ABSTRACT
A major challenge in organophosphate (OP) research has been the identification and utilization of reliable biomarkers for the rapid, sensitive, and efficient detection of OP exposure. Although Tyr 411 OP adducts to human serum albumin (HSA) have been suggested to be one of the most robust biomarkers in the detection of OP exposure, the analysis of HSA-OP adduct detection has been limited to techniques using mass spectrometry. Herein, we describe the procurement of two monoclonal antibodies (mAb-HSA-GD and mAb-HSA-VX) that recognized the HSA Tyr 411 adduct of soman (GD) or NI-(diisopropylamino)ethyl methylphosphonothioate (VX), respectively, but did not recognize nonphosphonylated HSA. We showed that mAb-HSA-GD was able to detect the HSA Tyr 411 OP adduct at a low level (i.e., human blood plasma treated with 180 nM GD) that could not be detected by mass spectrometry. mAb-HSA-GD and mAb-HSA-VX showed an extremely low-level detection of GD adducted to HSA (on the order of picograms). mAb-HSA-GD could also detect serum albumin OP adducts in blood plasma samples from different animals administered GD, including rats, guinea pigs, and monkeys. The ability of the two antibodies to selectively recognize nerve agents adducted to serum albumin suggests that these antibodies could be used to identify biomarkers of OP exposure and provide a new biologic approach to detect OP exposure in animals.

Introduction
Organophosphates (OPs) constitute a diverse class of chemicals that include insecticides, protease inhibitors, and chemical warfare agents. The use of OP toxicants as chemical warfare weapons is a threat to US military personnel as well as civilians. In addition, agricultural workers who handle OP pesticides are at risk of exposure because there are more than 100 different OP pesticides used worldwide, and it is estimated that globally there are 750,000 to 3 million human intoxications by OPs annually (Kwong, 2002). The worldwide level of OP exposure and the possible toxic consequences underscore the need to develop an efficient, portable, and inexpensive way to detect OPs (Chen and Mulchandani, 1998; Sogorb and Vilanova, 2002).

The acute toxic effects of OPs arise from the inhibition of acetylcholinesterase (AChE) to form a relatively stable OP-AChE adduct. AChE inhibition results in accumulation of excess acetylcholine that leads to stimulation of acetylcholine receptors, which can produce convulsions and other central nervous system toxicity (Marrs et al., 2007). In addition to AChE inhibition, OPs also covalently modify other proteins (Schopfer et al., 2005; Wieseler et al., 2006; Thompson et al., 2010), including butyrylcholinesterase (BChE) and serum albumin.

Analysis of biomedical samples (e.g., urine and blood) can provide not only qualitative but also quantitative information about OP exposure (Marrs et al., 2007). Numerous approaches have been used in the detection of OP exposure (Worek et al., 2005; Thompson et al., 2010), including determination of inhibition of serum AChE and BChE functional activities with biochemical assays and detection of unbound OPs in plasma or their decomposition products with mass spectrometry (MS) methods, including gas chromatography–MS (GC-MS) or tandem mass spectrometry; GC-MS, gas chromatography–mass spectrometry; GD, soman; GF, cyclosarin; HPLC, high-performance liquid chromatography; HRP, horseradish peroxidase; HSA, human serum albumin; HSA-GD, decapetide hapten that has a GB phosphonylated tyrosine; HSA-GD, decapetide hapten that has a GD phosphonylated tyrosine; HSA-GF, decapetide hapten that has a GF phosphonylated tyrosine; HSA-soman protein adduct, human serum albumin that has conjugated soman; HSA-VX, decapetide hapten that has a VX phosphonylated tyrosine; IS, internal standard; KLH, keyhole limpet hemocyanin; LC-MS/MS, liquid chromatography–tandem mass spectrometry; mAb, monoclonal antibody; mAb-HSA-GD, monoclonal antibody that recognizes HSA tyrosine 411 adduct of GD; mAb-HSA-VX, monoclonal antibody that recognizes HSA tyrosine 411 adduct of VX; MS, mass spectrometry; OP, organophosphate; PBS, phosphate-buffered saline; PDVF, polyvinylidene difluoride; RSA, rat serum albumin; OVAL, ovalbumin; SDS-PAGE, sodium dodecyl sulfate polyacrylamide gel electrophoresis; VX, S-[2-(diisopropylamino)ethyl]-O-ethyl methylphosphonothioate.

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liquid chromatography–tandem MS (LC-MS/MS). Other approaches include GC-MS detection of fluoride-induced reactivated AChE or BChE and detection of AChE-, BChE-, or albumin-OP adducts with capillary electrophoresis–MS or LC-MS-MS after proteolytic digestion of those proteins. Among these approaches, determination of inhibition of AChE and BChE functional activity in the blood is still the mainstay for quick initial screening, although determination of relative cholinesterase activity lacks sensitivity and specificity. MS approaches require sophisticated equipment and well trained operators (Worek et al., 2005).

Detection of protein-OP adducts (i.e., AChE, BChE, or serum albumin) as biomarkers for OP exposure has provided insight into the structural biology of OP action (Read et al., 2010; Thompson et al., 2010). Generally, the approach to detect protein-OP adducts has relied on MS (Thompson et al., 2010), although one report showed that antibodies could be raised against phosphorylated AChE (George et al., 2003). However, in that report, the antibodies reported to recognize the inhibited form of AChE did not distinguish OP-AChE adducts arising from different OPs (George et al., 2003). An attempt to develop antibodies against phosphorylated BChE showed that the chemical instability of phosphorylated decapeptide haptens designed to mimic phosphorylated BChE precluded using this approach (MacDonald et al., 2010, and unpublished observations). In addition, despite the fact that AChE and BChE are prominent targets of OP exposure, these enzymes are present at low concentrations (e.g., BChE is present at 4 μg/ml in plasma) and the OP-BChE adduct is labile (Li et al., 2007). In contrast, serum albumin is the most abundant protein in blood plasma (40,000 μg/ml), and after OP exposure, OPs form a stable adduct (e.g., with human serum albumin [HSA] at Tyr 411) on the basis of LC-MS analysis of proteolytic digests (Li et al., 2008). Thus, albumin is a functional scavenger of OPs, resulting in a stable OP adduct after exposure. It has been estimated that 1–2% of HSA forms a stable OP adduct after OP exposure (Ding et al., 2008). Based on the instability of phosphorylated adducts to AChE and BChE, the detection of HSA-OP adducts represent a more feasible strategy to detect in vivo biomarkers of OP exposure because the abundance of serum albumin is 10,000 times more than either AChE or BChE and the Tyr 411 adduct is extremely stable (Ding et al., 2008). Tyr 411 OP adducts of HSA have been reported in a number of cases of OP exposure (Peeples et al., 2005; Li et al., 2010; Lockridge and Schopfer, 2010; Marsillach et al., 2011).

Protein sequence alignment of serum albumin indicates that across different species, including humans, rats, and guinea pigs, the region surrounding Tyr 411 is highly conserved (Fig. 1). We therefore chose this region (i.e., amino acids 408–417, LVRYTKKVPQ, in HSA) as the epitope to synthesize phosphorylated antigens and haptens. Based on the three-dimensional structure of HSA, Tyr 411 lies on the surface of the protein (Ding et al., 2008) and Tyr 411 is apparently readily accessible and efficiently attacked by OPs to form stable covalent adducts of HSA. This may explain why Tyr 411 of HSA is preferentially phosphorylated if treated with OPs (Li et al., 2008).

In the present report, we discuss the rationale and the successful generation of five monoclonal antibodies (mAbs) (in which two were studied in detail) that were raised against four different phosphorylated decapeptides corresponding to the region surrounding Tyr 411 in HSA. Two of the mAbs selectively recognized soman (GD) or S-[2-(diisopropylamino)ethyl]-O-ethyl methylphosphonothioate (VX) adducts, respectively, at Tyr 411 on HSA. The two antibodies were shown to possess great sensitivity (i.e., as little as 10⁻¹² g of antigen detected in blood plasma samples) and detected OP-adducted serum albumin from various biologic samples prepared from in vitro and in vivo OP exposure experiments. The antibodies also stereoselectively recognized OP adducted to HSA. Because the OPs used in these studies possess two centers of chirality, four possible OP adducts could form. In the case of GD adducted to HSA, one of the adducts was selectively recognized over the other one. Two mAbs (i.e., mAb-HSA-GD and mAb-HSA-VX) were successfully used in studies to detect OP adducts of serum albumin in different species including humans, rats, guinea pigs, and monkeys. The availability of the two mAbs not only provides a powerful tool in basic research, but also could be used to identify biomarkers of OP exposure and provide a new approach to detect OP exposure in animals.

Materials and Methods

Biologic and Chemical Reagents. Buffers, reagents, and solvents were purchased from VWR Scientific, Inc. (San Diego, CA) in the highest purity commercially available. The nerve agent model compounds and their corresponding Sp and Rp isomers were synthesized as previously described (Berman and Leonard, 1989; Barakat et al., 2009). The nerve agent model compounds are toxic and should be handled with extreme care. Chemical wastes containing nerve agent model compounds were hydrolyzed by overnight incubation with 2.5 M NaOH and 10% ethanol before disposal. Goat anti-mouse antibodies conjugated with horseradish peroxidase (HRP) and SuperSignal West Pico Chemiluminescent Substrate was purchased from Pierce (Rockford, IL). Molecular biology reagents were purchased from Life Technologies (Carlsbad, CA) unless otherwise specified.

Synthesis of para-Nitrophenyl Esters of Nerve Agent Model Compounds. The synthesis of para-nitrophenyl esters of nerve agent model compounds was accomplished by combining the corresponding nerve agent monochloridates (50 mg, 0.25 mmol) with triethylamine (150 mg, 0.9 mmol) and dimethylaminopyridine (5 mg, 0.04 mmol) at 4°C in CH₂Cl₂. After addition of para-nitrophenol (150 mg, 1.1 mmol) and stirring at room temperature for 5 hours, the mixture was cooled to 4°C and made basic, and the separated organic fraction was evaporated to dryness and chromatographed to afford the desired products in 78%–91% yield. Each product was fully characterized by NMR.

Phosphorylation of HSA in the Presence of para-Nitrophenyl Esters of Sarin, GD, and VX. The incubation comprised 200 mM para-nitrophenyl esters of sarin (GB), GD, and VX (made up in a stock solution of CH₂CN) and 10 mM HSA in Tris-HCl buffer (10 mM, pH 8.0) in a total incubation volume of 0.6 ml. The incubation
was initiated after thorough mixing and continuously monitored at 400 nm with a Cary UV-Vis spectrophotometer (Agilent Technologies, Santa Clara, CA). The progress curves were plotted as a function of absorbance increase versus time. The half-life and rate constants were determined (n = 3).

**Characterization of Phosphonylated Peptide–Amino Capric Acid Hapten Purity and Aqueous Stability.** The purity of phosphonylated peptides was determined by high-performance LC (HPLC). Phosphonylated peptides were run on a Beckman Gold HPLC column using 0.1% trifluoroacetic acid in water-acetoni trile and a reverse-phase C18 column using a Hitachi 2000 HPLC (Hitachi Inc., Dallas, TX). HPLC data were reported for final, purified peptides. Final purities ranged from 92–99%. Peptide purity for each peptide is shown in Table 1. The stability of the phosphonylated peptides was determined by MS. Each phosphonylated peptide was prepared to a final concentration of 1 mg/ml and an internal standard (IS) was included to assist MS quantification (pH 7.0, 25°C). At the appropriate time point, an aliquot of each sample was injected into the MS (Hitachi M-8000; Hitachi Inc.) using electrospray ionization in the positive mode. Signal intensities for each phosphonylated peptide were tabulated and compared with the signal intensities of the IS peptide for each run (Peptide/IS). Peptide/IS for each day was normalized to the value obtained at day 0, and this value was multiplied by 100 to get percent remaining phosphonylated peptide. Plots of percent remaining versus time (days) were made in Excel. Half-lives were calculated for each phosphonylated peptide shown in Table 1.

**Preparation of Antigens and Conjugation of Antigens to Proteins.** Phosphonylated peptide antigens (408LVRY*TKKVPQ417-AcA, where AcA stands for ε-amino capric acid and * stands for a phosphonyl group) were synthesized as described previously (MacDonald et al., 2010). The purity of each phosphonylated peptide-AcA was >95% as judged by HPLC, and the structure was confirmed by HPLC-MS. Each phosphonylated peptide-AcA was conjugated to keyhole limpet hemocyanin (KLH; Pierce) and chicken egg white ovalbumin (OVA; Pierce) using ethyl-diisopropyl-carbodiimide and sulfo-N-hydroxysuccinimide (1:1). Highly purified conjugated phosphonylated peptide-AcA carrier protein was obtained after removing excess peptides and reagents by filtration and dialysis against phosphate-buffered saline (PBS) buffer (pH 7.4). The purity of the conjugated phosphonylated peptide-AcA was confirmed by HPLC-MS. Each phosphonylated peptide was purified to >95% as judged by HPLC, and the structure was confirmed by HPLC-MS. Each phosphonylated peptide-AcA was conjugated to keyhole limpet hemocyanin (KLH; Pierce) and chicken egg white ovalbumin (OVA; Pierce) using ethyl-diisopropyl-carbodiimide and sulfo-N-hydroxysuccinimide (1:1). Highly purified conjugated phosphonylated peptide-AcA carrier protein was obtained after removing excess peptides and reagents by filtration and dialysis against phosphate-buffered saline (PBS) buffer (pH 7.4) through repeated filtration with Amicon Ultra 4 centrifugation tubes (Millipore, Temecula, CA). Protein concentration was determined by absorbance increase versus time. The half-life and rate constants were determined (n = 3).

**Immunization.** The procedures and care of animals used in this work conformed to the Guiding Principles in the Care and Use of Animals provided by the American Physiological Society, as well as all federal and California regulations. Human BioMolecular Research Institute has an approved Assurance from the National Institutes of Health’s Office for Protection from Research Risks. Female Swiss Webster mice (5 weeks, 20 g) were purchased from Taconic Farms (Oxnard, CA) and used in immunization studies with the conjugated haptens. After a 1-week stabilization period, 10 mice were immunized with each KLH-conjugated hapten (i.e., HSA-VX-KLH, HSA-GD-KLH, HSA-GF-KLH, HSA-GB-KLH, and nonphosphonylated HSA-KLH). The immunization protocol consisted of an initial i.p. injection of 150 µg of conjugated hapten in 200 µl of an oil-in-water emulsion (PBS-containing Sigma Adjuvant System; Sigma, St. Louis, MO) followed by another boost 4 weeks later. After an additional 3 weeks, each animal was boosted with 100 µg OVAL-conjugated phosphonylated HSA decapetide (i.e., HSA-GD-OVAL, HSA-VX-OVAL, HSA-GF-OVAL, and HSA-GB-OVAL) in 200 µl PBS-containing alun (Pierce) as an i.p. injection. Mouse blood (50–100 µl) was collected from each animal from the tail vein 7–10 days after each boost. Heparin (Sigma), 1–3 IU, was added to each blood sample and mixed thoroughly, and plasma was separated from blood cells through centrifugation at 5000 rpm for 10 minutes at 4°C. Each plasma sample was tested for the presence of anti-phosphonylated peptide antibodies by enzyme-linked immunosorbent assay (ELISA) (see below). When the titer reached >6400 units, the mice were allowed to rest for at least 1 month. A final tail vein injection was made with KLH-conjugated phosphonylated peptides (25–50 µg in PBS) or nonphosphonylated HSA peptide–conjugated KLH hapten 3 days before the mice were taken for splenectomy and cell fusion.

**Cell Culture and Fusions.** Mouse myeloma cells NS-1 (American Type Culture Collection, Manassas, VA) were seeded at 50–100 × 10^3 cells/ml and grown in Medium A (Stemcell Technologies, Vancouver, BC, Canada). Three days after the final tail vein immunization, mice were sacrificed and 1–2 × 10^7 isolated spleen cells were fused with 1–2 × 10^7 myeloma cells using polyethylene glycol (Stemcell Technologies). After cell fusions, 500-mm dishes were plated with 2–4 × 10^5 cells/ml in methycellulose medium (Medium D; Stemcell Technologies) for 7–14 days. Single colonies of hybridoma cells were picked from methycellulose medium and cultured in Medium D (Stemcell Technologies) in 96-well plates. Cell culture supernatants were tested for the presence of antibodies to OVAL-conjugated HSA-OP peptides (i.e., HSA-GB-OVAL, HSA-GD-OVAL, HSA-GF-OVAL, and HSA-VX-OVAL) or nonphosphonylated HSA-OP using ELISA. Positive cultures were expanded and supernatants were tested against a panel of antigens to verify antibody selectivity before secondary cloning in methycellulose Medium D or by limited dilution in Medium E. After recloning of hybridoma cells for 2–3 rounds, the monoclonal cells afforded single heavy-chain and single light-chain clones and were deemed clones that generated mAbs and isotyped with isotyping kit (Pierce).

**Purification of Antibodies.** For hybridoma cells expressing the desired anti-HSA-OP mAbs, the cell culture was expanded. Briefly, culture medium of the hybridoma cells was centrifuged at 1000 rpm to separate cells from the supernatant, and the supernatant was filtered to remove residual debris. To the culture medium supernatant was added 0.01% NaNO_3_, and it was mixed thoroughly at room temperature prior to purification. For mAb purification, the protein was dialyzed against PBS (pH 7.4) with 0.01% NaNO_3_ and mixed 1:1 with binding buffer (Pierce) and then chromatographed on Protein G Sepharose 4B (Sigma) equilibrated with PBS (pH 7.4) containing 0.01% NaNO_3_. Culture medium was loaded onto the column at 1 ml/min. The Protein G column was washed extensively with PBS (pH 7.4) containing 0.01% NaNO_3_. Bound antibodies were eluted with low-pH elution buffer (pH 2.0; Pierce) and immediately neutralized to pH 7.5 with 1.0 M Tris. Protein-containing fractions were pooled based on optical density at wavelength of 280 nm (OD_280) absorbance and dialyzed against PBS (pH 7.4) containing 0.01% NaNO_3_ for routine application. Purified antibodies were quantified by

### TABLE 1

Characterization of phosphonylated peptide-AcA hapten purity and aqueous stability

<table>
<thead>
<tr>
<th>Name</th>
<th>Half-life</th>
<th>Mol. Wt.</th>
<th>Purity</th>
</tr>
</thead>
<tbody>
<tr>
<td>HSA-peptide-AcA-GB</td>
<td>12.0</td>
<td>1465.7</td>
<td>&gt;92.4</td>
</tr>
<tr>
<td>HSA-peptide-AcA-GD</td>
<td>16.3</td>
<td>1507.4</td>
<td>&gt;99.6</td>
</tr>
<tr>
<td>HSA-peptide-AcA-GF</td>
<td>19.3</td>
<td>1535.3</td>
<td>&gt;99.2</td>
</tr>
<tr>
<td>HSA-peptide-AcA-VX</td>
<td>8.5</td>
<td>1451.3</td>
<td>&gt;98.4</td>
</tr>
</tbody>
</table>

AcA, amino capric acid; GB, sarin; GD, soman; GF, cyclosarin; HSA, human serum albumin; VX, S-[(diisopropylamino)ethyl]-O-ethyl methylphosphonothioate.

* Peptide half-life was determined by mass spectrometric analysis (deionized H_2O, 25°C).
* Organothosphate-adducted peptide purity was determined by high-performance liquid chromatography.
OD280 absorbance (i.e., OD280 1.4 for 1 mg/ml IgG) and analyzed on sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) for purity.

Western Blot. Western blot analysis was conducted to determine levels of phosphorylated and nonphosphorylated OP-added serum albumin. The OP-added serum albumin was quantified in highly purified protein samples (or plasma protein samples) after removing cells by centrifugation at 5000 rpm at room temperature for 10 minutes. Blood plasma protein samples were prepared by addition to the plasma of a proteinase inhibitor cocktail (Sigma) in a 1:100 dilution containing 1 mM each of phenylmethylsulfonyl fluoride and Na3VO4 and 1 μM each of aprotinin, leupeptin, and pepstatin. Samples were run on 10% SDS-polyacrylamide gels. The primary antibodies were obtained from hybridoma culture supernatant or highly purified mAbs. Secondary antibodies used for Western blotting included HRP-conjugated goat anti-mouse antibodies (HRP-GAM) (Jackson ImmunoResearch Laboratories, Inc., West Grove, PA), and blots were developed using either SuperSignal West Pico Chemiluminescent (Thermo Fisher, Waltham, MA) for regular detection or Immobilon Western Chemiluminescent HRP Substrate (WBKLS0500; Millipore) for high-sensitivity detection. Polyvinylidene difluoride (PVDF) membranes were used in the experiment (IPVH00010; Millipore). For quantification, Western blots were scanned and densitometry was determined using Scion Image software (Scion Corporation, Frederick, MD).

ELISA. For quantification of immunoreactivity of serum from immunized mice or analysis of hapten binding of purified antibodies, ELISA was conducted. The wells of 96-well ELISA plates (Greiner, Neuburg, Germany) were coated with 0.1–2.0 μg of HSA-OP peptides or HSA-GB-OVAL, HSA-GD-OVAL, HSA-GF-OVAL, and HSA-VX-OVAL for 1 hour at room temperature in 50 μl of PBS (without calcium or magnesium) at pH 7.4. After rinsing with PBST (PBS with 0.1% Tween-20), the 96-well plates were blocked for 1 hour with 230 μl of blocking buffer [PBS containing 0.05% (v/v) Tween-20 (PBST) and 1% (w/v) bovine casein (Sigma) (PBSTC)]. After removing the blocking buffer, 50–100 μl of hybridoma culture medium or purified antibody in PBSTC was applied to each well and allowed to incubate at 37°C in a CO2 incubator for 1 hour. After rinsing off the primary antibody, a 50-μl aliquot of HRP-GAM secondary antibody (1:4000 dilution in PBSTC) (Jackson Laboratories) was added to each well and incubated at room temperature for 1 hour. After three rinses with PBST, bound antibody was detected by adding 100 μl of Pierce chemiluminescent substrate (Thermo Scientific, Brookfield, WI) to each well and reading the luminosity using a Wallac Victor2 plate reader (Perkin Elmer, Waltham, MA). For evaluation of serum titers, samples were diluted in PBSTC 1:100-fold and subsequently in twofold series. The diluted plasma assayed ranged generally from 1:100- to 1:128,000-fold dilutions depending on the titer.

Preparation of Rat Blood Plasma Samples. Male Sprague-Dawley rats (Crl:CD(SD); 250–300 g) were obtained from Charles River Laboratories (Wilmington, MA). Rats were housed individually in polycarbonate cages in a temperature-controlled (21 ± 2°C) and humidity-controlled (50% ± 10%) colony room and maintained on a reversed 12-hour light/dark cycle with lights off at 0900 hours. Food and water were available ad libitum in home cages. Blood samples were obtained from nerve agent–exposed animals as part of sample sharing from other studies conducted at the US Army Medical Research Institute of Chemical Defense. VX and GD were obtained from the US Army Edgewood Chemical Biologic Center (Aberdeen Proving Ground, MD) and diluted in sterile saline. For VX, 0.8 × LD50 (LD50 = 16 μg/kg) was administered s.c. into the right flank at a volume of 0.5 ml/kg body weight. For GD, 192 μg/kg (1.2 × LD50) was administered s.c. (0.5 ml/kg), followed 1 minute later with coadministration i.m. (0.5 ml/kg) of the oxime HI-6 (83.6 mg/kg) and atropine sulfate (2 mg/kg). Diazepam (10 mg/kg) was administered s.c. 30 minutes after seizure onset. Standard therapy of an oxime, atropine, and diazepam was given to maximize survival to a lethal dose of agent. Twenty-four hours after GD intoxication, the rats were given a wet mash of food pellets and sugar. S.c. injections of saline (3 ml) were administered as needed to prevent dehydration.

Effect of Sp-GD on Plasma Cholinesterase Functional Activity in Rats. From six 240-g male Sprague-Dawley rats (Harlan, Milpitas, CA) administered 250 μg/kg of Sp-GD, tail vein blood was collected at different times, including at 0, 2, 4, 6, 8, 24, 48, 72, 96, 144, and 240 hours, and immediately cooled to 4°C. Plasma was obtained from whole blood after separation by centrifugation at 2000g for 10 minutes. Cholinesterase functional activity was detected by the Ellman assay (Ellman et al., 1961). Typically, 20 μl of plasma was used in each assay [1 mM acetylthiocholine, 1 mM 5,5′-dithiobis-(2-nitrobenzoic acid) in 50 mM PBS buffer, pH 7.4]. Absorbance change of 5,5′-dithiobis-(2-nitrobenzoic acid) was monitored at 405 nm using a Cary UV-Vis spectrophotometer. Absorbance at 405 nm versus time was plotted to obtain initial rate measurements. Initial rate measurements were normalized to zero time (i.e., before Sp-GD administration, the change in absorbance at 405 nm was taken as 100% functional activity) and a plot of relative cholinesterase activity versus hours or days posttreatment was done with GraphPad Prism (GraphPad Software, La Jolla, CA).

Data Analysis. In the kinetic studies of antibody and ligand binding, the data analyses and statistical calculations were conducted with GraphPad Prism version 5.01. (GraphPad Software).

Results

Design and Synthesis of Antigens. Although there are numerous reports concerning the synthesis of phosphorylated peptides (McMurray et al., 2001), before our publication (MacDonald et al., 2010), there were very few reports in the literature concerning the synthesis of phosphorylated peptides (Fidder et al., 2002). As previously described (MacDonald et al., 2010), four phosphorylated peptides were used in the synthesis of the required haptons (Fig. 2), including peptide conjugates of GB 1, GD 2, cyclosarin (GF) 3, and VX 4. The chemical synthesis of the antigens and conjugated immunogens followed the general strategy described previously (MacDonald et al., 2010). Briefly, Fmoc-protected tyrosine, 5, was treated with monochloridates of nerve agent model compounds 1–4 to afford phosphorylated benzyl-protected Fmoc tyrosine 10–13, respectively (Fig. 3). The monochloridates 1–4 were prepared as described before (MacDonald et al., 2010). After purification by chromatography...
on silica gel, the benzyl group was removed by hydrogenolysis. All of the compounds were fully characterized by $^1$H NMR and $^{31}$P NMR, MS, and HPLC. The phosphorylated products 10–13 were incorporated into solid-phase peptide synthesis to afford the desired phosphorylated peptide 14–17 containing an AcA at the carboxyl terminus. These products were >95% pure on the basis of HPLC and MS. The carrier protein KLH was conjugated through the AcA to produce the desired conjugated hapten-KLHs that were used to immunize mice (Fig. 3). Alternatively, 14–17 was used to conjugate to OVAL or HSA, which were used in various bioassays. In total, three different phosphorylated peptide conjugates were synthesized (i.e., hapten-KLH, hapten-HSA, and hapten-OVAL) that were used to produce the desired conjugated hapten-KLHs that were used to immunize mice (Fig. 3). Alternatively, 14–17 was used to conjugate to OVAL or HSA, which were used in various bioassays. In total, three different phosphorylated peptide conjugates were synthesized (i.e., hapten-KLH, hapten-HSA, and hapten-OVAL) that were used to produce the desired conjugated hapten-KLHs that were used to immunize mice (Fig. 3). Alternatively, 14–17 was used to conjugate to OVAL or HSA, which were used in various bioassays. In total, three different phosphorylated peptide conjugates were synthesized (i.e., hapten-KLH, hapten-HSA, and hapten-OVAL) that were used to produce the desired conjugated hapten-KLHs that were used to immunize mice (Fig. 3). Alternatively, 14–17 was used to conjugate to OVAL or HSA, which were used in various bioassays. In total, three different phosphorylated peptide conjugates were synthesized (i.e., hapten-KLH, hapten-HSA, and hapten-OVAL) that were used to produce the desired conjugated hapten-KLHs that were used to immunize mice (Fig. 3).

**Procurement of Antibodies.** Tail vein blood was obtained from the immunized mice, and plasma obtained after centrifugation was tested for the presence of anti-phosphorylated HSA–peptide (i.e., anti-GB-, GD-, GF-, and VX-HSA antibodies) by ELISA. When the titer reached >6400 units, the mice were allowed to rest at least 1 month. A final tail vein injection was made with KLH-conjugated phosphorylated peptides before splenectomy and cell fusion. Spleens from immunized mice were obtained and finely minced for fusion with myeloma cells to give rise to primary hybridoma cells. Fusion of spleens from mice immunized against GB and GF yielded very few primary hybridomas, and thus, we focused our efforts on hybridoma cells from F41 and F46 (i.e., hybridomas that produced mAbs that selectively recognized HSA-GD and HSA-VX, respectively, named mAb41.5D5A2 and mAb46.5H10G7). After screening >200,000 primary cells, 40 clones were obtained. Recloning of these 40 positive hybridoma cells two or three more times provided homogenous cell lines expressing two lines of mAbs that selectively recognized HSA-GD or HSA-VX. Thus, mAb-HSA-GD and mAb-HSA-VX were obtained after cloning and recloning several generations of hybridoma cells. The mAbs were purified to homogeneity and characterized by SDS-PAGE and isotyping (Fig. 4). The isotypes for both mAb-HSA-GD and mAb-HSA-VX were determined to be IgG1k and their light chains were both k isotype.

**In Vitro Characterization of mAb-HSA-GD and mAb-HSA-VX.** Characterization of the selectivity of mAb-HSA-GD and mAb-HSA-VX was done using four different peptide haptens (i.e., HSA-GD, HSA-VX, HSA-GB, and HSA-GF) that were chemically synthesized to afford the same phosphonyl adduct as the actual nerve agents (Fig. 3). The nonphosphorylated decapeptide (i.e., HSA-NP) was also prepared and used for comparison. As shown in Fig. 5, ELISA data showed that mAb-HSA-GD had the greatest affinity for HSA-GD, while mAb-HSA-VX had the greatest affinity for HSA-VX. Compared with other mAbs obtained, mAb-HSA-GD showed...
selective recognition of HSA-GD and HSA-GF and to a much lesser extent the other HSA-OP adducts (i.e., GD > GF > GB > VX). mAb-HSA-GD did not recognize the HSA peptide that did not contain a phosphoryl group on Tyr 411 (i.e., HSA-NP). mAb-HSA-VX selectively recognized HSA-VX compared with the other HSA-OP peptide adducts (i.e., VX > GB > GF > GD). mAb-HSA-VX did not recognize the HSA peptide that did not contain a phosphoryl group (i.e., HSA-NP). The $K_d$ values for mAb-HSA-GD and mAb-HSA-VX were 30 nM and 0.4 nM, respectively (Fig. 5). The affinity of mAb-HSA-GD for HSA-GD or mAb-HSA-VX for HSA-VX, respectively, was confirmed with Western blot studies. As shown in Fig. 6, the immunoreactivity of mAb-HSA-GD for HSA-GD was observed for mAb-HSA-GD against HSA-GA (HSA-tabun) or HSA-NP at low immunogen levels.

In good agreement with the ELISA data, on the basis of Western blot analysis, the immunoreactivity of mAb-HSA-VX was HSA-VX >> HSA-GB > HSA-GF. No significant immunoreactive signal was observed for mAb-HSA-VX against HSA-GD, HSA-GA, or HSA-NP at low immunogen levels. The affinities of mAb-HSA-VX and mAb-HSA-GD suggested that the antibodies could have utility in the selective analysis of exposure to different nerve agent OPs because the two antibodies readily detected GD and VX, respectively, and to a lesser extent, GB and GF.

Because the antibodies were designed to be used in detection and analysis of OP exposure in vivo, we sought to increase the sensitivity of detection and develop a robust method to detect OPs at very low concentrations. To test the lowest level of detection of the antibodies, we conducted a titration experiment by serially diluting an authentic HSA-GD protein standard. Previously, the amount of phosphorylated HSA Tyr 411 adducted with GD had been determined by tryptic digestion and LC-MS studies and was available as a standard (Li et al., 2007; O. Lockridge, personal communication). An HSA-VX standard protein was not available, so the study was largely conducted with mAb-HSA-GD. As shown in Fig. 7B, the lowest level of detection of the GD adduct on Tyr 411 of HSA was on the order of $10^{-12}$ g with a femtogram-sensitive HRP substrate in a Western blot analysis format.

Because mAb-HSA-GD showed superb sensitivity in detecting GD-adducted serum albumin on the basis of serially diluted standard protein samples in PBS, we extended the studies to detection of OP-adducted serum albumin isolated from animal blood samples. Because it was estimated that only 1–2% of serum albumin would form Tyr 411 adducts during OP exposure (Ding et al., 2008), to recognize the serum albumin-GD adduct in blood plasma, it was required that the antibody not only have the ability to detect low levels of target protein as shown in Fig. 7, but also have the ability to distinguish serum albumin–GD adduct in the presence of large quantities of unmodified serum albumin. We therefore conducted experiments to simulate these conditions in blood plasma before embarking on more complex detection in blood samples from animals exposed to OPs. In these experiments, OP-adducted HSA standards previously quantified by LC-MS (O. Lockridge, personal communication) were serially diluted in the presence of unmodified HSA. The total protein in each dilution experiment was 1 μg, as shown by Ponceau staining in Fig. 7C. When an HSA solution was used in a serial dilution experiment, compared with dilution in PBS, the intensity of the signal was not significantly decreased (e.g., Fig. 7, B and C), and the antibody recognized far less than 1% of a theoretical amount of HSA-GD adduct in solution. For example, in an experiment with 1.6% of the serum albumin adducted with GD (determined by MS after tryptic digestion) at a typical blood plasma concentration, mAb-HSA-GD readily recognized HSA-GD adduct (i.e., the second lane of Fig. 7B represented a typical blood plasma that had 1–2% of HSA-GD adduct). Thus, using the methodology developed, detection of very low levels of HSA-adducted OPs was feasible even in the presence of large amounts of unmodified HSA.
Phosphonylation of HSA in the Presence of para-Nitrophenyl Esters of Nerve Agents. To study the kinetics of OP adduction of HSA in vitro, we synthesized para-nitrophenyl esters of nerve agent model compounds 18, 19, and 20 (Fig. 8). The para-nitrophenol (pNP) leaving group of these OP esters liberated when the para-nitrophenyl esters covalently modified HSA was continuously monitored at 400 nm. From the time course studies the half-life of adduction of covalently modified HSA was continuously monitored at 400 nm. We also examined the stereoselectivity of the adduction. Over the course of a 20-day incubation, based on the area of the product formation curves, Sp-GD methylthiocholine showed approximately a 1.3-fold greater stereoselectivity in HSA adduct formation recognition by mAb-HSA-GD compared with the amount of adduct formed from Rp-GD in the in vitro system (Fig. 9). Although the total amount of OP adduct recognized by the mAb-HSA-GD was significantly different for each OP enantiomer, the stability of the adducts appeared to be similar.

Immunodetection of GD in Human Blood Plasma Samples. As described above, we characterized mAb-HSA-GD and mAb-HSA-VX with highly purified OP-adducted HSA samples in vitro. mAb-HSA-GD was characterized for its ability to recognize nerve agent–adducted HSA in the presence of HSA. We next examined whether mAb-HSA-GD could recognize authentic HSA-adducted GD standards in the presence of human blood plasma. As shown in Fig. 10, mAb-HSA-GD detected extremely low levels of HSA-GD in the presence of human plasma that was treated with 0.18 μM GD for 72 hours. Under similar conditions, no detectable amount of OP adduct could be detected by LC-MS under standard tryptic digestion and analysis conditions (O. Lockridge, personal communication). After analysis by densitometry, the level of HSA-GD detected in human plasma treated with 0.18 μM GD was determined to be 0.1 ng/ml using an HSA-GD protein standard as a positive control.
Immunodetection of GD Adducts in Rat Serum Albumin from Plasma of Rats Administered GD. We examined whether mAb-HSA-GD could detect GD-adducted albumin in the presence of blood samples from animals treated with authentic nerve agents. Rat blood plasma samples were prepared from rats 72 hours after the animals had been administered GD s.c. at 1.2-fold the LD$_{50}$ for GD. As shown in Fig. 11, four different samples from plasma isolated from the blood of rats treated with GD (lanes 2, 3, 5, and 6) showed immunoreactive bands for GD–rat serum albumin (RSA-GD) with mAb-HSA-GD on the basis of Western blot analysis. Samples that showed no OP-adducted RSA signal (lanes 1 and 4) were derived from rats that were treated with PBS. Purified RSA and HSA were used as negative control samples in the same Western blot analysis (lanes 7 and 8). The positive control samples used in characterization of mAb-HSA-GD in vitro (Fig. 7) were also used as positive controls in the Western blot and a quantification standard (lane 9). The results showed that mAb-HSA-GD was suitable for low-level detection and sensitive analysis of authentic biologic samples of GD-adducted blood from rats. The RSA-GD detected was quantified by densitometry using HSA-GD as a standard to be 5.4 ± 1.1 ng/ml plasma.

Immunodetection of VX Adducted to RSA from Plasma of Rats Administered VX. We examined the plasma isolated from rats administered authentic VX with mAb-HSA-VX to detect VX-adducted serum albumin. In a similar fashion to the detection of GD-adducted RSA from rats administered GD described above, VX-adducted RSA was detected in plasma samples from rats treated with VX. As shown in Fig. 12, blood plasma samples isolated from rats 72 hours after administration of VX s.c. at 0.8-fold the LD$_{50}$ showed immunoreactive RSA-VX adducts. The selective immunodetection showed that VX adducted on Tyr 411 of RSA detected was quantified by densitometry using HSA-GD as a standard and was observed to be 5–50 ng/ml (quantification was done by comparing the VX band intensity in the Western blot with the HSA-GD band on the same blot because no independently verified HSA-VX was available).

In addition to GD and VX detected in RSA from rats administered the actual nerve agents, mAb-HSA-GD was able to detect GD-adducted serum albumin in the presence of plasma 30 minutes after African green monkeys were administered authentic GD (15 µg/kg). In addition, mAb-HSA-VX immunodetected VX-adducted serum albumin from the plasma of guinea pigs 72 hours after administration of VX s.c. at 0.6–0.8 × LD$_{50}$ (Supplemental Fig. 1). The immunodetection of GD adducted to monkey serum albumin under these conditions was determined to be 1.8 ± 0.4 ng/ml plasma.

Cholinesterase Activity Inhibition by GD Model Compound. One of the acute toxic effects of nerve agents is the cholinesterase inhibition in the blood. We therefore investigated the time course of cholinesterase inhibition with a GD model compound Rp-GD tertiary amine (Barakat et al., 2009) and compared it to one reported for GD. As shown in Fig. 13A, 80% of the functional activity of cholinesterase in rat blood plasma was inhibited by Sp-GD model compound at the 4-hour time point, but fully recovered to a normal level after 48 hours. At the 72-hour point, the cholinesterase activity was completely recovered to a normal level (Fig. 13B) while RSA-GD was still readily detectable (Fig. 11). Comparison of data for GD [i.e., 0–24 hours (Geller et al., 1987) and up to 10 days (Jovic, 1974)] with the Rp-GD model compound showed similar results in inhibition of cholinesterase functional activity (Fig. 13).

**TABLE 2**

<table>
<thead>
<tr>
<th>Nerve Agent Model Compound</th>
<th>Half-life (hr)</th>
<th>$k$</th>
<th>$k^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>para-Nitrophenyl-GB 18</td>
<td>10.9 ± 5</td>
<td>0.06 ± 0.02</td>
<td></td>
</tr>
<tr>
<td>para-Nitrophenyl-GB 19</td>
<td>6.1 ± 4</td>
<td>0.11 ± 0.07</td>
<td></td>
</tr>
<tr>
<td>para-Nitrophenyl-VX 20</td>
<td>16.4 ± 9</td>
<td>0.04 ± 0.02</td>
<td></td>
</tr>
</tbody>
</table>

GB, sarin; GD, soman; HSA, human serum albumin; VX, (S)-[2-(diisopropylamino)ethyl]-O-ethyl methylphosphonothioate. (Supplemental Fig. 1). The immunodetection of VX adducted to monkey serum albumin under these conditions was determined to be 1.8 ± 0.4 ng/ml plasma.

**Fig. 8.** Chemical structures of para-nitrophenyl esters of nerve agent as photometric model compounds. (A) Structure of para-nitrophenyl-GB. (B) Structure of para-nitrophenyl-GD. (C) Structure of para-nitrophenyl-VX.

**Fig. 9.** A plot of the amount of GD nerve agent model compound stereoisomers adducted to HSA as a function of time based on quantification of immunoactivity with mAb-HSA-GD. The amount of HSA-GD protein adduct was determined by Western blot analysis. After addition of nerve agent model compound Sp-GD or Rp-GD methylthiocholine to HSA, the incubation was sampled over time and compared with an HSA-GD-adducted protein standard. Quantification was done by densitometry. ○, Sp-GD methylthiocholine; □, Rp-GD methylthiocholine.
Discussion

Herein we describe the characterization of nerve agent– or nerve agent model compound–adducted serum albumin from large and small animals using mAbs raised against nerve agent–phosphonylated peptides of HSA. As shown in Fig. 13, nerve agent model compounds showed a similar time course of action compared with the actual nerve agent (i.e., Sp-GD versus GD itself), albeit at different doses. Consequently, data using nerve agent model compounds provide a useful approach in advance of studies using the actual nerve agents. The mAbs obtained showed remarkable sensitivity and selectivity in detecting phosphorylated serum albumin from both in vitro and in vivo samples. As shown in Fig. 7, A and B, in the case of in vitro samples, mAb-HSA-GD was able to recognize as little as 100 pg of HSA-GD. In contrast, detection of tryptic digests of a similar sample has a limit of detection of 250 ng (Ding et al., 2008). The mAbs described herein are remarkably selective and can distinguish phosphorylated HSA from nonphosphorylated HSA. Even minor structural changes of nerve agent–HSA adducts could be distinguished. As shown in Fig. 7B, mAb-HSA-GD was able to recognize HSA-GD in the presence of nonphosphorylated HSA at as low as a 213:1 molar ratio dilution without losing any sensitivity (i.e., compared with identical fold-dilutions in the presence of PBS or HSA solution). In the case of in vivo samples, mAb-HSA-GD was capable of detecting exposure to nerve agent model compounds or actual nerve agents from biologic samples at extremely low levels that were not detectable by MS methods (Fig. 10) (O. Lockridge, personal communication). Animals treated with nerve agents (i.e., GD or VX) at sublethal doses afforded plasma that showed immunoreactivity to the corresponding adduct at Tyr 411 of serum albumin.

Currently available approaches for the detection of OP exposure including functional enzyme assays and MS-based approaches have certain advantages and drawbacks (Worek et al., 2005; Thompson et al., 2010). For example, assays of

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Fig. 10. Western blot analysis of human blood plasma samples treated with GD. (Top) Western blot of human blood plasma samples treated with GD (0.18 μM) for 72 hours and probed with mAb-HSA-GD. (Bottom) Ponceau staining of the PVDF membrane used in the Western blot shows an equal amount of serum albumin. Lane 1, untreated human plasma (1 μl); lane 2, human plasma (1 μl) treated with 0.18 μM GD; lane 3, positive control protein HSA-GD-adducted protein standard (1 μg).

Fig. 11. Western blot using mAb-HSA-GD of plasma samples from rats treated with GD (1.2 × LD₅₀) for 72 hours. (A) Lanes 1 and 4, plasma (1 μl) from untreated rats; lanes 2, 3, 5, and 6, plasma from rats treated with GD; lane 7, untreated RSA (1 μg); lane 8, HSA (1 μg); lane 9, a standard of 1 μg of GD-adducted HSA (independently characterized by MS. (B) Ponceau staining of an identical gel as blot A run in parallel that showed equal protein loading of serum albumin.

Fig. 12. Western blot using mAb-HSA-VX of plasma samples from rats treated with VX (0.8 × LD₅₀) for 72 hours. (A) Lanes 1 and 3, plasma (1 μl) from untreated rats; lanes 2 and 4, plasma (1 μl) from treated rats. (B) Ponceau staining of an identical gel as blot A run in parallel that showed equal protein loading of serum albumin.

Fig. 13. Time course for inhibition of rat plasma cholinesterase functional activity by nerve agent model compound Sp-GD tertiary amine (Barakat et al., 2009) or GD after administration to rats. (A) Comparison of inhibition of rat plasma cholinesterase by nerve agent model compound Sp-GD (○) (250 μg/kg, i.p.) and GD (□) (88 μg/kg, i.p.) (modified from Geller et al., 1987). (B) Comparison of time course (0–10 days) for inhibition of rat plasma cholinesterase activity by nerve agent model compound Sp-GD tertiary amine (□) (250 μg/kg, i.p.) (Barakat et al., 2009) and GD (○) (25.5 μg/kg, i.p.) (modified from Jovic, 1974).
functional activity of AChE and BChE are quick and efficient but do not provide information about the type of nerve agent that is present and inhibiting the enzyme. MS-based approaches to detect protein adducts of OPs are quite sensitive but require expensive equipment and well trained operators (Worek et al., 2005). The approach described herein employs a biologic method using immunodetection that is efficient and robust.

Nerve agent adducts of serum albumin were chosen to develop immunodetection because previous studies of BChE showed that the phosphorylated peptide surrounding the active-site serine of BChE is prone to elimination and affords the corresponding dehydroalanine-containing peptide (Masson and Lockridge, 2010). The phosphorylated peptide that corresponds to the region surrounding Tyr 411 of HSA shows significant stability, and the adduct is not prone to elimination (Masson and Lockridge, 2010). The antibodies described herein, mAb-HSA-GD and mAb-HSA-VX, can distinguish phosphorylated Tyr 411 peptides from nonphosphorylated peptides. In particular, the antibodies selectively recognized phosphorylated Tyr 411 on serum albumin, (i.e., anti-HSA-GD recognizes GD-adducted HSA at Tyr 411 and anti-HSA-VX selectively recognizes VX-adducted HSA at Tyr 411) in Western blot analysis. To our knowledge, these antibodies are the first antibodies that can selectively recognize defined phosphorylated peptides of protein samples from in vivo serum samples.

The specificity of the antibodies reported herein was achieved using a defined epitope approach to raise antibodies that could distinguish native and phosphorylated peptide. Previously, a report showed that antibodies could distinguish phosphorylated AChE and inhibited AChE using this strategy (George et al., 2003), but despite their utility, these polyclonal anti–phosphorylated AChE antibodies did not have the ability to distinguish different OP adducts. The results of that report did suggest that it was possible to raise antibodies to individually recognize specific modifications of the dianionic phosphorus of OP-added proteins (George et al., 2003).

Another strategy reported in the literature to raise antibodies to selectively recognize OPs is to raise antibodies against different epitopes of the same hapten (Johnson et al., 2005). Although antibodies raised against OPs following this strategy did bind to different OP adducts with some affinity, their binding to the desired OP-adducted protein was reportedly relatively weak and not selective because most of the antibodies described had IC50 values to their desired antigen in the range of 10−4 to 10−6 M (Hunter and Lenz, 1982; Hunter et al., 1982, 1985, 1990; Brimfield et al., 1993; Johnson et al., 2005). For example, antibodies raised against GD that was conjugated to the carrier protein bovine serum albumin had an IC50 value of 10−4 M to its antigen (Lenz et al., 1992). Antibodies raised against GD that were conjugated to carrier protein KLH had a reported IC50 value of 10−6 M to its antigen (Hunter and Lenz, 1982). Compared with the mAbs described herein, the previously reported antibodies to phosphorylated haptens may be useful in qualitative analysis of biomedical samples, but may have less utility in quantitative studies.

Using a defined epitope approach, mAb-HSA-GD and mAb-HSA-VX, described herein, showed excellent detection of phosphorylated serum albumin using Western blot analysis across different species. The cross-species detection is undoubtedly due to the conserved nature of the region in the serum albumin protein sequence used in the hapten.

After exposure to nerve agents, numerous protein-OP adducts are formed in the blood, including cholinesterases and serum albumin. As shown in Fig. 13, following nerve agent model compound or actual nerve agent administration, rat cholinesterase functional activity is largely abrogated at 24 hours but functional activity is recovered to normal levels after 48 hours. Therefore, cholinesterase functional activity can be used as a biomarker in OP poisoning in the short term, but cholinesterase functional activity cannot be reliably used as a biomarker for OP exposure much beyond 48 hours. In contrast, serum albumin–GD adducts from animals exposed to nerve agents were still readily detectable by mAb-HSA-GD 72 hours after exposure, well after cholinesterase functional activity recovered back to normal levels. These data show that the detection of HSA-GD is a more robust biomarker than cholinesterase functional activity at longer time points post-administration. The availability of mAbs described herein also affords detection of nerve agent–adducted serum albumin and provides structural information to identify the chemical nature of the agent possibly long after the subject has been exposed to a nerve agent. In fact, some of the samples used in the studies described herein had been stored at −80°C for 3 years.

It is known that GD contains two centers of chirality. Although it has been reported that stereoisomers of GD have differential effects on their toxic potency in vivo, the potency of each isomer has largely been studied in detail for stereoselectivity of cholinesterase inhibition (Benschop and De Jong, 1988). Little has been reported about the reaction of GD stereoisomers with serum albumin. As shown in Fig. 9, mAb-HSA-GD stereoselectively recognized the Sp-GD isomer in preference to the Rp-GD isomer adducted to HSA. The result may suggest that Sp-GD more avidly binds to HSA compared with the Rp-GD isomer or it may suggest that the mAb-HSA-GD stereoselectively distinguishes GD–serum albumin adducts. If the nerve agent model compounds recapitulate the action of actual nerve agents (albeit at greater doses), the observation regarding stereoselective GD–serum albumin adduction may have some consequences for deconvoluting the mechanism of toxicity of GD (and other nerve agents). In conclusion, using a defined epitope approach, we obtained two mAbs that selectively recognize GD- or VX-adducted HSA at Tyr 411. The two mAbs showed unprecedented sensitivity and selectivity in detection of nerve agent biomarkers both in vitro and in vivo. The mAb approach presented herein provides a powerful tool in the detection of biomarkers for nerve agent OP detection.

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Authorship Contributions

 Participated in research design: Chen, Zhang, Cashman.
 Conducted experiments: Chen, Zhang, Cashman.
Contributed new reagents or analytic tools: Lumley.
Performed data analysis: Chen, Zhang, and Cashman.
Wrote or contributed to the writing of the manuscript: Chen, Zhang, Lumley, Cashman.

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