



# Immunization with lytic polysaccharide monoxygenase CbpD induces protective immunity against *Pseudomonas aeruginosa* pneumonia

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*Pseudomonas aeruginosa* (PA) CbpD belongs to the lytic polysaccharide monoxygenases (LPMOs), a family of enzymes that cleave chitin or related polysaccharides. Here, we demonstrate a virulence role of CbpD in PA pneumonia linked to impairment of host complement function and opsonophagocytic clearance. Following intratracheal challenge, a PA  $\Delta$ CbpD mutant was more easily cleared and produced less mortality than the wild-type parent strain. The x-ray crystal structure of the CbpD LPMO domain was solved to subatomic resolution (0.75Å) and its two additional domains modeled by small-angle X-ray scattering and AlphaFold2 machine-learning algorithms, allowing structure-based immune epitope mapping. Immunization of naive mice with recombinant CbpD generated high IgG antibody titers that promoted human neutrophil opsonophagocytic killing, neutralized enzymatic activity, and protected against lethal PA pneumonia and sepsis. IgG antibodies generated against full-length CbpD or its noncatalytic M2+CBM73 domains were opsonic and protective, even in previously PA-exposed mice, while antibodies targeting the AA10 domain were not. Preexisting antibodies in PA-colonized cystic fibrosis patients primarily target the CbpD AA10 catalytic domain. Further exploration of LPMO family proteins, present across many clinically important and antibiotic-resistant human pathogens, may yield novel and effective vaccine antigens.

lytic polysaccharide monoxygenase | vaccine antigen | pneumonia | mouse models | crystallography

*Pseudomonas aeruginosa* (PA) is an often highly multidrug-resistant bacterial pathogen that poses a serious threat to human health, particularly in immunocompromised and/or hospitalized patients. Chronic PA respiratory infections are seen in patients with cystic fibrosis (CF), bronchiectasis, and chronic obstructive pulmonary disease (1–3), where tracheobronchial colonization is linked to deterioration of pulmonary function, poor clinical outcome, and increased mortality (4–7). PA is also a leading agent of potentially fatal ventilator-associated pneumonia in intensive care units (8, 9) and of dangerous bloodstream infections and sepsis in patients with cancer, neutropenia, burns, and/or disruption of the microbiota from prior broad-spectrum antibiotic therapy (10–13). As PA strains have proven capable of deploying high-level intrinsic and acquired resistance mechanisms to counter all major classes of antibiotics in clinical use (14, 15), the development of vaccine-induced immunity against PA is considered an essential, albeit challenging pillar of future prevention strategies (16, 17).

Antigens possessing one or more of the following characteristics including strong immunogenicity, surface exposure, high abundance, and/or a role in bacterial virulence can serve in the foundation of an effective vaccine (18). Due to its medical importance, PA has been extensively studied in terms of its molecular epidemiology (19), fundamental biochemical composition (20, 21), gene regulation (22), and host–pathogen interactions (23, 24). A number of conserved PA surface molecules that proved promising in animal vaccination models later entered Phase I–III human clinical trials, including lipopolysaccharide (25–28), flagellins (29), alginate (30), and outer membrane proteins OprF–OprI (31, 32). Despite these many investigations, no licensed vaccines are available against PA, and the search for an optimal antigen or formulation of antigens continues.

Recently, we reported that chitin-binding protein D (CbpD), a highly conserved protein across PA clinical isolates (33, 34), contributed strongly to systemic virulence and immune evasion by interference with the terminal complement cascade (35). CbpD belongs to a class of ancient microbial copper-dependent redox enzymes, the lytic polysaccharide monoxygenases (LPMOs) (35), commonly known to degrade recalcitrant polysaccharides

## Significance

Bacterial lytic polysaccharide monoxygenases (LPMOs) are understudied in disease contexts. We show the trimodular *Pseudomonas aeruginosa* LPMO CbpD is a virulence determinant in pneumonia, and mice immunized with CbpD generate robust antibody responses that promote opsonophagocytosis and protect against lethal infection. Our findings can impact research on this broadly distributed enzyme class, their roles in bacterial pathogenesis, and their potential targeting by antivirulence therapies and vaccines.

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The authors declare no competing interest.

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such as chitin [ $\beta$ -1,4(GlcNAc) $_n$ ] and cellulose [ $\beta$ -1,4(Glc) $_n$ ] through oxidative cleavage of glycosidic bonds (36–39). In PA, CbpD is secreted by a type II secretion system (40, 41), regulated by quorum-sensing control (42, 43), and induced in the presence of human serum and respiratory mucus (35, 44). Abundant CbpD was reported in the secretome of a transmissible CF-associated PA strain under conditions mimicking CF lung physiology (45) and *cbpD* gene transcription was strongly induced by human respiratory tract mucus in a mucoid PA isolate from a CF patient (44).

Here, we show that the inducible expression and protein abundance of CbpD in the respiratory tract context is corroborated by an essential virulence role of the protein in PA in a murine pneumonia model. Combining structural biology and immunological studies, we demonstrate the potential of CbpD and its various domains as a protective vaccine antigen against PA lung infection. As similar LPMOs are present in many important human bacterial pathogens, their role(s) in virulence and immunogenicity merit further in-depth exploration.

## Results

**Loss of CbpD Reduces PA Virulence and Modifies Host Responses in Murine Lung Infection.** An isogenic CbpD knockout mutant ( $\Delta$ CbpD) of the well-characterized virulent wild-type (WT) PA clinical isolate PA14 (46) was kindly provided by Deborah Hogan (Dartmouth) and confirmed biochemically (immunoblotting and proteomics) and by whole genome sequencing in our earlier work (35). There we reported a preliminary finding of significantly reduced mortality of the  $\Delta$ CbpD mutant compared to the WT strain following intratracheal (IT) challenge of outbred CD-1 mice (35). Here, we show that complementation of the mutant with the *cbpD* gene expressed on a plasmid vector restored the virulence phenotype in the IT challenge model, with 100% mortality by 48 h while 50% of mutant mice survived through 96 h (*SI Appendix, Fig. S1A*). By 24 h post-IT challenge of both outbred CD-1 or inbred C57BL/6 mice, bacterial CFU recovered from the lungs were significantly lower than those infected with the WT parent strain (Fig. 1A). Reduced bacterial burdens in the lungs of  $\Delta$ CbpD-infected mice were accompanied by significantly lower levels of inflammatory serum cytokines and chemokines at the same 24 h time point, including IL-1 $\alpha$ , IL-3, G-CSF, and GM-CSF (FDR-adjusted  $P$  or  $q < 0.05$ ) (Fig. 1B).

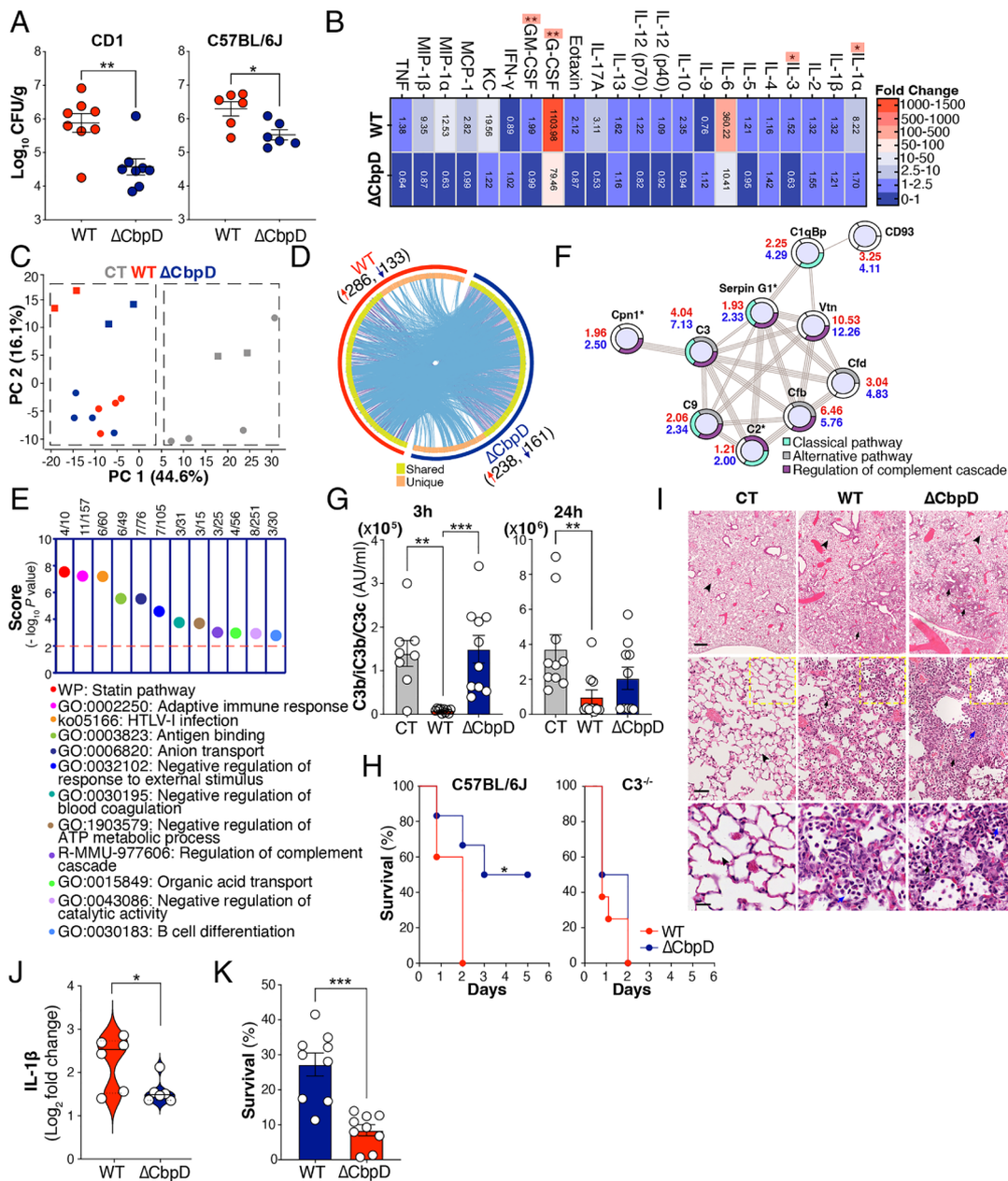
High-resolution tandem mass tag (TMT)-based quantitative proteomics was used to study the effect of CbpD deletion on lung responses to IT PA infection at the 24 h experimental endpoint; 3,944 proteins were quantified (*Datasets S1 and S2*). Principal component analysis (Fig. 1C) and similarity clustering (*SI Appendix, Fig. S2A*) segregated the lung proteomes into PA-infected (WT and  $\Delta$ CbpD) and control (mock-infected) clusters. Strong ontology term overlap was identified among the regulated proteins ( $\pi$  score  $\geq 2$ ) in WT-infected or  $\Delta$ CbpD-infected vs. mock-infected mice, particularly among proteins belonging to the shared proteome ( $n = 285$  proteins, Fig. 1D, *Dataset S3*). However, functional exploration of differentially regulated markers unique to the WT-infected (*Dataset S4*,  $n = 134$ ) or  $\Delta$ CbpD-infected lung (*Dataset S4*,  $n = 114$ ) identified more distinct strain-dependent immune responses (Fig. 1E and *SI Appendix, Fig. S2B*). The lung proteome data suggested effects on the complement system (Fig. 1E) or neutrophil responses (*SI Appendix, Fig. S2C*), which we therefore pursued through additional experimentation.

**CbpD and Complement Function in PA Lung Infection.** Among the 12 different immune-related categories that were differentially abundant in the unique proteome of  $\Delta$ CbpD-infected vs. WT-infected

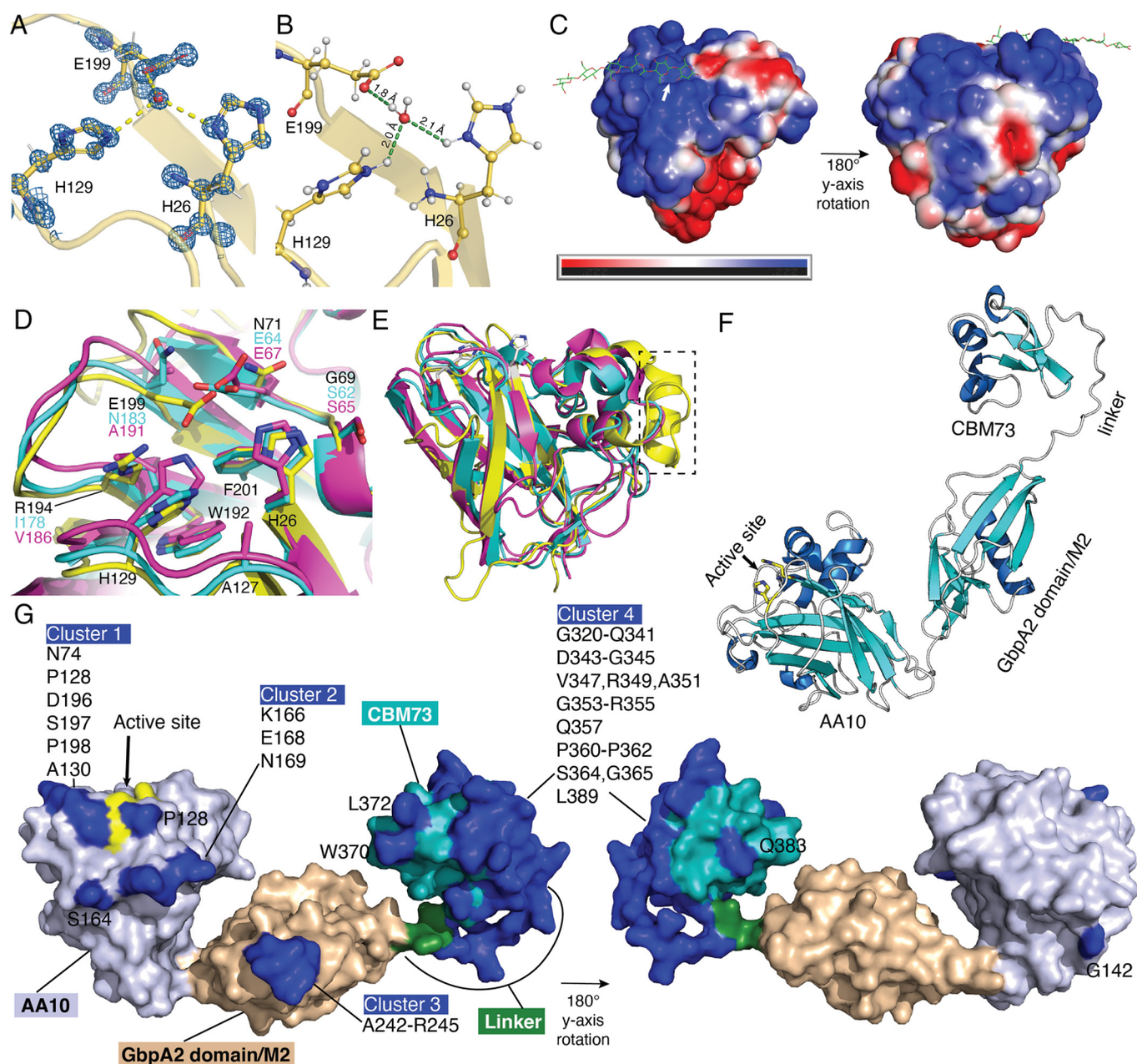
mice (Fig. 1E) were elements of the complement cascade (Serpin G1, C2, and Cpn1, Fig. 1F); proteins were further highly interconnected (Fig. 1F) to other regulated complement factors in the shared lung proteome (*Dataset S3*). Since CbpD reduced the deposition of lytic membrane attack complexes (MAC) on the bacterial surface (35), we explored serum complement cascade function both early (3 h) and later (24 h) post-IT PA infection in WT or  $\Delta$ CbpD strains vs. mock-infected controls. Plasma C3 activation (reviewed in refs. 47 and 48) as measured by C3b/iC3b/C3c (C3b-assay) was significantly reduced (below basal level) at both 3 h and 24 h time points in WT PA-infected mice compared to either mock- or  $\Delta$ CbpD mutant-infected mice ( $P < 0.05$ , Fig. 1G). Correspondingly, total C3 protein ranged lower in the lung proteome of WT-infected vs.  $\Delta$ CbpD-infected mice at the 24 h experimental endpoint (*SI Appendix, Fig. S1B*). We found that CbpD promotes PA resistance to complement-mediated killing, as the WT PA strain showed higher survival than the  $\Delta$ CbpD in normal human serum, but the two strains survived equally in serum that was heat-inactivated or depleted of key complement proteins (reviewed in refs. 47 and 48) C3 (49) ( $\Delta$ C3) or C3 and C5 (50) ( $\Delta$ C3/C5) (*SI Appendix, Fig. S1 C and D*). CbpD-mediated interference with C3-dependent complement activity contributed to PA pathogenicity in vivo, since the virulence defect of the  $\Delta$ CbpD mutant observed upon IT challenge of WT C57/BL6 mice was fully rescued in C3 knockout (C3 $^{-/-}$ ) mice (Fig. 1H).

**CbpD and Neutrophil Function in PA Lung Infection.** Strong infiltration of neutrophils was observed on histopathological examination of lungs of mice infected with either the WT or  $\Delta$ CbpD mutant PA strains (Fig. 1I and *SI Appendix, Fig. S1E*), and the importance of neutrophils in protection against mortality in the murine pneumonia model was confirmed by depletion of the cells with an anti-Ly6G antibody (*SI Appendix, Fig. S1 H and I*). Functional exploration of the top regulated proteins (*Dataset S3*,  $\pi$  score  $\geq 10$  at least in one group) in the shared lung proteome following infection with both PA strains revealed a strong association with leukocyte migration, neutrophil infiltration, and inflammation (*SI Appendix, Fig. S2C*). Among these, IL-1 $\beta$  was localized in the central part of the STRING network analysis and significantly reduced in the lung of  $\Delta$ CbpD-infected compared to WT-infected mice (Fig. 1J), fitting the overall pattern of diminished proinflammatory serum cytokines in the mutant-infected animals (Fig. 1B). As heavy lung neutrophil infiltration occurred in both groups, we hypothesized that the reduced bacterial loads seen in  $\Delta$ CbpD-infected animals (Fig. 1A) might reflect increased susceptibility to neutrophil opsonophagocytic killing. Indeed, while opsonized WT and  $\Delta$ CbpD mutant bacteria were killed with efficiency (*SI Appendix, Fig. S1G*) by murine alveolar macrophages (*SI Appendix, Fig. S1F*),  $\Delta$ CbpD mutants were more rapidly killed by freshly isolated human neutrophils ( $P < 0.05$ , Fig. 1K).

**Considerations for Vaccine Design—CbpD Structural Analysis.** The strong conservation and high prevalence of CbpD across PA clinical isolates (33–35), coupled with experimental animal studies indicating a virulence role in systemic (35) and lung infection (Fig. 1A and H and *SI Appendix, Fig. S1A*), suggests that the protein or its subdomain(s) could represent attractive vaccine antigens. Since CbpD catalytic activity is important for PA virulence (35), we determined the X-ray crystal structure of the CbpD catalytic domain (AA10) to subatomic resolution (0.75Å; PDB ID: 8C5N)—the highest-resolution LPMO structure to date, followed by 0.95Å structures with PDB ID: 4A02 (51) and 5O2X (51, 52) (Fig. 2A and *SI Appendix, Table S1*). The protein crystallized in space group  $P2_1$ , with a single chain in the asymmetric unit, and the model was



**Fig. 1.** CbpD virulence role in a murine pneumonia model. (A) Female CD-1 and C57BL/6J mice were inoculated IT with  $1 \times 10^6$  CFU PA WT or  $\Delta$ CbpD per mouse. Bacterial loads in the lung (CFU/g) were enumerated 24 h postinfection. Data are plotted as the mean  $\pm$  SEM, representing 8 (CD-1) and 6 (C57BL/6J) mice per group, and were analyzed by the two-tailed *t* test (CD-1: *P* = 0.0030; C57BL/6J: *P* = 0.0143). (B) Categorical heatmap of cytokines, chemokines, and growth factors in the serum of CD-1 mouse 24 h post IT infection (as described in A). The samples were examined using the Bio-Plex Pro™ mouse cytokine assay. Data depict mean fold-change of the cytokine values in WT- or  $\Delta$ CbpD-infected relative to mock-infected mice (10 mice/group). The data were analyzed by the multiple unpaired *t* test and the FDR-adjusted *P* value (or *q* value) was determined using the two-stage step-up method of Benjamini, Krieger, and Yekutieli. Significant differences between WT and  $\Delta$ CbpD are indicated by asterisks (\*). IL-1 $\alpha$ : *q* = 0.0181, IL-3: *q* = 0.0181, G-CSF: *q* = 0.0052, GM-CSF: *q* = 0.0054. (C) Principal component analysis (PCA) of identified proteins showing segregation of lung proteomes into infected (WT and  $\Delta$ CbpD) and uninfected groups; quantified proteins are plotted in two-dimensional principal component space by PC1 = 44.6% and PC2 = 16.1%. Each circle represents one mouse/biological replicate. One mouse sample per treatment was utilized as a technical replicate for each TMT group and depicted using square (*n* = 5 mice/group) (D) Dimension and overlap of differentially expressed proteins ( $\pi$  score  $\geq$  2) in the lung proteome of WT (red arc)- and  $\Delta$ CbpD (dark blue arc)-infected relative to mock-infected mice. The dark orange arc and purple lines reflect the regulated proteins shared in both datasets; light orange color represents regulated proteins unique to WT and  $\Delta$ CbpD-infected vs. control mice; blue lines indicate the ontology term overlap among the significantly regulated proteins; plot generated with Metascape. (E) Dot plot of enrichment score ( $-\log_{10}$  *P* value  $\geq$  2) showing pathways and cellular processes enriched in the lung proteome (Dataset S4) of  $\Delta$ CbpD-infected mice. Enrichment analysis was performed using the list of up-regulated proteins belonging to the unique category by Metascape; the *P* value was calculated based on cumulative hypergeometric distribution. (F) STRING network analysis showing connection of regulated proteins ( $\pi$  score  $\geq$  2) associated with complement activation in the lung proteome of WT- and/or  $\Delta$ CbpD-infected mice vs. control (Datasets S2–S4). The  $\pi$  score values are depicted next to the nodes in red (WT) and blue ( $\Delta$ CbpD). Regulated proteins unique to  $\Delta$ CbpD-infected versus control mice are marked with asterisks. Proteins without any interaction partners within the network are omitted from the graphics. GO:0006958: Complement activation, classical pathway (cyan), GO:0006957: Complement activation, alternative pathway (gray), MMU:977606: Regulation of complement cascade (purple). (G) Quantification of soluble complement factor C3b in EDTA-treated murine blood 3 h and 24 h post IT infection with WT or  $\Delta$ CbpD (as described in A); mock (PBS)-infected mice included as control (CT). Results are given in complement arbitrary units (AU) per mL; data are plotted as mean  $\pm$  SEM, representing 10 CD-1 mice/group. Data were analyzed by two-way ANOVA (Tukey's multiple comparisons). (Left) (3 h) CT vs. WT: *P* = 0.0025, CT vs.  $\Delta$ CbpD: *P* = 0.9893, WT vs.  $\Delta$ CbpD: *P* = 0.0009; (Right) (24 h) CT vs. WT: *P* = 0.0075, CT vs.  $\Delta$ CbpD: *P* = 0.1162, WT vs.  $\Delta$ CbpD: *P* = 0.3876. (H) Wild-type C57BL/6J (*n* = 5 to 6 mice/group) and C3 knock-out (C3<sup>-/-</sup>) mice (*n* = 8 mice/group) were inoculated IT with  $3 \times 10^6$  CFU PA WT or  $\Delta$ CbpD per mouse. Survival is represented by Kaplan-Meier survival curves and was analyzed with the log-rank (Mantel-Cox) test. (Left): *P* = 0.0445, (Right): *P* = 0.3464. (I) Representative hematoxylin and eosin-stained sections of spleen tissues from infected (WT or  $\Delta$ CbpD) and uninfected mice (*n* = 5 mice/group) were collected 24 h post IT infection (as described in A), analyzed by light microscopy. Arrowheads indicate normal alveolar structure characterized by patent airspaces lined by a thin border of alveolar epithelium with no inflammatory cells. Arrows show areas of the lung with an infiltration of neutrophils (black) and macrophages (blue) in the alveoli and the interstitium leading to loss of the pulmonary surface. (The scale bar represents 500, 50, and 20  $\mu$ m in the Upper, Middle, and Lower, respectively.) (J) Mean fold change values of IL-1 $\beta$  in the lung proteome of  $\Delta$ CbpD and WT infected compared to mock-infected (control) mice (as described in A); data are plotted as the mean ( $\log_2$  fold change)  $\pm$  SEM, representing 5 CD-1 mice/group (one mouse sample/treatment was utilized as technical replicate in each TMT group) and analyzed by the two-tailed *t* test. *P* = 0.004. (K) Survival of PA WT and  $\Delta$ CbpD upon incubation with freshly isolated human neutrophils. Bacterial survival was calculated relative to inoculum in percentage (%). Data are plotted as the mean  $\pm$  SEM, representing three experiments performed in triplicate, and analyzed by the two-tailed *t* test (*P* = 0.0001). Each data point represents an individual mouse, and the significant differences are marked with asterisks (\*): \**P*  $\leq$  0.05; \*\**P*  $\leq$  0.01; \*\*\**P*  $\leq$  0.001; \*\*\*\**P*  $\leq$  0.0001.



**Fig. 2.** Structural analysis of CbpD. (A) Electron density and protonation state of the LPMO domain.  $\sigma_a$ -weighted  $2mF_o$ -DFc map contoured at  $1.0 \sigma$  showing the histidine brace motif. (B) Protonation state of the histidine side chains of the active site at pH 7.3, calculated by H++ (56) to be fully protonated. (C) APBS-calculated surface potential for the CbpD LPMO domain, based on the protonated model generated by H++ for pH 7.3. The cellulose substrate (green) from the *Lentinus similis* LPMO9A (PDB ID: 5ACI) (57) was used to mark the putative substrate-binding region, and the active site is indicated by a white arrow. (D) Comparison of the active sites of CbpD (yellow; PDB ID: 8C5N, this work), GbpA from *Vibrio cholerae* (purple; PDB ID: 2XWX) and EfcBM33 from *Enterococcus faecalis* (cyan; PDB ID: 4ALE). Amino acids are labeled and color-coded, unless identical in all three proteins (in which case they are labeled by the residue number of CbpD, in black). (E) Superimposition of the three structures from panel (D) in cartoon representation. A dashed rectangle indicates the additional  $\alpha$ -helix and loop that extend the putative substrate-binding site of CbpD. (F) Ab initio model of full-length CbpD, generated using AlphaFold2 (UniProt entry Q02111 from the AlphaFold database). The three domains of CbpD and the linker region connecting the second domain [referred to as ‘GbpA2/M2’ domain] and third domain (‘CBM73’ domain) are labeled. An arrow points to the active site of the AA10 domain, where the two histidine residues are shown in stick representation. (G) Surface representation of the CbpD structure predicted with AlphaFold2, showing the AA10 LPMO domain (light purple; active site histidines colored yellow and indicated by arrow), the GbpA2/M2 domain (wheat), the linker connecting domains 2 and 3 (forest green), and the CBM73 domain (teal). The potential discontinuous structural epitopes predicted using Discotope 2.0 (58) are colored blue and labeled.

refined to final  $R_{work}/R_{free}$  values of 14.0%/15.2%. The structure closely resembles other AA10 LPMOs, such as those from *Tectaria macrodonta* (PDB code: 6IF7 (53); all-atom r.m.s.d.: 0.7Å) or *Vibrio cholerae* (PDB code: 2XWX) (54); all-atom r.m.s.d.: 1.1Å). All LPMOs have a pyramidal shape formed by a core  $\beta$ -sandwich fold with protruding loops, often hosting clusters of  $\alpha$ -helices, with an N-terminal histidine, which is a part of a characteristic histidine brace catalytic motif. The anomalous difference map showed no peak between His26 and His129 corresponding to the catalytically

essential copper ion; instead, a water molecule was modeled  $\sim 1\text{\AA}$  away from the putative copper position, additionally coordinated by a glutamate residue (Glu199), which may potentially be involved in proton delivery to the cosubstrate (55).

The CbpD LPMO domain protonation state was evaluated using the H++ server (56). Side chains of the histidine brace motif were fully protonated under neutral (pH 7.3; and acidic conditions (pH 5.0, 6.2, 6.7) conditions (Fig. 2B). Reduced pH during PA respiratory tract infection could theoretically impair activity

of epithelial-derived antimicrobial peptides (59) or enzymatic performance of CbpD. At pH 7.3, the adaptive Poisson–Boltzmann Solver (APBS)-calculated surface potential (60) of the catalytic site, and the CbpD enzyme surface as a whole, is dominated by positive charges (Fig. 2C). Compared with the structures of virulence-related LPMOs, such as GbpA from *Vibrio cholerae* and EfCBM33 from *Enterococcus faecalis*, the CbpD active site has several unique features (Fig. 2D), as also observed recently by Dade *et al.* (61). At the position where both GbpA and EfCBM33 carry nonpolar amino acids (Ile and Val), CbpD possesses an arginine residue (R194). The presumed catalytically important residue E199 in CbpD aligns with N183 and A191 in GbpA and EfCBM33, respectively. However, these two proteins have a glutamate residue (E64 and E67) spatially overlapping with the E199 side chain and likely play a similar role in catalysis. Finally, at a position where both GbpA and EfCBM33 exhibit a conserved serine residue, CbpD carries a glycine (G69). This makes CbpD more similar to the LPMO CelS2 (also known as ScAA10C) from *Streptomyces coelicolor* (39), which unlike CbpD oxidizes cellulose rather than chitin. CbpD also appears to have an extension of the putative substrate-binding surface, constituted by an  $\alpha$ -helix and a loop that is not present in GbpA or EfCBM33 (Fig. 2E, boxed). Finally, AlphaFold2 (62) was used to predict the full-length structure of CbpD, which is a three-domain protein. The top-ranked model (Fig. 2F) shows an elongated arrangement of the three domains, which is supported by the previously reported small-angle X-ray scattering (SAXS) data of the protein (35). A recent low-resolution (3Å) structure of the CbpD first and second domains (61) shows an interaction between the two domains. A similar interaction is seen for the *Vibrio cholerae* LPMO, GbpA (54), indicating that multidomain LPMOs may reorganize their domains upon substrate binding.

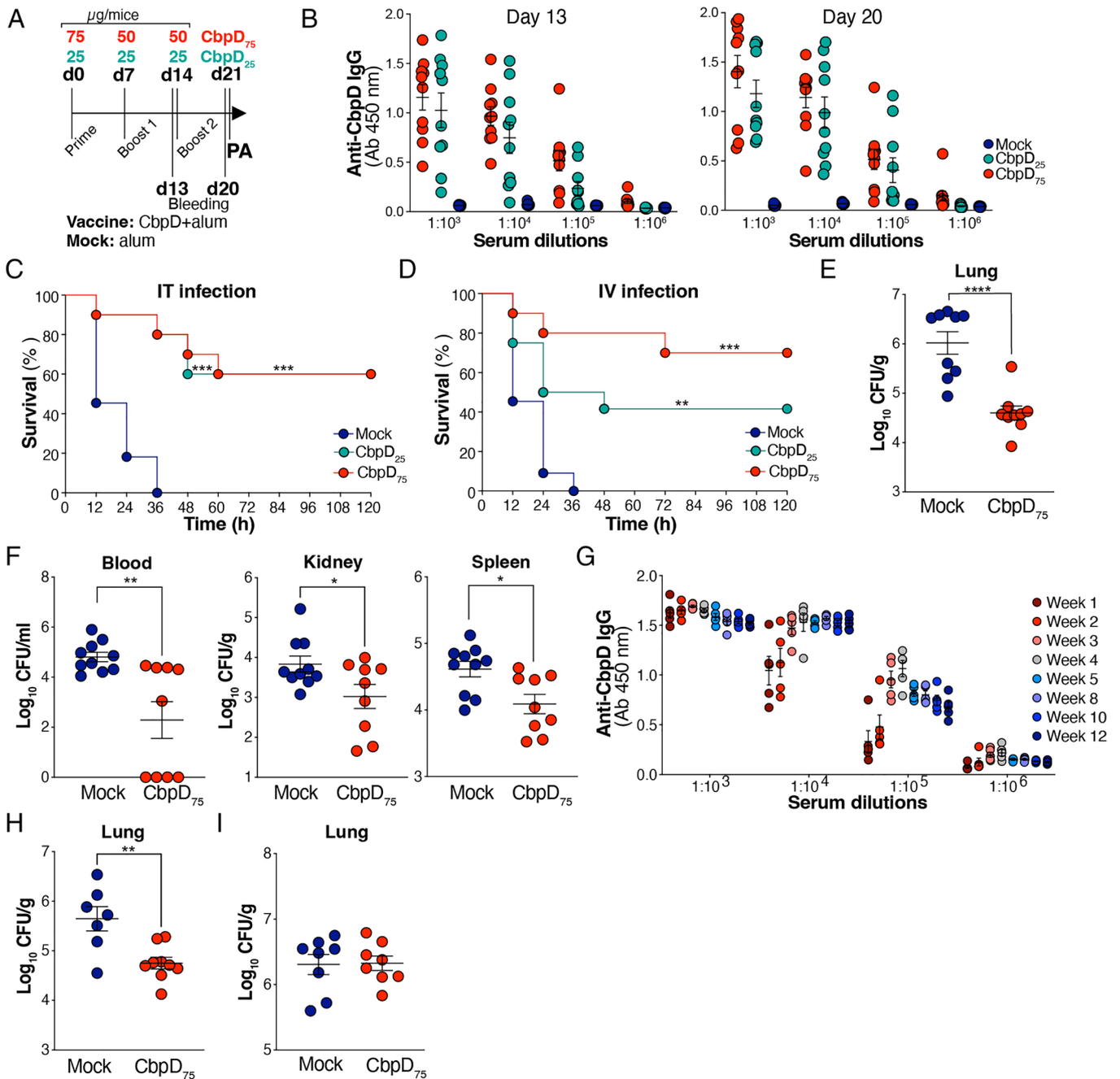
**Considerations for Vaccine Design—CbpD In Silico B Cell Epitope Prediction.** To gauge potential immunogenicity of CbpD, the Discotope 2.0 server (58) was used to predict continuous and discontinuous B-cell structural epitopes. Predicted epitopes were present in all three CbpD domains but were most abundant on the CBM73 domain (almost covering the complete domain) and the linker connecting the two domains (M2/CBM73) (Fig. 2G). In silico analysis revealed discontinuous epitopes that may be represented by four distinct clusters of amino acids: 1) N74, P128, A130, D196, S197, P198, and with P128 in close vicinity; 2) K166, E168, N169, and with S164 in close vicinity; 3) A242-R245 and 4) G320-Q341, D343-G345, V347, R349, A351, G353-R355, Q357, P360-P362, S364, G365, L389, and with L372 and W370 in close vicinity.

**Immunization with CbpD Provides Protection against Lethal PA Pneumonia and Sepsis.** We applied the IT challenge model of lethal PA pneumonia (Fig. 1A and H) to test the vaccine efficacy of CbpD adjuvanted with alum administered at two different dosing regimens (Fig. 3A): 75  $\mu$ g prime (day 0) + 50  $\mu$ g booster doses on days 7 and 14 (high dose); or 25  $\mu$ g prime (day 0) + 25  $\mu$ g booster doses on days 7 and 14 (low-dose). Blood was collected on days 13 and 20 postprime for antibody titer determination prior to experimental pneumonia (IT) or sepsis (intravenous, IV) challenge at day 21 postprime with a lethal dose of WT PA (Fig. 3A). CbpD immunization with either the high- or low-dose regimen induced high levels of anti-CbpD immunoglobulin G (IgG) measured by ELISA on days 13 and 20 postprime (Fig. 3B), which occurred without evidence of generalized inflammatory cytokine perturbations in the day 20 postprime sera comparing CbpD-immunized to mock-immunized

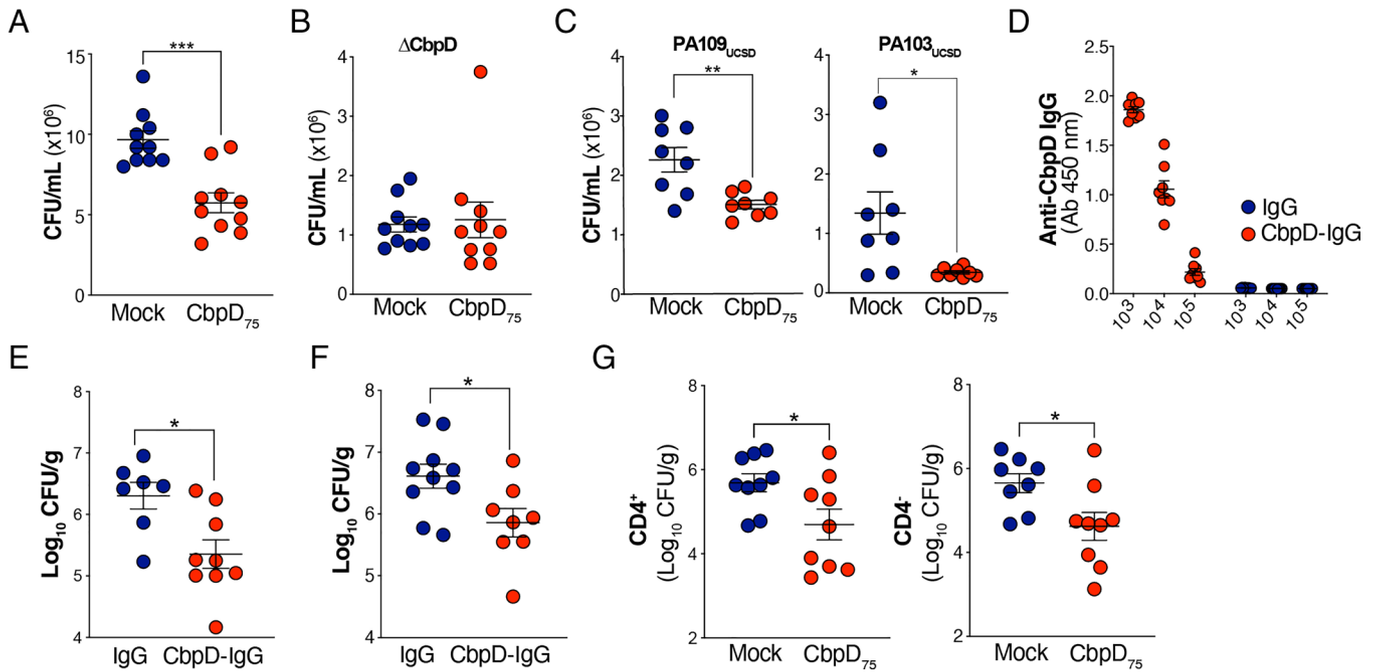
mice (SI Appendix, Fig. S3A). Both CbpD vaccine dosing regimens provided significant protection against lethal infectious challenges with PA. In the pneumonia model, all mock-immunized control mice died within 36 h of IT infection, whereas 60% of animals in the low- or high-dose CbpD vaccine groups survived (Fig. 3C). Likewise in the sepsis model, all mock-immunized control mice died within 36 h of IV infection, whereas the low-dose CbpD vaccination series led to 40% survival and the high-dose regimen of CbpD vaccination series yielded 70% survival (Fig. 3D). Given the superiority of the higher dose protocol in the systemic challenge, additional experiments were performed with high-dose (75  $\mu$ g prime + 50  $\mu$ g boosters) to assess bacterial clearance and cytokine responses. Here, we observed significantly reduced bacterial load in CbpD-immunized vs. control mice in the lung 24 h after IT challenge (Fig. 3E) and in the blood, kidney, and spleen 24 h after IV challenge (Fig. 3F). A simultaneous assessment by multiplexed immunoassay of 23 cytokines and chemokines in homogenates of lung tissue (post-IT challenge) or splenic tissue (post-IV challenge) showed no significant differences between CbpD-immunized and mock-immunized mice (SI Appendix, Fig. S3B), indicating that IgG-mediated capture of CbpD does not impair the acute cytokine response. Stability of anti-CbpD IgG titers was confirmed until 9 wk after the second booster in the high-dose vaccination regimen (Fig. 3G and SI Appendix, Fig. S3C). Additional intratracheal challenge experiments were performed to evaluate the vaccine efficacy of high-dose CbpD regimen against a hypervirulent clinical PA strain, PA103<sub>UCSD</sub>, isolated from a CF patient and shown to harbor *cbpD* in its genome (SI Appendix, Fig. S3D). The bacterial load in the lung of CbpD-immunized mice was significantly lower than that of mock-immunized mice 24 h post IT challenges (Fig. 3H). Furthermore, 24 h after IT challenge, the bacterial load of the isogenic PA  $\Delta$ CbpD mutant was unaffected by the high-dose vaccination protocol demonstrating the protective specificity to CbpD (Fig. 3I).

**Anti-CbpD Postvaccine Serum Promotes Neutrophil Opsonophagocytic Killing of PA Independent of Antibody Binding to the Bacterial Surface.** Given that CbpD is a secreted enzyme, CbpD-IgG binding to the WT PA14 surface was unaffected by sera from CbpD-immunized mice (SI Appendix, Fig. S3G). However, sera of mice fully immunized with the high-dose CbpD regimen (collected day 20 postprime) significantly boosted human neutrophil opsonophagocytic killing of WT PA compared to normal preimmune serum (Fig. 4A). Confirming antigen specificity of the antibody response to CbpD, the postimmune serum did not increase neutrophil opsonophagocytic killing of the isogenic PA  $\Delta$ CbpD (Fig. 4B). High-dose CbpD postimmune sera also improved the opsonophagocytic killing of two clinical PA strains (PA103<sub>UCSD</sub> and PA109<sub>UCSD</sub>) isolated from CF patients, and (Fig. 4C) that was confirmed to harbor *cbpD* in their genome (SI Appendix, Fig. S3D).

**Transfer of Anti-CbpD Postvaccine Serum Provides Passive Immunity to Naïve Animals.** Experiments were undertaken to determine whether anti-CbpD antibodies (humoral response) were the key factor explaining protection against PA challenge in immunized mice. High-titer anti-CbpD IgG was purified from mice immunized with the high-dose CbpD protocol and used in adoptive transfer to naïve mice, with an equivalent dose of normal purified mouse IgG serving as a control (Fig. 4D). When these antibodies were provided to naïve mice (single 70  $\mu$ g IV dose) followed by IT challenge with a sublethal inoculum of WT PA, mice that received the anti-CbpD postimmune IgG had a slightly lower bacterial CFU recovered from the lung (Fig. 4E) at



**Fig. 3.** CbpD-immunized mice are protected against PA infectious challenge. (A) The timeline depicts the immunization of CD-1 mice with CbpD plus alum (CbpD) or alum alone (Mock), bleeding and PA WT challenge. (B) Total serum anti-CbpD IgG levels following CbpD immunization with 0 µg/mouse (mock, alum), 25 µg/mouse (CbpD<sub>25</sub>) or 75 µg/mouse (CbpD<sub>75</sub>). The serum was collected on days 13 and 20 postprime (as described in A). The data are plotted as the mean ± SEM, representing 10 mice per group. (C) CbpD<sub>25</sub> (n = 10), CbpD<sub>75</sub> (n = 10), and mock (alum)-immunized (n = 11) CD-1 mice were inoculated IT with  $3 \times 10^6$  CFU/mouse PA WT at day 21 of the vaccination schedule (as described in A). Survival is represented by Kaplan–Meier survival curves and was analyzed by the log-rank (Mantel–Cox) test. CbpD<sub>25</sub> vs. Mock:  $P = 0.0001$ , CbpD<sub>75</sub> vs. Mock:  $P = 0.001$ . (D) CbpD<sub>25</sub> (n = 12 mice), CbpD<sub>75</sub> (n = 10 mice) and mock (alum)-immunized (n = 11 mice) CD-1 mice were inoculated IV with  $2 \times 10^7$  CFU/mouse PA WT at day 21 postprime. Survival is represented by Kaplan–Meier survival curves and was analyzed by the log-rank (Mantel–Cox) test. CbpD<sub>25</sub> vs. Mock:  $P = 0.0091$ , CbpD<sub>75</sub> vs. Mock:  $P = 0.0002$ . (E) CbpD<sub>75</sub> (n = 9) and mock (alum)-immunized (n = 9) CD-1 mice were IT infected with  $1 \times 10^6$  PA WT per mice at day 21 postprime. The bacterial burden in the lung (CFU/g) was enumerated 24 h postinfection. The data are plotted as the mean ± SEM, representing 9 to 10 mice per group, and were analyzed by the unpaired *t* test.  $P < 0.0001$ . (F) CbpD<sub>75</sub> (n = 9) and mock (alum)-immunized (n = 10) CD-1 mice were enumerated 4 h post IV infection with  $2 \times 10^7$  CFU WT PA per mice at day 21 postprime. Bacterial load in blood (CFU/mL) and tissues (CFU/g) was enumerated 24 h postinfection. The data are plotted as the mean ± SEM, representing 9 to 10 mice per group, and were analyzed by the unpaired *t* test. Blood:  $P = 0.0029$ ; spleen:  $P = 0.0112$ ; kidney:  $P = 0.0348$ . (G) Dynamics of changes in the titers of CbpD-IgG in CD-1 murine sera in the period from 1 to 12 wk postprime immunization. The timeline of the immunization is described in A. CbpD-IgG titers in the mock-immunized group are shown in *SI Appendix, Fig. S3C*. The data are plotted as the mean ± SEM, representing five mice per group. (H) CbpD<sub>75</sub> (n = 9) and mock (alum)-immunized (n = 7) CD-1 mice were enumerated 24 h post IT infection with  $0.5 \times 10^6$  PA103<sub>UCSD</sub> per mice at day 21 postprime. Bacterial load in the lung (CFU/g) was enumerated 24 h postinfection. The data are plotted as the mean ± SEM, representing seven to nine mice per group, and were analyzed by the unpaired *t* test.  $P = 0.0032$ . (I) CbpD<sub>75</sub> (n = 8) and mock (alum)-immunized (n = 8) CD-1 mice were enumerated 24 h post IT infection with  $2 \times 10^6$  CFU  $\Delta$ CbpD per mice at day 21 post prime, respectively. Bacterial load in the lung (CFU/g) was enumerated 24 h postinfection. The data are plotted as the mean ± SEM, representing eight mice per group, and were analyzed by the unpaired *t* test. Each data point represents an individual mouse, and the significant differences are marked with asterisks (\*):  $*P \leq 0.05$ ;  $**P \leq 0.01$ ;  $***P \leq 0.001$ ;  $****P \leq 0.0001$ .



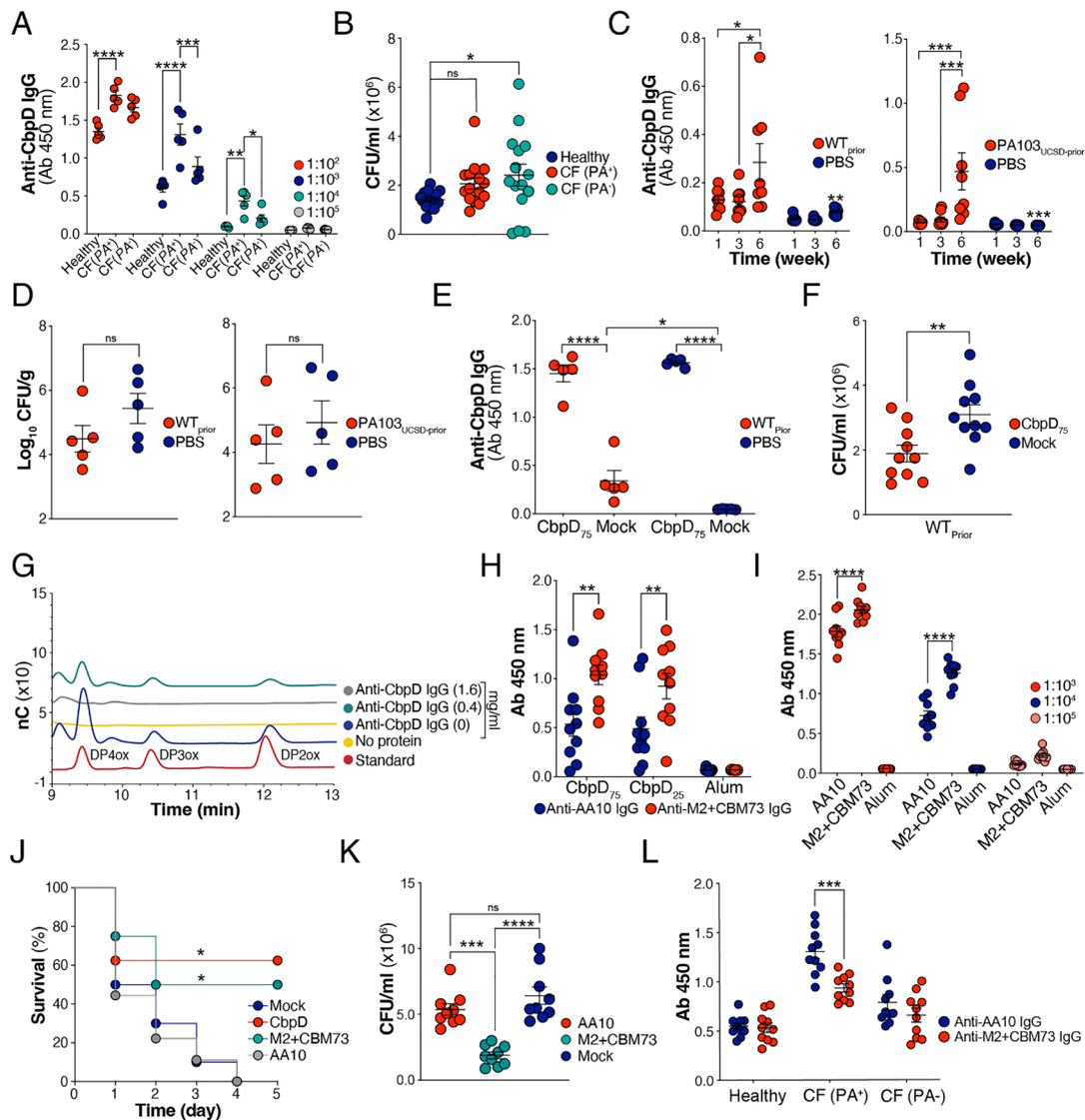
**Fig. 4.** Opsonophagocytic and protective activities of anti-CbpD IgG. (A) Opsonophagocytic killing of WT PA by freshly purified human neutrophils at MOI = 15 in the presence of 5% immunized murine sera (collected at day 20 postprime). The assay lasted 30 min. Data are plotted as the mean  $\pm$  SEM, representing 10 mice per group, and were analyzed by the unpaired *t* test.  $P = 0.0002$ . (B and C) Opsonophagocytic killing of (B)  $\Delta$ CbpD ( $n = 10$  mice), (C) PA109<sub>UCSD</sub> ( $n = 8$  mice), and PA103<sub>UCSD</sub> ( $n = 8$  mice) at MOI = 5 by freshly isolated human neutrophils in the presence of 5% fully immunized murine sera (collected at day 20 postprime). The assay lasted 30 min. The data are plotted as mean  $\pm$  SEM, representing 8 to 10 mice per group, and analyzed by the unpaired *t* test. PA109<sub>UCSD</sub>  $P = 0.0036$ , PA103<sub>UCSD</sub>  $P = 0.0140$ . (D and E) Anti-CbpD IgG (70  $\mu$ g/mouse,  $n = 8$  mice) or mouse IgG (70  $\mu$ g/mouse,  $n = 7$  mice) was IV transferred to naive CD-1 mice. The recipient mice were IT challenged with  $1 \times 10^6$  CFU/mouse WT PA 16 h after adoptive transfer. (D) The serum was collected 16 h post adoptive transfer for anti-CbpD IgG titer analysis. (E) The recipient mice were IT challenged with  $1 \times 10^6$  WT PA 16 h post adoptive transfer. Lungs were collected 24 h postinfection for enumeration. Data are plotted as the mean  $\pm$  SEM of two experiments, representing seven to nine mice per group. (F) Two doses of antibodies including 200  $\mu$ g/mouse (16 h prior infection) and 100  $\mu$ g/mouse (30 min postinfection)  $\mu$ g/mouse anti-CbpD IgG ( $n = 8$  mice) or mouse IgG ( $n = 10$  mice) were IV transferred to naive CD-1 mice. The recipient mice were IT challenged with  $2.5 \times 10^6$  CFU/mouse WT PA. The lung was collected 24 h postinfection for enumeration. The data are plotted as the mean  $\pm$  SEM of two experiments, representing 8 to 10 mice per group. (G) CbpD-immunized (CbpD<sub>75</sub>) and mock-immunized (alum) CD-1 mice were treated IV with anti-CD4 or isotype control ( $n = 9$  mice/group) 48 h and 24 h prior PA challenge. Next, 24 h post CD4 cells depletion, mice were inoculated IT with  $1 \times 10^6$  CFU/mouse PA WT. The bacterial load in the lung (CFU/g) was enumerated 24 h postinfection. Representative flow plots are presented in *SI Appendix, Fig. S4A*. The data are plotted as the mean  $\pm$  SEM, representing nine mice/group and were analyzed with the unpaired *t* test. Anti-CD4:  $P = 0.0311$ , Isotype control:  $P = 0.0246$ . Each data point represents an individual mouse, and the significant differences are marked with asterisks (\*): \* $P \leq 0.05$ ; \*\* $P \leq 0.01$ ; \*\*\* $P \leq 0.001$ ; \*\*\*\* $P \leq 0.0001$ .

a statistically significant level. A similar result was obtained using a two-dose passive transfer protocol (200  $\mu$ g + 100  $\mu$ g anti-CbpD vs. control IgG) followed by higher-dose IT PA challenge and reduced bacterial CFU in the lung observed with postvaccine IgG transfer (Fig. 4F). We confirmed no overall difference in lung inflammatory cytokine responses in post-CbpD vaccine IgG-treated mice vs. control IgG-treated mice, excluding a nonspecific mechanism of protection (*SI Appendix, Fig. S3E*). Finally, we performed an experiment in which two groups of mice underwent the high-dose CbpD vs. mock immunization protocol but were depleted of CD4<sup>+</sup> T cells using a monoclonal antibody (*SI Appendix, Fig. S4A*) prior to WT PA pneumonia challenge via examination of splenocytes. Though we did not specifically evaluate the efficiency of this treatment on depletion of tissue-resident memory C4 T cells in the lung, the CbpD-immunized mice showed reduced lung CFU compared to mock-immunized mice (Fig. 4G).

**Repeated Prior PA Infection Does Not Impair CbpD Vaccine Efficacy.** Human serum IgG recognizing CbpD was identified in healthy individuals (*SI Appendix, Fig. S3F* and Fig. 5A) and in CF patients colonized (PA<sup>+</sup>) or noncolonized (PA<sup>-</sup>) with PA at the time of blood collection (Fig. 5A). While the level of anti-CbpD IgG was significantly higher in CF(PA<sup>+</sup>) individuals compared to the two other groups, the serum failed to improve human neutrophil opsonophagocytic killing of WT PA (Fig. 5B). This observation raised a concern of whether preexisting IgG directed

against CbpD in chronically infected individuals might present a form of “original antigenic sin” (63) that interferes with effective anti-CbpD vaccine responses. To model chronic host exposure to PA and to determine whether preexisting antibodies could revoke protection conferred by CbpD immunization, 4-wk-old CD-1 mice were intratracheally infected with sublethal WT PA or PA103<sub>UCSD</sub> ( $3$  to  $5 \times 10^5$  CFU/mouse) vs. PBS control at weekly intervals for a total of six exposures. A gradual rise of anti-CbpD IgG titers in murine serum with substantial mouse-to-mouse heterogeneity was identified upon repeated IT infection with either PA strain compared to the mock-infected group (Fig. 5C). Next, we investigated the sensitization of the chronically exposed mice to a subsequent IT PA challenge. Regardless of the examined PA strain, the assessment of the lung bacterial loads revealed variation among individuals in the chronically exposed and nonexposed groups, with a general tendency to lower counts in the preexposed mice. (Fig. 5D). Nevertheless, prior PA infection did not interfere with the induction of robust anti-CbpD IgG titers in mice by the high-dose CbpD vaccination protocol (Fig. 5E) nor suppress the functional efficacy of the post-CbpD immune serum in boosting human neutrophil opsonophagocytic killing of WT PA (Fig. 5F).

**Antibodies Elicited by CbpD Immunization Block LPMO Activity of the Enzyme.** Sera from CbpD-immunized mice did not enhance IgG binding to WT PA surface (*SI Appendix, Fig. S3G*), consistent with the understanding that the enzyme is a secreted



**Fig. 5.** Immunization with CbpD elicits neutralizing antibodies. (A) Total plasma anti-CbpD IgG levels in healthy human individuals and cystic fibrosis (CF) patients who were colonized (PA<sup>+</sup>) or noncolonized (PA<sup>-</sup>) with PA at the time of blood collection. Data are plotted as the mean  $\pm$  SEM, representing 5 individuals per group. Data were analyzed by two-way ANOVA (Dunnnett's multiple comparisons test).  $10^2$ : CF (PA<sup>+</sup>) vs. Healthy:  $P = <0.0001$ ;  $10^3$ : CF (PA<sup>+</sup>) vs. Healthy:  $P = <0.0001$ ; CF (PA<sup>+</sup>) vs. CF (PA<sup>-</sup>):  $P = 0.0001$ ;  $10^4$ : CF (PA<sup>+</sup>) vs. Healthy:  $P = 0.0017$ , CF (PA<sup>+</sup>) vs. CF (PA<sup>-</sup>):  $P = 0.0360$ . (B) Opsonophagocytic killing of WT PA at MOI = 7 by freshly isolated human neutrophils in the presence of 5% human plasma obtained from healthy individuals or CF patients (as described in A). The assay lasted 30 min. Data are plotted as the mean  $\pm$  SEM, representing five individuals per group that were analyzed in triplicate and examined by the unpaired *t* test. Healthy vs. CF (PA<sup>-</sup>):  $P = 0.0319$ . (C) CD-1 mice were IT challenged with PA once per week for 6 wk (6x). Serum anti-CbpD IgG levels were measured (dilution:  $1:10^3$ ) at week 1, 3, and 6 postinfection at week 0 with WT (Left, WT<sub>Prior</sub>) or PA103<sub>UCSD</sub> (Right, PA103<sub>UCSD-Prior</sub>). Mock-infected mice (PBS) served as control. The data are plotted as the mean  $\pm$  SEM, representing 8 mice per group. Data were analyzed by two-way ANOVA (Tukey's multiple comparisons test). WT: week (1 vs. 6):  $P = 0.0240$ ; week (3 vs. 6):  $P = 0.0157$ ; PA103<sub>UCSD</sub>: week (1 vs. 6):  $P = 0.0004$ , week (3 vs.

6):  $P = 0.0009$ . (D) The lung bacterial burden in the mice that were recurrently exposed to WT PA (WT<sub>Prior</sub>) or PA103<sub>UCSD</sub> (PA103<sub>UCSD-Prior</sub>) and PBS (Mock-infected mice) as described in C. The mice were mock-immunized (alum) (as described in Fig. 3A) and rechallenged IT with a sublethal dose of  $1 \times 10^6$  WT PA or  $0.5 \times 10^6$  PA103<sub>UCSD</sub>. Bacterial load was enumerated 24 h postinfection. The data are plotted as the mean  $\pm$  SEM, representing 5 individuals per group. Data were analyzed by the two-tailed paired *t* test. (E) Total anti-CbpD IgG in sera (dilution:  $1:10^3$ ) from immunized (CbpD<sub>75</sub>) and mock-immunized mice that were recurrently exposed to WT PA (as described in C). The data are plotted as the mean  $\pm$  SEM, representing five mice per group. Data were analyzed by two-way ANOVA (Tukey's multiple comparisons test). WT<sub>Prior</sub>:  $P = <0.0001$ , PA103<sub>UCSD-Prior</sub>:  $P = <0.0001$ . (F) Opsonophagocytic killing of freshly isolated human neutrophils in the presence of 5% immunized (CbpD<sub>75</sub>) and mock-immunized mice that were recurrently exposed to WT PA (as described in C). The assay lasted 30 min at MOI = 5. Data are plotted as the mean  $\pm$  SEM of two experiments, representing five mice per group. Data were analyzed by the unpaired *t* test.  $P = 0.0075$ . (G) HPLC analysis of reaction products emerging from a mixture of copper-saturated recombinant CbpD,  $\beta$ -chitin, ascorbate as reducing agent (in all samples), and mouse anti-CbpD IgG, in buffer (20 mM Tris-HCl pH 7.0), for 3 h at 37  $^{\circ}$ C under shaking. The degrees of polymerization (DP) of chitoooligosaccharide aldonic acids in a standard sample are depicted. The reactions without 1 mM ascorbate, which resulted in generation of no oxidized product is shown in *SI Appendix, Fig. S3H*. (H) Total serum anti-AA10 IgG and anti-M2+CBM73 IgG levels (dilution:  $1:10^3$ ) following CbpD immunization with 0  $\mu$ g/mouse regimen (mock, alum), 25  $\mu$ g/mouse regimen (CbpD<sub>25</sub>), and 75  $\mu$ g/mouse regimen (CbpD<sub>75</sub>) as described in Fig. 3A. Serum was collected on day 20 postprime and recombinant AA10 and M2+CBM73 were immobilized on an ELISA plate. Data are plotted as the mean  $\pm$  SEM, representing 10 mice per group, and analyzed by two-way ANOVA (Tukey's multiple comparisons test). CbpD<sub>75</sub>:  $P = 0.0018$ ; CbpD<sub>25</sub>:  $P = 0.0078$ . (I) Total serum anti-AA10 IgG and anti-M2+CBM73 IgG level following immunization with truncated variants of CbpD (25  $\mu$ g/mouse regimen) as described in Fig. 3A. The serum was collected on day 20 postprime and recombinant CbpD were immobilized on an ELISA plate. The data are plotted as the mean  $\pm$  SEM, representing nine mice per group. Data were analyzed by two-way ANOVA (Tukey's multiple comparisons test).  $1:10^3$ :  $P = <0.0001$ ;  $1:10^4$ :  $P = <0.0001$ . (J) CbpD- (n = 8 mice, 25  $\mu$ g/mouse regimen), AA10- (n = 9 mice, 25  $\mu$ g/mouse regimen), M2+CBM73 (n = 8 mice, 25  $\mu$ g/mouse regimen), and mock (alum)-immunized (n = 10 mice) were inoculated IT with  $2 \times 10^6$  CFU/mouse PA WT. Survival is represented by Kaplan-Meier survival curves and analyzed by the log-rank (Mantel-Cox) test. CbpD vs. Mock:  $P = 0.0130$ , M2+CBM73 vs. Mock:  $P = 0.0360$ . (K) Opsonophagocytic killing of WT PA by freshly purified human neutrophils at MOI = 12 in the presence of 5% immunized mouse sera (collected at day 20 postprime). The assay lasted 30 min. Data are plotted as the mean  $\pm$  SEM, representing nine mice per group, and were analyzed by two-way ANOVA (Tukey's multiple comparisons test). AA10 vs. M2+CBM73:  $P = 0.0008$ ; Mock vs. M2+CBM73:  $P = <0.0001$ . (L) Total plasma anti-AA10 and M2+CBM73 IgG levels in healthy human individuals and cystic fibrosis (CF) patients (dilution:  $1:10^2$ ) who were colonized (PA<sup>+</sup>) or noncolonized (PA<sup>-</sup>) with PA at the time of blood collection. Recombinant AA10 and M2+CBM73 were immobilized on an ELISA plate. The data are plotted as the mean  $\pm$  SEM of two independent ELISA, representing five individuals per group. Data were analyzed by two-way ANOVA (Šidák's multiple comparisons test). AA10 vs. M2+CBM73 CF (PA<sup>+</sup>):  $P = 0.0002$ . Each data point represents an individual mouse/human, and the significant differences are marked with asterisks (\*): \* $P \leq 0.05$ ; \*\* $P \leq 0.01$ ; \*\*\* $P \leq 0.001$ ; \*\*\*\* $P \leq 0.0001$ .

protein. To assess the neutralizing characteristics of CbpD IgG, we evaluated the enzymatic activity of CbpD on a model substrate,  $\beta$ -chitin, in the presence or absence of the anti-CbpD IgG (Fig. 5G)

purified from the serum of immunized mice with a higher dose of CbpD (Fig. 3A). Copper-saturated and ascorbate-reduced full-length CbpD was fully active toward  $\beta$ -chitin in the absence of

anti-CbpD IgG, leading to the release of chitooligosaccharide aldonic acids with a degree of polymerization ranging from four to six (Fig. 5G). Incubation of the recombinant CbpD with a lower dose (0.4 mg/mL) anti-CbpD antibodies reduced activity against the chitin substrate, while a higher dose (1.6 mg/mL) abolished activity almost completely (Fig. 5G). No oxidized products were observed in the non-ascorbate-driven control reactions (SI Appendix, Fig. S3H).

**The Noncatalytic Domains of CbpD Are Important for Protective Immunogenicity.** The trimodular CbpD structure consists of an N-terminal catalytic (AA10) domain followed by a module with an unknown function (M2) and a C-terminal CBM domain (CBM73)(35). Antibodies elicited by CbpD immunization at the low- and high-dose series (Fig. 3A) bound more broadly to M2+CBM73 domains than to the AA10 domain (Fig. 5H). Immunization with truncated variants of CbpD, either the AA10 domain or the M2+CBM73 domains, showed a significantly higher level of antibody response to the latter (Fig. 5I). While mice vaccinated with the CbpD AA10 domain were not protected (full mortality within 96 h post IT infection), 50% of those immunized with the CbpD M2+CBM73 domains survived the lethal challenge (Fig. 5J). Additionally, sera of mice fully immunized with the M2+CBM73 regimen (collected at day 20 postprime), but not those immunized with AA10, significantly increased human neutrophil opsonophagocytic killing of WT PA compared to mock-immunized serum (Fig. 5K). Of note, preexisting antibodies in CF patients with PA respiratory tract colonization (Fig. 5L and SI Appendix, Fig. S3I) were primarily directed against the AA10 domain, which may in part explain their failure to improve the opsonophagocytic killing of WT PA (Fig. 5B) or impede the ongoing susceptibility to PA pneumonia in this patient population.

## Discussion

Here, we provide evidence that the LPMO CbpD contributes to PA virulence in the lower respiratory tract, as shown by reduced mortality and increased clearance of an isogenic PA  $\Delta$ CbpD mutant in a murine pneumonia model (Fig. 1). While a comparable inflammatory cytokine response and robust neutrophil recruitment occurred in the lungs of both WT PA- and  $\Delta$ CbpD-infected mice, the reduced bacterial burden seen with  $\Delta$ CbpD challenge mirrored enhanced opsonophagocytic killing of the isogenic mutant by human neutrophils (Fig. 1). C3b is a pivotal opsonin for PA (64) and increased levels of C5 convertase assembly and deposition of C3b on the  $\Delta$ CbpD surface were demonstrated in our previous study (35). Increased soluble C3b, as measured by a neoepitope expressed in C3b, iC3b, and C3c, thus reflecting specific activation of C3 in the plasma of intratracheally challenged CbpD-infected mice (Fig. 1) and vulnerability of the isogenic mutant to complement-mediated killing (SI Appendix, Fig. S1) could also contribute to accelerated lung bacterial clearance. Neutrophils (65, 66) and the complement system (67, 68) are crucial immune defenses against PA, and neutropenic (69) and complement-deficient mice (70–72) suffer more severe PA infections. In contrast to normal mice, we found comparable mortality upon  $\Delta$ CbpD- and WT PA IT challenge of C3<sup>-/-</sup> (Fig. 1) or neutropenic mice (SI Appendix, Fig. S1), indicating that complement activation and neutrophil functions are involved in CbpD-mediated virulence functions.

Both cell-mediated immunity and opsonizing antibodies have been linked to protective immunological responses against PA (26, 73–77). Immunization with CbpD elicited protective antibodies

that promoted neutrophil opsonophagocytic killing, independent of their binding to the bacterial surface, and exhibited neutralizing characteristics (Figs. 3–5). The encoding *cbpD* gene is highly conserved and prevalent across multiple PA clinical isolates (33–35), an important characteristic of a vaccine candidate. Immunization of naive mice with recombinant CbpD provided antibody-mediated protection against murine acute pneumonia and systemic infections (Fig. 3). Given CbpD is a secreted virulence factor (40, 41), enhanced opsonophagocytic killing of PA (Fig. 4) is likely related to the potency of anti-CbpD IgG in immunized murine sera to block enzyme activity (Fig. 5) and mitigate CbpD-mediated immune subversion (Fig. 3) to the advantage of the host. In contrast to the biomass-degradation potential of LPMOs (reviewed in refs. 78 and 79), their role in bacterial pathogenesis (reviewed in refs. 80 and 81) and protective antigenicity are important but understudied aspects of LPMO biology. Among other LPMO family members, passive immunization of mice with rabbit antisera raised against *Vibrio cholerae* GbpA reduced intestinal colonization with the pathogen (82). The results of our in silico discontinuous epitopes analysis (Fig. 2) corroborate our observation that CbpD-immunized mice developed significant amounts of protective antibodies against the M2+CBM73 domains and that immunization of CD-1 mice with these two domains provided nearly comparable protection to full-length CbpD in IT-challenged mice and promoted opsonophagocytic mediated killing of the bacterium (Fig. 5).

Previously, a recombinant-based vaccine candidate (IC43) containing recombinant OprF and OprI, two conserved and prevalent PA outer membrane proteins, progressed to clinical trials (31, 83–85). Although IC43 elicited a significant level of immunogenicity, it expressed no therapeutic advantage over the sham immunized group in terms of overall mortality and invasive PA infections in severely ill ICU patients, who were on mechanical ventilation (31). Recently, prior *Staphylococcus aureus* exposure was shown to render several protective recombinant-based vaccine candidates against this bacterium, such as IsdB, MntC, and FhuD2, ineffective in vivo (86). Importantly, despite offering substantial protection against invasive *S. aureus* infection during preclinical animal evaluation, these vaccines underperformed in human clinical trials (87, 88). In contrast, we observed immunization with CbpD-induced robust IgG titers with high levels of opsonophagocytic killing even in mice that were recurrently exposed to PA (Fig. 5). Protective immunogenicity is not evenly distributed through the trimodular architecture of CbpD. Our studies suggest that prior exposure to PA in healthy individuals or chronic colonization with the bacterium in CF patients may lead to a propensity for the development of antibodies against the nonprotective AA10 domain of CbpD. Immunization with recombinant CbpD in naive mice generates high titers of antibodies against the protective noncatalytic domains (M2+CBM73), and these domains combined function effectively as an antigen for vaccine protection (Fig. 5). This could offer a theoretical advantage for immunization since the AA10 domain carries a potentially toxic copper ion in its holo form or may act as a copper scavenger in its apo-form.

Chitin or chitosan polymers can be converted into oligosaccharides (COS) that contain  $\beta$ -(1,4)-linked d-glucosamine monomers, *N*-acetyl-glucosamine monomers, or a combination of these. Due to the absence of such biopolymers in the mammalian host, they are able to evoke host humoral and cellular immunological responses (reviewed in ref. 89), and many variables including the degree of polymerization and chain length may alter their immunostimulatory properties (reviewed in ref. 89). Despite the low

likelihood of such molecules arising from CbpD activity in the mammalian host, it cannot be ruled out that products arising from CbpD activity could have properties similar to COS and thereby provide an immunopotentiating activity. The study of such effects would be of great interest once the CbpD substrate is defined. PA lung infections are a major concern in vulnerable populations, and we show the involvement of the conserved LPMO CbpD in exacerbation of PA pneumonia through interference with complement and neutrophil-mediated killing. Our immunization studies provide evidence on protective characteristics of CbpD as a potential PA vaccine candidate and suggest that further exploration of the functional immunogenicity provided by the LPMO family of proteins, represented across many clinically important and antibiotic-resistant human pathogens, is warranted.

## Materials and Methods

Materials and methods, describing in detail reagents, bacterial strains, constructs, cells, and serum collection; protein production and purification; murine model of immunization; murine model of intratracheal (IT) PA infection; lung histopathology analysis; murine model of systemic PA infection; isolation and infection of murine alveolar macrophages (AMs); fluorescence microscopy analysis of AMs; serum killing assay; neutrophil killing assay; CbpD-specific antibody response in human and murine sera; cytokine profiling in murine sera or organ homogenates; complement analysis (C3b-assay); in vivo depletion of neutrophils; in vivo depletion of CD4 cells; adoptive transfer of anti-CbpD IgG in vivo; CbpD-IgG neutralization assay; flow cytometric analysis of CbpD antibody binding to PA cell surface; analysis of lung proteome; X-ray crystallography of CbpD catalytic domain; modeling the protonation state of CbpD catalytic domain (AA10); epitope prediction; and statistical analysis are provided in *SI Appendix*.

**Ethics Declarations.** Animal experiments followed all ethical regulations for animal research and were carried out in accordance with the rules and regulations of the Institutional Animal Care and Use Committee, which was approved by the UC San Diego IRB protocol S00227M. Blood was collected from several healthy volunteers (male and female) and cystic fibrosis (CF) patients in accordance with the Helsinki Declaration's ethical principles. The two blood collection processes were

approved by the UC San Diego Human Research Protection Program under IRB protocol numbers 131,002 and 160,078, respectively. All blood donors signed a written informed consent form.

**Data, Materials, and Software Availability.** Mass spectrometry proteomics data are deposited to the ProteomeXchange Consortium (90) via the PRIDE (91) partner repository with dataset identifier [PXD039298](https://proteomecentral.proteomex.org/submitter/PXD039298). The crystal structure of CbpD is provided by publicly available protein structures with PDB identifiers [8C5N](https://www.rcsb.org/structure/8C5N). All study data are included in the article and/or [supporting information](#).

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