of TBI.
Fig 1. Peripheral expression of selected olfactory receptors in blast-induced rat model of TBI. Rat blood RNAs were extracted at 4, 10 and 22 weeks following blast. Quantitative PCR was performed using ABI 7900HT Fast Real-Time PCR System. *p<0.05, ***p<0.001 by two-tailed t-test.

There is no information on how TBI exposure might induce long-term changes in the regulation of select ORs. In ongoing studies using the blast-induced rat model of TBI, we are exploring potential epigenetic changes, such as changes in DNA methylation and histone modifications associated with TBI injury, which might lead to down regulation of select OR gene expression long-term.

Investigations on the relationships between OR signaling and tau neuropathogenic mechanisms.
These studies are contingent on the availability of high-affinity, select ligands for individual ORs affected by TBI. As a first step in establishing high-affinity, selective OR ligands, we have identified low-affinity ligands for specific ORs. Using a low-affinity OR4M1 ligand at a concentration of 10µM, we found that exogenous expression and pharmacological activation of OR4M1 reduced tau phosphorylation at Ser396/Ser404 (PHF-1 epitope), coincidental with reduced c-Jun NH2-terminal kinase (JNK) signaling in primary neuron cultures (Fig 2) (Zhao et al., 2013). Our observation validated our hypothesis that long-term down-regulation of ORs in the brain following TBI exposure promotes tau neuropathological responses that potentially contribute to the development of neurologic complications, such as AD. As we described in more details below, our ongoing studies are continuing to isolate and characterize selective, high-affinity OR ligands that we could use to investigate the impact of OR downstream tau signaling on tau neuropathogenic mechanisms.

Fig 2. Activation of OR4M1 resulted in reduced tau phosphorylation via JNK signaling pathway. Primary cortico-hippocampal neuron culture was transduced with lentiviral particles with overexpression of OR4M1 or control lentiviral particles by spin infection (250 g×90 min at 30°C). JNK and tau phosphorylation were measured by western blot analysis using antibodies recognizing phospho-JNK, phospho-tau (PHF-1 epitope) and quantified. *p<0.05, ***p < 0.001 by two-tailed t-test.
Identification of candidate high-affinity, selective OR4M1 ligands via virtual receptor modeling and in silico ligand screening

Construction of a three-dimensional OR receptor model: We constructed a three-dimensional (3D) structure model for OR4M1 by comparative modeling based on the crystal structure of bovine rhodopsin (Okada et al., 2004). Because the sequence similarity between OR4M1 and bovine rhodopsin is below 25%, alignment of the two sequences was built using profiles of homologous olfactory receptors and other G protein-coupled receptors (GPCRs). The profiles were matched to each other using the SALIGN routine of program MODELLER (Martí-Renom et al., 2004). The alignment was then manually edited using predictions of transmembrane helices for OR4M1. A preliminary model of OR4M1 was built using MODELLER version 9.12 (Eswar et al., 2003), using the alignment and the rhodopsin structure as input. The quality of the modeling alignment was evaluated by mapping, which predicted olfactory receptor binding site residues (Man et al., 2004c) on the preliminary model. Most of the predicted binding site residues are spatially close in the model, suggesting that the alignment is reliable. The alignment shows that the region between Ala 142 and Pro 175 of OR4M1 is highly divergent from the equivalent region in rhodopsin, hence this region was further refined using the loop modeling routine in MODELLER (Fiser et al., 2000). A total of 100,000 conformations for the loop were generated and the best scoring one, based on the DOPE potential (Shen and Sali, 2006), was selected as the final model. In the final model, most of the predicted binding site residues from Man et al. (Man et al., 2004b) cluster into one region (Fig. 3), which is consistent with a binding pocket identified in the model using program SiteHound (Ghersi and Sanchez, 2009b).

Virtual screening of the OR4M1 binding pocket: The pocket identified in the 3D OR4M1 model was used as a target for virtual screening of small molecule compounds. A set of ~5 million lead-like compounds (Teague et al., 1999), derived from the ZINC library of commercially available compounds (Irwin and Shoichet, 2005), was docked into the binding site using program DOCK version 6.5 (Moustakas et al., 2006). Lead-like compounds were selected to facilitate optimization of validated hits. The top 700 hits from the DOCK screening (ranked by docking score) were visually inspected, and 57 compounds that showed good occupancy of the predicted binding pocket and at least two hydrogen bonds with the protein were selected. The selected compounds were clustered based on similarity, as measured by the Tanimoto coefficient. Single-linkage clustering was used using a Tanimoto coefficient of 0.7 as the cutoff. This resulted in 32 clusters of structurally similar compounds. The representatives from each cluster (the highest ranking compound in each cluster based on the DOCK score) conform a set of 32 compounds for experimental validation. Ongoing studies are now testing the selectivity and affinity of these 32 compounds as an OR4M1 ligand.
KEY RESEARCH ACCOMPLISHMENTS
1. We assessed blood OR content in a rat model of TBI, found significant decrease of select olfactory receptors, and established a cause-and-effect relationship between TBI exposure and long-term down-regulation of select ORs.
2. We generated evidence supporting that down-regulation of ORs observed in the periphery (in PBMC) following TBI exposure may reflect similar down-regulation of ORs in the brain.
3. We have published one manuscript on decreased level of olfactory receptors in blood cells following TBI and its potential association with tauopathy (Appendix 1).
4. We have found that activation of OR4M1 by a low affinity ligand resulted in reduced tau phosphorylation via JNK signaling pathway.
5. We have constructed a virtual 3D structure for OR4M1.
6. The pocket identified in the OR4M1 model was used as a target for virtual screening of small molecule compounds. 57 compounds were identified and clustered into 32 clusters based on their structural similarities.

REPORTABLE OUTCOMES
We have published one manuscript on decreased level of olfactory receptors in blood cells following traumatic brain injury and potential association with tauopathy (Appendix 1). Additionally, we presented this evidence in our presentation entitled, “Attenuation of neuropathological features shared by TBI and AD through olfactory receptor activation,” at the Arrowhead’s 3rd Annual TBI Conference in Washington D.C., March 6, 2013.

CONCLUSION
We established a cause-and-effect relationship between TBI exposure and down-regulation of ORs. Our evidence also implicates that down-regulation of ORs observed in PBMC following TBI reflects similar down-regulation of ORs in the brain that may contribute to neuropathogenic mechanism, such as tau-mediated neuropathology. Consistent with this, we found that activation of OR4M1 by a low-affinity ligand resulted in reduced tau phosphorylation via JNK signaling pathway; a manuscript was published based on this finding. We constructed a virtual 3D structural model for OR4M1 to screen for potential high-affinity ligands. The pocket identified in the OR4M1 model was used as a target for virtual, high-throughput screening of small molecule compounds. 57 compounds were identified and clustered into 32 clusters based on their structural similarities. In ongoing studies, we are testing the ability of compounds, identified from virtual screening, in activation of OR4M1 in a cell culture system. Once confirmed in vitro, high affinity ligands will be used to assess downstream tau signaling. We found a significant decrease of the blood OR content in a rat model of TBI. We are continuing to monitor the blood OR content in the rat model of TBI. We will also assess the brain OR...
content and tau phosphorylation in this rat model of TBI. Outcomes from our study will provide a better understanding of TBI complications and how they are related to AD.

REFERENCES


APPENDIX
Decreased Level of Olfactory Receptors in Blood Cells Following Traumatic Brain Injury and Potential Association with Tauopathy

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Abstract. Traumatic brain injury (TBI) is a leading cause of death and disability among children and young adults in the United States. In this study, we explored whether changes in the gene expression profile of peripheral blood mononuclear cells (PBMC) may provide a clinically assessable “window” into the brain, reflecting molecular alterations following TBI that might contribute to the onset and progression of TBI clinical complications. We identified three olfactory receptor (OR) TBI biomarkers that are aberrantly down-regulated in PBMC specimens from TBI subjects. Down-regulation of these OR biomarkers in PBMC was correlated with the severity of brain injury and TBI-specific symptoms. A two-biomarker panel comprised of OR11H1 and OR4M1 provided the best criterion for segregating the TBI and control cases with 90% accuracy, 83.3% sensitivity, and 100% specificity. We found that the OR biomarkers are ectopically expressed in multiple brain regions, including the entorhinal-hippocampus system known to play an important role in memory formation and consolidation. Activation of OR4M1 led to attenuation of abnormal tau phosphorylation, possibly through JNK signaling pathway. Our results suggested that addition of the two-OR biomarker model to current diagnostic criteria may lead to improved TBI detection for clinical trials, and decreased expression of OR TBI biomarkers might be associated with TBI-induced tauopathy. Future studies exploring the physiological relevance of OR TBI biomarkers in the normal brain and in the brain following TBI will provide a better understanding of the biological mechanisms underlying TBI and insights into novel therapeutic targets for TBI.

Keywords: Biomarker, olfactory receptor, peripheral blood mononuclear cell, tauopathy, traumatic brain injury

Supplementary data available online: http://www.j-alz.com/issues/34/vol34-2.html#supplementarydata04

INTRODUCTION

Traumatic brain injury (TBI) is a leading cause of death and disability among children and young adults in the United States [1]. TBI is an acquired injury caused by a sudden trauma to the head that disrupts
normal brain functioning, which leads to either transient or chronic impairments in physical, cognitive, emotional, and/or behavioral functions. In the civilian population, TBI is typically associated with direct, closed impact mechanical trauma to the brain due to falls, motor vehicle accidents, sports, etc. [2]. In contrast, TBI among military personnel, particularly among veterans returning from the Persian Gulf region, is primarily due to exposure to blast pressure waves stemming from blast-producing weaponry, leading to prototypical cognitive deficits including impairments in attention, memory, processing speed, and executive functioning [2–4]. Both civilians and veterans who suffer from TBI exhibit symptoms that range in severity from mild to very severe, with a minimal to profound impact on daily functioning. The reasons why TBI induces different clinical symptoms among affected individuals are not yet known.

New evidence has highlighted defects in neural circuit and synapses, and the plastic processes controlling these functions, in TBI [5–11]. While genes relevant to these processes are expressed in the brain, some of these genes are also expressed in circulating blood cells, such as peripheral blood mononuclear cells (PBMCs) [12–15]. Consistent with this, recent studies have illustrated that PBMC-associated biomarkers may provide insights into the pathogenesis of neurological disorders such as Alzheimer’s disease and can be used to monitor disease diagnosis and progression [16, 17]. Thus, PBMC may also provide an ideal, clinically assessable “window” into the brain, reflecting molecular alterations following TBI which might contribute to the onset and progression of clinical TBI phenotypes.

Evidence suggests that appropriate interventions can reduce functional impairment after TBI [18–20]. In order to demonstrate the efficacy of clinical interventions, research must identify the biological, clinical, and neurological indexes that are sensitive to the detection of functional impairments after TBI. Therefore, in this study we explored the feasibility of identifying clinically assessable TBI biomarkers and the potential function of identified TBI biomarkers in TBI neuropathology.

MATERIALS AND METHODS

Civilian TBI and control subjects

Eleven individuals with TBI (five male and six female), with TBI severity ranging from mild to severe, and nine control participants (four male and five female) were recruited from the Brain Injury Research Center at the Mount Sinai School of Medicine (MSSM). Medical documentation of TBI was reviewed for each participant. The age- and education-matched controls were recruited using advertisements placed in local media and flyers, and through word of mouth. Following a capacity screening to assess the individual’s ability to comprehend the purpose and procedures of the study and provide informed consent, the informed consent process was performed in accordance with MSSM IRB policies and procedures.

Veteran TBI and control subjects

For this biomarker validation study, we obtained banked PBMCs from a cohort of five OEF/OIF veterans and a control cohort of seven age- and gender-matched veterans without TBI (non-TBI veterans). TBI diagnosis is based on confirmation according to Defense and Veterans Brain Injury Center (DVBIC) criteria of sustained injury to the head plus subsequent alteration of consciousness, and Repeatable Battery for Neuropsychological Testing (RBANS) scores of one standard deviation below the norms established for age and education of individuals in question. Non-TBI control classification is based on DVBIC confirmation of no sustained injury to the head and RBANS scores less than one standard deviation below the norms established. Most of the veteran TBI cases exhibit co-morbid post-traumatic stress disorder (PTSD). Thus, our non-TBI control cases are also matched for PTSD.
diagnosis. In this study, diagnosis of PSTD is based on a score of 50 or more in the PTSD Checklist – Civilian Version.

Postmortem brain specimens

Human postmortem brain samples from 4 neurologically normal cases (characterized by a Clinical Dementia Rating (CDR) of 0) were obtained from the Alzheimer’s Disease Brain Bank of the Mount Sinai School of Medicine. A multistep approach based on cognitive and functional status during the last 6 months of life was applied to the assignment of CDR [23] as previously reported [24, 25].

PBMC and RNA isolation

Blood specimens were collected into BD Vacutainer CPT Cell Preparation Tubes. PBMC were isolated following manufacturer’s instructions (Becton, Dickinson and Company) and were stored at −80°C until use. Total RNA was isolated from approximately 10–50 mg of PBMC using RNA STAT-60 (Tel-Test) according to the manufacturer’s instructions. The purity and concentration of RNA samples were determined from OD260/280 readings using a dual beam UV spectrophotometer and RNA integrity was determined by capillary electrophoresis using the RNA 6000 Nano Lab-on-a-Chip kit and the Bioanalyzer 2100 (Agilent Technologies).

Microarray study and analysis

Total RNA was directly labeled using the FlashTag™ HSR Biotin RNA Labeling Kit according to the manufacturer’s instructions (Genisphere). Verification of biotin labeling was obtained by an enzyme-linked oligoabsorbant assay (ELOSA) using Immobilon™ Amino – 8 well strips (Nunc/Thermo Fisher Scientific) according to instructions supplied by Genisphere. Labeled RNA (1.0 μg) was hybridized for 16 h at 48°C to Affymetrix HuGene 1.0 ST arrays containing probe sets for 28,869 genes. Arrays were washed and stained on a Fluidics Station 450 (Affymetrix) according to the manufacturer’s recommended procedures. The arrays were stained with phycoerythrin-conjugated streptavidin (Life Technologies) and the fluorescence intensities were determined using a GCS 3000 TG high-resolution confocal laser scanner and AGCC software (Affymetrix). The scanned images were analyzed with the RNA QC tool (Affymetrix) using RNA global background correction, quantile normalization and median polish summarization to generate quantified data (as recommended by Genisphere). Quality control metrics for arrays included normalized signal values >1000 for five spike-in control oligo probe sets (Genisphere). RNA probe sets exhibiting significant differential expression (SDE) were identified using the following steps in GeneMaths XT (Applied Maths): 1) Probe sets with array detection p-values ≤0.05 for all samples in at least one experimental group were selected for further analysis; 2) Performed Discriminant Analysis (DA) and determined the largest percentage of remaining probe sets that permitted correct group assignment of samples in unsupervised hierarchical clustering by the Unweighted Pair-Group Method using Arithmetic averages (UPGMA) based on cosine correlation of row mean centered log2 signal values; this was the top 50 percentile; and 3) in the DA top 50 percentile, selected probe sets with absolute signal log2 fold changes ≥1.0 and independent t-test p-values ≤0.05 adjusted for multiple testing error by the Benjamin-Hochberg FDR correction method. Unsupervised hierarchical clustering of probe sets and heat map generation were performed in GeneMaths XT following row mean centering of log2 transformed MA5.0 signal values; probe set clustering was performed by the UPGMA method using Cosine correlation as the similarity metric. For comparative purposes, clustered heat maps included probe sets for spike-in controls (Genisphere), or endogenous small RNAs exhibiting: 1) Array detection p-values ≤0.05, and 2) either a) a log2 signal value standard deviation ≤0.025 for all samples or b) in the DA top 50 percentile with a FC >1.3 in the opposite direction of the selected SDE profile.

Confirmatory quantitative PCR

First strand cDNA was synthesized from 1 μg of total RNA using Superscript III Super-mix for qRT-PCR (Invitrogen). Quantitative RT-PCR was performed using Maxima SYBR Green master mix (Fermentas) in ABI Prism 7900HT in four replicates with primers amplifying the seven olfactory receptor candidates: OR4Q3 (Forward: CACCTGCTCCAATCTCATG, Reverse: TCCCCTAACATCTTTGGCAC); OR51L1 (Forward: TTCCCACACCTTTGCTACTG, Reverse: AATACTGTTGGTCCTGGCATC); OR4D10 (Forward: CCACTGCTCCAATCTCATG, Reverse: ATGGCTGACTTCATCTCATGG); OR4M1 (Forward: TCTGTTAATGTCCTATGCTTCC, Reverse: AATGTGGGAATAGCAGGTGG);
OR52N5 (Forward: ATGCTACCACTCTACCAAC, Reverse: CATCAGCAATACCCCTCAG); OR2J3 (Forward: CCTCTCATCCTCTACCTTC, Reverse: CAAACACTTTCTGAAGCCCG); OR11H1 (Forward: AACTGGTCATACTGTGCTGG, Reverse: GGCGAAGGATCCATTTTGAA). Human TATA-binding protein (TBP, Forward: TGCACAGGAGCCAAAGTGAA, Reverse: CTGGAACGGTGAAGGTGACA) expression level was used as an internal control. Data were normalized using the 2^{−ΔΔCt} method [26]. Levels of olfactory receptor mRNAs were expressed relative to those in control groups and plotted in GraphPad Prism.

**Lentiviral plasmid construction and lentivirus packaging**

We obtained pCMV6-XL5-OR4M1 cDNA clone from Origene and subcloned the OR4M1 ORF into lentiviral plasmid pLVX-IRES-ZsGreen (Clontech). The inserted OR4M1 sequence was verified by sequencing. For lentivirus packaging, we transfected Lenti-X 293T cells with either pLVX-OR4M1-IRES-ZsGreen or pLVX-IRES-ZsGreen using Lentiviral Packaging System (Clontech). Medium was collected 48 h after transfection and a ten-fold concentration step was performed using Lenti-X Concentrator (Clontech).

**Primary neuron culture, lentiviral transduction, and cAMP assay**

Embryonic day 15 cortico-hippocampal neuronal cultures were prepared from C57BL6 mouse (Jackson Laboratory) as previously described [27]. Cells were seeded onto poly-D-lysine-coated 12-well plates at 5 × 10^5 cells per well and cultured in Neurobasal medium supplemented with 2% B27, 0.5 mM L-glutamine, and 1% penicillin-streptomycin (Life Technologies). On Day 5 of culture, primary cortico-hippocampal neuron cultures were transduced with lentiviral particles overexpressing OR4M1 or control lentiviral Packaging System (Clontech). Medium was collected 48 h after transfection and a ten-fold concentration step was performed using Lenti-X Concentrator (Clontech).

Luminex multiplex assay and western blot analysis

Primary cortico-hippocampal neurons were infected with OR4M1 lentiviral particles and stimulated with acetophenone (10 μM) for 1 h. Multiplex luminex assay was performed using the Milliplex xMAP 8-plex multipathway signaling-phosphoprotein kit (Millipore) as previously described [28]. Protein concentrations were determined using the BCA protein assay kit (Thermo Scientific). 30 μg of total protein were loaded onto 12% SDS-PAGE and subjected to western blot analysis with antibody recognizing phospho-JNK (T183/Y185) (Cell Signaling), phospho-ERK (T185/Y187) (Cell Signaling), and PHF-1 (phospho-tau (S396/S404), a generous gift from Peter Davies at Albert Einstein College of Medicine). Blots were quantified in ImageJ (NIH), normalized with glyceraldehyde-3-phosphate dehydrogenase (GAPDH) (Santa Cruz Biotechnologies) and then plotted in GraphPad Prism.

**RESULTS**

**Identification of select olfactory receptors as clinically accessible candidate TBI biomarkers**

We analyzed the gene expression profile in PBMC derived from subjects with TBI and age-, gender-, and education-matched healthy controls. Our data showed that female subjects had a more robust profile than males probably due to a rather smaller number of male subjects in the study cohort. Taking this into consideration, further analysis and confirmatory studies were all performed in a subset of the study cohort which only contains female subjects. Demographic information for the female TBI and control participants is summarized in Table 1. Using microarray gene expression profile analysis, we identified a panel of 102 candidate biomarker genes (36 annotated and 66 non-annotated genes) that are differentially regulated (t-test p < 0.05) following Benjamini-Hochberg false discovery rate correction [29] by >1.5-fold in PBMC specimens of TBI cases compared to control cases (Fig. 1A). More importantly, we were able to correctly separate all cases into TBI and non-TBI groupings using this panel of 102 genes in an unsupervised hierarchical clustering analysis (Fig. 1A). Interestingly, among the panel of 102 candidate TBI biomarkers that we identified from microarray data, 7 were olfactory receptors (Fig. 1B). Our microarray studies revealed that each of the 7 ORs were down-regulated in PBMC.
Table 1

Demographic information for the civilian study cohort. TBI severity was classified using a 7-point scale ranging from 1 (no loss of consciousness, no confusion (i.e., no TBI)) to 7 (loss of consciousness greater than 4 weeks in duration) [23]. Time post injury is the time frame between the occurrence of brain injury and volunteer’s participation in this biomarker study.

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Confirmatory qPCR studies

Using independent qPCR assays, we confirmed that three of the ORs (OR11H1, OR4M1, and OR52N5) were significantly down-regulated in PBMC of TBI cases (Fig. 2B). For the remaining four ORs (OR4D10, OR2J3, OR4Q3, OR51L1), our qPCR evidence showed lower levels of these ORs in PBMC of TBI compared to control cases, but these differences did not reach statistical significance (Fig. 2B). This particular result is conceivably due to a high degree of gene expression variability for these 4 ORs among healthy control cases and the relatively small sample sizes that we used in our exploratory study.

We continued to explore the sensitivity and specificity of an individual or combined role of OR11H1, OR4M1, and OR52N5 in distinguishing TBI cases from normal healthy controls in our study cohort (Fig. 2C). Using unsupervised clustering analyses, we found that a two-biomarker panel comprised of OR11H1 and OR4M1 provides the best criterion for segregating the TBI and control cases with 90% accuracy, 83.3% sensitivity, and 100% specificity (Fig. 2C, D).

Fig. 1. Development of blood biological indices (biomarkers) capable of correctly segregating TBI and control cases. Gene expression profile analysis of PBMC specimens from TBI and 5 age-, gender-, and education-matched control cases using a microarray platform (Affymetrix) led to the identification of a panel of 102 candidate biomarker genes. A) The 108 differentially-regulated genes identified were subjected to unsupervised hierarchical clustering analysis using the UPGMA algorithm with cosine correlation as the similarity metric. Results are presented as a heat map (left panel) demonstrating that the content of the 108 biomarker panel is able to correctly segregate TBI from control cases. B) 7 ORs and one OR pseudogene that are down-regulated in PBMC of TBI cases.
Fig. 2. Down-regulation of OR genes in PBMC of TBI cases provide a sensitive and specific criterion for distinguishing TBI from control cases. mRNA contents for each of the 7 candidate OR biomarker genes in TBI and control cases were analyzed by microarray (A) or independent qPCR (B). Bar graphs represent mean ± SEM. *p < 0.05, **p < 0.01, ***p < 0.001 by student t-test, TBI versus control. The efficacy of using biomarker contents from PBMC as a criterion to correctly segregate TBI and control cases was tested by unsupervised clustering analysis using the UPGMA algorithm with cosine correlation as the similarity metric. C) The accuracy, sensitivity, and specificity of OR11H1, OR4M1, OR52N5, or panels of ORs to distinguish TBI from control cases. D) A heat map graphically depicting the efficacy of using the two biomarker panel to distinguish TBI cases and control cases by unsupervised clustering analysis.

TBI biomarker validation studies: Testing the validity of OR11H1, OR4M1, and OR52N5 to distinguish TBI from non-TBI control cases from a study cohort of OEF/OIF veterans

The pathophysiological mechanisms underlying mechanical and blast-related TBI may differ in some ways, but they share important pathophysiological features. Similarities between the pathophysiologies of mechanical and blast-related TBI [2] suggested that information gathered from TBI cases in the civilian population may also be relevant to combat-related TBI. Based on this, we tested the ability of OR11H1, OR4M1, and OR52N5 to distinguish TBI cases from non-TBI controls from a study cohort of OEF/OIF veterans. For this biomarker validation study, we obtained banked PBMCs from a cohort of five OEF/OIF veterans and a control cohort of seven age- and gender-matched veterans without TBI (non-TBI veterans) (Table 2). Similar to our above-mentioned observations regarding OR biomarkers from the civilian study cohort, we found that OR4M1 (Fig. 3A), OR11H1 (Fig. 3B), and OR52N5 (Fig. 3C) are also significantly down-regulated in PBMC specimens of veteran TBI cases compared to that of non-TBI control cases. The two biomarker panel of OR11H1 and OR4M1 we identified from the civilian TBI study cohort is also capable of distinguishing veteran TBI from control veteran cases with 83% accuracy, 80% sensitivity, and 86% specificity (Fig. 3D & E).

OR biomarker contents in PBMCs are correlated with TBI severity & long-term clinical neuropsychological complications

Continuing to explore the potential relevance of OR11H1, OR4M1, and OR52N5 to TBI clinical symptoms, we found that the contents of the three OR biomarkers in PBMC was significantly and inversely correlated with TBI severity: lower OR mRNA
Table 2

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>TBI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of cases</td>
<td>7</td>
<td>5</td>
</tr>
<tr>
<td>Gender</td>
<td>57% male</td>
<td>80% male</td>
</tr>
<tr>
<td>Age (mean ± SD)</td>
<td>30.1 ± 9.4</td>
<td>31.0 ± 7.1</td>
</tr>
<tr>
<td>Interval since last deployment (mean ± SD)</td>
<td>2.8 ± 2.3</td>
<td>4.0 ± 3.8</td>
</tr>
<tr>
<td>Education (mean ± SD)</td>
<td>13.3 ± 2.8</td>
<td>13.8 ± 1.5</td>
</tr>
<tr>
<td>Percentage of cases with co-morbid PTSD</td>
<td>85.7%</td>
<td>100%</td>
</tr>
<tr>
<td>Ethnicity composition</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Black-African American</td>
<td>14.3%</td>
<td>40%</td>
</tr>
<tr>
<td>Hispanic</td>
<td>85.7%</td>
<td>60%</td>
</tr>
</tbody>
</table>

Demographic information for the veteran study cohort. TBI diagnosis is based on confirmation according to Defense and Veterans Brain Injury Center (DVBIC) criteria of sustained injury to the head plus subsequent alteration of consciousness, and Repeatable Battery for Neuropsychological Testing (RBANS) scores of one standard deviation below the norms established for age and education of individuals in question. Non-TBI control classification is based on DVBIC confirmation of no sustained injury to the head and RBANS scores less than one standard deviation below the norms established. Most of the veteran TBI cases exhibit co-morbid post-traumatic stress disorder (PTSD). Thus, our non-TBI control cases are also matched for PTSD diagnosis. In this study, diagnosis of PTSD is based on a score of 50 or more in the PTSD Checklist – Civilian Version.

Contents in PBMCs are associated with exposure to more severe head injury (Fig. 4A–C). We also found that OR TBI biomarker contents in PBMC were significantly correlated with select aspects of self-reported clinical TBI symptoms. In particular, PBMC contents of OR11H1 or OR4M1 were significantly associated in an inverse fashion with TBI-specific symptoms, a summation of 25 cognitive symptoms that are sensitive and specific to TBI [22]. OR11H1 or OR4M1 content within PBMC were not associated with self-assessments of mood (Fig. 4D). Expression level of the third OR biomarker, OR52N5, was not correlated with TBI-specific symptoms or self-assessment of mood (Fig. 4D).

Ectopic expression of OR TBI biomarkers in multiple brain regions outside of the olfactory bulb

The brain represents a key target tissue for understanding TBI clinical complications and for the development of clinical interventions. In order to explore the potential physiological relevance of OR11H1, OR4M1, and OR52N5 in the brain, we assessed the RNA contents of OR biomarkers in the brain. We found that the three TBI biomarker ORs were expressed in multiple regions of the brain from post-mortem brain specimens, including temporal gyrus (BM22), entorhinal cortex (BM36), occipital cortex (BM17), and the hippocampal formation (Fig. 5A & B). In agreement with our RT-PCR data, we also
found the expression of OR TBI biomarkers in the hippocampal formation region from the human brain genome-wide microarray database [30] (Fig. 5C–E). Our observation that the three OR biomarkers were ectopically expressed in multiple brain regions outside of the olfactory bulb suggested potential function(s) of these ORs in the brain unrelated to the detection and processing of olfactory information.

Activation of OR TBI biomarker modulates tau neuropathology-related phenotypes in vitro

Olfactory receptors are members of the class A rhodopsin-like family of G protein-coupled receptors (GPCRs). Once the odorant has bound to the odor receptor, the receptor undergoes structural changes and it binds and activates the olfactory-type G protein on the inside of the olfactory receptor neuron. The G protein (Golf and/or Gs) in turn activates adenylate cyclase, which converts ATP into cyclic AMP (cAMP). So we first examined the expression level of Golf and β-arrestin in the PBMC of TBI and control subject and no change was detected (Supplementary Figure 1; available online: http://www.j-alz.com/issues/34/vol34-2.html#supplementarydata04).

Considering that tau neuropathology is a main pathological feature following TBI, we continued to explore whether OR4M1 activation might influence tau processing mechanisms. We transduced primary cortical-hippocampal neuron culture with OR4M1 lentiviral particles and screened several odorants for their ability to activate OR4M1 using cAMP ELISA assay. In Fig. 6A, we presented some representative odorants that were capable or not capable of activating OR4M1. Positive ligands were used to stimulate OR4M1-overexpressing primary cortico-hippocampal neurons and the potential impact of activation of OR4M1 on signaling pathways and aberrant tau phosphorylation were assessed. Using luminex multiplex technology, we found that upon acetophenone stimulation, phosphorylation of c-Jun N-terminal kinase
Fig. 5. Olfactory receptor TBI biomarkers expression in the brain. OR11H1, OR4M1, and OR52N5 mRNA expression by (A) RT-PCR and (B) relative expression level in postmortem superior temporal gyrus (BM22), hippocampal formation (HF), occipital cortex (BM17), and entorhinal cortex (BM36) specimens from neurologically normal cases. Expression pattern of (C) OR4M1, (D) OR11H1, and (E) OR52N5 in the brain by genome-wide microarray from the Allen Human Brain Atlas [30].

(JNK) at Thr183/Tyr185 was significantly reduced in neurons overexpressing OR4M1 (Fig. 6B), indicating the activation of OR4M1 might influence JNK signaling pathway. The phosphorylation of ERK1/2 (Fig. 6B), among others (STAT3, MEK, p70S6, IKBα, and CREB, see Supplemental Figure 2), did not change upon acetophenone stimulation, suggesting that this inhibition of JNK signaling is rather specific. The stress-activated kinase JNK belongs to the mitogen-activated protein kinase family and takes part in signaling cascades initiated by various forms of stress. Its targets include the microtubule-associated protein tau [31]. Hyperphosphorylation of tau at Ser396/Ser404 is a known feature associated with tau neuropathology in neurodegenerative disorders such as Alzheimer’s disease. Encouragingly, we observed that overexpression and activation of OR4M1 in primary neurons significantly reduced cellular content of abnormally phosphorylated tau at Ser396/Ser404 (PHF1 epitope) (Fig. 6C), suggesting that activation of OR4M1 might result in protection against abnormal tau processing.

Collectively, our data demonstrated that select olfactory receptors (e.g., OR11H1, OR4M1, and OR52N5) were down-regulated in the PBMC of TBI cases and could serve as TBI biomarkers. Activation of OR4M1 resulted in protection against tau neuropathological features possibly through the JNK signaling pathway (Fig. 7).

DISCUSSION

Consistent with accumulation evidence suggesting that PBMC-associated biomarkers may provide
insights into the pathogenesis of neurological disorders, results from our studies revealed that expression of select ORs in PBMC may serve as clinically assessable surrogate biological indices of TBI. ORs are G protein-coupled receptors known to be expressed in nasal epithelium olfactory neurons, where they are responsible for the detection of odorants [32]. However, ectopic expression of select ORs, defined as a biological event or process that occurs in an abnormal location or position in the brain or other tissues, has also been described [33]. Our high throughput microarray evidence, followed by data from independent qPCR confirmatory studies, led to the identification of three OR TBI biomarkers (OR11H1, OR4M1, and OR52N5) that are ectopically expressed in PBMC and are aberrantly down-regulated in PBMC specimens from subjects with a history of TBI. Among the TBI cases in our study cohort, there was a significant time lag between the occurrence of brain injury and the volunteers’ participation in this biomarker study, the average post-injury interval among the TBI cases was 5.4 ± 5.3 years (Table 1). Thus, down-regulation of the three ORs in PBMC among TBI cases was reflections of long-term physiological consequences of TBI.

We found that down-regulation of the three OR biomarkers in PBMC were directly correlated with the severity of brain injury in our TBI participants. Among the three OR biomarkers, we found that a two-biomarker panel, comprised of OR11H1 and OR4M1, provides the best criterion for segregating the TBI and control cases with 90% accuracy, 83.3% sensitivity, and 100% specificity. Interestingly, we found that PBMC contents of OR11H1 and OR4M1 were inversely correlated with cognitive-related symptoms.
While additional studies will be required to clarify the mechanisms underlying the inter-relationships between changes in OR biomarker contents in PBMC, the initial severity of brain injury, and long-term clinical consequences of TBI outcomes from our studies suggest that additional applications of this two-OR biomarker panel to current diagnostic criteria may lead to improved TBI detection and more sensitive outcome measures for clinical trials.

Initial and persistent cognitive deficits are the most common complaints after TBI [34, 35]. The entorhinal-hippocampus system is known to play an important role in the formation and consolidation of memories, particularly spatial memories [36–38]. Among the brain regions surveyed, we found the highest content of OR TBI biomarkers in the hippocampal formation, with relatively lower levels of OR TBI expression in the entorhinal cortex. Based on this and on our observation that the down-regulation of OR11H1 and OR4M1 in the PBMC are inversely correlated with self-reported indexes of cognitive functions, it is likely that ectopic expression of OR biomarkers in the entorhinal-hippocampus circuitry might be relevant to long-term deficits in cognitive functions following TBI.

It is well known that TBI may be a risk factor for dementia [39], but recent evidence suggests that Alzheimer’s disease-related neuropathological mechanisms may contribute to cognitive dysfunction in TBI. The two characteristic neuropathologies of Alzheimer’s disease are the abnormal accumulation and deposition of amyloid-β peptides and tau proteins in the brain. Evidence from humans [40–42] and experimental animal models [43] has also revealed abnormal accumulations of amyloid-β peptides and tau proteins in the brain and in cerebrospinal fluid following TBI. Interestingly, elevation of plasma tau levels has been associated with increasingly severe outcomes following TBI. Multiple signaling pathways have been demonstrated to be involved in TBI pathology, including the AKT signaling pathway [45], glycogen synthase kinase-3 signaling pathway [46], STAT3 signaling pathway [47], ERK signaling pathway [48], and JNK signaling pathway [49]. We found that activation of OR4M1 could lead to attenuation of tau neuropathology through OR activation.

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REFERENCES


