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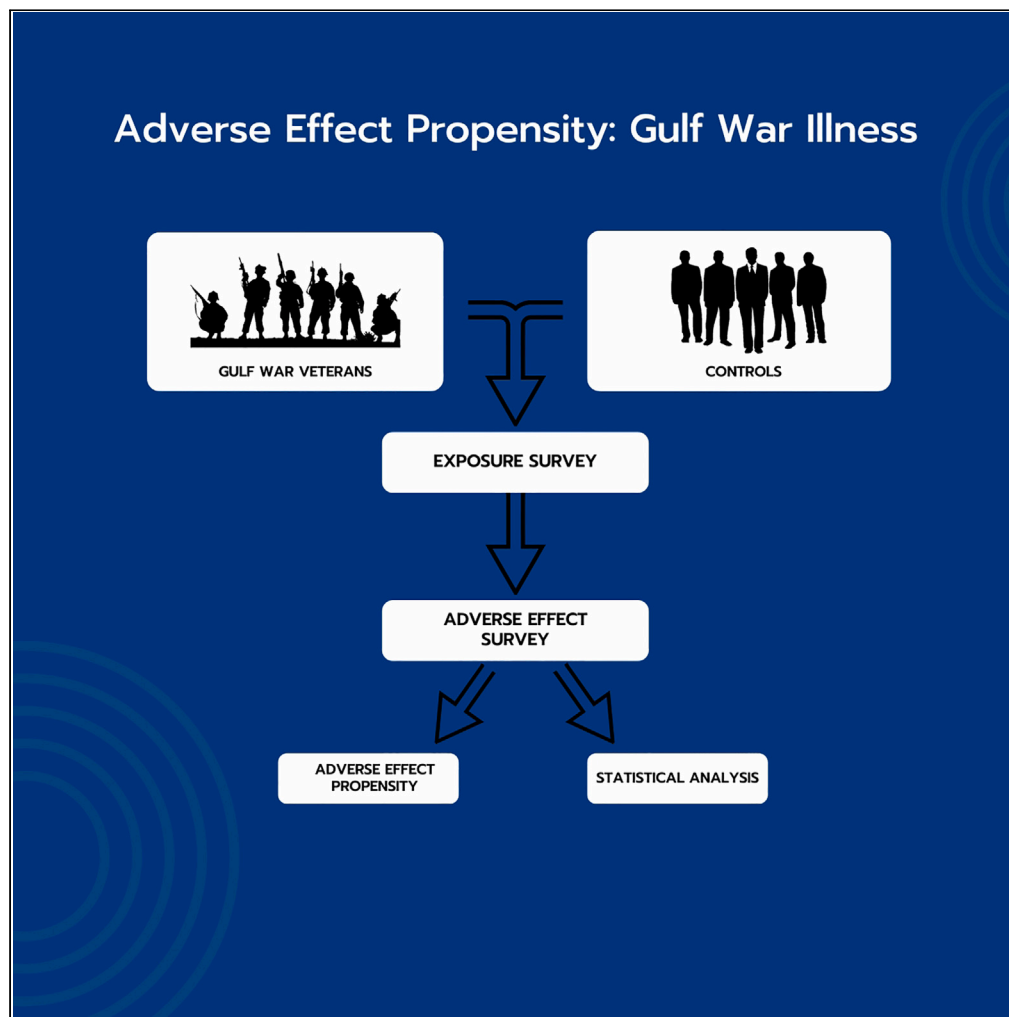
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## Article

## Adverse effect propensity: A new feature of Gulf War illness predicted by environmental exposures



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**Highlights**

Some people experience adverse effects (AEs) to many drugs/chemicals of differing class

Total AE Propensity (TAEP) was greater in veterans with Gulf War illness (VGWI)

Pesticides and radiation predicted greater TAEP, while copper appeared protective

Exposures reproducibly predicted TAEP in all, VGWI, controls, and age strata

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## Article

## Adverse effect propensity: A new feature of Gulf War illness predicted by environmental exposures

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## SUMMARY

**A third of 1990-1 Gulf-deployed personnel developed drug/chemical-induced multisymptom illness, “Gulf War illness” (GWI). Veterans with GWI (VGWI) report increased drug/exposure adverse effects (AEs). Using previously collected data from a case-control study, we evaluated whether the fraction of exposures that engendered AEs (“AE Propensity”) is increased in VGWI (it was); whether AE Propensity is related to self-rated “chemical sensitivity” (it did); and whether specific exposures “predicted” AE Propensity (they did). Pesticides and radiation exposure were significant predictors, with copper significantly “protective”—in the total sample (adjusted for GWI-status) and separately in VGWI and controls, on multivariable regression. Mitochondrial impairment and oxidative stress (OS) underlie AEs from many exposures irrespective of nominal specific mechanism. We hypothesize that mitochondrial toxicity and interrelated OS from pesticides and radiation position people on the steep part of the curve of mitochondrial impairment and OS versus symptom/biological disruption, amplifying impact of new exposures. Copper, meanwhile, is involved in critical OS detoxification processes.**

## INTRODUCTION

Gulf War illness (GWI) is a chronic multisymptom illness affecting approximately one-third of the ~700,000 U.S. troops deployed to the 1990-1 Persian Gulf theater of operations.<sup>1,2</sup> It is a symptom-characterized condition, typified by multiple symptoms spanning protean domains, including fatigue/sleep, pain, neurological, respiratory, gastrointestinal, dermatological, and autonomic.<sup>3</sup> Numerous objective markers have been found to be altered in GWI,<sup>4–7</sup> though there is no pathognomonic test. Alterations with replicated documentation include increased inflammation,<sup>8,9</sup> mitochondrial impairment,<sup>6,10–12</sup> altered eicosanoids,<sup>4</sup> altered hormone status,<sup>13,14</sup> altered autonomic function,<sup>7,15,16</sup> and increased autoantibodies<sup>5,17,18</sup>—among others.

Veterans with GWI (VGWI) have repeatedly been found to be at increased risk of developing multiple chemical sensitivity (MCS).<sup>3,19–24</sup> Many also report intolerance to numerous medications and some cite intolerance to electromagnetic radiation (EMR).<sup>25</sup> In a recent meeting of the Department of Veterans’ Affairs Research Advisory Committee on Gulf War Veterans’ Illnesses,<sup>26</sup> in the “Public Comment” session, a veteran inquired whether/what additional work was being done on chemical sensitivity in VGWI.

Adverse effects (AEs) of many drugs, chemicals, and radiation involve intertwined mechanisms of oxidative stress and mitochondrial impairment—irrespective of the nominal specific mechanism of action of the agent (or frequency of the radiation).<sup>27–43</sup> MCS, nonionizing radiation sensitivity, and ionizing radiation sensitivity involve oxidative and mitochondrial mechanisms.<sup>44–49</sup> Thus, a relationship between propensity to experience AEs to environmental factors, including radiation, and self-reported chemical sensitivity is plausible. Since mitochondrial injury may lead to ongoing oxidative stress/free radical production,<sup>50–52</sup> leading to increased competition for antioxidant defenses—and positioning mitochondrial injury further along the path toward clinical mitochondrial “threshold effects”<sup>53,54</sup> that may be triggered by a new exposure, it is plausible that specific past exposures may have contributed to enhanced risk of AE vulnerability in VGWI.

The UCSD Gulf War illness study is an observational study that has produced findings on Gulf War illness relations to metabolomics, prostaglandins and leukotrienes, malondialdehyde, citric acid cycle markers, and the vaccine experience.<sup>4,12,55,56</sup> Data from that study were here used to examine two questions: first, have VGWI indeed manifested increased apparent vulnerability to AEs of exposures received (case-control

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analysis). Second, to what extent are specific exposures associated with vulnerability to AEs spanning exposures to many drugs and chemicals—and how does this AE Propensity relate to self-rated chemical sensitivity (cross-sectional analysis). This was not prespecified but considering the challenges to study participation for this compromised group, we consider there to be an ethical desideratum to glean the greatest possible information from the sacrifices made by veterans to participate; and to accelerate science and hypothesis generation related to concerns of this group, while VGWI remain alive to potentially benefit from the knowledge gained.

## RESULTS

**Participants:** The study involved a one-time visit. Because there was only one visit, no dropouts between visits occurred. All eligible/interested qualifying cases and matching eligible/interested controls were seen and evaluated, and all eligible parties included in analysis. One extra case was recruited to expand demographic options for identifying a matching control. However, for this study, all participants, including the “extra” (unmatched) case, are included in the analysis.

As [Table 1](#) shows, GWI cases and controls were (by selection) closely similar on age, sex, and ethnicity. VGWI were more likely to be married, perhaps reflecting that for those with this condition, those who have social support are more likely to be able to add research study participation to their challenging lives.

[Table 1](#) also shows symptom scores based on Kansas criteria, for all, and for cases and controls separately. Different numbers of symptom queries are asked for the different domains, providing large differences in maximum possible score, and accounting for material differences in mean scores across domains, in all, in GWI cases, and in controls. The mean summed symptom score for affected veterans is almost 40, and is approximately 75 times higher than the mean score for controls. (Cases are selected for/defined by the presence of symptoms in these categories, as described previously; while controls are selected for a dearth of such symptoms.)

The focus of our analysis was on reported AEs, rather than whether participants considered themselves to be chemically sensitive. However, some data are available for the latter. [Table 1](#) shows self-rated chemical sensitivity, based on a Kansas criteria question and our own question.

“chemsenssx” was assessed as a binary (and also as an ordinal variable) variable, “chemsensbinary”, rated 1 if the chemsenssx variable was rated 1, 2, or 3; and 0 if it was rated 0. These variables cannot be assessed against individual exposures in controls (because only one control reported chemical sensitivity), a further impetus to focus on predictors of actual AE Propensity.

[Table 1](#) further shows the summed AE score, the summed exposure score, and the ratio of the summed AE score to the summed exposure score—the total AE Propensity—in all, in cases and in controls. Affected Gulf War veterans’ summed exposure scores were about 4 times higher, while summed AE scores were close to 10 times higher than controls’ scores. AE Propensity was approximately three times higher in GWI cases than in controls.

A key purpose of this study is to examine whether there is a relationship between nature of exposures experienced and propensity to develop AEs.

To maximize prediction in multivariable models, while limiting the number of included variables, composite variables were generated by summing potentially important exposures within a class. The pesticide variable included pesticides on clothes or bedding and pyrethroids. The composite fuel-fume variable included burning fuel, diesel fumes, diesel on skin, petroleum products, degreasers, and carbon monoxide. (The relation of diesel fumes to carbon monoxide and carbon monoxide to fume exposures may be difficult to disentangle, so these were included in a single variable.) A composite radiation variable included radiation therapy, X-ray radiation, radioactive chemicals, and other radiation. [Table 1](#) shows the mean values of these variables in all, in cases and controls, and the case-control difference. Mean values and ranges assist in interpreting coefficients in regression.

Total AE Propensity showed modest relationships to Kansas domains with the strongest relationship being to skin symptoms in cases and to gastrointestinal (GI) symptoms in controls ([Table 2](#)).

**Table 1. Participant characteristics**

	N	All N = 81 Mean (SD) <sup>a</sup>	Case N = 41 Mean (SD)	Control N = 40 Mean (SD)	p case vs. control
Age (Years)	81	49.8 (7.5)	50.1 (7.6)	49.5 (7.5)	0.68
Male %	81	92.6	92.7	92.5	0.98
Ethnicity %	81				
Caucasian		54.3	53.7	55.0	0.90
African American		21.0	22.0	20.0	0.83
Latino		14.8	14.6	15.0	0.96
Asian		7.41	7.32	7.50	0.98
Native American		2.47	2.44	2.50	0.99
Married %	81	53.1	65.9	40.0	0.020
Kansas symptom subscales	81				
Total fatigue		3.88 (4.33)	7.49 (3.20)	0.175 (0.501)	<0.0001
Total pain		2.73 (3.14)	5.27 (2.48)	0.125 (0.404)	<0.0001
Total neurological		9.47 (11.0)	18.5 (8.31)	0.175 (0.501)	<0.0001
Total skin		0.95 (1.67)	1.85 (1.96)	0.025 (0.158)	<0.0001
Total GI		1.86 (2.44)	3.66 (2.29)	0.025 (0.158)	<0.0001
Total respiratory		1.19 (1.85)	2.32 (2.04)	0.025 (0.158)	<0.0001
Total Kansas symptoms		20.1 (22.5)	39.1 (16.1)	0.550 (0.959)	<0.0001
Chemsensk (0–3)	81	0.593 (0.946)	1.15 (1.06)	0.025 (0.158)	<0.0001
Rated 0, 1, 2, 3		54, 11, 11, 5	15, 10, 11, 5	39, 1, 0, 0	<0.001
Chemsenssx	80	0.575 (1.02)	1.13 (1.20)	0.025 (0.158)	<0.0001
Rated 0, 1, 2, 3		57, 8, 7, 8	18, 7, 7, 8	39, 1, 0, 0	<0.001
Chemsensbinary	81	0.284 (0.454)	0.537 (0.505)	0.025 (0.158)	<0.0001
Rated 0, 1		58, 23	19, 22	39, 1	<0.001
totAE: Summed AE score	81	12.0 (14.9)	21.4 (14.8)	2.31 (6.33)	<0.0001
totExp: Summed exposure score	81	38.4 (28.7)	60.9 (21.5)	15.4 (12.0)	<0.0001
AE Propensity <sup>b</sup>	81	0.216 (0.209)	0.330 (0.172)	0.100 (0.176)	<0.0001
Composite pesticide	81	0.51 (0.72), 0-2	0.90 (0.75)	0.10 (0.38)	<0.0001
Composite fuel	81	2.00 (2.02), 0-6	3.20 (1.82)	0.78 (1.39)	<0.0001
Total radiation	81	1.78 (2.03), 0-8	2.60 (2.27)	0.95 (1.32)	0.0002

GI = gastrointestinal. chemsensbinary = a binarized version of chemsenssx (The missing value for chemsenssx was assigned a value of 0). chemsensk = having physical or mental symptoms after breathing in certain smells or chemicals. chemsenssx = chemical sensitivity (e.g., unusual sensitivity to smells). AE = adverse effect. totAE = summed adverse effect score. totExp = summed exposure score. Composite pesticide: sum of pesticides on clothes or bedding, and pyrethroids. Composite fuel: sum of burning fuel, diesel fumes, diesel on skin, petroleum products, degreasers, and carbon monoxide. Total radiation: sum of radiation therapy, X-ray radiation, radioactive chemicals, other radiation, SD = standard deviation. N = number. p = probability (two-sided).

<sup>a</sup>Except where “%” is signified in characteristics column.

<sup>b</sup>Summed AE score/summed exposure score.

Table 3 shows the relation of AE Propensity to self-rated chemical sensitivity, by the various measures, in GWI cases. (This is not meaningful in controls, with only one participant designating themselves as chemically sensitive.) The strongest relation of AE Propensity is to the UCSD binary chemical sensitivity measure. This analysis uses correlation, which though not the optimal assessment when a variable is binary, provides a more intuitive index of the relationship, for comparison across measures. This affirms that AE Propensity is related to self-rated chemical sensitivity but is not isomorphic to it.

On a similar theme, Table 4 shows the mean AE Propensity, by chemical sensitivity rating, for the two binary ratings and for the Kansas rating. Those who consider themselves to be chemically sensitive, on average

**Table 2. Relation of AE propensity to Kansas GWI symptom domains**

Kansas domain	All		Cases		Controls	
	r	p	r	p	r	p
Fatigue	0.52	<0.0001	0.14	0.38	0.10	0.52
Pain	0.57	<0.0001	0.35	0.024	-0.13	0.43
Neurological	0.58	<0.0001	0.35	0.024	-0.15	0.35
GI	0.51	<0.0001	0.22	0.16	0.32	0.046
Respiratory	0.41	0.0001	0.15	0.35	-0.050	0.76
Skin	0.50	<0.0001	0.40	0.0093	-0.054	0.74
Total symptoms	0.59	<0.0001	0.36	0.0195	-0.044	0.79

r = correlation coefficient. p = probability (two-sided).

reported AEs to a higher fraction of exposures. A negative relationship of AE Propensity to neurologic symptoms especially may be because controls could only be included if they did not have multiple symptoms in a category. Chemical sensitivity which related to AE Propensity relatively excludes controls with other neurological symptoms and with any symptoms with which those were correlated.

Table S1a shows the relationships between individual exposures and the AE Propensity score, for all participants, and for GWI cases and controls separately. For exposures that are materially more common in GWI cases, they may relate to AE Propensity in the total sample simply because of this. The relations in GWI cases and controls separately are not subjected to this limitation. The far-right column shows the relationship of an exposure to AE Propensity in all participants adjusted for GWI case status, via regression with robust standard errors. Particularly strong relationships occur in categories of fuels-solvents and pesticides. Controls show an apparent relationship of AE Propensity to some metals. Several individual exposures show relationships. One variable related to radiation is significant in the case-adjusted regression, with added interest based on shared direction and p values <0.2 for two of the other three variables in that category.

Negatively signed exposure correlations to AE Propensity, significant or otherwise, were not common. Among them were several exposures in the vaccine and immune globulin category; and also both copper and selenium (each of which is central to a key antioxidant system).

Table S1b shows Gulf-specific exposure queries, asked only of cases. Of note, combat and combat-stress related exposures (such as seeing Americans killed or seeing dismembered bodies) were not included in totExp, and AEs to them were not included in totAE, as the purpose was to examine predictors of drug-environment AE Propensity. These variables were, however, examined to see if they were predictors of AE Propensity. The most highly significant variable is pesticide related, with significance also present for two additional pesticide variables. Gas mask use was significant, which could cohere with pesticide use findings since gas mask use occurred in some settings of chemical alarm sounding, which could potentially reflect either actual organophosphate nerve gas (to which exposure occurred in the Gulf Conflict<sup>57,58</sup>), or

**Table 3. Correlations among AE propensity and chemical sensitivity scores: GWI cases**

	AE Propensity r (p)
AE Propensity	1.0
Kansas: chemsensk	0.31 (0.047)
UCSD: chemsenssx	0.41 (0.0081)
UCSD Binary	0.44 (0.0037)
Kansas Binary	0.26 (0.10)

chemsensk, having physical or mental symptoms after breathing in certain smells or chemicals; chemsenssx, chemical sensitivity (e.g., unusual sensitivity to smells); UCSD, University of California, San Diego; r, correlation coefficient; p, probability (two-sided).

**Table 4. Mean AE propensity by chemical sensitivity rating**

Value of measure (ordered category)	0	1	2	3	
<b>Chemical sensitivity</b>					
<b>Self-rating measure</b>	Mean (SD)	Mean (SD)	Mean (SD)	Mean (SD)	p <sup>a</sup>
UCSD Binary Chemical Sensitivity rating	0.15 (0.19)	0.39 (0.16)	N/A	N/A	<0.0001
Kansas Binary Chemical Sensitivity rating	0.15 (0.19)	0.35 (0.18)	N/A	N/A	<0.0001
Kansas Full Chemical Sensitivity rating	0.15 (0.19)	0.29 (0.21)	0.38 (0.14)	0.41 (0.17)	<0.01 <sup>b</sup>

Chem, chemical; SD, standard deviation; r, correlation coefficient; p, probability (two-sided); UCSD, University of California, San Diego.

<sup>a</sup>For difference across categories.

<sup>b</sup>Some statistical commands in Stata afford different numbers of significant digits. This p value is listed as "0.00."

possibly organophosphate or other pesticide or chemical classes. (A recent study found a gene environment interaction, in which paraoxinase 1 (PON1) variants adverse to sarin detoxification coupled with reported hearing of chemical alarms were associated with a 9-fold increase in GWI.<sup>59</sup>) A possible consideration is that if gas masks were reused, as evidently they were, near-nose inhalation of chemical residue on the mask is a possibility. Inhalation of smoke (potentially from oil fires) was a significant predictor. Eating local food and drinking bad water were significant on univariable analyses.

A number of combat related exposures show an unadjusted relationship to AE Propensity. Factors such as forward deployment that correlated with combat were also correlated with higher environmental exposures, producing (unadjusted) appearance of a link of combat to exposure-associated outcomes. For variables with correlation p values <0.15, the right hand column reports if the variable was significant in multivariable adjustment, and whether it was included in the main model for cases (Table 6); or reports the loss of significance with the variable added to the main model. That is, combat stressors lost significance with adjustment for primary model environmental exposures. For each candidate variable not included in a multivariable model (shown later), neither significance nor borderline significance was retained when the variable was incorporated into the multivariable assessment. (In contrast, in each such instance, the variables of the main model retained significance with addition of the candidate variable.)

To maximize prediction in multivariable models, while limiting the number of included variables, composite variables were generated by summing potentially important exposures within a class. The pesticide variable included pesticides on clothes or bedding and pyrethroids. The composite fuel-fume variable included burning fuel, diesel fumes, diesel on skin, petroleum products, degreasers, and carbon monoxide. A composite radiation variable included radiation therapy, X-ray radiation, radioactive chemicals, and other radiation. Table 1 shows the mean values of these variables in all, in cases and controls, and the case-control difference. Mean values and ranges assist in interpreting coefficients in regression.

Table 5 shows the unadjusted relationships of these variables to the AE Propensity variable, in all, in GWI cases and in controls. Although the presence of these exposures is far greater in GWI cases, shown in

**Table 5. Composite variables: Univariable relation to total AE per exposure**

	All		GWI case		Control		All adjusted for GWI case	
	r	p	r	p	r	p	β (SE)	p
Composite pesticide	0.69	<0.0001	0.52	0.0005	0.69	<0.0001	0.16 (0.032)	<0.001
Composite fuel-fume	0.64	<0.0001	0.44	0.0038	0.51	0.0008	0.050 (0.013)	<0.001
Total radiation	0.54	<0.0001	0.47	0.0019	0.34	0.0298	0.038 (0.011)	0.001

Composite pesticide: sum of pesticides on clothes or bedding, and pyrethroids. Composite fuel: sum of burning fuel, diesel fumes, diesel on skin, petroleum products, degreasers, and carbon monoxide. Total radiation: sum of radiation therapy, X-ray radiation, radioactive chemicals, other radiation. The regressions use robust (heteroskedasticity-independent) standard errors.<sup>60</sup>

AE, adverse effect; r, correlation coefficient; β, regression coefficient; SE, standard error; p, probability (two-sided).

**Table 6. AE propensity: Multivariable model assessed across groups**

Exposure	All adjusted for case		Case		Control	
	b (SE)	p	b (SE)	p	$\beta$ (SE)	p
Composite pesticide	0.11 (0.026)	<0.001	0.083 (0.025)	0.002	0.32 (0.085)	0.001
Total radiation	0.028 (0.0073)	<0.001	0.028 (0.0092)	0.005	0.031 (0.012)	0.012
Copper	-0.14 (0.030)	<0.001	-0.15 (0.033)	<0.001	-0.16 (0.070)	0.026
Composite fuel	0.027 (0.012)	0.030	0.021 (0.010)	0.045	0.0082 (0.025)	0.74
Case	0.070 (0.045)	0.12	N/A		N/A	
	R <sup>2</sup> = 0.67, p < 0.0001		R <sup>2</sup> = 0.60, p < 0.0001		R <sup>2</sup> = 0.58, p = 0.0002	

Composite pesticide: sum of pesticides on clothes or bedding, and pyrethroids. Composite fuel-fume: sum of burning fuel, diesel fumes, diesel on skin, petroleum products, degreasers, and carbon monoxide. Total radiation: sum of radiation therapy, X-ray radiation, radioactive chemicals, other radiation.

All regressions use robust (heteroskedasticity-independent) standard errors.<sup>60</sup>  $\beta$  = regression coefficient. SE = standard error. p = probability (two-sided). R<sup>2</sup> = “coefficient of determination,” a measure of the proportion of the variance that is explained by the variables in the regression model.

Table 2, the relationship to the AE vulnerability variable is replicated in controls for each of the exposure classes.

Table 6 shows multivariable models, in which the exposure relationship is adjusted for other potentially relevant exposures. Table 6 shows a single multivariable model, implemented in all participants (adjusted for GWI case status), and separately in VGWI and in controls. This shows that the composite pesticide variable, the composite radiation variable, and copper exposure (inverse predictor i.e., “protective”) are significant predictors in each. The composite fume-fuel exposure is significant in the total sample, and in cases, but not in controls. Its significance level is less than for the other predictors. In each model, a fairly strong R<sup>2</sup> is produced (0.58 or greater), with high-model significance. Of note, the R<sup>2</sup> remains 0.58 in the control model, omitting the non-significant composite fume-fuel variable.

Table S2 shows alternative models. Immune globulin was a significant protective predictor.

Table 7 shows candidate models specific to cases that incorporate Gulf-specific exposures. Inhaling smoke in the Gulf is substituted for the composite fuel-fume variable (Oil fire smoke was a significant problem in the Gulf.). Aerosolized depleted uranium (DU) exposure occurred when munitions struck vehicles (e.g., “friendly fire” episodes), and either contact with these or proximity to them (data not shown for the latter) served as independent predictors, beyond other radioactive chemical exposure. As aforementioned, gas mask use may serve as a relative proxy for organophosphate chemical exposure—either organophosphate nerve agents or pesticides that may have triggered chemical alarms and caused symptoms or other features adding concern. (As aforementioned, based on comments by veterans, there may also be a question of whether gas masks, which were reused, could have thereby served as a source of recurrent chemical inhalation in some settings.)

An unexpected finding was the apparent positive relation of sunscreen use in the Gulf to AE Propensity. The relationship was strong. (There was no relationship of sunscreen use *outside* of the Gulf to AE Propensity, either in controls or in Gulf War veterans; see discussion.) Of note, no combat-related or stress-related variables were independent predictors (except seeing or directly contacting destroyed enemy vehicles—a potential proxy for inhalational and/or dermal exposure to DU), often switching to a negative coefficient (non-significant); whereas model variables retained strong significance.

Table 8 shows control-specific models. These retain the composite pesticide exposure, total radiation, and copper exposures present in the models for all, and for cases. Immune globulin was a negative predictor and measles mumps rubella vaccine (MMR) a (weaker) positive predictor, as in analyses in the total sample (Table S2). (“Total vaccines” was not a predictor.) A modest positive relationship to metal exposure is suggested; adding iron (which is ferromagnetic, prooxidant, and relates to “ferroptosis”) increases significance of all but one of the other exposures in the model, suggesting it is addressing some of the variance that curbs significance for other variables. However, the variable itself does not reach significance.



**Table 7. AE propensity: Multivariable regression models, VGWI**

"Main" model: Model 1			Alternative model: Model 2		
Exposure	$\beta$ (SE)	p	Exposure	$\beta$ (SE)	p
Composite pesticide	0.087 (0.021)	<0.001	Composite pesticide	0.11 (0.022)	<0.001
Total radiation	0.030 (0.0071)	<0.001	Total radiation	0.035 (0.0068)	<0.001
Copper	-0.13 (0.033)	<0.001	Copper	-0.12 (0.031)	0.001
Inhaled smoke from oil-well fires (in Gulf)	0.17 (0.038)	<0.001	Inhaled smoke from oil-well fires (in Gulf)	0.13 (0.035)	0.001
Gas mask	0.16 (0.051)	0.004	Sunscreen	0.099 (0.034)	0.006
$R^2 = 0.71, p < 0.0001$			$R^2 = 0.72, p < 0.0001$		

Models 3–5 each add one variable to "Main Model"

Model 3		
Exposure	$\beta$ (SE)	p
Composite pesticide	0.085 (0.020)	<0.001
Total radiation	0.029 (0.0069)	<0.001
Copper	-0.12 (0.031)	<0.001
Inhaled smoke from oil-well fires	0.14 (0.035)	<0.001
Gas mask	0.15 (0.047)	0.003
Direct contact destroyed enemy vehicles (proxy for DU)*	0.070 (0.027)	0.014
$R^2 = 0.74, p < 0.0001$		

DU = depleted uranium  
 \*Direct contact with destroyed enemy vehicles – a proxy for potential depleted uranium inhalation exposure (since DU-girded munitions were used to penetrate/destroy enemy vehicles).  
 Note that substituting for this say, combat injury, air combat, ground combat, danger, saw death, saw dismemberment, these other variables have no relationship, and the strong relationships of the other variables are preserved.

Model 4			Model 5		
Sprayed pesticides in theater add separate significance to the composite pesticide exposure			Includes both gas mask (main model) and sunscreen (alternative model)		
Exposure	$\beta$ (SE)	p	Exposure	$\beta$ (SE)	p
Composite pesticide	0.068 (0.021)	0.003	Composite pesticide	0.096 (0.022)	<0.001
Total radiation	0.029 (0.0064)	<0.001	Total radiation	0.032 (0.0068)	<0.001
Copper	-0.12 (0.031)	<0.001	Copper	-0.13 (0.031)	<0.001
Inhaled smoke	0.15 (0.035)	<0.001	Inhaled smoke	0.14 (0.034)	<0.001
Gas mask	0.14 (0.054)	0.013	Gas mask	0.12 (0.058)	0.042
Saw pesticides sprayed in theater	0.075 (0.034)	0.033	Sunscreen	0.077 (0.033)	0.024
$R^2 = 0.75, p < 0.0001$			$R^2 = 0.76, p < 0.001$		

Model 6			Model 7		
Expanded model			Limited model		
Exposure	$\beta$ (SE)	p	Exposure	$\beta$ (SE)	p
Composite pesticide	0.094 (0.020)	<0.001	Composite pesticide	0.10 (0.025)	<0.001
Total radiation	0.031 (0.0070)	<0.001	Total radiation	0.033 (0.007)	<0.001
Copper	-0.11 (0.031)	0.001	Copper	-0.13 (0.034)	0.001
Inhaled smoke	0.12 (0.028)	<0.001	Inhaled smoke	0.15 (0.041)	0.001
Direct contact with destroyed enemy vehicles*	0.066 (0.024)	0.008			
Gas mask	0.12 (0.047)	0.017			

(Continued on next page)

**Table 7. Continued**

Model 6			Model 7		
Expanded model			Limited model		
Exposure	$\beta$ (SE)	p	Exposure	$\beta$ (SE)	p
Sunscreen	0.075 (0.031)	0.021			
$R^2 = 0.78, p < 0.0001$			$R^2 = 0.64, p < 0.0001$		

Composite pesticide: sum of pesticides on clothes or bedding, and pyrethroids; DU: depleted uranium; total radiation: sum of radiation therapy, X-ray radiation, radioactive chemicals, other radiation. \*Direct contact with destroyed enemy vehicles – a proxy for potential DU inhalation exposure (since DU-girded munitions were used to penetrate/destroy enemy vehicles). Note that substituting in, say, combat-related injury, air combat, ground combat, danger/direct combat, saw Americans or Allied troops badly wounded or killed, saw dismembered bodies, these variables have no relationship, and the strong relationships of the other variables are preserved.

$\beta$ , regression coefficient; SE, standard error; p, probability (two-sided);  $R^2$ , “coefficient of determination,” a measure of the proportion of the variance that is explained by the variables in the regression model.

Table 9 shows results stratified at the mean participant age of 50.

In Table 9, for the total sample stratified by age using the common model, pesticides, radiation, and copper are supported in independent samples. Based on coefficients, the pesticide relationship appeared stronger in younger age, the radiation relationship stronger in older age. For other variables, coefficients are a bit stronger in younger age but variance greater and significance lower. Power is lower in age groups assessed separately. Despite this, a role for pesticides, radiation, and copper, at least, is supported in each age group separately.

Table 10 shows age-stratified results in GWI cases, using the main and alternative case-specific models. Again, based on the coefficient, the pesticide relationship appeared stronger in younger age, the radiation relationship in older age. Gas mask (main model) but not sunscreen (alternative model) relationships are preserved across stratification. Both age groups contribute to significance of the copper variable, though with a smaller sample size, significance is borderline in both groups.

Table 11 shows age-stratified results in controls. Age stratification supports the primacy of pesticides as a contributor. Copper and radiation are also supported, via significance or borderline significance in each analysis.

Given the relationship of the AE Propensity variable to skin domain symptoms in VGWI and the relation of sunscreen exposure in the Gulf to the AE Propensity variable, we examined (on an exploratory basis) the relation of AE Propensity to the Kansas skin symptom total, in those reporting and not reporting sunscreen use in theater (Table 12); and then those reporting and not reporting sunscreen use in general. Those who did not use sunscreen in theater have markedly lower AE Propensity and skin symptoms than those who did; and the relationship between skin symptoms and AE Propensity is lost in this group. There is no similar relationship for sunscreen use outside of the theater. Those without sunscreen use have lower AE Propensity than overall for cases, i.e., those with sunscreen use have higher AE Propensity. However, the relation of skin symptoms to AE Propensity is particularly low (essentially nonexistent) in those with no sunscreen exposure.

As a final step, to further assess whether predictors of AE Propensity (which can be assessed also in controls) were also predictors of self-rated chemical sensitivity (which could be examined here only in cases since only one control self-designated as having any chemical sensitivity), we assessed the key predictors in cases against the binary UCSD chemical sensitivity measure, using logistic regression with robust standard errors (Table 13).

The three primary case predictors—the pesticide variable, radiation, and inhaled smoke—were each significant predictors of the binary chemical sensitivity variable, on logistic regression. Copper and sunscreen use were not predictors. (The *sign* for the sunscreen variable was not reproduced—not shown.) Adding either copper or sunscreen, the three “primary” variables retained significance. In general, a role in cases for pesticides, and especially radiation, and smoke inhalation as predictors of chemical sensitivity is supported.

**Table 8. AE propensity prediction: Control-specific model**

Model 1: 4-variables			Model 2: 5-variables		
Exposure	$\beta$ (SE)	P	Exposure	$\beta$ (SE)	p
Composite pesticide	0.37 (0.073)	<0.001	Composite pesticide	0.38 (0.065)	<0.001
Total radiation	0.032 (0.019)	0.019	Total Radiation	0.031 (0.012)	0.012
Copper	-0.13 (0.050)	0.012	Copper	-0.12 (0.050)	0.023
Immune globulin	-0.16 (0.056)	0.006	MMR	0.076 (0.031)	0.021
			Immune globulin	-0.15 (0.050)	0.004
R <sup>2</sup> = 0.62, p = 0.0008			R <sup>2</sup> = 0.64, p = 0.0001		
If exclude the five participants who cited immune globulin as “unsure” (value 0.5), all are still significant but coefficients are stronger for all variables (0.41, 0.040, -0.18, -0.25) and significance is stronger for the other variables: <0.001, 0.008, 0.001; but is 0.031 for immune globulin.			If exclude the five participants who cited immune globulin exposure as “unsure”, all are still significant but coefficients are strengthened for all variables, especially total radiation ( $\beta = 9.938$ , $p = 0.006$ ) and copper ( $\beta = -0.16$ , $p = 0.001$ ); and p values are strengthened for all except MMR which changes little ( $\beta = 0.082$ , $p = 0.023$ ).		
“Total vaccines” was not a predictor.					
Model 3: 6-variables			Model 4: 6-variables		
Exposure	$\beta$ (SE)	p	Exposure	$\beta$ (SE)	p
Composite pesticide	0.28 (0.063)	<0.001	Composite pesticide	0.25 (0.073)	0.002
Total Radiation	0.030 (0.011)	0.009	Total Radiation	0.027 (0.010)	0.012
Copper	-0.14 (0.055)	0.013	Copper	-0.15 (0.055)	0.011
MMR	0.082 (0.030)	0.011	MMR	0.086 (0.030)	0.008
Immune globulin	-0.12 (0.042)	0.009	Immune globulin	-0.12 (0.039)	0.004
Iron	0.16 (0.089)	0.089	Iron + cobalt	0.12 (0.067)	0.084
R <sup>2</sup> = 0.68, p = 0.0001			R <sup>2</sup> = 0.69, p = 0.0001		
If exclude the five participants who cited immune globulin exposure as “unsure”, all are still significant but coefficients are strengthened for all variables, especially total radiation ( $\beta = 9.938$ , $p = 0.006$ ) and copper ( $\beta = -0.16$ , $p = 0.001$ ); and p values are strengthened for all except MMR which changes little ( $\beta = 0.082$ , $p = 0.023$ ).					
Composite pesticide: sum of pesticides on clothes or bedding, and pyrethroids. MMR, measles, mumps, rubella. Total radiation: sum of radiation therapy, X-ray radiation, radioactive chemicals, other radiation. $\beta$ = regression coefficient. SE = standard error. p = probability (two-sided). R <sup>2</sup> = “coefficient of determination,” a measure of the proportion of the variance that is explained by the variables in the regression model.					

## DISCUSSION

### Key results

Our data affirm that AE Propensity is markedly elevated in VGWI relative to healthy controls, identifying AE Propensity as a *new feature* of GWI. Specific exposure classes that are more prevalent in VGWI predicted AE Propensity, and did so in both VGWI and controls, both validating findings in VGWI and adding relevance beyond VGWI. Predictors of AE Propensity include not only pesticides and radiation, but also combustion products. Pesticides and radiation were significant predictors in the full sample; separately in cases and controls; and separately in younger and older age groups, providing robust internal replication. Unexpectedly, reported copper exposure appeared significantly protective, in all, separately in cases and controls, and separately in younger and older age groups, in age-stratified analysis.

Additional variables were possible predictors in one or two of the “three” groups, considering cases, controls, and the total sample (adjusted for case status). Immune globulin appeared protective in the total group and in controls. Gulf-theater sunscreen appeared to be promoting in affected veterans, while sunscreen outside of this setting was not (and the sign was negative, not shown).

**Table 9. AE propensity prediction. Combined VGWI and controls: Stratified by age (at mean age of 50)**

Exposure	Age ≤50 N = 47		Age >50 N = 34	
	β (SE)	p	β (SE)	p
Composite pesticide	0.15 (0.043)	0.001	0.082 (0.028)	0.006
Total Radiation	0.021 (0.0093)	0.036	0.045 (0.011)	<0.001
Copper	−0.14 (0.041)	0.002	−0.11 (0.030)	0.001
Composite fuel	0.027 (0.019)	0.17	0.022 (0.010)	0.044
Case	0.071 (0.084)	0.40	0.066 (0.042)	0.13
	R <sup>2</sup> = 0.65, p < 0.0001		R <sup>2</sup> = 0.78, p < 0.0001	

Composite pesticide: sum of pesticides on clothes or bedding, and pyrethroids. Total radiation: sum of radiation therapy, X-ray radiation, radioactive chemicals, other radiation. Pesticide relationship is stronger in younger age; radiation relation appears stronger in older age.

Adding immune globulin, it was significant in younger age (p = 0.03), but not older (p = 0.56).

Adding MMR with immune globulin, it was not significant in younger, p = 0.52; but approached significance in older age, p = 0.05 (and was significant, p = 0.03, if stratification is at age 48, the median age).

N = number. β = regression coefficient. SE = standard error. p = probability (two-sided). R<sup>2</sup> = “coefficient of determination,” a measure of the proportion of the variance that is explained by the variables in the regression model.

### Interpretation/fit with existing literature

One lens with which to view findings is through twin intertwined considerations of oxidative stress and mitochondrial impairment, which are common mechanisms of toxicity for many exposure classes—irrespective of nominal mechanisms of action, for drugs/chemicals,<sup>27–35</sup> and extending also to radiation.<sup>36–43</sup> These in turn contribute to relevant downstream mechanisms of autoantibody induction,<sup>61,62</sup> apoptosis,<sup>63,64</sup> and inflammation<sup>65</sup>—mechanisms relevant to GWI.<sup>4,5,8,9,17</sup>

Pesticide exposure predicted AE Propensity in each group. Pesticides were previously shown to relate to development of multiple chemical sensitivity (MCS) in Gulf War veterans, OR = 12.3(95%CI:5.1–30.0).<sup>24</sup> In GWI cases, gas mask use and pesticides relatively attenuated the significance of each other: gas masks might serve as a proxy for organophosphate nerve gas exposure or other chemicals that cause these alarms to sound (potentially including organophosphate pesticides).

Radiation exposure predicted AE Propensity. This coheres with reports to us by nonveterans and veterans, who cite onset of chemical and electrical sensitivity following a significant EMR exposure. (Many who develop electrosensitivity do so on a backdrop of existing chemical sensitivity. Some cite *de novo* sensitivity to both chemicals and EMR, concurrently.) While this often refers to nonionizing radiation, toxicity mechanisms involving oxidative stress are shared for nonionizing and ionizing radiation (and conversely antioxidant protections are shared).<sup>66,67</sup> Further aligned with a role for radiation, contact with destroyed enemy vehicles—which were destroyed via DU-girded munitions, that produce aerosol/inhalational exposure and potentially dermal exposure to radioactive chemicals when they hit a target—was a possible independent predictor. Other combat and combat stress-related exposures bore no independent relationship.

Fuels-fumes were not a material independent predictor in controls, though these were a predictor in the total sample, and inhaling smoke (e.g., from burning oil fires) was a predictor in cases.

Pesticides,<sup>68–77</sup> across major classes, as well as radiation across the spectrum<sup>78,79</sup> can depress activity of critical endogenous antioxidant systems, like glutathione, superoxide dismutase and catalase, perhaps by consuming antioxidants. (Results vary according to specifics of the study: such systems can in certain conditions, be adaptively upregulated.<sup>80</sup>) This could be one factor contributing to their role. Moreover, radiation can also depress levels of melatonin,<sup>81–94</sup> which though better known for its relation to sleep, has critical antioxidant functions, and has been shown to defend against toxicity not only from radiation itself<sup>66,95–115</sup> (across the electromagnetic spectrum—nonionizing and ionizing<sup>66,67</sup>) but from toxicity of many drugs and chemicals,<sup>116–151</sup> including toxicity by pyrethroids<sup>152</sup> and organophosphates<sup>153–155</sup>—which was present in nerve gas to which some personnel were exposed, and in key pesticides used in the Persian Gulf. In a French study including >700 persons with nonionizing-radiation sensitivity

**Table 10. Total AE propensity, GWI Case regression models: Stratified by age (at mean age of 50)**

Exposure	Main Model				
	Age ≤ 50 N = 23		Age >50 N = 18		
	β (SE)	p	β (SE)	p	
Composite pesticide	0.13(0.033)	0.001	0.044 (0.031)	0.18	
Total radiation	0.022 (0.0089)	0.022	0.048 (0.011)	0.001	
Inhaled smoke from oil-well fires	0.21 (0.069)	0.007	0.088 (0.050)	0.103	
Copper	−0.13 (0.063)	0.055	−0.080 (0.042)	0.082	
Gas mask	0.19 (0.052)	0.002	0.13 (0.061)	0.036	
	R <sup>2</sup> = 0.75, p not calculated		R <sup>2</sup> = 0.82, p < 0.0001		
Exposure	Alternative Model				
	Age ≤ 50 N = 23		Age >50 N = 18		
	β (SE)	p	β (SE)	p	
Composite pesticide	0.14 (0.036)	0.001	0.083 (0.020)	0.001	
Total radiation	0.024 (0.008)	0.007	0.057 (0.012)	0.001	
Inhaled smoke from oil-well fires	0.27 (0.066)	0.023	0.098 (0.079)	0.24	
Copper	−0.11 (0.067)	0.12	−0.086 (0.030)	0.013	
Sunscreen	0.070 (0.056)	0.23	0.11 (0.039)	0.013	
	R <sup>2</sup> = 0.74, p < 0.0001		R <sup>2</sup> = 0.84, p = 0.0004		

Composite pesticide: sum of pesticides on clothes or bedding, and pyrethroids. Total radiation: sum of radiation therapy, X-ray radiation, radioactive chemicals, other radiation.  
 In cases too, the pesticide relationship (and gas mask relationship) appears stronger in younger age; radiation relationship (best on the coefficient) appears stronger in older age.  
 Inhaled smoke is far stronger in younger age. If the split is at the median rather than the mean, gas mask use is significant in younger (p = 0.003) and older (p = 0.04) age; copper is significant in older age, p = 0.019 and – with a stronger coefficient – almost significant in younger age (p = 0.06). Pesticides still lose significance in older age (younger personnel may have been involved in actual pesticide application in theater), and with the smaller sample and younger age, significance for radiation is lost (p = 0.02).  
 N = number. β = regression coefficient. SE = standard error. p = probability (two-sided). R<sup>2</sup> = “coefficient of determination,” a measure of the proportion of the variance that is explained by the variables in the regression model.

(electrosensitivity), many with chemical sensitivity, a depressed ratio of 24-h excretion of a urine melatonin metabolite (6-hydroxymelatonin sulfate) relative to creatinine was a biomarker.<sup>156</sup> Radiation and radioactive chemicals<sup>157</sup> can also depress activity of other endogenous antioxidant systems, such as the glutathione system,<sup>158–165</sup> which also defends against radiation and drug-chemical-environmental oxidative stress (OS) injury<sup>160,166–178</sup> (including from pesticides-solvents).<sup>179</sup> (In turn, low glutathione peroxidase is linked to drug and chemical intolerance.<sup>175</sup>)

Depressed antioxidant defenses may explain why pesticide and radiation exposures that can promote development of MCS also can set people up for developing electrosensitivity following chemical or especially significant radiation exposures. Indeed, chemical-induced depression in antioxidant systems has been shown to be involved in increased vulnerability to toxicity of radiation.<sup>180</sup> A role for antioxidant defenses in chemical sensitivities is supported by genetic evidence relating an adverse polymorphism of superoxide dismutase 2 (SOD2) to chemical sensitivity,<sup>49,181</sup> and evidence of depressed glutathione levels, depressed catalase and glutathione S transferase activities in patients with MCS,<sup>156,182</sup> and the aforementioned findings with respect to melatonin metabolite excretion.

Both pesticides<sup>183–203</sup> and radiation are reported to adversely affect mitochondria,<sup>40,41,46,202–211</sup> as do radioactive chemicals, including DU.<sup>212–214</sup> Impaired mitochondria amplify free radical production<sup>51,52</sup> and risks with many exposures. Again, AEs often involve mitochondrial compromise and OS.<sup>27–35</sup>

**Table 11. Total AE Propensity, control regression models: Stratified by age (at mean age of 50)**

Exposure	Controls			
	Age ≤50 N = 24		Age >50 N = 16	
	β (SE)	p	β (SE)	p
Composite pesticide	0.55 (0.089)	<0.001	0.26 (0.025)	<0.001
Total Radiation	0.029 (0.014)	0.053	0.050 (0.020)	0.032
Copper	-0.083 (0.038)	0.040	-0.13 (0.033)	0.002
Immune globulin	-0.050 (0.085)	0.56	-0.13 (0.050)	0.028
	R <sup>2</sup> = 0.71, p < 0.0001		R <sup>2</sup> = 0.69, p not defined	

Composite pesticide: sum of pesticides on clothes or bedding, and pyrethroids. Total radiation: sum of radiation therapy, X-ray radiation, radioactive chemicals, other radiation.

Pesticide relationship is stronger in younger age, other relationships are stronger in older age.

β = regression coefficient. SE = standard error. p = probability (two-sided). R<sup>2</sup> = "coefficient of determination," a measure of the proportion of the variance that is explained by the variables in the regression model.

(Perhaps not surprisingly given potential for mitochondrial injury and impaired antioxidant defenses, both pesticides<sup>76,215–222</sup> and radiation produce OS at least acutely.<sup>39,41,66,96,98–101,105,107,109,210,223–256</sup>)

Pesticides and radiation have also each been shown to alter membrane properties,<sup>184,201,257–265</sup> the latter finding replicated across many frequencies and lifeforms.<sup>205,211,230,266–274</sup> This extends to radioactive chemicals (including uranium).<sup>275</sup>

Inhaled combustion products were predictors in VGWI, who had more prospects for significant smoke inhalation exposure, via exposure to oil fire smoke. Smoke/combustion products and inhaled particulates are also linked to OS<sup>276–280</sup> and mitochondrial injury.<sup>277,281</sup> Delta psi, the mitochondrial membrane potential, which relates to apoptosis risk, is affected by combustion products/smoke,<sup>281</sup> radiation,<sup>281</sup> and pesticides.<sup>188,201,282–285</sup>

**Table 12. Relation of AE propensity to skin symptoms (Kansas criteria), as a function of sunscreen use in and out of theater, in GWI cases**

Sunscreen in Gulf Theater	No Sunscreen in Theater N = 16		Yes Sunscreen in Theater N = 23	
	AE Propensity: Mean (SD)	0.28 (0.16)		0.37 (0.18)
Skin symptoms: Mean (SD)	1.2 (1.6)		2.4 (2.1)	
Correlation of skin symptoms to AE Propensity	r	p	r	p
	0.015	0.96	0.48	0.020

Sunscreen, Not Gulf	Sunscreen use, Not Gulf-specific			
	No Sunscreen, General N = 20		Sunscreen, General N = 19	
AE Propensity: Mean (SD)	0.35 (0.20)		0.33 (0.15)	
Skin symptoms: Mean (SD)	1.9 (2.4)		1.7 (1.4)	
Correlation of skin symptoms to AE Propensity	r	p	r	p
	0.49	0.029	0.46	0.046

Those citing no Gulf-theater sunscreen exposure have lower AE Propensity, half the Kansas skin symptoms score; and no correlation of AE Propensity to skin symptoms. In contrast, in those with Gulf sunscreen exposure, there is a strong correlation between these, nearly 0.5 (with p = 0.020). Of note, there is no such disparity with sunscreen use outside of the Gulf (in which with and without are associated with a significant correlation of skin symptoms to AE Propensity).

N, number; r, correlation coefficient; p, probability (two-sided); SD, standard deviation.

**Table 13. Multivariable assessment of chemical sensitivity variables in cases using the model optimized for Total AE Propensity**

UCSD binary chemical sensitivity			Kansas, full chemical sensitivity		
Exposure	$\beta$ (SE)	p	Exposure	$\beta$ (SE)	p
Composite pesticide	0.94 (0.45)	0.037	Composite pesticide	-0.15 (0.42)	0.72
Total radiation	0.38 (0.19)	0.048	Total radiation	0.47 (0.20)	0.021
Inhale smoke	3.3 (1.4)	0.016	Inhale smoke	3.0 (0.85)	<0.001
$R^2 = 0.25, p = 0.013$			Pseudo $R^2 = 0.14, p = 0.0056$		
UCSD binary			Kansas, full		
Exposure	$\beta$ (SE)	p	Exposure	$\beta$ (SE)	p
Composite pesticide	0.96 (0.47)	0.039	Composite pesticide	-0.14 (0.42)	0.74
Total radiation	0.39 (0.20)	0.044	Total radiation	0.48 (0.22)	0.024
Inhale smoke	3.2 (1.3)	0.016	Inhale smoke	2.9 (0.82)	0.001
Copper	0.59 (0.81)	0.47	Copper	-0.48 (0.66)	0.47
$R^2 = 0.25, p = 0.013$			Pseudo $R^2 = 0.15, p = 0.012$		
Adding gas mask use, it is not a predictor: $p = 0.76$ .			Substituting the saw-pesticides-sprayed variable, the pesticide becomes 0.88 (0.60) 0.14; the variables retain similar coefficients, with $p = 0.26$ and 0.002.		

Composite pesticide: sum of pesticides on clothes or bedding, and pyrethroids. Total radiation: sum of radiation therapy, X-ray radiation, radioactive chemicals, other radiation.

Regression approach: Logistic regression (robust standard errors) for the UCSD Binary Chemical Sensitivity variable; ordinal logit regression (robust standard errors) for the Kansas, Full Chemical Sensitivity measure.

$\beta$ , regression coefficient; SE, standard error; p, probability (two-sided);  $R^2$ , "coefficient of determination," a measure of the proportion of the variance that is explained by the variables in the regression model.

Thus, implicated agents may markedly enhance vulnerability to toxicity from many exposures, encompassing many drugs, chemicals and radiation exposures that can further impair mitochondrial function, and/or cause OS, which can promote apoptosis, or alter membrane properties (involved in barrier function, mitochondrial function, and apoptosis regulation).

OS, induced by the above exposures, can enhance triggering of immune and autoimmune reactions<sup>286–298</sup> (hence effectiveness of adjuvants in vaccines), and some triggered AEs may have immune mediation; Gulf War veterans have been shown to have increased autoantibodies,<sup>5,17</sup> as do persons with radiation AEs,<sup>156</sup> and epidemiological evidence associates pesticide exposure with autoimmune conditions.<sup>299,300</sup> (It is a prediction of the present paper that if radiation has not already been shown to do so, it will. Nonionizing radiation sensitivity has been tied to increased autoantibodies.<sup>156</sup>) This might account for the (tentatively supported) protective association of immune globulin (gamma globulin), which was given to some Persian Gulf War personnel,<sup>301,302</sup> to AE Propensity—a finding supported in controls and the total sample. Its use has been associated with improvement in experimental and naturally occurring autoimmune conditions.<sup>303,304</sup> The association with protection must be considered tentative—but could suggest a candidate treatment to try.

An unexpected finding was the apparent protective association of copper-exposure against AE Propensity, seen in cases, controls, the full sample and on age-stratified analysis. Excess copper can cause prooxidant injury; however, copper intake in US adolescents and adults has been reported to be low<sup>305</sup>; perhaps copper-zinc SOD may depend, in this setting, on adequacy of copper. Moreover, though able to cause prooxidant effects, copper can also increase activity of catalase and increase gene expression for multiple critical antioxidants and antioxidant regulators, including SOD1 (copper-zinc SOD), glutathione peroxidase, glutathione-S-transferase, Nrf2 (viewed as a master antioxidant regulator), and Kelch-like ECH associated protein 1a (Keap-1a)—albeit, here assessed in a nonmammal organism.<sup>306</sup> In a seminar for medical providers focused on genetic predictors of health problems,<sup>307</sup> a family with extreme multiple chemical intolerance was described, members of which were found to lack remarkable findings in other antioxidant systems but showed strong multiple adverse polymorphisms related to copper-associated detoxification systems (copper chaperone antioxidant-1, "Atox-1").

Cell copper accumulation is reported to be higher in linoleate supplemented cells<sup>308</sup> (however, this study was done in microbes). Linoleic acid is the precursor for arachidonic acid, and we have found many arachidonic acid products to be markedly depressed in VGWI<sup>4,309</sup>—suggesting, but not confirmative for, the possibility of depressed levels of linoleate. Following a speculative line of reasoning, if veterans have reduced linoleic acid, leading to reduced copper retention, those affected with GWI may have relative shortfalls in cellular copper—potentially affecting antioxidant activity,<sup>310</sup> as earlier, and mitochondrial function (which is low in VGWI,<sup>6</sup> as below). Frank deficiency may be unlikely; copper deficiency is linked to peripheral neuropathy (many VGWI indeed cite neuropathic symptoms), ataxia (a feature in some VGWI<sup>311</sup>)—and sometimes motor neuron problems (which three studies reported to be present in excess in GWI, in the first decade or so following Gulf deployment<sup>312–314</sup>). But it is also linked to myelopathy and anemia, which are not reported to be characteristic in VGWI. Of note, the fatty acid changes consistent with those in GWI (potentially opposed to copper uptake) are also reportedly opposed to accumulation of radioactive chemicals (in microbes),<sup>315</sup> suggesting the possibility that observed alterations in VGWI might be adaptive.

The redox activity of copper is essential for mitochondrial enzyme activity,<sup>310</sup> which is depressed in GWI.<sup>6</sup> Additionally, copper deficiency may enhance vulnerability to apoptosis<sup>316</sup>; altered apoptosis regulation has repeatedly emerged as altered in GWI.<sup>317–319</sup> On the flip side, copper is a source of endogenous free radicals<sup>310</sup>; endogenous copper may be mobilized during periods of OS, and may be “particularly dangerous” in settings of mitochondrial dysfunction.<sup>320</sup> A number of agents that cause OS act synergistically with copper to depress mitochondrial membrane potential.<sup>320</sup> Though no definite inferences are possible, assessment of copper status may be merited in VGWI.

Gulf exposure to sunscreen was apparently adverse, an intriguing finding with both mundane and interesting potential bases. This could be a chance finding. Breakdown products arising from desert heat (and nighttime cold) exposure could have conferred problems themselves or with other exposures. The specific sunscreen(s) that used in theater could have had components, or breakdown products, that were a problem themselves or with other exposures. The sunscreen could have afforded a depot and conduit for other chemicals to persist and continue to penetrate the skin. Other oxidative stressor exposures combined with sunscreen application could have led to an immune reaction by mechanisms discussed above.

More interesting hypotheses relate to reduced UV-B exposure and its implications for production of vitamin D and also melanin—which confers antioxidant protection operating locally in the skin.<sup>321,322</sup> UV-B induces production of vitamin D,<sup>323</sup> which is critical for its benefits to OS, apoptosis, autoimmunity, muscle strength, and mitochondrial function. Vitamin D protects against radiation injury including radiation-induced cell senescence and cell/keratinocyte apoptosis.<sup>324,325</sup> It may decrease oxidative injury in GWI-relevant organs including the gastrointestinal tract.<sup>326</sup> It is linked to reduced development of autoimmunity,<sup>327–330</sup> which is increased in VGWI and related to neurological and pain symptoms.<sup>5,17</sup> Improved vitamin D status enhances mitochondrial energy production in muscle,<sup>331</sup> and is linked to muscle strength including in RCTs<sup>332,333</sup> and reduced pain in some studies,<sup>334,335</sup> potentially contributing to the greater impact of Gulf-sunscreen-abstinence-presence on the relation of AE Propensity to symptoms in the neurological and pain domains, relative to others (besides skin). Activation of the vitamin D receptor (e.g., by vitamin D) inhibits energy loss-restoration injury to tissue (reperfusion injury), by “reducing oxidative stress, and inhibiting apoptosis and autophagy dysfunction-mediated cell death”.<sup>336</sup> Vitamin D activates the Nrf2-Keap1 (“master”) antioxidant pathway.<sup>337</sup> It is reported to reverse age-related increases in microglial activation that may contribute to brain effects such as those in aging (also relevant to the neurological domain).<sup>338</sup> So having endogenous UV-B triggered vitamin D on board at the time of other Gulf-relevant exposures—many known to trigger OS, cell energy compromise, autoimmune triggering, and apoptosis—may have protected against injury at the time. This may protect against later symptoms either from exposure-induced worsening of oxidative and mitochondrial mechanisms, e.g., already impaired by Gulf-relevant exposures—and/or some AEs may involve immune-autoimmune systems as suggested by apparent protection by immune globulin.

Sunscreen (and associated disruption of skin access to UV-B) also reduces production of melanin which confers antioxidant protections to the skin.<sup>321,322</sup> This would account for why sunscreen exposure was apparently detrimental selectively in theater, the setting with the high load of toxins against which UV-induced products protect. AE Propensity in VGWI was most strongly related to the skin domain among Kansas GWI domains.



### Limitations of the study

This analysis has limitations. Findings are cross-sectional and causality and in some cases temporality is difficult to infer, particularly for nonveteran participants. (Virtually all veterans describe themselves as having had health that was “very good” or “excellent” at the time of their participation in the Gulf War.<sup>6</sup> Early Gulf experiences are relatively likely to precede health problems.) Information is based on self-report, which may be subject to recall and recording bias.

Selection of participants was not contingent on any specific exposure, so the power to assess the impact of different exposures differs. Of note, participants did not know that we would examine the relation of reported exposures to reported AE Propensity, so at least that expectation will not have influenced reporting.

VGWI are a comparatively heavily exposed group, which benefits ability to see exposure relations for a number of exposures. However, the number of participants, though sizable for a study of VGWI that also assessed objective markers (presented separately)<sup>4,12,55,309</sup> and with excellent power for paired analyses, is limited in the number of exposures that can be meaningfully included in a regression, particularly where subgroup analyses are also conducted. As for all new findings, authority of findings will rest on replication. In this vein, internal replication of major findings—shown in GWI cases and controls, younger and older participants—materially enhances confidence in key findings. The fit of the pesticide findings with chemical sensitivity literature; and the radiation finding with emerging data in the electrosensitivity and radiosensitivity sphere,<sup>156,339,340</sup> further reinforce the likelihood that these are genuine risk factors. Combustion products, which were weaker and less consistent predictors here, have been highlighted in MCS.<sup>341</sup>

Relevant risk factors may have been missed. Insufficient participants with (or without) an exposure, or existence of a vulnerable subgroup *vis-à-vis* that exposure, may necessitate a larger sample or different analysis approach to see effects. Though the exposure assessments were extensive, they are not all-inclusive.

Evaluation of AE propensity should be extended in future studies to other military and veteran groups, as well as to potentially exposed civilian subgroups. However, replication within this study of key findings in nonveteran controls as well as GWI cases, and in younger and older age strata, suggests these findings likely generalize.

Heightened AE Propensity may be viewed as a new feature of GWI, distinct from presence of ongoing symptoms. Concordance of exposure relationships to AE Propensity between VGWI and controls serves as a reminder that findings in the unique Gulf War veteran population can offer vital information of relevance to the rest of us. Exposure to pesticides, radiation, and perhaps combustion products may render some people more susceptible to AEs spanning many classes of exposure. Additional studies are needed to determine whether exposures subsequent to the Gulf, in affected veterans, may not only precipitate time-limited worsening, but may actually magnify severity of the overall illness in VGWI, as some veterans report.

### STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- RESOURCE AVAILABILITY
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- EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS
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## SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.isci.2023.107363>.

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## AUTHOR CONTRIBUTIONS

B.G. designed and oversaw conduction of the study, performed the initial analyses, drafted the initial paper, and participated in the investigation, supervision, administration, and funding acquisition for the project. J.H. participated in validation of statistical results, review and editing of the manuscript, as well as administrative aspects of manuscript submission. All authors read and approved the final manuscript.

## DECLARATION OF INTERESTS

The authors declare no competing interests.

## INCLUSION AND DIVERSITY

We support inclusive, diverse, and equitable conduct of research.

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## STAR★METHODS

### KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Software and algorithms		
Stata Versions 9.0 and 12.0	College Station, Texas	<a href="https://www.stata.com">https://www.stata.com</a>
G*Power Version 3.1.9.7	Faul et al. <sup>345,346</sup>	<a href="https://www.psychologie.hhu.de/arbeitsgruppen/allgemeine-psychologie-und-arbeitspsychologie/gpower">https://www.psychologie.hhu.de/arbeitsgruppen/allgemeine-psychologie-und-arbeitspsychologie/gpower</a>

### RESOURCE AVAILABILITY

#### Lead contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Beatrice A. Golomb, MD, PhD ([bgolomb@ucsd.edu](mailto:bgolomb@ucsd.edu)).

#### Materials availability

This study did not generate new unique reagents.

#### Data and code availability

- All data reported in this paper will be shared by the [lead contact](#) upon request.
- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the [lead contact](#) upon request.

### EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS

#### Human participants

**Table 1** shows the breakdown of age, gender, and ethnicity. The majority of participants were male and Caucasian, consistent with the demographic composition of those deployed to the Persian Gulf. Socioeconomic status was not separately assessed. 81 participants included 41 veterans with Gulf War illness and 40 controls matched 1:1 to 40 of the cases on sex, age, and ethnicity (designated by the participant). An additional case completed the study; recruitment had continued until there were 40 matched pairs, and for this individual case, a matched control had not at that time been identified. Since analyses here are cross-sectional (the primary purpose is not to compare cases to controls, but to look at relationships across and within these groups), the additional case provides additional relevant information on the relation of exposures to outcomes, and is included. Biological sex best describes the data. The dearth of female participants (consistent with the small fraction of Gulf deployed personnel that were female) precludes meaningful sex-stratified analysis. The parent study from which the data were secured was approved by the UC San Diego Human Research Protections Program (HRPP) and the Department of Defense (DoD) Human Research Protection Office (HRPO). All participants gave written informed consent.

### METHOD DETAILS

This study encompasses case-control and cross-sectional design elements. We assess propensity to experience adverse events to exposures (AE Propensity) in VGWI vs. matched healthy controls. We assess the cross-sectional relationships between individual exposures and AE Propensity, in the total sample and in GWI cases and healthy controls separately. Participants were obviously not randomized to exposures in the Gulf. As this is not a treatment study, randomization did not occur. All participants and study staff were blinded to the intent to assess adverse effect propensity, or to relate it to Gulf exposures.

To qualify as a GWI case, veterans must have been deployed to the Persian Gulf theater of operations between August 1, 1990 and July 31, 1991, inclusive. VGWI were additionally required to meet both CDC and Kansas symptom inclusion criteria for GWI.<sup>3,342</sup> CDC criteria require presence of symptoms for at least

6 months, arising during or after Gulf War participation, in at least two of the three domains of fatigue/sleep, mood-cognitive, and musculoskeletal.<sup>342</sup> The more discriminating, more specific Kansas criteria require that symptoms have been present for at least 6 months, arising during or after Gulf deployment, in at least three of a suite of six categories comprising fatigue/sleep, pain, neurological/cognitive/mood, respiratory, gastrointestinal, and dermatologic.<sup>3</sup> For a symptom to qualify toward Kansas symptom criteria, the component symptoms must be at least moderate in severity (not mild) and/or there must be multiple symptoms within the domain.<sup>3</sup>

To qualify as a control, it was required that participants were nonveterans, meeting neither Kansas nor CDC symptom inclusion criteria for GWI, and additionally not meeting Kansas exclusion criteria (that is, they could not have other health conditions such as lupus or multiple sclerosis that could produce symptoms that could be confused for those of GWI). Controls were selected to match 1:1 to enrolled cases on sex, ethnicity, and age. A half-match for ethnicity was deemed to be qualifying, in recognition of the prevalence of mixed ethnicities. Age matching for matched pairs was within 4 years.

Nonveteran and veteran controls each have advantages and disadvantages. We preferred nonveteran controls. 1. Veterans of the 1990-1 Gulf War who remain healthy despite exposures may have specific protective genetics and/or engage distinct protective adaptations that may render them unsuitable as controls. Gulf era veterans who were not deployed are known to differ significantly: unhealthier persons are not selected for deployment to high threat areas and it has been affirmed that nondeployed era veterans were materially less healthy.<sup>343</sup> Veterans from other conflicts may have experienced other problematic exposures (e.g. antimalarials like mefloquine that may compromise suitability). All military personnel experienced some exposures (e.g. mandatory vaccinations, depleted uranium, permethrin, impregnated uniforms) that may contribute toward shared mechanisms with GWI, reducing statistical power to observe differences. For these reasons, our preference is for nonveteran healthy controls. Controls were drawn from rosters of healthy controls used in prior GWI studies, supplemented by outreach primarily through ResearchMatch.

## QUANTIFICATION AND STATISTICAL ANALYSIS

### Measurements

Demographic characteristics including age, sex, ethnicity, and marital status were assessed. Military information was collected for VGWI.

GWI-relevant symptoms were gauged by Kansas criteria symptom scores for each Kansas symptom domain.

Exposures were elicited via inquiries for a list of general exposures (non-Gulf specific – see Table 1a); and for veterans, a list of Gulf-specific exposures (see Table 1b). Whether the exposure had been experienced was rated no, unsure, or yes – which were coded as 0, 0.5, and 1, respectively. A summed exposure score (**totExp**) summed responses on queried exposures.

Adverse effects: Those who rated an exposure as “yes” were asked if they had experienced an AE or health effect to the exposure. Response options were again no, unsure, or yes, coded as 0, 0.5, and 1, respectively. A summed adverse effect score (**totAE**) summed responses on the AE queries.

AE Propensity: To gauge a proxy for AE Propensity, a ratio was calculated, of totAE divided by totExp – roughly, the fraction of assessed exposures to which AEs were reported to have been experienced.

Chemical sensitivity: The study assessed self-rated chemical sensitivity via the chemical sensitivity question from the widely-employed Kansas GWI questionnaire, as well as via our single-item UCSD GWI chemical sensitivity self-rating, analyzed as a binary assessment (0 if absent, 1 if present). The Kansas criteria question states: “Having physical or mental symptoms after breathing in certain smells or chemicals.” The timeframe for the Kansas query is the prior 6 months, with a 4-point Likart scale as absent, mild, moderate, severe rated 0, 1, 2, 3, respectively. The UCSD self-rating states: “Chemical sensitivity (e.g., unusual sensitivity to smells).” The timeframe is 2 weeks. This was assessed as a binary rating, 0 if absent, 1 if present. The Kansas and UCSD measures show convergent validation against one another:  $r=0.57$ ,  $p=0.0001$ . The single-item UCSD chemical sensitivity measure was further validated by affirming a relation to SOD2 a16v, an

adverse polymorphism of the major mitochondrial antioxidant SOD2,<sup>344</sup> that had elsewhere related to chemical sensitivity in a Japanese cohort.<sup>49</sup>

Missing data in exposure (and AE) tallies, a missing value was treated as absence of the corresponding exposure (or AE).

### Analysis

Descriptive statistics characterized participant characteristics, Kansas symptom ratings, totAE, totExp, AE Propensity and the chemical sensitivity measures for all participants, and for cases and controls separately. T-tests and chi-squared tests compared characteristics of cases to those of controls for continuous and categorical variables respectively. Ethnicity was used for matching of controls to cases for case-control assessments, but was otherwise not a focus of the present study.

Summed AEs, summed exposures, and AE Propensity were compared in (all) cases vs. controls (unpaired t-test), The correlations of AE Propensity to Kansas symptom domains and to chemical sensitivity indices were assessed.

We evaluated how individual exposures correlated to AE Propensity, in the total sample, and in cases and controls separately. Correlation in each group was complemented by regression in the total sample, adjusted for case status. All regression analyses used robust standard errors (aka heteroskedasticity-independent standard errors).<sup>60</sup>

Shared relationships among exposures within an exposure class were used to guide generation of composite variables for inclusion in multivariable models (to limit the number of predictors included).

A shared set of predictors were identified for use in multivariable models, spanning all participants (adjusted for case status), and cases and controls separately. These models could not include Gulf-specific exposures, as these were not elicited for controls.

Additional models were individually optimized for all participants, adjusted for case status; for cases; and for controls. For the model specific to cases, Gulf-specific exposures could be included.

After identification of a robust "main" model, a number of variables were assessed for candidacy as added variables (particularly those with  $p \leq 0.15$  on univariable assessment), and the best identified candidates are shown.

Main models for the combined case-control samples were reassessed in age-stratified analysis (stratified at the mean sample age) to assess internal replication in independent groups.

Key predictors were then reassessed via logistic regression in GWI cases, as predictors of the binary UCSD chemical sensitivity variable.

Analyses used Stata 9.0 and Stata 12.0 (College Station, Texas). Two-sided  $p < 0.05$  designated statistical significance. Adjustment for multiple comparisons was not undertaken.

### Sample size and power

Analyses capitalized on data procured for a different primary purpose though from the outset, the intention was to assess other exposure/outcome relationships, to enhance yield per dollar expended and more importantly, per Gulf War veteran contribution/burden. The sample size of 40 per group (matched pairs) provides power of 99% to identify a 0.5 SD difference with two-sided alpha of 0.05 (G\*Power 3.1.9.7).<sup>345,346</sup> Alternatively, the study has 80% power at two-sided alpha of 0.05 to detect an effect size of  $\geq 0.32$  standard deviations (strong power for a modest effect size). Power for regression analyses varies by group/subset assessed, but significance for key exposure outcome relationships, upheld in stratified analyses, affords post hoc affirmation of adequate study power for these assessments.





### Efforts to reduce bias

- (1) Key limitations include prospects for recall and reporting bias, with a material span from Gulf exposures to survey completion. However, an “unsure” response helped capture uncertainty in responses. Moreover, there is not strong rationale for considering that specific exposure classes would be disproportionately reported *in alignment with* adverse effect propensity spanning exposure classes.
- (2) Primary models were assessed in all participants, in VGWI and controls separately, and in analysis stratified by age to protect against spurious findings. Assessment of calculated AE Propensity against self-rated chemical sensitivity further reinforces validity of the AE Propensity construct assessed.