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Deletion of the Zinc Transporter Lipoprotein AdcAII Causes Hyperencapsulation of Streptococcus pneumoniae Associated with Distinct Alleles of the Type I Restriction-Modification System

Claire Durmort,^a Giuseppe Ercoli,^b Elisa Ramos-Sevillano,^b Suneeta Chimalapati,^b Richard D. Haigh,^c Megan De Ste Croix,^c Katherine Gould,d Jason Hinds,d Yann Guerardel,e [Thierry Vernet,a](https://orcid.org/0000-0003-2034-7944) Marco Oggioni,c Jeremy S. Brownb

alnstitut de Biologie Structurale (IBS), Univ. Grenoble Alpes, CEA, CNRS, Grenoble, France

^bCentre for Inflammation and Tissue Repair, Department of Medicine, Royal Free and University College Medical School, Rayne Institute, London, United Kingdom

c Department of Genetics and Genome Biology, University of Leicester, Leicester, United Kingdom

dInstitute for Infection and Immunity, St. George's University of London, London, United Kingdom

^eUniv. Lille, CNRS, UMR 8576—UGSF-Unité de Glycobiologie Structurale et Fonctionnelle, Lille, France

Giuseppe Ercoli and Elisa Ramos-Sevillano contributed equally to the manuscript.

ABSTRACT The capsule is the dominant Streptococcus pneumoniae virulence factor, yet how variation in capsule thickness is regulated is poorly understood. Here, we describe an unexpected relationship between mutation of adcAII, which encodes a zinc uptake lipoprotein, and capsule thickness. Partial deletion of adcAII in three of five capsular serotypes frequently resulted in a mucoid phenotype that biochemical analysis and electron microscopy of the D39 adcAII mutants confirmed was caused by markedly increased capsule thickness. Compared to D39, the hyperencapsulated AadcAII mutant strain was more resistant to complement-mediated neutrophil killing and was hypervirulent in mouse models of invasive infection. Transcriptome analysis of D39 and the Δ adcAII mutant identified major differences in transcription of the Sp_0505-0508 locus, which encodes an SpnD39III (ST5556II) type I restrictionmodification system and allelic variation of which correlates with capsule thickness. A PCR assay demonstrated close linkage of the SpnD39IIIC and F alleles with the hyperencapsulated Δ adcAII strains. However, transformation of Δ adcAII with fixed SpnD39III alleles associated with normal capsule thickness did not revert the hyperencapsulated phenotype. Half of hyperencapsulated Δ adcAII strains contained the same single nucleotide polymorphism in the capsule locus gene cps2E, which is required for the initiation of capsule synthesis. These results provide further evidence for the importance of the SpnD39III (ST5556II) type I restriction-modification system for modulating capsule thickness and identified an unexpected linkage between capsule thickness and mutation of Δ adcAII. Further investigation will be needed to characterize how mutation of adcAII affects SpnD39III (ST5556II) allele dominance and results in the hyperencapsulated phenotype.

IMPORTANCE The Streptococcus pneumoniae capsule affects multiple interactions with the host including contributing to colonization and immune evasion. During infection, the capsule thickness varies, but the mechanisms regulating this are poorly understood. We have identified an unsuspected relationship between mutation of adcAII, a gene that encodes a zinc uptake lipoprotein, and capsule thickness. Mutation of adcAII resulted in a striking hyperencapsulated phenotype, increased resistance to complement-mediated neutrophil killing, and increased S. pneumoniae virulence in mouse models of infection. Transcriptome and PCR analysis linked the hyperencapsulated phenotype of the Δ adcAII strain to specific alleles of the

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Address correspondence to Claire Durmort, [claire.durmort@ibs.fr.](mailto:claire.durmort@ibs.fr)

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SpnD39III (ST5556II) type I restriction-modification system, a system which has previously been shown to affect capsule thickness. Our data provide further evidence for the importance of the SpnD39III (ST5556II) type I restriction-modification system for modulating capsule thickness and identify an unexpected link between capsule thickness and Δ adcAII, further investigation of which could further characterize mechanisms of capsule regulation.

KEYWORDS Streptococcus pneumoniae, capsule expression, virulence, AdcAII, restriction modification, SpnD39III

*S*treptococcus pneumoniae (the pneumococcus) is a Gram-positive bacterial commensal of the human nasopharynx (1) and also a common invasive pathogen causing pneumonia, septicemia, and meningitis (2). S. pneumoniae has multiple virulence factors which facilitate disease pathogenesis (3), the most important of which is the capsule. The capsule is an extracellular polysaccharide layer which plays a crucial role in S. pneumoniae immune evasion by inhibiting complement recognition, phagocytosis, and bacterial entrapment by mucus (4). Variation in S. pneumoniae capsule structure results in multiple different biochemical and antigen structures, with at least 98 distinct capsule polysaccharide serotypes recognized at present (5). This diversity is mainly related to genetic variation in the multigene cps locus (6) and correlates closely with strain phenotypes such as invasive potential, duration of colonization, and ability to evade complement-mediated neutrophil phagocytosis (7 , 8). The degree of capsule expression by S. pneumoniae is also affected by phase variation at different sites of infection (9, 10). Opaque-phase S. pneumoniae has increased thickness of the capsule layer and is associated with invasive infections such as septicemia, whereas transparent-phase S. pneumoniae has thinner capsule layers and is associated with colonization and biofilm formation (11–13). Despite the importance of capsule expression during S. pneumoniae interactions with the host, the molecular mechanisms underpinning phase variation and capsule thickness remain relatively poorly understood.

One mechanism that has been recently described to control capsule expression is epigenetic regulation by phase-variable control of DNA methylation driven by the type I restriction-modification system SpnD39III (ST5556II) (14). The SpnD39III (ST5556II) system consists of multiple genes that can be shuffled by recombination on inverted repeats to create enzymes capable of methylation at six different recognition sites. Capsule expression and thickness (opaque versus transparent) have been correlated with different SpnD39III alleles (14–16), and this system may be involved in regulating at least some aspects of S. pneumoniae phase variation. As yet, both the environmental conditions influencing allele distribution and how the effects of methylation patterns on gene expression lead to changes in capsule thickness have not been resolved.

Within mammalian hosts, the available concentrations of several cations are strictly controlled. As a consequence, cation ABC transporters of iron, manganese, and zinc are essential for S. pneumoniae growth and survival in the host (17–19). ABC transporters consist of a membrane-attached lipoprotein substrate binding protein and membrane permease(s) and ATPase proteins. Zinc acquisition is mediated by two ABC transporters identified by their lipoprotein components as AdcA and AdcAII (20 , 21). Adjacent to adcAII is phtD, which encodes the surface protein PhtD, a member of the Pht histidine triad surface protein family that are involved in S. pneumoniae virulence. The histidine triad motifs of Pht proteins have a high affinity for zinc, and these proteins may provide a surface reservoir of zinc for import into S. pneumoniae via AdcA and AdcAII ABC transporters (22–24). We have previously demonstrated that deletion of adcA partially attenuates virulence, and deletion of both adcA and adcAII had a profound effect on S. pneumoniae physiology under low zinc conditions and strongly attenuated virulence (19, 25). In contrast, the virulence of the single adcAII deletion mutant was significantly increased. Here, we describe this unexpected consequence of partial deletion of adcAII in detail and show that the hypervirulence of the D39 Δ adcAll mutant strains is

FIG 1 Creation and macroscopic phenotype of the ΔadcAII mutant. (A) Gene map of the adcAII locus showing the bp 201 to 750 deletion and replacement with an antibiotic resistance cassette (cat or kana) present in the ΔadcAII mutant. (B) Colony morphology on Columbia blood agar plates of wild-type (WT) D39 strain and the ΔadcAII mutant. (C) Relative amount of monosaccharides in capsule extracts of WT D39 strain and ΔadcAII mutant determined by GC-MS. All monosaccharide derivatives were identified according to their specific retention times and EI-MS fragmentations, as described in reference 26. (D) Example of measuring the volume of D39 and ΔadcAII bacterial pellets using microcapillary tubes. (E) Height (mm) of bacterial pellets for the WT and mutant strains in the indicated strains measured using microcapillary tubes. Each point represents data for independent clones containing the indicated mutation, and bars represent mean values for independently derived colonies for each mutant strain. P values were calculated using unpaired t test. **, $P < 0.01$; ***, $P < 0.001$.

associated with a mucoid phenotype and increased capsule expression and is correlated closely with specific SpnD39III alleles and a point mutation in the csp2E capsule locus gene.

RESULTS

Deletion of *adcAII* **in the** *S. pneumoniae* **D39 strain results in a markedly increased expression of the capsule.** During our previous investigation of the functional roles of the AdcA and AdcAII zinc ABC transporter systems, a single deletion mutant of the adcAII gene was made by partial replacement of the adcAII gene with the chloramphenicol resistance cassette cat (Fig. 1A). The resulting Δ adcAll mutant strains displayed a visibly increased mucoid colony morphology (Fig. 1B). Capsule thicknesses

FIG 2 Microscopic phenotype of the ΔadcAII mutant. (A, D, and G) Wild type. (B, E, and H) ΔadcAII mutant. (C and F) ΔcpsD mutant. (I) R6 unencapsulated strain. (A to F) Electron microscopy of WT D39 strain and mutants, showing ultrathin sections of pneumococcus after capsule spring fixation using lysine-acetate-based ruthenium red-osmium protocol. Scale bars and mean capsule width in nm (SD) are given in the closeup views of selected examples of each strain in the right-hand column. (G to I) Confocal microscopy of wild-type D39, R6, and ΔadcAII mutant strains showing the capsule in yellow (anti-type 2 capsule antibody and Alexa Fluor 546 anti-rabbit antibody) and DAPI in blue.

were compared between the D39 and Δ adcAll mutant strains using a range of assays. Initially, colony volume was assessed by transferring single colonies to a capillary tube and measuring the height of visible bacterial material. This demonstrated an increased volume of the Δ adcAII strain compatible with a thicker capsule layer (Fig. 1D and E). Capsule width was then directly visualized for the Δ adcAll and wild-type D39 strains using electron microscopy (EM), which demonstrated that the bacterial cells of the AadcAII mutant had a considerably enlarged capsule layer compared to D39 (Fig. 2A to F). The mean capsule radius indicated that the ΔadcAII mutant expressed a capsule 5.6 times thicker than the wild-type (WT) D39 (capsule width of 61 \pm 1.8 nm versus 343 \pm 8.3 nm for the D39 and ΔadcAll strains, respectively; $n = 30$ for each strain). Monosaccharide composition of capsule extracts for the ΔadcAII mutant and WT D39 strain extracts were assessed biochemically using gas chromatography-mass spectrometry (GC-MS). Total polysaccharide in capsule extracts demonstrated a 1.5-fold increase in the ΔadcAII mutant compared to the wild-type strain, largely due to a 2-fold increase in rhamnose content (Fig. 1C). Despite these changes in polysaccharide content, the hyperencapsulated ΔadcAII strain was still recognized by serotype-specific antisera (Fig. 2G to I). The small amount of GalNac detected was probably from teichoic acids extracted with the capsular polysaccharide. Overall, these data demonstrated that partial deletion of adcAII modified the polysaccharide content of the capsule with overexpression of rhamnose-containing polysaccharides. To assess whether the ΔadcAII mutant phenotype was serotype specific, additional ΔadcAII mutant strains were obtained in capsular serotype 4, 6A, 6B, and 17F strains. Partial deletion of adcAII in the 6A and 6B serotypes also resulted in a mucoid phenotype suggestive of increased capsule thickness but did not affect capsule thickness in the serotype 4 and 17F strains (Fig. 1E).

TABLE 1 ΔadcAII mutant method of construction/source of DNA for the targeted deletion related to the capsule phenotype for multiple transformants

	No. of clones	Capsule phenotype			
DNA source for transformation	analyzed	Absent ^a	Normal	Thick	
PCR fragment adcAll::kana	100	14(1)	32	44	
PCR fragment adcAll::cat	100	45 (4)	13	42	
Genomic DNA R6 AadcAll::cat1 1st	4	0	0		
Genomic DNA R6 AadcAll::cat1 2nd	4		0		
Genomic DNA R6 ΔadcAll::cat2	15				

aNumbers in parentheses are numbers of absent capsule strains sequenced all of which contained the Q308 stop codon mutation in cps2E.

Consistent association of the *adcAII* **mutation with increased capsule expression by D39.** To characterize further the relationship between partial deletion of adcAII and increased capsule thickness, additional transformation and phenotyping experiments were performed. Increased capsule expression was also detected in Δ adcAll strains made using the kanamycin resistance cassette kana instead of cat and if the deletion included the immediate downstream gene (phtD) (Fig. 1E and Table 1). Combined deletion of adcA and adcAII did not result in an increased capsule thickness phenotype. When the adcAII mutation was created in an unencapsulated D39 strain (ΔcpsD), colony volumes were similar to the parental strain and markedly lower than with Δ adcAII mutations in the WT D39 strain (Fig. 1E). The frequency with which deletion of adcAII resulted in a strain with an increased capsule thickness was investigated using multiple transformants made using the adcAII deletion constructs or by transformation with genomic DNA extracted from a Δ adcAll strain mutant. Of the 100 transformants, 44% (kana) or 42% (cat) had increased capsule thickness when transformed with the PCR construct and 78% (18 out of 23) when transformed with genomic DNA (Table 1). The remaining mutant clones either had a normal capsule thickness or were unencapsulated. Growth of the Δ adcAll strain in chemically defined medium (CDM) supplemented with 33 μ M cations (Mn²⁺ or Zn²⁺), 5% sucrose, or recombinant PhtD (50 μ g/ml) or in CDM depleted of cations by treatment with 1 mM EDTA did not reduce increased capsule expression (data not shown; measured using capillary tube colony volume). The increased capsule thickness phenotype was stable, with 100% of 100 colonies retaining a thick capsule after a single mucoid colony was cultured in THY (Todd-Hewitt broth supplemented with yeast extract) liquid medium followed by plating on blood agar plates over five generations. These data show that transformation of the S. pneumoniae D39 strain with a deletion construct affecting adcAII frequently results in transformants with a marked increase in capsule quantity.

The hyperencapsulated D39 Δ adcAII strain is resistant to complement**mediated phagocytosis.** The capsule is an essential virulence factor that prevents opsonophagocytosis of S. pneumoniae but at a metabolic cost during S. pneumoniae growth (7 , 26). We therefore investigated the phenotypes of the hyperencapsulated D39 Δ adcAll strain in vitro and in murine infection models. Growth of the Δ adcAll strain was similar to the WT D39 in complete medium THY and in CDM (supplemented with 33 μ M zinc to overcome effects of loss of adcAII on zinc transport) (Fig. 3A and B). In contrast, in blood approximately 1 log₁₀ more Δ adcAll bacteria were recovered after 4 h of incubation compared to the D39 WT strain, with large differences in CFU persisting at 6 h (Fig. 3C). Flow cytometry demonstrated increased resistance to opsonization with complement and macrophage phagocytosis of the D39 Δ adcAll strain compared to the D39 WT strain (Fig. 4A to C). The D39 Δ adcAll strain also had increased resistance to killing by neutrophils compared to the WT strain; these differences were lost if bacteria were opsonized in heat-treated (i.e., complement-deficient) sera or in phosphatebuffered saline (PBS) alone, demonstrating that the differences were largely complement dependent (Fig. 4D). Adhesion assays showed there was no defect for the D39 AadcAII strain in binding to the respiratory epithelium cell line Detroit 562 compared to the WT strain (Fig. 5A). Hence, increased capsule expression by the Δ*adcAll s*train was

FIG 3 Growth phenotype of the WT D39 strain and ΔadcAII mutant. (A and B) Bacteria were inoculated at an OD₅₉₅ of 0.01 in THY (A) or CDM (B) supplemented with 33 μ M Zn and incubated at 37°C for 8 h. Black circles, WT D39; white circles, ΔadcAII mutant. Two independent assays were performed using triplicate wells. Each point is the mean (SD) for the results of a representative experiment. (C) Mean (SD) WT D39 or Δ adcAII mutant CFU after culture in blood (1 ml inoculated with 1 \times 10⁶ CFU) for 4 and 6 h. P values were calculated using unpaired Student's t test. $*$, $P < 0.05$.

associated with resistance to complement-mediated phagocytosis but did not inhibit adhesion to a human nasopharyngeal cell line.

The hyperencapsulated Δ adcAII strain has increased virulence. Both colony forming units (CFU) in nasal washes at day 5 and competitive infection experiments demonstrated that the hyperencapsulated D39 AadcAII strain colonized the nasopharynx to a similar degree as the WT D39 (Fig. 5B; Table 2), results which are consistent with the lack of a difference between the strains for adhesion to Detroit 562 cells. In contrast, the hyperencapsulated $\Delta a d c A$ II strain had increased virulence during systemic or pneumonic infection. In competitive infection experiments using a sepsis model (intraperitoneal [i.p.] inoculation), the D39 adcAII strain strongly outcompeted the WT strain (Table 2), and in a murine sepsis model using pure inocula of each strain, 80% of

FIG 4 The ΔadcAII mutant has increased resistance to complement and phagocytosis. (A) Mean fluorescence index (MFI; measured in arbitrary units) of C3b/iC3b deposition on WT D39 or ΔadcAII mutant measured using flow cytometry in 25% human serum. Error bars represent SDs. ***, $P < 0.001$, unpaired t test. (B) Examples of flow cytometry histograms for C3b/iC3b deposition on WT D39 or ΔadcAII mutant in 100% human serum. Gray shadowing indicates the results for bacteria incubated in PBS alone. (C) Flow cytometry quantification of macrophage (THP-1 cells) phagocytosis of isothiocyanate fluorescein-labeled WT D39, R6 (unencapsulated derivative of D39), and the ΔadcAII mutant for 1 h at 37°C (50 CFU/cell). The percentage of fluorescent macrophages was quantified by flow cytometry, and the data are expressed as means (SD) of the percentage of the results for the WT D39 strain. **, P < 0.01, unpaired Student's t tests. (D) Mean proportions of WT D39 (white columns) and the ΔadcAII mutant (black columns) surviving incubation with fresh human neutrophils for 45 min (MOI of 500 bacteria/neutrophil). Data are given for bacteria preincubated in PBS, 25% normal human serum, or 25% heat-inactivated human serum (no complement activity). Error bars represent SDs, and P values were obtained using unpaired t tests.

mice infected with the D39 Δ adcAll strain progressed to fatal infection by 40 h compared to 40% of mice infected with WT D39 (Fig. 6A). Finally, in a pneumonia model higher CFU was recovered in both the lungs and blood from mice infected with the AadcAII mutant compared to wild-type D39 (Fig. 6B and C).

Transcriptome analysis of wild-type and hyperencapsulated Δ **adcAII strains.** To investigate mechanisms causing increased capsule production by the Δ adcAll strain, a transcriptome microarray analysis was performed on WT D39, one hyperencapsulated Δ adcAII and one Δ adcAII/phtD strain clone, and one Δ adcAII::cat unencapsulated clone (Cl44) (Table 3). Three independent RNA extracts for each strain were submitted to transcriptomic analysis. In total, 89 genes showed significant changes in expression $(>= 1.5$ -fold, $P < 0.05$) between the wild-type D39 and hyperencapsulated Δ adcAll strain (78 with reduced and 11 with increased expression in the mutant strain including the deleted adcAII and downstream phtD genes), 96% (86/89) of which also showed comparable changes in expression in the thick-capsule Δ adcAII/phtD strain. In contrast, 11% (10/89) of these genes showed similar changes in expression in the Cl44 Δ adcAll strain without increased capsule expression, suggesting that the gene expression changes were linked to the capsule phenotype. Expression of the D39 capsule locus genes was not significantly different between the strains. Genes showing increased expression in the hyperencapsulated Δ adcAII and Δ adcAII/phtD strains included genes related to zinc uptake (adcR, adcA, phtA, and phtE), suggesting compensatory effects

FIG 5 Effects of ΔadcAII on adhesion to epithelial cells. (A) Bacterial CFU recovered from the Detroit 562 cell adhesion assay (duration 1 h) with WT D39 or ΔadcAII mutant. There were no statistical differences in CFU recovered for each strain (unpaired t tests). (B) WT D39 or ΔadcAII mutant CFU in nasal washes recovered from mice 5 days after inoculation of either strain with 107 CFU under light halothane general anesthesia. Each symbol represents data from a single mouse, bars represent medians, and error bars represent the upper interquartile range. There were no statistically significant differences in nasal wash CFU.

due to loss of the AdcAII zinc transporter. The other genes showing increased expression in the hyperencapsulated strains encode proteins of unknown function or containing LysM domains predicted to be involved in cell wall metabolism (27). Three of the operons that showed reduced expression in the hyperencapsulated strains are predicted to be involved in pyrimidine synthesis: SPD_0608-09, encoding a predicted orotate decarboxylase and phosphoribosyltransferase and being part of a larger operon encompassing SPD_0608 to SPD_06187 (28); SPD_0851-52, predicted to encode a dihydroorotate dehydrogenase electron transfer subunit (29); and SPD_1131, predicted to encode a carbamoylphosphate synthase large subunit required for pyrimidine synthesis from glutamine (30). Other genes showing reduced expression in the hyperencapsulated strains have roles in iron uptake (SPD_0224, -0226, and -1650), carbohydrate uptake (SPD_0279, 0362, 1050-1053, 1501, and 1987-95), and riboflavin synthesis (SPD_0166-69). Of particular interest, the hyperencapsulated strains showed reduced expression of SPD_0450, SPD_0452, and SPD_0453, creX (psrA), hdsS' (hsdS2), and hdsS (hsdS1), from the SpnD39III (ST5556II) type I restriction-modification system, respectively; this is discussed in detail below.

Increased capsule thickness of the Δ adcAII strains correlated closely with **specific** *hsd* **alleles.** The S. pneumoniae SpnD39III (ST5556II) type I restriction-modification locus undergoes genetic variation due to recombination within the locus between pairs of inverted repeats, generating six allelic variants which are linked to

TABLE 2 Competitive index data for infection models using a mixed inoculum of 50% WT D39 and 50% D39 Δ adcAII hyperencapsulated strain

		P value
1.04 (0.15)		0.58
4.6(0.62)		< 0.0001
	Nasal washes (5 days)	

FIG 6 The ΔadcAII mutant has increased virulence in mouse models of sepsis and pneumonia. (A) For the septicemia model, 5×10^4 CFU of each strain was injected intraperitoneally and the progress of infection was followed over time. Empty circles represent data for the WT D39 strain, and filled black circles show the ΔadcAII mutant strain. P values were obtained using the log rank test. (B and C) Blood (B) and lung (C) CFU for the pneumonia model determined by plating serial dilutions on Columbia blood agar recovered 48 h after infection for mice inoculated by intranasal instillation of 5×10^6 CFU. Each symbol represents data from a single mouse, bars represent medians, and P values were calculated using unpaired t tests. \overline{P} / 0.05.

opaque (increased capsule expression) and transparent (reduced capsule expression) colony morphology (14–16). This suggests that the detected changes in expression of genes within the SpnD39III (ST5556II) locus could reflect differences in the proportions of the allelic variants between the WT and Δ adcAll strains, and these differences could underpin the hyperencapsulated phenotype of the latter. Hence, the proportion of each of the six SpnD39III (ST5556II) variants was obtained for multiple individual Δ adcAII strains expressing either thick or normal-size capsules using a previously described assay based on PCR followed by restriction digestion of the products (14) (Table 4). This showed a clear correlation between capsule phenotype and the dominant SpnD39III (ST5556II) variant. The WT D39 strain contained a mixture of the SpnD39III (ST5556II) variants, mainly SpnD39IIIC with also a significant proportion of the SpnD39IIID and F variants. With one exception, SpnD39IIIC (3 strains) and F (5 strains) were the dominant variants found in the hyperencapsulated Δ adcAII strains, whereas SpnD39IIID (7 strains) or A (1 strain) was the dominant variant found in the Δ adcAll strains with normal capsule thickness. To try to link increased capsule formation by some Δ adcAII mutants to changes in the dominant alleles of the SpnD39III (ST5556II) type I restriction-modification locus, the hyperencapsulated Δ adcAII strain was transformed with genomic DNA from D39 mutant strains with locked SpnD39III (ST5556II) alleles due to an inactivated *creX* gene. Flow cytometry analysis of complement sensitivity was used to rapidly assess capsular phenotype for 10 transformants for each allele (A to F). All transformants retained the complement-resistant phenotype of the hyperencapsulated Δ adcAII strain, even those made using the SpnD39III (ST5556II) alleles associated with a normal capsule width in Δ adcAII transformants (A and D) (Fig. 7), suggesting they all remained hyperencapsulated.

Genome sequence data for *AadcAll* strains. Genome sequencing of one *AadcAll* and one Δ adcAll/pht strain confirmed they contained the expected partial deletion of adcAII or adcAII and phtD, respectively, with insertion of the antibiotic resistance

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^aFor comparison, the fold differences compared to WT D39 strain for the thick-capsule ΔadcAII/phtD strain (AII + Pcl14) and a normal-capsule-thickness ΔadcAII strain (Cl144) are provided alongside. The adcAII, phtD, and SpnD39III (ST5556II) type I restriction-modification system genes are indicated in bold, and gene expression profile differences in the ΔadcAII/phtD or the ΔadcAII (Cl144) compared to the ΔadcAII AIIL hyperencapsulated strain are indicated in italics. Abbreviations: CoA, coenzyme A; PTS, phosphotransferase.

cassette with no additional mutations elsewhere in the genome. Sequencing of the cps locus in additional Δ adcAII strains found that a nonsynonymous single nucleotide polymorphism (SNP) affecting the capsule locus gene cps2E (E to K at amino acid 322) was present in five out of 10 clones (Table 5), suggesting that this SNP may be relevant for the hyperencapsulated phenotype at least for a proportion of Δ adcAII strains. However, the same SNP was also present in one of eight Δ adcAll strains with normal capsule thickness. All five of the unencapsulated Δ adcAII strains investigated had cps2E genes containing a stop codon at amino acid 308, which could explain their unencapsulated phenotype (Table 5).

DISCUSSION

In this work, we have described that mutation of the zinc transporter lipoprotein gene adcAII in the S. pneumoniae D39 strain leads to an unexpected and striking increase in capsule expression in 42% of the resulting mutants. This phenotype occurred with Δ adcAll mutations made by transformation either with a PCR construct or with genomic DNA from another Δ adcAll mutant and was stable over many bacterial generations. A similar mucoid phenotype was also observed with the Δ adcAll mutation in two of the four other S. pneumoniae capsular serotypes investigated. The increased capsule quantity was very marked, with EM showing a greater-than-5-fold increase in capsule width and nuclear magnetic resonance (NMR) showing a 60% increase in the quantity of monosaccharides in purified capsule. This level of increase in capsule expression is markedly greater than that seen between opaque and transparent TIGR4

TABLE 4 Proportions of variants (identified by PCR analysis) for the SpnD39III (ST5556II) type I restriction-modification system for selected Δ adcAII mutant strains divided into those with thick and normal capsule thicknesses

			Proportion (%) of SpnD39III (ST5556II) variant:				
Phenotype	Strain	A	B	C	D	E	F
Wild type	D39	2.2	$\mathbf{0}$	67.2	15.6	$\mathbf{0}$	15.0
Thick capsule	C182 CI72	1.3 1.7	0 Ω	3.8	7.1 9.2	1.0 0	86.8
	C ₁₀	1.3	Ω	4.3 4.0	8.8	Ω	84.8 85.9
	Cl38 C13 2P	2.0 1.0	Ω Ω	1.9 2.5	8.4 6.4	2.3 1.6	85.4 88.6
	$C15$ 1P $C13$ 1P	0.7 1.7	Ω Ω	87.8 84.8	9.8 10.3	Ω Ω	1.7 3.3
	Cl1 1G Cl7 1G	2.2 Ω	$\mathbf{0}$ $\mathbf{0}$	83.1 7.7	11.1 92.3	Ω Ω	3.7 Ω
	AIIL	3.8	0.6	74.9	17.8	0.0	2.8
Normal capsule	C188	Ω	0	6.0	92.7	0.00	1.3
	Cl28 Cl35	0 Ω	0 Ω	6.7 6.92	93.3 93.1	0 0	0 Ω
	Cl6	0	0.7	5.71	93.6	0	0
	CI73	Ω	0	9.88	90.1	Ω	Ω
	Cl ₂₀ Cl17	0 Ω	Ω 9.0	9.93 2.9	90.1 88.2	0 Ω	0 Ω
	Cl1	0.4	1.7	16.4	80.8	0	0.6
	Cl36	94.3	5.8	0	0	0	0

capsular switched (less than 2-fold) (31) and 6B strains (32), justifying describing the D39 *AadcAII* strain as hyperencapsulated. The phenotypic consequence of the increased capsule expression was a high degree of resistance to complement-mediated immunity and hypervirulence in mouse models of pneumonia and sepsis. These

FIG 7 Flow cytometry analysis of complement sensitivity of the hyperencapsulated ΔadcAII strain after transformation with locked SpnD39III (ST5556II) alleles (A to F) containing an inactivated creX gene. ΔFP441, ΔFP442, ΔFP443, ΔFP444, ΔMRO559, and ΔMRO560 are all double mutant strains carrying the adcAII mutation and an extra one in allele SpnIIIB, allele SpnIIIC, allele SpnIIIA, allele SpnIIID, allele SpnIIIE, and allele SpnIIIF, respectively. (A) Fluorescence index (MFI measured in arbitrary units multiplied by proportion of bacteria positive for C3b/iC3b) of C3b/iC3b deposition on ΔadcAII mutants and ΔadcAII fixed SpnD39III allele transformants (alleles A to F) as a proportion of the fluorescence index for the wild-type normal-capsule-thickness D39 strain. The data were measured using flow cytometry after preincubation in 30% human serum. Error bars represent SDs, and 10 transformants were tested for each double mutant strain. For all mutant strains, the P value for results compared to D39 was \leq 0.001 (unpaired t tests). (B) Examples of flow cytometry histograms for C3b/iC3b deposition on WT D39 (dark gray line) and one ΔadcAII/SpnD39IIID allele (light gray line) double mutant transformant. Gray shading indicates the results for bacteria incubated in PBS alone.

TABLE 5 Mutation construction, capsule phenotype, and (where available) cps2E gene genome sequence data for S. pneumoniae strains

aaa, amino acid position.

phenotypes are exaggerated versions of the well-described effects of the capsule on S. pneumoniae evasion of host immunity (7), demonstrating that under a normal level of expression the capsule effects on immune evasion have not reached maximal potential. Previous data have shown that capsule expression comes at a metabolic cost which inhibits growth when cultured in defined medium and that the capsule prevents adhesion by respiratory epithelium (26, 33, 34). However, surprisingly, these negative aspects of capsule expression were not identified with the hyperencapsulated Δ adcAll strain. The serotype 2 S. pneumoniae capsule repeating unit is a hexasaccharide consisting of one glucuronic acid, two glucoses, and three rhamnoses (6, 35). NMR demonstrated that the relative proportion of glucose to rhamnose was altered in the Δ adcAII strain compared to WT D39, shifting from almost 1 to 1 in the latter to closer to the expected 2-to-3 ratio. This would be compatible with an increased proportion of the total S. pneumoniae glucose pool being used for capsule production. The larger comparative increase in capsule width compared to changes in monosaccharide quantity suggests the organization of the capsule may have been altered, perhaps with more loosely packed but longer capsule strands in the Δ adcAll strain compared to D39.

Why there is increased expression of the capsule in the Δ adcAll strain is not clear. The close linkage to adcAII suggests a role for disruption of zinc utilization, yet the hyperencapsulated phenotype did not occur with mutation of the other S. pneumoniae zinc uptake lipoprotein gene adcA (19) and was not affected by zinc availability. Combined deletion of adcA and adcAII was also not associated with the hyperencapsulated phenotype, but the double mutation had major effects on S. pneumoniae

physiology (19) which could have obscured or suppressed the mucoid phenotype. Overall regulation of S. pneumoniae capsule expression is poorly understood and is further complicated by the large number of different capsular carbohydrate structures with potentially significant differences in regulatory mechanisms. Factors affecting thickness of the capsule layer include regulation of cps locus gene expression by RitR (an orphan two-component signal transduction component) (36), CpsR (a GntR family regulator) (37), and RegM (38), as well as the conserved 5' cpsABCD (also termed wzg, wzh, wzd, and wze) genes of the cps locus (39–41). Two S. pneumoniae quorum-sensing systems (LuxS/AI-2 and the Rgg/small hydrophobic peptide system) increase capsule thickness (42–44), which can also be regulated independently of gene transcription by the supply of capsule monosaccharide precursors (45) or by increased capsule shedding mediated by LytA (12). However, our transcriptome analysis did not identify increased cps locus gene expression or any effects on the abovementioned known regulators of capsule expression in the AadcAll strain.

Another potential mechanism causing the hyperencapsulated phenotype in the ΔadcAII mutant was identified by effects on transcription of the SPD_0450-0453 locus. This encodes the SpnD39III (ST5556II) type I restriction-modification system, allelic variants of which correlate with capsule thickness for several serotypes (14–16). We found that the hyperencapsulated phenotype of Δ adcAll mutants was associated with a predominance of either the SpnD39IIIC or F allelic variant, whereas SpnD39IIID was the dominant allele for the majority of Δ adcAII mutants with normal capsule thickness. This link between the hyperencapsulated phenotype of the Δ adcAII strain and specific alleles of the SpnD39III (ST5556II) system seems unlikely to be coincidental given the known effects of this restriction-modification system on capsule expression. However, transformation with fixed SpnD39III (ST5556II) alleles, including those associated with normal capsule thickness (A and D), did not alter the hyperencapsulated phenotype of the *AadcAII* mutant, showing that any effects of SpnD39III alleles on the capsule thickness of the Δ adcAII mutation are not readily reversed by switching alleles. This situation is further confused by the similarity in allele composition of the wild-type D39 strain and the AIIL AadcAII mutant and by differences between our data and published papers in which SpnD39III alleles are linked to thick or thin capsule phenotypes. Manso et al. found that A, E, and F allele strains were largely opaque but C strains were more transparent, Li et al. found that only E allele strains (termed hsdSa in their paper) were opaque, and Oliver et al. found that the A and B alleles were opaque and the others transparent (14–16). The presumed mechanism of capsule regulation by SpnD39III is differential methylation of genes or regulatory regions (14 , 15), but the genes involved remain undetermined. Our transcriptome data have identified multiple additional genes showing differential expression between hyperencapsulated Δ adcAll strains and wild-type D39 or a normal-capsule-width Δ adcAII mutant, some of which could be involved in mediating increased capsule expression. These include three operons annotated as being involved in pyrimidine metabolism, suggesting a potential role for pyrimidine in controlling capsule expression. Which genes showing differential expression between the Δ adcAII strains and WT D39 strains are involved in the capsular phenotype and whether differential regulation is related to differences in methylation will require considerably more detailed genetic studies.

Interestingly, 50% of independently obtained hyperencapsulated AadcAll strains contained an identical nonsynonymous SNP affecting the cps locus gene csp2E. The SNP is predicted to affect the cytoplasmic tail of Csp2E, a glucose phosphate transferase that initiates the assembly of capsule components on the cell membrane and is partially conserved among most capsular serotypes (39). Point mutations of cps2E that affect capsule expression have been previously described (32 , 39), suggesting a causative role for this SNP for the Δ adcAII-related capsule phenotypes. However, the same SNP was not present in one lineage of Δ adcAII with increased capsule thickness (the original transformant and four back-crossed derivatives) and was also identified in one out of eight normal-capsule-thickness Δ adcAII strains. All the unencapsulated Δ adcAII transformants also contained the same SNP in $csp2E$ predicted to introduce a stop codon.

This high frequency of $cspE2$ stop codon mutations suggests that partial deletion of adcAII causes significant physiology stress to S. pneumoniae that may induce loss of capsule production as an escape mutation.

To conclude, we have identified that in the S. pneumoniae D39 strain a hyperencapsulated phenotype is an unexpected consequence of targeted mutation of adcAII, which encodes a zinc ABC transporter lipoprotein. This strain will be a useful tool for investigating how the capsule affects S. pneumoniae interactions with the host. The hyperencapsulated phenotype partially correlated with both a nonsynonymous SNP in cps2E and changes in allelic dominance within the SpnD39III (ST5556II) restrictionmodification system. Further investigation of genes showing differential expression between normal and hyperencapsulated D39 strains could help to further identify the underlying mechanism(s) controlling S. pneumoniae capsule thickness.

MATERIALS AND METHODS

Bacterial strains and growth conditions. The ΔadcAII, ΔphtD, ΔadcA/adcAII, and ΔadcAII/phtD mutant strains were created either in the wild type or in the ΔcpsD D39 strain as well as in wild-type serotype 4 (TIGR4), 6A, 6B (strains 6Aa and 6Ba, respectively, from the work of Hyams et al. [31]), and 17F (46) strains by gene replacement using genomic DNA or PCR-amplified fragments obtained from the corresponding R6 mutants and standard transformation protocols for S. pneumoniae (19). The cat and kana genes were inserted in the reverse orientation without promoter or terminator sequences to avoid affecting expression of adjacent genes. Mutant identities were verified by PCR with primers flanking the cloned regions. S. pneumoniae was grown at 37°C with 5% CO₂ in air in THY or on Columbia agar containing 5% blood. Working stocks grown to an optical density (OD) of 0.4 (\sim 10⁸ CFU/ml) were made using THY and stored at -80°C in 10% glycerol as single-use aliquots. CFU were confirmed by colony counting of log_{10} serial dilutions of bacteria cultured overnight on 5% Columbia blood agar. Growth curves were determined by measuring OD₅₉₅ for bacteria cultured in 2.5 ml of THY or chemically defined medium (CDM) supplemented with 33 μ M Zn in 24-well plates sealed with a transparent film and incubated at 37°C in a FLUOstar reader. To measure blood growth, 1×10^6 CFU/ml of S. pneumoniae was inoculated into 1 ml of heparinized human blood and incubated at 37°C, with plating of serial dilutions at 0, 4, and 6 h to assess bacterial CFU.

Capsule size measurement and microscopy. An indirect method was developed to measure capsule size by determining the size of the bacterial pellet. Briefly, 12 ml of culture was centrifuged, the pellet was resuspended in 120 μ l of PBS, and 35 μ l was loaded in a microcapillary tube. After centrifugation for 15 min at 800 \times g , the height of the pellet within the tube was measured with a ruler. Electron microscopy of mid-log-phase S. pneumoniae fixed in 3% paraformaldehyde (PAF) was performed using a ruthenium red and London resin capsule-preserving protocol as previously described (33). Capsule thickness was calculated by direct measurement of the surface layer for 30 randomly chosen S. pneumoniae bacteria/strain using ImageJ software.

Confocal microscopy on bacteria was performed using an Olympus FV1000 confocal laser scanning microscope with a 63 \times objective. Bacteria were fixed for 30 min with 4% PFA (Sigma) on slides (Thermo Scientific; SuperFrost Plus 10149870) and subsequently stained with anti-serotype 2 antibody (Statens Serum Institute) plus Alexa Fluor 546-conjugated anti-rabbit antibody. DNA was stained with 4',6diamidino-2-phenylindole (DAPI).

Capsular polysaccharide extraction and quantification. Capsular polysaccharides were extracted from 1 liter of culture, and bacteria were resuspended in 10 ml of 0.15 M Tris buffer (pH 8) supplemented with 0.1% deoxycholate and incubated for 10 min at 37°C and then for 35 min at 50°C. Cell debris was removed by centrifugation under acidic condition. Proteins were eliminated from the supernatant by two successive extractions using a 5:1 ratio of chloroform and butanol, before precipitating capsular polysaccharides in 80% ethanol. Pellets were dried, resuspended in 0.1 M phosphate buffer (pH 7.2), and incubated with DNase and RNase for 1 h at 37°C, and then trypsin was added for 2 h at 37°C before purification of capsular polysaccharide by ion exchange on a column of DEAE Sepharose. Monosaccharide composition was established by GC and GC-MS as alditol acetate derivatives. Briefly, samples were hydrolyzed in 4 M trifluoroacetic acid (TFA) for 4 h at 100°C and reduced with sodium borohydride in 0.05 M NaOH for 4 h. Reduction was stopped by dropwise addition of acetic acid until pH 6 was reached, and borate salts were codistilled by repetitive evaporation in dry methanol. Peracetylation was performed in acetic anhydride at 100°C for 2 h. All monosaccharide derivatives were identified according to their specific retention times and electron ionization MS (EI-MS) fragmentation patterns (47).

Phagocytosis, neutrophil killing, complement deposition, and adhesion assays. Flow cytometry phagocytosis and complement deposition assays were performed as previously described (7 , 48) using S. pneumoniae incubated for 30 min with human serum (25%), human heat-inactivated serum (25%), or just Hanks balanced salt solution (HBSS) medium. For macrophage phagocytosis, THP-1 monocytes cultured in suspension in RPMI medium supplemented with 10% fetal bovine serum (FBS) were treated for 24 h with 10 nM phorbol 12-myristate 13-acetate (PMA) to induce cell adhesion and macrophage differentiation. Flow cytometry was performed using a fluorescence-activated cell sorting (FACS) Verse machine (BD), and the data were analyzed with FlowJo software. For neutrophil killing assays, fresh human neutrophils were purified using a magnetically activated cell sorting (MACS) neutrophil isolation kit (Miltenyi Biotec) and resuspended in HBSS medium at a concentration of 1×10^6 cells/ml. S.

pneumoniae previously incubated with human sera was incubated with the neutrophils at a multiplicity of infection (MOI) of 1:500 (bacteria to neutrophils) in a 48-well plate for 45 min at 37°C. Adhesion assays were performed on Detroit 562 human nasopharyngeal cells (ATCC CCL-138) as previously described (34, 49) using 3×10^5 cells/well seeded into 24-well plates, infected with MOIs of 25 and 50, and incubated for 1 h before being washed three times with PBS, followed by addition of Dulbecco's modified Eagle's medium (d-MEM)-1% saponin for 10 min and plating of serial dilutions to count bacterial CFU.

Genome sequencing and transcriptional microarray analysis. Mutant strains were genome sequenced by the Wellcome Trust Centre for Human Genetics (Oxford, United Kingdom) using an Illumina MiSeq sequencer. Sequences were assembled using Velvet, annotated using Prokka, and mapped to the published D39 (R00000036) reference genome. Bases and single-nucleotide variants were identified using the SAMtools "mpileup" command and Bcftools. Sites were filtered to a minimum depth of five reads at each and a single-nucleotide variant quality of 25, and the Integrated Genome Viewer was used to visualize mapping and coverage. Gene transcriptome microarrays were performed as described previously (34). Briefly, RNA was extracted with the RNeasy minikit (Qiagen), and labeled cDNA was prepared using Cy3-dCTP (GE Healthcare, United Kingdom) and SuperScript II reverse transcriptase with random hexamer primers (Life Technologies). Labeled cDNA was hybridized overnight to the $B\mu G@S$ SPv1.4.0 Agilent SurePrint platform (Agilent Technologies) microarray designed by the Bacterial Microarray Group at St. George's, University of London. After hybridization, washed slides were scanned immediately, using an Agilent high-resolution microarray scanner, at $5-\mu m$ resolution; scanned images were quantified using Feature Extraction software v 10.7.3.1 (Agilent Technologies); and statistical analysis of raw intensity data was performed in GeneSpring v14.9.1 (Agilent Technologies). Data for 3 independent biological replicate experiments were analyzed and normalized using a 75th percentile shift plus baseline normalized to the median for the related control sample for each biological replicate. Statistically significant ($P < 0.05$) differences between strains were identified in an unpaired t test with Benjamini and Hochberg false-discovery rate correction.

Assessing allelic variants of SpnD39III. Primers AMRE74L (5' 6-carboxyfluorescein [FAM] label, FAM-GGAAACTGAGATATTTCGTGGTGATGATGGGA) and AMRE59 (CCTGATCGAGCGGAAGAATATTTCTGCC GAGGTTGCC) were used to PCR amplify a 4.2-kb fragment from S. pneumoniae under the following conditions: denaturation at 95°C for 5 min, followed by 40 cycles of 1 min of denaturation at 95°C, 1 min of annealing at 68°C, and 5 min of extension at 68°C, with a final extension of 10 min at 68°C. PCR products were digested using 1 U Dral, 2 U PleI, and $1 \times$ CutSmart Buffer (all from New England Biolabs, United Kingdom) in a 20-µl volume. Each FAM-labeled SpnIII variant has a unique size that can be distinguished through capillary electrophoresis on an ABI Prism gene analyzer (Applied Biosystems, USA) and analyzed using Peak Scanner v1.0 software. Genomic DNA for transformation using locked SpnD39III (ST5556II) alleles due to an inactivated creX gene was obtained from preexisting strains (14).

Animal models of infection. All animal experiments conformed to institutional and United Kingdom Home Office guidelines and regulations. Outbred CD1 sex-matched white mice were used for infection experiments using established models of infection (50–52). For the nasopharyngeal colonization model, 10⁷ CFU of bacteria in 10 μ l was administered by intranasal inoculation under light halothane general anesthesia, and nasal washes were obtained after 5 days. Mice were inoculated by intraperitoneal (i.p.) injection of 5×10^4 CFU for the sepsis model and by intranasal (i.n.) inoculation under isoflurane inhalational anesthesia of 5 \times 10⁶ CFU for the pneumonia model. Mice were sacrificed after 24 h (i.p.) or 48 h (i.n.), and serial dilutions of blood and lung homogenates were plated to enumerate target organ CFU. For the sepsis model, disease development was also monitored by observing mice three times a day $(n = 8)$. For colonization and sepsis competitive infection models, mice were inoculated with a 50/50 ratio of D39 wild type and ΔadcAII strain to determine the competitive index (CI; ratio of mutant to WT strain recovered from mice divided by the ratio of mutant to WT strain in the inoculum).

Statistical analysis. Statistical analyses were conducted using Prism 7 (Graph Pad, USA). Parametric data are presented as means, and error bars represent standard deviations. Nonparametric date were analyzed using the Mann-Whitney U test. For the disease development model, data were analyzed using the log rank (Mantel-Cox) test.

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REFERENCES

- 1. Yahiaoui RY, den Heijer C, van Bijnen EM, Paget WJ, Pringle M, Goossens H, Bruggeman CA, Schellevis FG, Stobberingh EE, APRES Study Team. 2016. Prevalence and antibiotic resistance of commensal Streptococcus pneumoniae in nine European countries. Future Microbiol 11:737–744. [https://doi.org/10.2217/fmb-2015-0011.](https://doi.org/10.2217/fmb-2015-0011)
- 2. Gillespie SH1, Balakrishnan I. 2000. Pathogenesis of pneumococcal in-

fection. J Med Microbiol 49:1057–1067. [https://doi.org/10.1099/0022](https://doi.org/10.1099/0022-1317-49-12-1057) [-1317-49-12-1057.](https://doi.org/10.1099/0022-1317-49-12-1057)

- 3. Weiser JN, Ferreira DM, Paton JC. 2018. Streptococcus pneumoniae: transmission, colonization and invasion. Nat Rev Microbiol 16:355–367. [https://](https://doi.org/10.1038/s41579-018-0001-8) [doi.org/10.1038/s41579-018-0001-8.](https://doi.org/10.1038/s41579-018-0001-8)
- 4. Nelson AL, Roche AM, Gould JM, Chim K, Ratner AJ, Weiser JN. 2007.

Capsule enhances pneumococcal colonization by limiting mucusmediated clearance. Infect Immun 75:83–90. [https://doi.org/10.1128/IAI](https://doi.org/10.1128/IAI.01475-06) [.01475-06.](https://doi.org/10.1128/IAI.01475-06)

- 5. Henrichsen J. 1995. Six newly recognized types of Streptococcus pneumoniae. J Clin Microbiol 33:2759-2762. [https://doi.org/10.1128/JCM.33](https://doi.org/10.1128/JCM.33.10.2759-2762.1995) [.10.2759-2762.1995.](https://doi.org/10.1128/JCM.33.10.2759-2762.1995)
- 6. Yother J. 2011. Capsules of Streptococcus pneumoniae and other bacteria: paradigms for polysaccharide biosynthesis and regulation. Annu Rev Microbiol 65:563–581. [https://doi.org/10.1146/annurev.micro](https://doi.org/10.1146/annurev.micro.62.081307.162944) [.62.081307.162944.](https://doi.org/10.1146/annurev.micro.62.081307.162944)
- 7. Hyams C, Camberlein E, Cohen JM, Bax K, Brown JS. 2010. The Streptococcus pneumoniae capsule inhibits complement activity and neutrophil phagocytosis by multiple mechanisms. Infect Immun 78:704 –715. [https://doi.org/10.1128/IAI.00881-09.](https://doi.org/10.1128/IAI.00881-09)
- 8. Domenech A, Ardanuy C, Grau I, Calatayud L, Pallares R, Fenoll A, Brueggemann AB, Liñares J. 2014. Evolution and genetic diversity of the Spain23F-ST81 clone causing adult invasive pneumococcal disease in Barcelona (1990 –2012). J Antimicrob Chemother 69:924 –931. [https://](https://doi.org/10.1093/jac/dkt473) [doi.org/10.1093/jac/dkt473.](https://doi.org/10.1093/jac/dkt473)
- 9. Kim JO, Weiser JN. 1998. Association of intrastrain phase variation in quantity of capsular polysaccharide and teichoic acid with the virulence of Streptococcus pneumoniae. J Infect Dis 177:368-377. [https://doi.org/](https://doi.org/10.1086/514205) [10.1086/514205.](https://doi.org/10.1086/514205)
- 10. Oggioni MR, Trappetti C, Kadioglu A, Cassone M, Iannelli F, Ricci S, Andrew PW, Pozzi G. 2006. Switch from planktonic to sessile life: a major event in pneumococcal pathogenesis. Mol Microbiol 61:1196 –1210. [https://doi.org/10.1111/j.1365-2958.2006.05310.x.](https://doi.org/10.1111/j.1365-2958.2006.05310.x)
- 11. Kohler S, Voß F, Gómez Mejia A, Brown JS, Hammerschmidt S. 2016. Pneumococcal lipoproteins involved in bacterial fitness, virulence, and immune evasion. FEBS Lett 590:3820 –3839. [https://doi.org/10.1002/](https://doi.org/10.1002/1873-3468.12352) [1873-3468.12352.](https://doi.org/10.1002/1873-3468.12352)
- 12. Kietzman CC, Gao G, Mann B, Myers L, Tuomanen EI. 2016. Dynamic capsule restructuring by the main pneumococcal autolysin LytA in response to the epithelium. Nat Commun 7:10859. [https://doi.org/10](https://doi.org/10.1038/ncomms10859) [.1038/ncomms10859.](https://doi.org/10.1038/ncomms10859)
- 13. Sanchez CJ, Hinojosa CA, Shivshankar P, Hyams C, Camberlein E, Brown JS, Orihuela CJ. 2011. Changes in capsular serotype alter the surface exposure of pneumococcal adhesins and impact virulence. PLoS One 6:e26587. [https://doi.org/10.1371/journal.pone.0026587.](https://doi.org/10.1371/journal.pone.0026587)
- 14. Manso AS, Chai MH, Atack JM, Furi L, De Ste Croix M, Haigh R, Trappetti C, Ogunniyi AD, Shewell LK, Boitano M, Clark TA, Korlach J, Blades M, Mirkes E, Gorban AN, Paton JC, Jennings MP, Oggioni MR. 2014. A random six-phase switch regulates pneumococcal virulence via global epigenetic changes. Nat Commun 5:5055. [https://doi.org/10.1038/ncomms6055.](https://doi.org/10.1038/ncomms6055)
- 15. Li J, Li JW, Feng Z, Wang J, An H, Liu Y, Wang Y, Wang K, Zhang X, Miao Z, Liang W, Sebra R, Wang G, Wang WC, Zhang JR. 2016. Epigenetic switch driven by DNA inversions dictates phase variation in Streptococcus pneumoniae. PLoS Pathog 12:e1005762. [https://doi.org/10.1371/](https://doi.org/10.1371/journal.ppat.1005762) [journal.ppat.1005762.](https://doi.org/10.1371/journal.ppat.1005762)
- 16. Oliver MB, Basu Roy A, Kumar R, Lefkowitz EJ, Swords WE. 2017. Streptococcus pneumoniae TIGR4 phase-locked opacity variants differ in virulence phenotypes. mSphere 2:e00386-17. [https://doi.org/10.1128/mSphere.00386](https://doi.org/10.1128/mSphere.00386-17) [-17.](https://doi.org/10.1128/mSphere.00386-17)
- 17. Brown JS, Gilliland SM, Holden DW. 2001. A Streptococcus pneumoniae pathogenicity island encoding an ABC transporter involved in iron uptake and virulence. Mol Microbiol 40:572–585. [https://doi.org/10](https://doi.org/10.1046/j.1365-2958.2001.02414.x) [.1046/j.1365-2958.2001.02414.x.](https://doi.org/10.1046/j.1365-2958.2001.02414.x)
- 18. Johnston JW, Myers LE, Ochs MM, Benjamin WH, Briles DE, Hollingshead SK. 2004. Lipoprotein PsaA in virulence of Streptococcus pneumoniae: surface accessibility and role in protection from superoxide. Infect Immun 72:5858 –5867. [https://doi.org/10.1128/IAI.72.10.5858-5867.2004.](https://doi.org/10.1128/IAI.72.10.5858-5867.2004)
- 19. Bayle L, Chimalapati S, Schoehn G, Brown J, Vernet T, Durmort C. 2011. Zinc uptake by Streptococcus pneumoniae depends on both AdcA and AdcAll and is essential for normal bacterial morphology and virulence. Mol Microbiol 82:904 –916. [https://doi.org/10.1111/j.1365-2958.2011.07862.x.](https://doi.org/10.1111/j.1365-2958.2011.07862.x)
- 20. Dintilhac A, Claverys JP. 1997. The adc locus, which affects competence for genetic transformation in Streptococcus pneumoniae, encodes an ABC transporter with a putative lipoprotein homologous to a family of streptococcal adhesins. Res Microbiol 148:119 –131. [https://doi.org/10](https://doi.org/10.1016/S0923-2508(97)87643-7) [.1016/S0923-2508\(97\)87643-7.](https://doi.org/10.1016/S0923-2508(97)87643-7)
- 21. Loisel E, Jacquamet L, Serre L, Bauvois C, Ferrer JL, Vernet T, Di Guilmi AM, Durmort C. 2008. AdcAII, a new pneumococcal Zn-binding protein homologous with ABC transporters: biochemical and structural analysis. J Mol Biol 381:594 – 606. [https://doi.org/10.1016/j.jmb.2008.05.068.](https://doi.org/10.1016/j.jmb.2008.05.068)
- 22. Loisel E, Chimalapati S, Bougault C, Imberty A, Gallet B, Di Guilmi AM, Brown JS, Vernet T, Durmort C. 2011. Biochemical characterization of the histidine triad protein PhtD as a cell surface zinc-binding protein of pneumococcus. Biochemistry 50:3551–3558. [https://doi.org/10.1021/bi200012f.](https://doi.org/10.1021/bi200012f)
- 23. Plumptre CD, Hughes CE, Harvey RM, Eijkelkamp BA, McDevitt CA, Paton JC. 2014. Overlapping functionality of the Pht proteins in zinc homeostasis of Streptococcus pneumoniae. Infect Immun 82:4315– 4324. [https://](https://doi.org/10.1128/IAI.02155-14) [doi.org/10.1128/IAI.02155-14.](https://doi.org/10.1128/IAI.02155-14)
- 24. Eijkelkamp BA, Pederick VG, Plumptre CD, Harvey RM, Hughes CE, Paton JC, McDevitt CA. 2016. The first histidine triad motif of PhtD is critical for zinc homeostasis in Streptococcus pneumoniae. Infect Immun 84: 407– 415. [https://doi.org/10.1128/IAI.01082-15.](https://doi.org/10.1128/IAI.01082-15)
- 25. Brown LR, Gunnell SM, Cassella AN, Keller LE, Scherkenbach LA, Mann B, Brown MW, Hill R, Fitzkee NC, Rosch JW, Tuomanen EI, Thornton JA. 2016. AdcAII of Streptococcus pneumoniae affects pneumococcal invasiveness. PLoS One 11:e0146785. [https://doi.org/10.1371/journal.pone](https://doi.org/10.1371/journal.pone.0146785) [.0146785.](https://doi.org/10.1371/journal.pone.0146785)
- 26. Hathaway LJ, Brugger SD, Morand B, Bangert M, Rotzetter JU, Hauser C, Graber WA, Gore S, Kadioglu A, Mühlemann K. 2012. Capsule type of Streptococcus pneumoniae determines growth phenotype. PLoS Pathog 8:e1002574. [https://doi.org/10.1371/journal.ppat.1002574.](https://doi.org/10.1371/journal.ppat.1002574)
- 27. Giefing-Kröll C, Jelencsics KE, Reipert S, Nagy E. 2011. Absence of pneumococcal PcsB is associated with overexpression of LysM domain-containing proteins. Microbiology 157:1897–1909. [https://doi.org/10.1099/mic.0](https://doi.org/10.1099/mic.0.045211-0) [.045211-0.](https://doi.org/10.1099/mic.0.045211-0)
- 28. Donini S, Ferraris DM, Miggiano R, Massarotti A, Rizzi M. 2017. Structural investigations on orotate phosphoribosyltransferase from Mycobacterium tuberculosis, a key enzyme of the de novo pyrimidine biosynthesis. Sci Rep 7:1180. [https://doi.org/10.1038/s41598-017-01057-z.](https://doi.org/10.1038/s41598-017-01057-z)
- 29. Rowland P, Nørager S, Jensen KF, Larsen S. 2000. Structure of dihydroorotate dehydrogenase B: electron transfer between two flavin groups bridged by an iron-sulphur cluster. Structure 8:1227–1238. [https://doi](https://doi.org/10.1016/s0969-2126(00)00530-x) [.org/10.1016/s0969-2126\(00\)00530-x.](https://doi.org/10.1016/s0969-2126(00)00530-x)
- 30. Arioli S, Monnet C, Guglielmetti S, Mora D. 2009. Carbamoylphosphate synthetase activity is essential for the optimal growth of Streptococcus thermophilus in milk. J Appl Microbiol 107:348 –354. [https://doi.org/10](https://doi.org/10.1111/j.1365-2672.2009.04213.x) [.1111/j.1365-2672.2009.04213.x.](https://doi.org/10.1111/j.1365-2672.2009.04213.x)
- 31. Hyams C, Yuste J, Bax K, Camberlein E, Weiser JN, Brown JS. 2010. Streptococcus pneumoniae resistance to complement-mediated immunity is dependent on the capsular serotype. Infect Immun 78:716 –725. [https://doi.org/10.1128/IAI.01056-09.](https://doi.org/10.1128/IAI.01056-09)
- 32. Kim JO, Romero-Steiner S, Sørensen UB, Blom J, Carvalho M, Barnard S, Carlone G, Weiser JN. 1999. Relationship between cell surface carbohydrates and intrastrain variation on opsonophagocytosis of Streptococcus pneumoniae. Infect Immun 67:2327–2333. [https://doi.org/10.1128/IAI.67](https://doi.org/10.1128/IAI.67.5.2327-2333.1999) [.5.2327-2333.1999.](https://doi.org/10.1128/IAI.67.5.2327-2333.1999)
- 33. Hammerschmidt S, Wolff S, Hocke A, Rosseau S, Muller E, Rohde M. 2005. Illustration of pneumococcal polysaccharide capsule during adherence and invasion of epithelial cells. Infect Immun 73:4653– 4667. [https://doi](https://doi.org/10.1128/IAI.73.8.4653-4667.2005) [.org/10.1128/IAI.73.8.4653-4667.2005.](https://doi.org/10.1128/IAI.73.8.4653-4667.2005)
- 34. Schaffner TO, Hinds J, Gould KA, Wüthrich D, Bruggmann R, Küffer M, Mühlemann K, Hilty M, Hathaway LJ. 2014. A point mutation in cpsE renders Streptococcus pneumoniae nonencapsulated and enhances its growth, adherence and competence. BMC Microbiol 14:210. [https://doi](https://doi.org/10.1186/s12866-014-0210-x) [.org/10.1186/s12866-014-0210-x.](https://doi.org/10.1186/s12866-014-0210-x)
- 35. Bentley SD, Aanensen DM, Mavroidi A, Saunders D, Rabbinowitsch E, Collins M, Donohoe K, Harris D, Murphy L, Quail MA, Samuel G, Skovsted IC, Kaltoft MS, Barrell B, Reeves PR, Parkhill J, Spratt BG. 2006. Genetic analysis of the capsular biosynthetic locus from all 90 pneumococcal serotypes. PLoS Genet 2:e31. [https://doi.org/10.1371/journal.pgen.0020031.](https://doi.org/10.1371/journal.pgen.0020031)
- 36. Ulijasz AT, Andes DR, Glasner JD, Weisblum B. 2004. Regulation of iron transport in Streptococcus pneumoniae by RitR, an orphan response regulator. J Bacteriol 186:8123– 8136. [https://doi.org/10.1128/JB.186.23](https://doi.org/10.1128/JB.186.23.8123-8136.2004) [.8123-8136.2004.](https://doi.org/10.1128/JB.186.23.8123-8136.2004) (Erratum, 188:6045, 2006, [https://doi.org/10.1128/JB](https://doi.org/10.1128/JB.00861-06) [.00861-06.](https://doi.org/10.1128/JB.00861-06))
- 37. Wu K, Xu H, Zheng Y, Wang L, Zhang X, Yin Y. 2016. CpsR, a GntR family regulator, transcriptionally regulates capsular polysaccharide biosynthesis and governs bacterial virulence in Streptococcus pneumoniae. Sci Rep 6:29255. [https://doi.org/10.1038/srep29255.](https://doi.org/10.1038/srep29255)
- 38. Giammarinaro P, Paton JC. 2002. Role of RegM, a homologue of the catabolite repressor protein CcpA, in the virulence of Streptococcus pneumoniae. Infect Immun 70:5454 –5461. [https://doi.org/10.1128/iai.70](https://doi.org/10.1128/iai.70.10.5454-5461.2002) [.10.5454-5461.2002.](https://doi.org/10.1128/iai.70.10.5454-5461.2002)
- 39. James DB, Gupta K, Hauser JR, Yother J. 2013. Biochemical activities of

Streptococcus pneumoniae serotype 2 capsular glycosyltransferases and significance of suppressor mutations affecting the initiating glycosyltransferase Cps2E. J Bacteriol 195:5469 –5478. [https://doi.org/10.1128/JB](https://doi.org/10.1128/JB.00715-13) [.00715-13.](https://doi.org/10.1128/JB.00715-13)

- 40. Geno KA, Hauser JR, Gupta K, Yother J. 2014. Streptococcus pneumoniae phosphotyrosine phosphatase CpsB and alterations in capsule production resulting from changes in oxygen availability. J Bacteriol 196: 1992–2003. [https://doi.org/10.1128/JB.01545-14.](https://doi.org/10.1128/JB.01545-14)
- 41. Morona JK, Miller DC, Morona R, Paton JC. 2004. The effect that mutations in the conserved capsular polysaccharide biosynthesis genes cpsA, cpsB, and cpsD have on virulence of Streptococcus pneumoniae. J Infect Dis 189:1905–1913. [https://doi.org/10.1086/383352.](https://doi.org/10.1086/383352)
- 42. Trappetti C, McAllister LJ, Chen A, Wang H, Paton AW, Oggioni MR, McDevitt CA, Paton JC. 2017. Autoinducer 2 signaling via the phosphotransferase FruA drives galactose utilization by Streptococcus pneumoniae, resulting in hypervirulence. mBio 8:e02269-16. [https://doi.org/](https://doi.org/10.1128/mBio.02269-16) [10.1128/mBio.02269-16.](https://doi.org/10.1128/mBio.02269-16)
- 43. Yadav MK, Vidal JE, Go YY, Kim SH, Chae SW, Song JJ. 2018. The LuxS/AI-2 quorum-sensing system of Streptococcus pneumoniae is required to cause disease, and to regulate virulence- and metabolismrelated genes in a rat model of middle ear infection. Front Cell Infect Microbiol 8:138. [https://doi.org/10.3389/fcimb.2018.00138.](https://doi.org/10.3389/fcimb.2018.00138)
- 44. Junges R, Salvadori G, Shekhar S, Åmdal HA, Periselneris JN, Chen T, Brown JS, Petersen FC. 2017. A quorum-sensing system that regulates Streptococcus pneumoniae biofilm formation and surface polysaccharide production. mSphere 2:e00324-17. [https://doi.org/10.1128/mSphere.00324-17.](https://doi.org/10.1128/mSphere.00324-17)
- 45. Ventura CL, Cartee RT, Forsee WT, Yother J. 2006. Control of capsular polysaccharide chain length by UDP-sugar substrate concentrations in Streptococcus pneumoniae. Mol Microbiol 61:723–733. [https://doi.org/10](https://doi.org/10.1111/j.1365-2958.2006.05259.x) [.1111/j.1365-2958.2006.05259.x.](https://doi.org/10.1111/j.1365-2958.2006.05259.x)
- 46. Domenech de Cellès M, Opatowski L, Salomon J, Varon E, Carbon C, Boëlle PY, Guillemot D. 2011. Intrinsic epidemicity of Streptococcus pneumoniae depends on strain serotype and antibiotic susceptibility

pattern. Antimicrob Agents Chemother 55:5255–5261. [https://doi.org/](https://doi.org/10.1128/AAC.00249-11) [10.1128/AAC.00249-11.](https://doi.org/10.1128/AAC.00249-11)

- 47. Faille C, Lequette Y, Ronse A, Slomianny C, Garénaux E, Guerardel Y. 2010. Morphology and physico-chemical properties of Bacillus spores surrounded or not with an exosporium. Consequences on their ability to adhere to stainless steel. Int J Food Microbiol 143:125–135. [https://doi](https://doi.org/10.1016/j.ijfoodmicro.2010.07.038) [.org/10.1016/j.ijfoodmicro.2010.07.038.](https://doi.org/10.1016/j.ijfoodmicro.2010.07.038)
- 48. Wilson RM, Galvin AM, Robins RA, Reeves WG. 1985. A flow cytometric method for the measurement of phagocytosis by polymorphonuclear leucocytes. J Immunol Methods 76:247–253. [https://doi.org/10.1016/](https://doi.org/10.1016/0022-1759(85)90301-1) [0022-1759\(85\)90301-1.](https://doi.org/10.1016/0022-1759(85)90301-1)
- 49. Sanchez CJ1, Kumar N, Lizcano A, Shivshankar P, Dunning Hotopp JC, Jorgensen JH, Tettelin H, Orihuela CJ. 2011. Streptococcus pneumoniae in biofilms are unable to cause invasive disease due to altered virulence determinant production. PLoS One 6:e28738. [https://doi.org/10.1371/](https://doi.org/10.1371/journal.pone.0028738) [journal.pone.0028738.](https://doi.org/10.1371/journal.pone.0028738)
- 50. Brown JS, Ogunniyi AD, Woodrow MC, Holden DW, Paton JC. 2001. Immunization with components of two iron uptake ABC transporters protects mice against systemic Streptococcus pneumoniae infection. Infect Immun 69:6702– 6706. [https://doi.org/10.1128/IAI.69.11.6702-6706](https://doi.org/10.1128/IAI.69.11.6702-6706.2001) [.2001.](https://doi.org/10.1128/IAI.69.11.6702-6706.2001) (Erratum, 72:6755, 2004, [https://doi.org/10.1128/IAI.72.11.6755](https://doi.org/10.1128/IAI.72.11.6755.2004) [.2004.](https://doi.org/10.1128/IAI.72.11.6755.2004))
- 51. Brown JS, Hussell T, Gilliland SM, Holden DW, Paton JC, Ehrenstein MR, Walport MJ, Botto M. 2002. The classical pathway is the dominant complement pathway required for innate immunity to Streptococcus pneumoniae infection in mice. Proc Natl Acad Sci U S A 99:16969-16974. [https://doi.org/10.1073/pnas.012669199.](https://doi.org/10.1073/pnas.012669199)
- 52. Wilson R, Cohen JM, Reglinski M, Jose RJ, Chan WY, Marshall H, de Vogel C, Gordon S, Goldblatt D, Petersen FC, Baxendale H, Brown JS. 2017. Naturally acquired human immunity to pneumococcus is dependent on antibody to protein antigens. PLoS Pathog 13:e1006259. [https://doi.org/](https://doi.org/10.1371/journal.ppat.1006259) [10.1371/journal.ppat.1006259.](https://doi.org/10.1371/journal.ppat.1006259)