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Alga-Produced Cholera Toxin-Pfs25 Fusion Proteins as Oral Vaccines

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Infectious diseases disproportionately affect indigent regions and are the greatest cause of childhood mortality in developing countries. Practical, low-cost vaccines for use in these countries are paramount to reducing disease burdens and concomitant poverty. Algae are a promising low-cost system for producing vaccines that can be orally delivered, thereby avoiding expensive purification and injectable delivery. We engineered the chloroplast of the eukaryotic alga *Chlamydomonas reinhardtii* to produce a chimeric protein consisting of the 25-kDa *Plasmodium falciparum* surface protein (Pfs25) fused to the β subunit of the cholera toxin (CtxB) to investigate an alga-based whole-cell oral vaccine. Pfs25 is a promising malaria transmission-blocking vaccine candidate that has been difficult to produce in traditional recombinant systems due to its structurally complex tandem repeats of epidermal growth factor-like domains. The noncatalytic CtxB domain of the cholera holotoxin assembles into a pentameric structure and acts as a mucosal adjuvant by binding GM1 ganglioside receptors on gut epithelial cells. We demonstrate that CtxB-Pfs25 accumulates as a soluble, properly folded and functional protein within algal chloroplasts, and it is stable in freeze-dried alga cells at ambient temperatures. In mice, oral vaccination using freeze-dried algae that produce CtxB-Pfs25 elicited CtxB-specific serum IgG antibodies and both CtxB- and Pfs25-specific secretory IgA antibodies. These data suggest that algae are a promising system for production and oral delivery of vaccine antigens, but as an orally delivered adjuvant, CtxB is best suited for eliciting secretory IgA antibodies for vaccine antigens against pathogens that invade mucosal surfaces using this strategy.

There is an urgent need to develop vaccines for practical use in developing countries, both in terms of cost and delivery. Infectious and parasitic diseases killed more than 1.7 million children and adolescents in low-income countries in 2008 (1), which represents nearly half of all deaths in that age bracket. In contrast, infectious and parasitic diseases caused 4,500 deaths (4%) in high-income countries in the same age bracket (1). This disparity highlights one of the great challenges in human health today, which is to reduce the disease burden in the poorest regions of the world, especially for the young. Much of the inequality is the result of disparate vaccine coverage in the developing world (2), mainly brought about by the high cost and logistical difficulty of large-scale vaccination campaigns in countries with underdeveloped health infrastructures. Additionally, pathogens for which no traditional vaccine exists, like malaria, disproportionately affect the poorest countries (3), and some have argued that these countries are unable to climb out of poverty because of this disease burden (4). Thus, feasibility of implementation must be considered when developing vaccines for such regions of the world. Heat-stable oral vaccines could overcome the largest obstacles that deter widespread vaccination in low-income countries. Thermostability would eliminate the need for cold-chain storage, and oral delivery would be safer, simpler, and cheaper than injectable vaccines (5–8). Indeed, the National Institutes of Health, the World Health Organization, and others have emphasized the need for needle-free vaccination strategies.

Oral vaccines are not a new idea; indeed, oral vaccines for a few human pathogens, including *Vibrio cholerae* (Dukoral), *Salmonella typhi* (Vivotif), rotavirus (Rotarix and RotaTeq), and polio (Sabin vaccine), are commercially available. Despite these few examples, the potential of oral vaccines remains largely unrealized. The complexity of the mucosal immune system, which must discriminate between detrimental and innocuous antigens in the gut, has slowed the development of oral vaccine candidates (9, 10).

This is especially true for subunit vaccines, because the response to most antigens at mucosal surfaces is one of nonresponsiveness or tolerance (11, 12). Recent studies suggest that this may be overcome by proteins that act as adjuvants at mucosal surfaces. The best-characterized mucosal adjuvants are the ADP-ribosylating enterotoxins from *V. cholerae* (cholera toxin [CT]) and *Escherichia coli* (heat labile toxin [LT]) (13, 14). Other potential mucosal adjuvants include cytokines, which have been investigated for intranasal vaccines (15), and Toll-like receptor (TLRs) or other pattern recognition receptor (PRR) agonists (16–18). Alternatively, antigens could be specifically targeted to microfold (M) cells, which overlay Payer’s patches and are often the site of entry for pathogens or particles from the gut lumen to immune cells (19, 20).

One major challenge for oral vaccines is to overcome antigen degradation by commensal bacteria, proteases, and the acidic stomach environment in order to safely deliver intact antigens and mucosal adjuvants to the gut-associated lymphoid tissue. Plant delivery systems are particularly attractive for this purpose for two reasons. First, oral delivery is restricted to Food and Drug Administration (FDA)-approved organisms that are generally regarded as safe (GRAS) (21). Unlike conventional organisms used for recombinant protein production, GRAS organisms pose little risk of
harmed by viral, prion, toxin, or bacterial contaminants. Second, plant cells have a rigid cell wall that can protect encapsulated vaccine antigens, allowing ambient temperature storage. The cell wall can also provide a layer of protection from proteolysis in the gastrointestinal tract.

Advances in molecular genetics have made recombinant protein production possible in many plants (22); the first example of a plant-based oral vaccine was accomplished in tobacco and potatoes more than 15 years ago (23). Plants also afford an unprecedented economy of scale that results in a robust and inexpensive platform for producing recombinant proteins (24). Oral delivery via an edible plant also eliminates the need for antigen purification and drives costs down even further. Several experimental plant-based oral vaccines elicit an immune response, although not all demonstrate protective efficacy (25). There are inherent drawbacks of terrestrial plants, however, which undoubtedly contribute to the absence of commercially available plant-based vaccines despite a few that have been approved for use (26). First, terrestrial plants grow relatively slowly. It can take years from the initial transformation to validate an antigen from a stable plant line, although newer methods using transient transcription in tobacco allow a more rapid turnaround (27). Both methods often result in antigen production variability between plants. Second, the environmental impact of gene flow or lateral gene transfer remains unclear (28). Thus, from a time and regulatory perspective, transgenic food crops may not be ideal. Lastly, many plants require extensive processing to become palatable, which reduces antigen immunogenicity (29).

The unicellular eukaryotic green alga *Chlamydomonas reinhardtii* is an attractive alternative to terrestrial plants for producing oral vaccines. Eukaryotic green algae are considered GRAS organisms by the FDA (30). *C. reinhardtii* can be rapidly transformed into stable transgenic strains and grown in enclosed bioreactors or outdoor ponds. Importantly, several algae are already grown at commercial scales for consumption (e.g., *Earthrise* and *CyanoTech*), and the interest in algal-based biofuel is further driving innovation for low-cost biomass production. The impending industrial utility of algae is also accelerating the development of genetic tools for metabolic engineering and recombinant protein production as potential coproducts (31). Thus far, *C. reinhardtii* chloroplasts have been used to make vaccine antigens (32, 33), industrial enzymes (34), antibodies (35), human therapeutics (36), and immunotoxins (37). The versatility of the chloroplast is owed to the unique combination of eukaryotic chaperones (38), protein disulfide isomerases (39), and peptidylprolyl isomerases (40) coupled with prokaryotic-like ribosomal machinery (41).

Here, we describe the production and characterization of a complex chimeric protein consisting of the CTβ subunit (CtxB), a mucosal adjuvant, and Pfs25, a malaria transmission-blocking vaccine (TBV) candidate (42), in *C. reinhardtii* chloroplasts. CtxB assembles into a pentameric structure and binds the GM1 ganglioside receptor found on gut epithelial cells, which facilitates an immune response via the gut mucosa but lacks the toxic ADP-ribosylating activity of the CT α subunit (CtxA) (43). Pfs25 is a structurally complex protein found on the surface of sexual stage *Plasmodium falciparum* parasites (44); antibodies to Pfs25 block malaria transmission (45). We previously demonstrated that alga-produced Pfs25 resembles native Pfs25 structure and elicits transmission-blocking antibodies (46). In these studies, we characterized the chimeric CtxB-Pfs25 fusion protein and the immune response in mice that were administered lyophilized whole algae that produce CtxB-Pfs25.

**Materials and Methods**

Plasmid and *C. reinhardtii* strain construction. The peptide sequence of CtxB (Thr23, Asn13, Asp14, UniProt identifier P01556) was reverse translated using the Gene Designer software (DNA2.0, Menlo, CA) using the *C. reinhardtii* chloroplast codon table (www.kazusa.or.jp) as a reference table. The codon-optimized *C. reinhardtii* ctxB nucleotide sequence (Cr.ctxB) was validated by calculating the codon adaptation index (CAI; 0.94) (47) and synthesized by Integrated DNA technologies (San Diego, CA). *C. reinhardtii* codon-optimized *Plasmodium falciparum* surface protein 25 (Cr.pfs25) was previously described (46). Cr.ctxB and Cr.pfs25 were separately amplified and fused together with uracil-specific excision reagent (USER) cloning (48) such that it contained a linker composed of a flexible hinge (G-P-G-P), a furin protease recognition sequence (R-A-R-R) (49), and a BglII restriction site. Flanking Cr.ctxB-pfs25 are Ndel and AgeI restriction sites on the 5′ and 3′ ends, respectively. The chimeric gene (Cr.ctxB-pfs25) was digested with Ndel and AgeI and cloned into the chloroplast transformation vector pJAG15 (46), a pD1-KanR (50) derivative, also digested with Ndel and AgeI. The resulting plasmid, pJAG101, encodes Cr.CtxB-Pfs25 with a C-terminal FLAG affinity tag driven by the psbA promoter and flanked by the psbA 5′ and 3′ untranslated regions for message stabilization. Integration of pJAG101 into the psbA locus of the chloroplast is directed by 5′ and 3′ psbA homology regions and confers kanamycin resistance. All constructs were verified by sequencing (Regeneron, San Diego, CA). Mid-log-phase *C. reinhardtii* strain W1.1 (51) cells were transformed with pJAG101 by particle bombardment using DNA-coated gold particles as previously described (46, 52) to make strain JAG101. Transformants were selected on Tris-acetate-phosphate (TAP) agar plates supplemented with 100 μg/ml kanamycin and propagated with 150 μg/ml kanamycin. Gene-specific primers were used to confirm integration of Cr.ctxB-Pfs25, and homoplasy was confirmed as previously described (36). A *C. reinhardtii* strain that is isogenic to JAG101 (Cr.ctxB-Pfs25), except that it lacks Cr.ctxB-Pfs25 and the kanamycin resistance cassette, was constructed (here referred to as the ΔpsbA strain) using previously described methods to remove the selectable marker (53). The ΔpsbA strain has a seamless deletion that extends from −300 bp to +1,525 bp, relative to the psbA start codon and stop codon, respectively.

**Western blotting of algal lysates.** Initial screens for the Cr.CtxB-Pfs25 protein were performed on homoplasy JAG101 strains. JAG101 and ΔpsbA cultures were grown in liquid TAP medium on a rotary shaker in low light. Cultures were shifted to high light (~5,000 lx) at mid-log phase (~1 × 10⁶ cells/ml), and samples were harvested at various time points after light shift and stored at −20°C. Alga pellets were resuspended in lysis buffer (50 mM Tris [pH 8.0], 400 mM NaCl, 0.1% Tween 20) with protease inhibitor cocktail (Roche, Mannheim, Germany) and lysed by sonication. Lysates were cleared by centrifugation at 30,000 × g and quantitated with the DC Bio-Rad protein assay. Soluble protein samples were prepared in SDS buffer with urea and β-mercaptoethanol (50 mM Tris, 2% SDS, 10% glycerol, 2 M urea, and 10% β-mercaptoethanol), heated at 37°C for 10 min, resolved on RUNCLEAR precast SDS-PAGE gels (Expe- deon, San Diego, CA), and transferred to nitrocellulose membranes. The nitrocellulose membrane was probed with mouse anti-FLAG monoclonal antibodies (MAbs) (Sigma catalog no. F3165) and detected with goat anti-mouse IgG antibodies conjugated to alkaline phosphatase (Sigma catalog no. A1682) using nitro-blue tetrazolium (NBT) and 5-bromo-4-chloro-3-indolylphosphate (BCIP) p-toluidine salt as the substrates in alkaline phosphatase buffer.

**Affinity purification of alga-produced CtxB-Pfs25.** *C. reinhardtii* strain JAG101 (Cr.CtxB-Pfs25) and JAG9 (Cr.Pfs25) were grown in 20-ml photobioreactors constructed from Nalgene carboys (part number 2251-0050) fitted with Nunc bulkhead adapters (part number 6149-002). Photobioreactors were inoculated with 250 ml of stationary-phase JAG101 grown in TAP and grown to mid-log phase in the dark. Mixing...
was achieved by bubbling air through a 0.1-μm-pore-size filter. Cultures were then illuminated with 32-W 6500K T8 fluorescent light bulbs and harvested 24 h later using a Lavin L2 continuous-flow centrifuge (AML Industries, Hatboro, PA) fed by a peristaltic pump. Cells were snap-frozen with liquid nitrogen and stored at −80°C. Cells were resuspended and lysed as described above, followed by the addition of M2 anti-FLAG affinity resin (Sigma-Aldrich) and end-over-end mixing for 2 to 4 h at 4°C. Resin was washed twice with 20 column volumes of lysis buffer or until washes were no longer green, followed by one wash with lysis buffer without Tween 20. Bio-Rad Econo-pac columns were loaded with the washed resin, and bound protein was eluted using 100 mM glycine (pH 3.5) and 400 mM NaCl. Eluted fractions were neutralized with 1 M Tris (pH 8.0) to a final concentration of 50 mM Tris and tested for protein by Western blotting. Fractions containing protein were combined and buffer exchanged and concentrated into phosphate-buffered saline (PBS) using a 10-kDa cutoff Vivaspin centrifugal concentrator (GE Healthcare) and quantitated using the DC Bio-Rad protein assay. Pure protein was prepped with 4× native buffer (Expe) for nonreduced samples or SDS buffer with β-mercaptoethanol for reduced samples and resolved by SDS-PAGE and transferred onto nitrocellulose membranes. The membranes were probed using rabbit anti-CT antibodies (Sigma catalog no. C3062), mouse anti-FLAG MAb (Sigma catalog no. F3165), or anti-Pfs25 (4B7) MAb and detected with goat anti-rabbit IgG (Sigma catalog no. A6367) or goat anti-mouse IgG (Sigma catalog no. A1682) secondary antibodies conjugated to alkaline phosphatase. Bands were visualized as described above. For gel staining, samples were resolved on 4 to 20% gradient SDS-native buffer (Expedeon) for nonreduced samples or SDS buffer with 0.1% Tween 20. Bio-Rad Econo-pac columns were loaded with the washed resin, and bound protein was eluted using 100 mM glycine (pH 3.5) and tested for protein by Western blotting. Fractions containing protein were combined and buffer exchanged and concentrated into phosphate-buffered saline (PBS) using a 10-kDa cutoff Vivaspin centrifugal concentrator (GE Healthcare) and quantitated using the DC Bio-Rad protein assay. Pure protein was prepped with 4× native buffer (Expe) for nonreduced samples or SDS buffer with β-mercaptoethanol for reduced samples and resolved by SDS-PAGE and transferred onto nitrocellulose membranes. The membranes were probed using rabbit anti-CT antibodies (Sigma catalog no. C3062), mouse anti-FLAG MAb (Sigma catalog no. F3165), or anti-Pfs25 (4B7) MAb and detected with goat anti-rabbit IgG (Sigma catalog no. A6367) or goat anti-mouse IgG (Sigma catalog no. A1682) secondary antibodies conjugated to alkaline phosphatase. Bands were visualized as described above. For gel staining, samples were resolved on 4 to 20% gradient SDS-PAGE precast gels (Bio-Rad) and stained using Coomassie blue.

Determination of alga-produced CtxB-Pfs25 accumulation. The percent total soluble protein that constitutes the Cr.CtxB-Pfs25 fraction was determined by enzyme-linked immunosorbent assay (ELISA) as previously described (46). Briefly, soluble lysates from JAG101 and the ΔpsbA mutant were prepared as described above and diluted to 0.5 mg/ml in PBS. A standard curve was prepared by adding known amounts of purified CtxB (Sigma catalog no. C9903) to ΔpsbA lysates. JAG101 and ΔpsbA lysates with purified CtxB were applied to MaxiSorp plates in triplicate, detected with rabbit anti-CT primary antibodies (Sigma catalog no. C3062) and goat anti-rabbit IgG secondary antibodies conjugated to alkaline phosphatase. Plates were then visualized with p-nitrophenyl phosphate disodium salt (PNPP) in 1 M diethanolamine buffer (pH 9.8; Thermo Scientific catalog no. A4937) conjugated to alkaline phosphatase for 2 h at room temperature. Plates were visualized with PNPP and measured at 405 nm using an Infinite M200 Pro plate reader. Figures were assembled in JMP version 9.02.

GM1 ganglioside binding ELISA. MaxiSorp plates (Nunc, Rochester, NY) were coated with 130 μg/ml GM1 ganglioside (Sigma, G7641) in PBS, incubated overnight at 4°C, and blocked with 5% milk in PBS with 0.1% Tween 20 (PBS-T). Total soluble protein lysates from JAG101 and ΔpsbA algae were prepared as described above. Lysates were diluted to 20 μg/ml soluble protein in PBS and applied to the GM1-coated plate for 1 h at room temperature in triplicate. GM1 binding by Cr.CtxB-Pfs25 in lysates was detected with rabbit anti-CT antibodies followed by goat anti-rabbit IgG antibodies conjugated to horseradish peroxidase (HRP; Thermo Scientific). Binding was then visualized with the TMB substrate kit (Thermo Scientific) and read with an Infinite M200 Pro plate reader (Tecan, Switzerland) at 405 nm.

Stability of alga-produced CtxB-Pfs25 in lyophilized algae. The stabilities of Cr.CtxB-Pfs25 in lyophilized algae stored at 4°C, 22°C (room temperature), and 37°C were determined by GM1 ganglioside binding as described above with minor modifications. The JAG101 and ΔpsbA strains were grown in 20-liter photobioreactors as described above and stored at −80°C in aliquots, several of which were lyophilized. Total soluble protein lysates from the JAG101 and ΔpsbA strains were prepared from frozen and lyophilized samples and diluted to 20 μg/ml. Two-fold serial dilutions were prepared in PBS and applied to GM1 ganglioside-coated plates in triplicate. Purified CtxB was used as a positive control. Plates were washed with PBS-T, and binding was detected with rabbit anti-CT primary antibodies and goat anti-rabbit secondary antibodies conjugated to horseradish peroxidase. Plates were visualized with the TMB substrate kit (Thermo Scientific). Endpoint titers of GM1 binding activity in JAG101 lysates were defined as the last dilution that gave a reading that was three standard deviations above the signal from ΔpsbA lysates at a concentration of 10 μg/ml.

Laboratory mice and animal protocols. Animal experiments were approved by the Institutional Animal Care and Use Committee (IACUC) of the University of California, San Diego, under protocol S-12026. Female BALB/c mice 4 to 6 weeks of age were purchased from Charles River Labs and housed in a UCSD vivarium. Mice were given ad libitum access to food and water. Experimental groups were composed of at least five mice.

Immunization regimes and blood and fecal sample collection. For comparing the immunogenicity of Cr.CtxB-Pfs25 to that of Cr.Pfs25, mice were vaccinated by intraperitoneal injection with 20 μg of affinity-purified Cr.CtxB-Pfs25 or an equimolar amount of Cr.Pfs25 in a 1:1 mix of 2% Alhydrogel (Ivivogen). Three injections were administered at 3-week intervals, and blood was collected prior to each immunization and 1 week following the final injection.

Oral vaccinations were administered to naive and primed mice. Mice were primed by intraperitoneal injection with 40 μg of affinity-purified Cr.CtxB-Pfs25 using complete Freund’s adjuvant; naive mice were not injected. Primed and naive mice were each orally immunized by intragastic gavage with 10 mg lyophilized JAG101 or the ΔpsbA strain for a total of four groups of five mice: (i) primed with ΔpsbA algae, (ii) primed with JAG101 algae, (iii) naive with ΔpsbA algae, and (iv) naive with JAG101 algae. Oral immunizations were prepared by resuspending lyophilized algae in PBS and were administered using a bulb-tipped gastric gavage needle every 7 days. In total, 12 oral immunizations were administered over a 12-week period. From each mouse, fecal pellets were collected weekly and blood was collected prior to vaccinations (prebleed) and periodically thereafter by submandibular bleed; final bleeds were performed by cardiac puncture at the completion of the immunization regime.

Determination of antigen-specific IgG and IgA endpoint titers. Pfs25- and CtxB-specific serum IgG antibody titers were determined by ELISA as follows. Purified CtxB and Cr.Pfs25 were diluted to 1 μg/ml in PBS and applied to MaxiSorp or Immulon 2HB ELISA plates, which were incubated overnight at 4°C on a rocker, blocked with 5% milk in PBS-T for 2 h at room temperature, and washed three times with PBS-T. Serum samples from each mouse were prepared by 1:4 serial dilutions starting at 1:100 in PBS; diluted samples were added to the plate and incubated overnight at 4°C. After three washes with PBS-T, antigen-specific IgG antibodies were detected with goat anti-mouse IgG antibodies (Sigma; A3562) conjugated to alkaline phosphatase for 2 h at room temperature. Plates were visualized with PNPP and measured at 405 nm using an Infinite M200 Pro plate reader. Endpoint titers were defined as the lowest dilution with a signal that was three standard deviations above that of prebleed sera. Statistics were calculated in Excel using a paired Student t test, and figures were assembled in JMP version 9.02 (SAS, Cary, NC).

Pfs25- and CtxB-specific fecal IgA antibody levels were determined by ELISA as follows. ELISA plates were prepared in an identical manner to IgG antibody detection. Fecal extracts were prepared by pooling fecal pellets from each experimental group and resuspending in PBS with 0.01% Tween 20. Samples were homogenized with a 19-gauge needle followed by sonication. Lysates were cleared by centrifugation and quantitated using the DC protein assay (Bio-Rad). Fecal extracts were added to antigen-coated ELISA plates at 0.5 mg/ml in triplicate and incubated overnight at 4°C on a rocker. Plates were washed three times with PBS-T, and IgA antibodies were detected with goat anti-mouse IgA antibodies (Sigma catalog no. A4937) conjugated to alkaline phosphatase at room temperature for 2 h. Following three washes with PBS-T, plates were visualized with PNPP and measured at 405 nm with an Infinite M200 Pro plate reader. Figures were assembled in JMP version 9.02.
**RESULTS**

Protein engineering and construction of transgenic algal chloroplasts expressing ctxB-pfs25. We synthesized the β subunit of the *Vibrio cholerae* toxin (ctxB) gene using a *C. reinhardtii* chloroplast codon bias (here referred to as *Cr.ctxB*). Synthetic *Cr.ctxB* was fused to the 5′ end of a codon-optimized *pfs25* (*Cr.pfs25*). We placed the *Pfs25* domain on the carboxy side of the *ctxB* domain, because N-terminal peptide additions to *ctxB* reduce oral immunogenicity (54). *Pfs25* is a malaria transmission-blocking vaccine candidate protein that lacks glycosylation and contains tandem repeats of structurally complex epidermal growth factor (EGF)-like domains. The codon adaptation index (CAI), which quantitates the codon bias of a transgene against a reference set, of *Cr.ctxB-pfs25* is 0.909 (47). A CAI score of 1 indicates that every instance of an amino acid is encoded by the most prevalent codon of the reference table for that amino acid (55). *CtxB* and *Pfs25* are separated by a flexible glycine linker and a furin protease recognition sequence, which facilitates antigen separation following endocytosis by gut epithelial cells. A nine-amino-acid flag epitope was placed on the carboxy end of the protein to facilitate purification, with a TEV protease site between the flag epitope and *Pfs25* to allow for removal of the flag epitope following purification. *Cr.ctxB-pfs25* was cloned into a chloroplast expression vector containing the light-dependent *psbA* promoter and 5′ and 3′ untranslated regions (Fig. 1A; pJAG101). *C. reinhardtii* strain W1.1 (51) was transformed with plasmid pJAG101 by DNA-coated gold particle bombardment; integration of *Cr.ctxB-pfs25* and chloroplast homoplasmicity were confirmed by PCR as previously described (data not shown;36), and protein accumulation was confirmed by Western blotting (Fig. 1B). *Cr.CtxB-Pfs25* accumulates in chloroplasts and reaches levels of 0.09% total soluble protein as determined by ELISA (see Materials and Methods).

**Characterization of purified CtxB-Pfs25.** Purified *Cr.CtxB-Pfs25* was prepared using anti-FLAG M2 affinity resin (see Materials and Methods) and evaluated by Western blotting (Fig. 2A). As expected, *Cr.CtxB-Pfs25* migrates more slowly than *Cr.Pfs25* when analyzed by SDS-PAGE, near the predicted size of 34 kDa when probed with anti-FLAG MAbs (Fig. 2A, arrow). There is also a slightly larger band that migrates around 40 kDa (Fig. 2A, asterisk), which is similar to the band in JAG101 algae lysates (Fig. 1B). The absence of the smaller band in lysates is unclear but could be due to incomplete reduction of *Cr.CtxB-Pfs25*, because both *CtxB* and *Pfs25* contain multiple disulfide bonds.

We previously demonstrated that *Cr.Pfs25* resembles native *Pfs25* when produced in algal chloroplasts (46). To test whether *Cr.Pfs25* retains its structure when produced as a fusion with *CtxB*, we probed both reduced and nonreduced *Cr.CtxB-Pfs25*. Reduced *Cr.CtxB-Pfs25* lacks the necessary disulfide bridges that are essential to *Pfs25* structure, while nonreduced *Cr.CtxB-Pfs25* should maintain conformational epitopes. We first probed with conformation-specific anti-*Pfs25* monoclonal antibodies (4B7 MAbs). 4B7 MAbs block malaria transmission and are specific to an epitope found on properly folded *Pfs25* (56). Consistent with conformation-dependent binding of *Pfs25*, 4B7 MAbs recognized nonreduced but not reduced *Cr.CtxB-Pfs25* (Fig. 2B). Thus, *Cr.CtxB-Pfs25* contains the 4B7 epitope found only on native *Pfs25*, suggesting that *CtxB* does not disrupt *Cr.Pfs25* folding. Interestingly, nonreduced *Cr.CtxB-Pfs25* migrates as a larger-molecular-weight species. This could be due to *Cr.Pfs25*, which was previously shown to migrate at a larger molecular weight in native and nonreduced gels (46), or because native *CtxB* assembles into a pentameric structure (57). Similar to 4B7 MAbs, anti-cholera toxin (CT) antibodies recognize larger but less well-defined molecular weight species in nonreduced samples. Stronger recognition of nonreduced *Cr.CtxB-Pfs25* by anti-CT antibodies suggests that *Cr.CtxB* is folding correctly when fused to *Pfs25*. Additionally, the wide distribution suggests that *Cr.CtxB-Pfs25* is assembling into quaternary structures, including but not limited to pentamers. These data suggest that both *Cr.Pfs25* and *Cr.CtxB* are folded correctly when produced together as a chimeric protein.

Purified *Cr.CtxB-Pfs25* separated by SDS-PAGE and stained with Coomassie blue (Fig. 2D) is consistent with the Western blots. Reduced and nonreduced *Cr.CtxB-Pfs25* migrate near 33 kDa (Fig. 2D, arrowhead) and 100 to 135 kDa (Fig. 2D, arrow), respectively. However, the quantitation of purified *Cr.CtxB-Pfs25* determined by the DC Bio-Rad protein assay (see Materials and Methods) appears to overestimate the amount of recombinant protein recovered compared to bovine serum albumin (BSA).
standards. Thus, where purified Cr.CtxB-Pfs25 was used in animal studies (described below), we likely used less than the indicated amounts.

Alga-produced CtxB-Pfs25 binds the GM1 ganglioside receptor. The mucosal adjuvant properties of CtxB are dependent on GM1 ganglioside receptor binding and subsequent internalization into the gut-associated lymphoid tissue. GM1 binding was evaluated in soluble lysates prepared from Cr.CtxB-Pfs25-producing algae (strain JAG101) and an isogenic control strain lacking psbA (the /H9004psbA strain) collected at 0, 24, 48, and 72 h after light induction. Light induction is necessary because Cr.ctxB-pfs25 is controlled by the psbA promoter, which is light dependent. Lysates were applied to ELISA plates coated with GM1 ganglioside and tested for Cr.CtxB-mediated binding using anti-CT antibodies by ELISA (Fig. 3). Significant binding was detected in JAG101 samples collected at 24, 48, and 72 h after light induction compared to control algae lacking psbA. Thus, fusing Cr.Pfs25 to the C terminus of Cr.CtxB does not disrupt GM1 ganglioside binding by Cr.CtxB-Pfs25.

Effect of lyophilization and storage temperature on GM1 ganglioside binding. Cold-chain storage increases vaccine distribution complexity and costs, which could be avoided if antigens were temperature stable. We evaluated GM1 binding by Cr.CtxB-Pfs25 by ELISA from both pre- and postlyophilized JAG101 and /H9004psbA C. reinhardtii strains collected at 0, 24, 48, and 72 h after light induction (JAG101 is driven by the light-dependent psbA promoter) and tested for Cr.CtxB-mediated binding using anti-CT antibodies by ELISA (Fig. 3). Significant binding was detected in JAG101 samples collected at 24, 48, and 72 h after light induction compared to control algae lacking psbA. Thus, fusing Cr.Pfs25 to the C terminus of Cr.CtxB does not disrupt GM1 ganglioside binding by Cr.CtxB-Pfs25.

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Cr.CtxB-Pfs25 is relatively stable in lyophilized algae at moderate temperatures.

**Immunogenicity of CtxB-Pfs25.** We previously demonstrated that injecting mice with Cr.Pfs25 elicits significant levels of Pfs25-specific IgG antibodies. To compare the immune response to Cr.CtxB-Pfs25 and Cr.Pfs25 alone, we vaccinated BALB/c mice via intraperitoneal injection with 20 μg of affinity-purified Cr.CtxB-Pfs25 and an equimolar amount of Cr.Pfs25 formulated with Alhydrogel. Injections were given at 3-week intervals, and blood samples were collected prior to each vaccination and 1 week after the last injection. Sera were then tested for Pfs25-specific IgG antibodies by ELISA (Fig. 5). Surprisingly, Pfs25-specific IgG titers were significantly higher in mice injected with Cr.Pfs25 alone after the first two injections, while both proteins had similar high titers after the third injection. Thus, Cr.CtxB-Pfs25 elicits Pfs25-specific antibodies, which reach levels that are similar to Cr.Pfs25 alone after two boosts using Alhydrogel.

**Oral vaccination of primed mice with lyophilized algae.** We hypothesized that oral vaccination with whole algae that produce CtxB-Pfs25 could act to boost antigen-specific serum IgG antibody levels of mice previously injected with a single antigen dose (58). We also wanted to determine whether oral vaccination with CtxB-Pfs25 elicits a secretory IgA response not normally associated with injectable vaccines (59). To test this hypothesis, BALB/c mice were injected with affinity-purified Cr.CtxB-Pfs25 using complete Freund’s adjuvant followed by weekly oral vaccinations with 10 mg of JAG101 or ΔpsbA algae, starting 2 weeks after injection (Fig. 6; oral vaccinations are indicated by arrows). Soluble protein was measured as 23% of the total algal biomass, and of that 0.09% is Cr.CtxB-Pfs25, which results in an effective dose of approximately 31 μg of Cr.CtxB-Pfs25 for each immunization. Pfs25- and CtxB-specific IgG and IgA antibodies in sera and feces, respectively, were detected by ELISA. Pfs25- and CtxB-specific IgG antibodies were similar for mice fed with JAG101 or control ΔpsbA algae (Fig. 6A and B). Pfs25-specific IgG antibody titers were also similar when mice were injected with Cr.Pfs25 (rather than Cr.CtxB-Pfs25) followed by six doses of JAG101 or ΔpsbA algae over 3 days (see Fig. S1 in the supplemental material). Thus, oral vaccination appears to have no effect on serum IgG antibodies on a primed immune system when using relatively low doses of antigen (microgram quantities). However, CtxB- and Pfs25-specific IgA antibodies were significantly higher in fecal extracts of mice that were fed JAG101 algae than in those fed ΔpsbA algae (Fig. 6C and D). Significant levels of CtxB-specific IgA antibodies were seen after a single dose of JAG101 algae, whereas four doses were required for Pfs25-specific IgA antibodies. Together, these data suggest that lyophilized algae sufficiently protect Cr.CtxB-Pfs25 from degradation in the gastrointestinal tract and that Cr.CtxB-Pfs25 binds the GM1 ganglioside receptor and elicits an antigen-specific IgA immune response to both Cr.CtxB and its fusion partner, Cr.Pfs25.

**Oral vaccination of naive mice with lyophilized algae.** We characterized the immune response of naive BALB/c mice during a 12-week oral vaccination schedule to evaluate whether antigen-specific IgA antibodies were dependent on prior injections of the antigen. Lyophilized JAG101 or ΔpsbA algae were administered once per week starting at day 0 by oral gavage in PBS; blood and
fecal samples were collected periodically and tested for antigen-specific IgG and IgA antibodies, respectively, by ELISA. Pfs25-specific IgG antibodies in sera were not detected at any point during the vaccination regime (Fig. 7A). CtxB-specific IgG antibodies in sera increased after three immunizations (~21 days) and reached significant levels after five immunizations (35 days) relative to those in preimmune sera (Fig. 7B). Similar to primed mice, significant levels of CtxB- and Pfs25-specific IgA antibodies were detected in mice fed JAG101 algae compared to those in mice fed /H9004 psbA C. reinhardtii (Fig. 7C and D). Antigen-specific IgA antibodies appeared to increase more slowly in naive mice than in primed mice, which indicates that priming by injection may enhance the IgA antibody response to oral vaccinations. Thus, oral vaccination using Cr.CtxB-Pfs25-producing algae elicits both CtxB-specific IgG and IgA antibodies and Pfs25-specific IgA antibodies.

DISCUSSION

In this study, we demonstrated that C. reinhardtii chloroplasts can produce a structurally complex chimeric protein consisting of CtxB, the β subunit from the V. cholerae toxin, and Pfs25, a eukaryotic protein from P. falciparum and promising malaria TBV candidate. To date, algae are the only recombinant system to produce an unmodified aglycosylated Pfs25 protein that resembles native Pfs25 structure (46). Like alga-produced Pfs25, the Cr.CtxB-Pfs25 chimera is recognized by conformation-specific anti-Pfs25 transmission-blocking monoclonal antibodies, suggesting that Cr.Pfs25 is similar in structure to native Pfs25 even when produced as a chimeric protein. Furthermore, GM1 ganglioside binding by Cr.CtxB-Pfs25 in soluble alga extracts indicates that Cr.CtxB also resembles its native counterpart from V. cholerae. Thus, Cr.CtxB-Pfs25 accumulates as a soluble, correctly folded, and functional protein within algal chloroplasts.

Bifunctional chimeric proteins, like the CtxB-Pfs25 fusion produced here, are becoming increasingly useful for vaccine formulation, cancer therapeutics, immunotherapy (60, 61), and industrial enzymes (62). Ultimately, algae could be an inexpensive platform for producing many of these complex proteins, because they can be grown photosynthetically and have minimal growth requirements. Economic manufacturing of recombinant proteins using algae will further improve as new methods to increase yields are identified. In the near term, proteins that are recalcitrant to production in conventional systems or otherwise biologically suited for production in algae will help drive development of the algae platform. Indeed, algae were recently used to make immunotoxins in a single step (37). Immunotoxins are currently made by chemically conjugating a human antibody, made and purified from CHO cells, to a toxin molecule or protein, followed by a second purification of the chimeric molecule. Another significant advantage for an alga platform is oral delivery of a protein, thereby eliminating all purification steps. Unlike bacterial and mammalian expression platforms, eukaryotic green algae are considered safe and can therefore be consumed, making oral delivery of unpurified protein a possibility.

In this work, we demonstrated that Cr.CtxB-Pfs25 is protected from degradation in the gastrointestinal tract within lyophilized algae. Cr.CtxB-Pfs25 is eventually released from the alga cell in the gut, whereupon it binds the GM1 ganglioside receptor found on gut epithelial cells and elicits an immune response. Thus, lyophilized algae could be a cost-effective and efficient means of storing

FIG 6 Immune response to whole-cell alga oral immunization in primed mice. Mice were injected with Cr.CtxB-Pfs25 using complete Freund’s adjuvant followed by weekly doses of 10 mg of JAG101 or /ΔpsbA C. reinhardtii starting 2 weeks postinjection. Serum IgG antibodies for Pfs25 (A) and CtxB (B) were detected by ELISA. Error bars indicate the standard errors of the means. Pfs25 (C)- and CtxB (D)-specific IgA antibodies in fecal extracts diluted to 0.5 mg/ml were detected by ELISA. Error bars indicate 1 standard deviation. Arrows indicate when oral doses were administered.

FIG 7 Immune response to whole-cell alga oral immunization in naive mice. Mice were administered weekly doses of JAG101 algae or /ΔpsbA C. reinhardtii. Serum IgG antibodies for Pfs25 (A) and CtxB (B) were detected by ELISA. Error bars indicate standard errors of the means. Pfs25 (C)- and CtxB (D)-specific IgA antibodies in fecal extracts diluted to 0.5 mg/ml were detected by ELISA. Error bars indicate 1 standard deviation. Arrows indicate when oral doses were administered. ^, P < 0.05; ^^, P < 0.01.
and delivering proteins to the gut. In this case, Cr.CtxB-Pfs25 did not elicit the prerequisite IgG antibodies that would block malaria transmission. However, a similar strategy using the D2 fibronectin-binding domain from Staphylococcus aureus fused to CtxB did elicit IgG antibodies (63); eliciting an antigen-specific IgG response therefore appears to be dependent on the antigen itself and not just on CtxB. Hence, CtxB is not the appropriate mucosal adjuvant for an oral Pfs25-based transmission-blocking vaccine. The predominantly IgA antibody response in this study may be more appropriate for Helicobacter pylori, Shigella sp., rotaviruses, or pathogens that invade host mucosal surfaces (64).

Whole-cell oral vaccines would greatly simplify vaccine delivery and eliminate costly purification steps, both of which would enable affordable vaccination in poor countries. Eukaryotic green algae are devoid of endotoxins, human pathogens, or other known toxic compounds and provide a stable storage and delivery mechanism. Furthermore, lyophilized whole cells are a significant cost advantage over injectable vaccines and alternative oral vaccination strategies that use formulated nanoparticles (65, 66). For example, algal biomass can be produced for about $3/kg (67), meaning that a therapeutic dose could cost just a few cents. A more thorough understanding of mucosal adjuvant proteins is paramount to developing oral vaccines moving forward. Ideally, antigens could be paired with mucosal adjuvants that elicit an appropriate immune response that confers long-lived protection from disease. If this can be achieved, the gap in vaccine coverage between wealthy and poor regions of the world could be closed, and the disease burdens could be reduced in areas of the world where this is needed most.

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