Chimeric Plantibody Passively Protects Mice against Aerosolized Ricin Challenge

Erin K. Sully, Kevin J. Whaley, Natasha Bohorova, Ognian Bohorov, Charles Goodman, Do H. Kim, Michael H. Pauly, Jesus Velasco, Ernie Hiatt, Josh Morton, Kelsi Swope, Chad J. Roy, Larry Zeitlin, Nicholas J. Mantis

Division of Infectious Disease, Wadsworth Center, New York State Department of Health, Albany, New York, USA; Mapp Biopharmaceutical, Inc., San Diego, California, USA; Kentucky BioProcessing, Owensboro, Kentucky, USA; Infectious Disease Aerobiology, Division of Microbiology, Tulane National Primate Research Center, Covington, Louisiana, USA; Department of Biomedical Sciences, School of Public Health, University at Albany, Albany, New York, USA

Recent incidents in the United States and abroad have heightened concerns about the use of ricin toxin as a bioterrorism agent. In this study, we produced, using a robust plant-based platform, four chimeric toxin-neutralizing monoclonal antibodies that were then evaluated for the ability to passively protect mice from a lethal-dose ricin challenge. The most effective antibody, c-PB10, was further evaluated in mice as a therapeutic following ricin exposure by injection and inhalation.

Ricin toxin is a member of the medically important A-B family of plant and bacterial ribosome-inactivating proteins (RIPs) (1). In its mature form, ricin is a 65-kDa heterodimeric glycoprotein that is a natural constituent of the seeds of the castor plant (*Ricinus communis*). The 267-amino-acid A subunit (RTA) of ricin is an RNA N-glycosidase that depurinates a conserved adenine residue within the so-called sarcin-ricin loop (SRL) of eukaryotic 28S rRNA, which is required for activation of the elongation factor EF-Tu (2, 3). RTA is linked via a single disulfide bond to the 262-amino-acid B subunit (RTB), a 262-amino-acid lectin that is specific for glycoproteins and glycolipids terminating in galactose and N-acetylgalactosamine (Gal/GalNAc). In addition to its role in attachment, RTB also mediates the retrograde transport of ricin to the trans-Golgi network (TGN) and endoplasmic reticulum (ER), where RTA is ultimately delivered across the ER membrane and into the cytoplasm (4, 5). The extraordinary capacity of RTA to inactivate ribosomes (*k*<sub>cat</sub> 1,500/min) makes ricin one of the most potent known RIPs (6, 7). For example, in rodents and nonhuman primates, the 50% lethal dose (LD<sub>50</sub>) of ricin by injection is approximately 5 µg/kg, while the LD<sub>50</sub> of ricin by inhalation is estimated to be as low as 3 µg/kg (8). The respiratory mucosa is especially sensitive to ricin, as even trace amounts of toxin are known to elicit widespread necrosis in the airways and alveoli, peribronchovascular edema, mixed inflammatory cell infiltrates, and massive pulmonary alveolar flooding (9–15).

The U.S. Departments of Defense (DOD) and Health and Human Services (HHS) have ongoing initiatives to develop antibody-based products capable of providing passive protection against systemic and mucosal ricin exposure (16, 17). In addition to protective efficacy, issues related to platform technology, scalability, and speed of manufacturing will ultimately dictate which product(s) will be pursued for advanced development and are consistent with the unique needs for biodefense. In this regard, the *Nicotiana benthamiana*-based rapid antibody-manufacturing platform (RAMP) provides the potential for extremely fast and high-yield monoclonal antibody (MAb) production that is already proven to be applicable to biodefense (18–21). The technology entails mass infiltration of mature *Nicotiana* plants with an *Agrobacterium tumefaciens* suspension carrying T-DNA encoding viral replicons and results in high MAb recovery from original DNA constructs within days (22). The *Nicotiana*-based RAMP is being used for manufacturing recombinant MAb s or “plantibodies” for clinical studies under good manufacturing practices (GMP) (20, 23).

In this study, we sought to exploit the *Nicotiana*-based RAMP for the purpose of identifying and advancing potential immunotherapeutic agents for ricin toxin. In a proof-of-concept study, we recently characterized chimeric GD12 (c-GD12), a partially humanized murine monoclonal antibody (MAb) directed against an immunodominant, linear B cell epitope on RTA (24). c-GD12, produced in *N. benthamiana*, protected mice from the effects of a systemic ricin toxin challenge when administered before or up to 4 h after ricin injection. However, for as of yet underdetermined reasons, c-GD12 ultimately proved difficult to express in high quantities via traditional cell culture or the RAMP technologies. For that reason, we have now extended our studies to include four additional recently identified toxin-neutralizing MAb s (Table 1), two MAb s (SyH7 and PB10) directed against RTA (25, 26), and two MAb s (SyH3 and JB4) directed against RTB (27, 28).

The murine V<sub>H</sub> and V<sub>λ</sub> domains of SyH7, PB10, SyH3, and JB4 were amplified by PCR from cDNA derived from the respective murine B cell hybridomas (26–28). PCR amplicons were sequenced, and consensus contigs for each domain were generated based on the Kabat and International ImMunoGeneTics (IMGT) databases (29). The codon-optimized V<sub>H</sub> and V<sub>λ</sub> regions of each MAb were then synthesized commercially (GeneArt; Life Technologies, Grand Island, NY) and fused to human IgG1 and κ constant regions as described previously (24). Mouse-human chimeric antibodies were produced using the *N. benthamiana*-based RAMP. For this study, production was done with a transgenic N.
benthamiana line devoid of xylosyl transferase and fucosyl transferase activities, which results in immunoglobulins with N-glycans that are mammalian and generally more homogeneous than those produced in mammalian cell lines used for MAb production (e.g., CHO and NS0) (30).

All four of the chimeric antibodies were successfully expressed and purified from the N. benthamiana-based system. Briefly, 24- to 26-day-old plants were vacuum infiltrated with A. tumefaciens strains carrying viral vectors encoding the cognate VH and VL of SyH7, PB10, SylH3, and JB4. Seven days postinfiltration, leaf tissues were extracted, clarified using a plate-and-frame filter press (Ertel-Alsop), and then subjected to a MabSelect SuRe protein A column (GE Healthcare Biosciences, Pittsburgh, PA), Capto Q (GE Healthcare), and final polishing via a ceramic hydroxyapatite type II (CHT) column (80 μm) (Bio-Rad). Chimeric MAbs were fully assembled, as determined by SDS-PAGE, and had <1% aggregate, as determined by high-pressure liquid chromatography (HPLC)-size exclusion chromatography (SEC) (see Fig. S1 in the supplemental material). Yields were in the range of 100 to 200 mg per kg of fresh leaf biomass. Antibodies were placed into an appropriate formulation buffer and sterile filtered into crystal zenith vials (West Pharma, Exton, PA) and stored at −80°C.

We found that, as determined by enzyme-linked immunosorbent assay (ELISA), the chimeric MAbs retained antigen specificity and apparent affinities (50% effective concentration [EC50]) compared to the parental murine MAbs (see Fig. S2 in the supplemental material). We next performed ricin toxin-neutralizing assays using Vero cells, as previously described (24). With respect to functional activity, all four chimeric MAbs (c-MAbs) had 50% inhibitory concentrations (IC50s) that were indistinguishable or, in the case of c-PB10 and c-SylH7, slightly better than their murine counterparts (Table 1 and Fig. 1). c-PB10 had the lowest IC50.

**TABLE 1 Characteristics of murine and chimeric MAbs used in this study**

| MAb | Target | $K_D$ (M) | m-IC50 (μg/ml) | c-IC50 (μg/ml) | No. of mice survived/no. of mice challenged | Reference |
|-----|--------|----------|---------------|---------------|------------------------------------------|-----------|
| PB10 | RTA     | $4 \times 10^{-4}$ | 0.015 | 0.03 | 10/10 | 26 |
| SyH7 | RTA     | $4 \times 10^{-4}$ | 0.125 | 0.10 | 3/10 | 26 |
| SylH3 | RTB    | $3.3 \times 10^{-9}$ | 1.8 | 1.8 | 5/10 | 27 |
| JB4  | RTB     | $2 \times 10^{-10}$ | 1.1 | 1.1 | 10/10 | 28 |

| a | Equilibrium dissociation constant as determined by BIAcore analysis and reported previously. |
| b | Inhibitory concentration (IC50) of murine (m) or chimeric (c) MAbs, as determined in Vero cytotoxicity assay. |
| c | BALB/c mice were administered 20 μg of each MAb and then challenged with ricin (10× LD50) 24 h later, as described in the text. |

**FIG 1** In vitro neutralizing capabilities of chimeric MAbs. Dilutions of indicated murine (m) or chimeric (c) MAbs were mixed with ricin (10 ng/ml), added to Vero cells for 2 h, washed, and then incubated with Dulbecco modified Eagle medium (DMEM) for an additional 48 h, after which cell viability was assessed. (A) SyH7, (B) PB10, (C) SylH3, (D) JB4. Treatments were performed in triplicate. Cells treated with media only were used as controls, with 100% viability. Variability of cell viability around 100% (±10%) is not uncommon in this assay.
of the four c-MAbs and its IC50 was >5-fold lower than that of c-GD12 (24).

We next compared c-PB10, c-SyH7, c-SylH3, and c-JB4 to determine which chimeric MAb was the most effective at ricin neutralization in vivo. BALB/c mice received 20 μg of chimeric MAb 24 h prior to being subjected to a 10 × LD50 ricin challenge (~2 μg per mouse) by the i.p. route 24 h before the animals were challenged with 10 × LD50 ricin, as described for panel A. Survival was monitored over a 5-day period. (D and E) c-PB10 (25 μg/mouse) was administered to mice (n = 8 mice per group) at indicated time points before or after the animals were challenged with 10 × LD50 ricin. Survival (D) and blood glucose levels (E) were recorded as described above. Data are displayed as the mean ± SEM. Statistical significance was determined using the Student t test. **, P < 0.01.

FIG 2 Evaluation in mice of chimeric plantibodies in conferring passive immunity to ricin toxin. BALB/c mice (female, 6 to 8 weeks of age) were purchased from Taconic Labs (Hudson, NY) and housed under conventional, specific pathogen-free conditions. (A and B) Chimeric MAbs were administered to mice (n = 10 mice per group) by intraperitoneal (i.p.) injection 24 h prior to i.p. challenge with 10 × LD50 ricin. Survival (A) was monitored over a period of 6 days. Morbidity (B) was assessed based on the onset of hypoglycemia, as done previously (36). (C) c-PB10 and c-JB4 were administered at indicated amounts to mice (n = 6 mice per group) by the i.p. route 24 h before the animals were challenged with 10 × LD50 ricin, as described for panel A. Survival was monitored over a 5-day period. (D and E) c-PB10 (25 μg/mouse) was administered to mice (n = 8 mice per group) by the i.p. route 24 h before the animals were challenged with 10 × LD50 ricin. Survival (D) and blood glucose levels (E) were recorded as described above. Data are displayed as the mean ± SEM. Statistical significance was determined using the Student t test. **, P < 0.01.

To more fully compare the efficacy of c-PB10 and c-JB4, we performed a dose step-down experiment in which groups of BALB/c mice were administered the chimeric MAbs at 5 μg (~0.2 mg/kg), 2.5 μg, 1.25 μg, or 0.6 μg per mouse 24 h prior to being challenged with 10 × LD50 ricin (Fig. 2C). c-PB10 demonstrated a dose-dependent capacity to protect mice against ricin intoxication; 100% protection was achieved with 5 μg, 75% protection with 2.5 μg, and 40% protection was achieved with 1.25 μg, while...
Before ricin exposure, at the time of ricin exposure, or 4 h after ricin challenge (Fig. 2D and E). Control mice that did not receive c-PB10 at 2 or 3 h after toxin challenge survived. Mice treated with c-PB10 at the 2- to 3-h time points experienced a transient reduction in blood glucose levels between 24 and 48 h, but those levels trended to baseline levels thereafter. Administration of c-PB10 4 or 5 h after ricin intoxication challenge proved sufficient to protect 90% of the test animals, whereas antibody treatment at 6 h afforded only 30% protection. Reductions in blood glucose levels were correspondingly severe in the mice given c-PB10 at 4 to 6 h after ricin challenge. Overall, these data indicate that the therapeutic window in mice for antibody-mediated rescue from ricin intoxication is between 3 and 4 h, a result which is in accordance with what we observed previously with c-GD12 (24).

The respiratory tract is the compartment most vulnerable to ricin intoxication, and assessing the efficacy of potential therapeutic agents to protect against toxin exposure by this route is critical (8). To assess c-PB10 as a protectant against aerosolized ricin, BALB/c mice were administered a single intraperitoneal injection of ~250 μg (10 mg/kg) of c-PB10 at one of three time points: 24 h before ricin exposure, at the time of ricin exposure, or 4 h after ricin exposure. As it is not known what concentrations of serum antibody are required to confer protective immunity to aerosol challenge, c-PB10 was administered to mice at a dose (i.e., 10 mg/kg) that would be considered toward the upper limit of what would be acceptable for a therapeutic antibody. Ricin exposure was performed in a whole-body inhalation exposure chamber (10). Aerosols were generated into the exposure system using a three-jet Collision nebulizer (BGI, Inc., Waltham, MA) operated at 20 pressure square inch gauge with an output flow of 7.5 l/min. Time-of-flight aerodynamic particle size distribution measured prior to the exposure using an aerodynamic particle sizer (APS) (model 3321; TSI, Inc., St. Paul, MN) indicated a highly respirable particle size distribution (mass median aerodynamic diameter [MMAD], 1 μm; geometric standard deviation (σ_g) = 1.4). Control mice that were exposed to ricin but not treated with c-PB10 experienced a rapid decrease in core body temperature and body weight, and then died within 48 h (Fig. 3A through C). Conversely, mice that received c-PB10 (irrespective of the time point of c-PB10 administration) experienced weight loss and a slight decrease in core body temperature but ultimately survived challenge. Histopathologic analysis of lung tissues revealed that systemic c-PB10 administration greatly damped neutrophil cell infiltration and toxin-induced tissue damage in ricin-challenged mice compared to control toxin-challenged mice (Fig. 3D). These experiments demonstrate that c-PB10 is sufficient in mice to serve as a protectant and therapeutic agent against aerosolized ricin challenge. However, we recognize that more comprehensive dose-response studies are necessary to determine the effectiveness of c-PB10 at lower concentrations. Such experiments are currently planned but are beyond the scope of this current article.
In summary, we generated mouse-human chimeric IgG1 derivatives of four ricin toxin-neutralizing MAbs and then expressed and purified them using a Nicotiana-based platform. The chimeric plantibodies, two directed against RTA (c-PB10 and c-SyH7) and two against RTB (c-JB4 and c-SyH3), were tested for their capacity to neutralize ricin in vitro and in vivo. We found that c-PB10 was the most effective of the four chimeric MAbs (and more effective than the previously described c-GD12) in that (i) it had the lowest IC_{50} in a Vero cell-based toxin-neutralizing assay, (ii) it was sufficient to passively protect mice against systemic and aerosol toxin challenge, and (iii) it had demonstrable therapeutic potential, in that it rescued mice from the effects of ricin when administered up to 3 to 4 h after toxin challenge. In addition, preliminary studies indicated that scale-up production of c-PB10 using the N. benthamiana-based RAMP is readily achievable (J. Morton and L. Zeitlin, unpublished observations). Although not investigated in this study, we have tested c-PB10 in combination with the other three chimeric MAbs; no enhanced toxin-neutralizing activity was observed in any of the combinations tested, indicating that c-PB10 alone affords maximal antiricin activity in vitro (E. Sully and N. Mantis, unpublished results). Collectively, these data make c-PB10 an ideal candidate for further development as a fully humanized immunoprotectant against ricin toxin, either as a stand-alone countermeasure or as part of a cocktail with other category B toxin-specific antibodies.

While active vaccination of mice and rabbits has been shown to elicit serum antibody titers that are sufficient to confer immunity against aerosolized ricin challenge, our study is the first to demonstrate that immunity in the respiratory tract can be achieved by MAb therapy (14, 31, 32). PB10 is known to recognize a conformational immunodominant linear epitope (residues 98 to 106) -based RAMP is readily achievable (J. Doebler et al., manuscript in preparation). We postulate, therefore, that in the aerosol challenge model, c-PB10 likely neutralizes ricin on the epithelial surfaces of the respiratory tract, although it remains to be determined whether c-PB10 is found in mucosal secretions at the time of challenge and, if so, whether transudation or active transport is responsible for delivery of the antibody into this compartment (35). Therefore, moving forward it will also be important to resolve whether a secretory IgA (sIgA) derivative of c-PB10 affords any benefit over the IgG form of c-PB10 in mucosal protection.

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