Molecular Epidemiology and Dynamics of Pseudomonas aeruginosa Populations in Lungs of Cystic Fibrosis Patients


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The ability to establish lifelong persistent infections is a fundamental aspect of the interactions between many pathogenic microorganisms and their mammalian hosts. One example is chronic lung infections by the opportunistic pathogen Pseudomonas aeruginosa in cystic fibrosis (CF) patients. This infection process is associated with extensive genetic adaptation and microevolution of the infecting bacteria. Through investigations of P. aeruginosa populations and infection dynamics in a group of CF patients followed at the Danish CF Clinic in Copenhagen, we have identified two distinct and dominant clones that have evolved into highly successful colonizers of CF patient airways. A significant component of the evolutionary success of these two clones has been their efficient transmissibility among the CF patients. The two clones have been present and transmitted among different CF patients for more than 2 decades. Our data also suggest that the P. aeruginosa population structure in the CF patient airways has been influenced by competition between different clones and that the two dominant clones have been particularly competitive within the lungs, which may add to their overall establishment success. In contrast, we show that adaptive traits commonly associated with establishment of chronic P. aeruginosa infections of CF patients, such as transition to the mucoid phenotype and production of virulence factors, play minor roles in the ability of the two dominant clones to spread among patients and cause long-term chronic infections. These findings suggest that hitherto-unrecognized evolutionary pathways may be involved in the development of successful and persistent P. aeruginosa colonizers of CF patient lungs.

Pseudomonas aeruginosa, the most common pathogen associated with morbidity and mortality in patients suffering from cystic fibrosis (CF), causes chronic lung infections that, once established, are difficult to eradicate even with intensive antibiotic treatment (15).

The development of chronic lung infections follows a characteristic pattern. Typically, a period of intermittent colonization with P. aeruginosa early in the life of the patient precedes the establishment of chronic infections (5, 17). These early-infesting strains typically resemble those found in the environment, being nonmucoid, fast growing, and relatively susceptible to antibiotics (16). This stage is temporary, and eventually the airways of nearly all patients become permanently colonized by P. aeruginosa.

During chronic infection, the bacterial population adapts to the environment in the CF patient airways. This process correlates with considerable genetic adaptation and accumulation of loss-of-function mutations in specific P. aeruginosa genes (40). One common mutation is in the mucA gene, which causes a transition from a nonmucoid to a mucoid, alginate-overproducing phenotype (24). Mucoid P. aeruginosa is of special interest as an indicator of chronic infection and, because of the association with poor prognosis, deteriorating lung function and tissue damage (33). Early intervention and aggressive antimicrobial chemotherapy after initial detection of P. aeruginosa have been shown to delay the transition from intermittent to chronic infection in the majority of patients (11, 27, 42).

The prevailing view of the infection process is that most of the infections are clonal and that individual CF patients acquire unique P. aeruginosa strains independently, presumably from diverse environmental sources (16, 22, 35, 36). However, several recent reports have shown the occurrence of aggressive and transmissible strains of P. aeruginosa in patients attending CF centers in Europe and Australia (2, 3, 7, 8, 10, 19, 25, 29–31, 34, 39), and some transmissible clones may in fact be more virulent than the usual infecting types, potentially resulting in poor prognoses for the patients (1, 3, 28, 37).

We have investigated these different aspects of P. aeruginosa infections in a defined group of CF patients from the CF Center in Copenhagen, Denmark. In particular, we have used molecular epidemiological tools to better understand how the bacteria infect and establish persistent infections in CF patient airways, from the early intermittent colonization in children and young adults to the chronic state of infection in older patients.

MATERIALS AND METHODS

Bacterial isolates. The P. aeruginosa strain collection examined consisted of the following: 122 isolates from 7 CF patients with long-term chronic infection...
Chronic status, pulmonary function, and microbiology of lower-airway secretions. All patients were monitored on a monthly basis by evaluation of their clinical shown in Table 1. As part of the general management structure at the CF Center, phoresis as previously described (14). Serum from all included patients was determined by crossed immunoelectrophoresis. The number of precipitating antibodies against _P. aeruginosa_ in sputum for 6 consecutive months or less when persistence was further analysis. All ing of 96 colonies per sample for differences in antibiotic resistance profiles samples taken in 2005, we furthermore routinely performed phenotypic screen- ing of 96 colonies per sample for differences in antibiotic resistance profiles and/or colony appearance. Colonies with different phenotypes were chosen for further analysis. All _P. aeruginosa_ isolates collected were frozen at –80°C.

**CF patients.** Patient data for the 22 CF patients included in this study are shown in Table 1. As part of the general management structure at the CF Center, all patients were monitored on a monthly basis by evaluation of their clinical status, pulmonary function, and microbiology of lower-airway secretions. Chronic _P. aeruginosa_ infection was defined as the persistent presence of _P. aeruginosa_ in sputum for 6 consecutive months or less when persistence was combined with the presence of two or more precipitating antibodies against _P. aeruginosa_ (18). The number of precipitating antibodies against _P. aeruginosa_ in serum from all included patients was determined by crossed immunoelectro- phoresis as previously described (14).

**P. aeruginosa genotyping.** Identification of _P. aeruginosa_ genotypes was done by single-nucleotide polymorphism (SNP) typing using AT biochips (Clondiag Chip Technologies, Germany). The biochip allows simultaneous detection of up to 77 SNPs in different genomic regions such as _oriC_ (12). Sequencing of _mucA_ was mucA1 (5′–GGTAGGCCGATGCACAATCCATTTCT 0.3′), mucA2 (5′–GGGATGCTTCAAGCAGAGACG-3′), and mucA2rev (5′–GAAGAATGACGCTTTGGAACAG-3′). MucA1 and mucA2 were used as positive controls. The accuracy of using these SNPs for genotyping is 99.7% (26) (Clondiag Chip Technologies, Germany). In addition to SNP typing, the AT chip allows detection of differences in antibiotic resistance profiles to confirm their origin from the lower airways by the presence of mucus and leukocytes (15). The sputum samples were obtained by expectoration or were previously described (15). The sputum samples were obtained by expectoration or endotracheal suction, followed by Gram staining and microscopic examination to confirm their origin from the lower airways by the presence of mucus and leukocytes (15). _P. aeruginosa_ was identified by conventional biochemical tests. _P. aeruginosa_ isolates from samples obtained during 2005 were isolated on _Pseudomonas_ isolation agar (Difco) containing ampicillin (100 μg/ml). At least one colony of each morphotype present in a sputum sample was analyzed. For samples taken in 2005, we furthermore routinely performed phenotypic screening of 96 colonies per sample for differences in antibiotic resistance profiles and/or colony appearance. Colonies with different phenotypes were chosen for further analysis. All _P. aeruginosa_ isolates collected were frozen at –80°C.

**TABLE 1. Characteristics of patients included in this study**

<table>
<thead>
<tr>
<th>Patient type</th>
<th>Patient no.</th>
<th>Yr of birth</th>
<th>Gender</th>
<th>Yr chronic infection began (duration [yr])</th>
<th>Median P. aeruginosa-specific precipitants (range) last 5 yr</th>
<th>Bacteriology(^a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intermittently colonized</td>
<td>B1</td>
<td>1990</td>
<td>M</td>
<td>—</td>
<td>1 (0–1)</td>
<td>NM</td>
</tr>
<tr>
<td></td>
<td>B4</td>
<td>2004</td>
<td>M</td>
<td>—</td>
<td>0 (0–0)</td>
<td>NM</td>
</tr>
<tr>
<td></td>
<td>B6</td>
<td>2002</td>
<td>F</td>
<td>—</td>
<td>1 (0–1)</td>
<td>NM</td>
</tr>
<tr>
<td></td>
<td>B7</td>
<td>1995</td>
<td>M</td>
<td>—</td>
<td>0 (0–1)</td>
<td>NM</td>
</tr>
<tr>
<td></td>
<td>B9</td>
<td>2001</td>
<td>F</td>
<td>—</td>
<td>0 (0–1)</td>
<td>NM</td>
</tr>
<tr>
<td></td>
<td>B10</td>
<td>1999</td>
<td>F</td>
<td>—</td>
<td>1 (0–2)</td>
<td>NM</td>
</tr>
<tr>
<td></td>
<td>B12</td>
<td>1998</td>
<td>M</td>
<td>—</td>
<td>0 (0–1)</td>
<td>M</td>
</tr>
<tr>
<td></td>
<td>B14</td>
<td>1982</td>
<td>M</td>
<td>—</td>
<td>0 (0–1)</td>
<td>NM</td>
</tr>
<tr>
<td></td>
<td>B18</td>
<td>1988</td>
<td>F</td>
<td>—</td>
<td>0 (0–0)</td>
<td>NM</td>
</tr>
<tr>
<td></td>
<td>B22</td>
<td>1996</td>
<td>F</td>
<td>—</td>
<td>0 (0–0)</td>
<td>NM</td>
</tr>
<tr>
<td></td>
<td>B23</td>
<td>1998</td>
<td>F</td>
<td>—</td>
<td>0 (0–1)</td>
<td>M</td>
</tr>
<tr>
<td></td>
<td>B29</td>
<td>1999</td>
<td>F</td>
<td>—</td>
<td>0 (0–0)</td>
<td>NM</td>
</tr>
<tr>
<td></td>
<td>B36</td>
<td>1993</td>
<td>F</td>
<td>—</td>
<td>1 (0–2)</td>
<td>NM</td>
</tr>
<tr>
<td>Recently chronically infected</td>
<td>B3</td>
<td>2002</td>
<td>F</td>
<td>2005 (&lt;1)</td>
<td>5 (0–16)</td>
<td>M/NM</td>
</tr>
<tr>
<td></td>
<td>B19</td>
<td>1979</td>
<td>F</td>
<td>2005 (&lt;1)</td>
<td>1 (0–23)</td>
<td>NM</td>
</tr>
<tr>
<td></td>
<td>p6</td>
<td>1975</td>
<td>F</td>
<td>1984 (21)</td>
<td>27 (22–29)</td>
<td>NM(^b)</td>
</tr>
<tr>
<td></td>
<td>p7</td>
<td>1986</td>
<td>F</td>
<td>1990 (15)</td>
<td>17 (12–20)</td>
<td>NM(^b)</td>
</tr>
<tr>
<td></td>
<td>p10</td>
<td>1963</td>
<td>M</td>
<td>1980 (25)</td>
<td>25 (20–30)</td>
<td>NM(^b)</td>
</tr>
<tr>
<td></td>
<td>p11</td>
<td>1983</td>
<td>M</td>
<td>1993 (12)</td>
<td>18 (13–22)</td>
<td>M/NM</td>
</tr>
</tbody>
</table>

\(^a\) M, mucoid; NM, nonmucoid; M/NM, mucoid and nonmucoid.

\(^b\) The bacteriology records showed that mucoid variants have been found (but not stored) in occasional samples in the history of the patients.

and 28 isolates from 15 CF patients who were intermittently colonized or recently chronically infected and who were attending the Danish CF Center, Rigshospi- talaet, Copenhagen; 11 isolates from non-CF patients obtained from six different departments at Rigshospitalet, Copenhagen; 1 environmental isolate from a sanitary facility in Copenhagen; and 1 reference strain (PA01).

Isolation and identification of _P. aeruginosa_ from sputum were done as previ- ously described (15). The sputum samples were obtained by expectoration or endotracheal suction, followed by Gram staining and microscopic examination to confirm their origin from the lower airways by the presence of mucus and leukocytes (15). _P. aeruginosa_ was identified by conventional biochemical tests. _P. aeruginosa_ isolates from samples obtained during 2005 were isolated on _Pseudo- monas_ isolation agar (Difco) containing ampicillin (100 μg/ml). At least one colony of each morphotype present in a sputum sample was analyzed. For samples taken in 2005, we furthermore routinely performed phenotypic screening of 96 colonies per sample for differences in antibiotic resistance profiles and/or colony appearance. Colonies with different phenotypes were chosen for further analysis. All _P. aeruginosa_ isolates collected were frozen at –80°C.

The clonal relatedness of selected _P. aeruginosa_ isolates was also assessed by pulsed-field gel electrophoresis (PFGE) as described previously (20).

**Sequenceing of mucA and algT genes.** A 687-bp fragment covering the entire mucA region and an 831-bp fragment covering the entire algT region were amplified by standard PCR. Purified PCR products were sequenced by Macrogen’s (South Korea) sequencing service. Primers used for PCR amplification and sequencing of mucA were mucA1 (5′–CTCTGCAAGCCGTTTGTGCGAAG-3′), mucA1rev (5′–CTGCCAGAAGCCCCAGAGGAGG-3′), mucA2 (5′–GTGCGTCGGCTGACACCCAGACGCG-3′), and mucA2rev (5′–GTCGGTCTGACACCCAGACGCG-3′). MucA1, MucA2, MucA1rev, and MucA2rev were used as positive controls.

**Caenorhabditis elegans killing assay.** The killing of the _C. elegans pha-1(e2123ts) strain by _P. aeruginosa_ isolates was measured on NG agar (slow-killing
FIG. 1. Genotyping of *P. aeruginosa* isolates. (A) Isolates from young CF patients with intermittent colonization. (B) Clinical isolates from non-CF patients. Boxes below the names of the isolates indicate specific genotype classes. White boxes specify unique genotypes found only once in the data set. Labeled boxes specify that a genotype was found in another patient. The asterisk indicates mucoid isolates. Genotyping was based on SNP typing. In the SNP panel, light gray and black boxes indicate similar and different SNP sequences, respectively, compared to the PA01 sequence. Dark gray boxes indicate SNPs that were not measured. Below the SNP panel, the presence or absence of particular genes or regions in different isolates is shown by black and light gray boxes, respectively.
assay) essentially as previously described (43). The temperature-sensitive sterile C. elegans pha-1(e2123ts) strain used here grows normally at 15°C, but the allele is embryonic lethal at 25°C (38). Mutation in the pha-1 gene does not affect the immune function of C. elegans (44). The Escherichia coli and C. elegans strains used in this work were provided by the Caenorhabditis Genetics Center (University of Minnesota, Minneapolis).

RESULTS

High P. aeruginosa genotype diversity in intermittently colonized children and young adults with CF. In order to understand the P. aeruginosa population dynamics during the early stages of CF lung infection, we first determined the genotypes of a set of P. aeruginosa strains isolated from 13 Danish CF patients with intermittent P. aeruginosa colonization and with a mean age of 10.3 years (Table 1). None of the patients had elevated levels of precipitating antibodies against P. aeruginosa, and 12 of the P. aeruginosa isolates were nonmucoid and susceptible to standard antipseudomonas antibiotics (data not shown). Mucoid variants were found only in patient B23. Genotyping of the infecting isolates was performed by analyzing SNPs of conserved genomic regions using ClonDiag AT biochips, which provide strain discrimination up to a specificity of 99.7% (Materials and Methods).

Our data show that 8 of the 13 young CF patients were colonized with unique clones, whereas identical P. aeruginosa genotypes were isolated from patients B1 and B12 and from B9 and B18 (Fig. 1A). In addition, the same P. aeruginosa genotype was isolated from patient B23 and the recently chronically infected patient B3 (cf. Fig. 1A and 3A). We also genotyped a collection of 11 clinical P. aeruginosa isolates from non-CF patients admitted to the hospital as well as the reference strain PA01 and an environmental isolate. Again, 11 of these 13 strains represented unique genotypes, whereas the same clone was found in two different non-CF patients; interestingly, the clone was identical to the one found in CF patients B1 and B12 (Fig. 1A and B). Thus, despite some examples of possible cross-infections, our data show that intermittently colonized CF patients have an overall high level of P. aeruginosa genotype diversity equivalent to the diversity found in settings other than the CF patient airways. In agreement with previous findings, our data thus show that there is not a specific genotype of P. aeruginosa responsible for colonization of young CF patients (5, 27).

Recolonization by P. aeruginosa can be caused by identical or different clones. Our genotyping data suggest that most intermittently colonized CF patients acquire unique clones from the environment. Early aggressive antibiotic chemotherapy temporarily eradicates these bacteria; eradication is followed by new colonization events (11, 27, 42). To further understand the dynamics of these reemerging P. aeruginosa colonizations, we genotyped strains isolated from patients in which P. aeruginosa was again isolated after a period of no detectable P. aeruginosa in sputum samples. From patients B7 and B12 the same P. aeruginosa genotypes as those originally isolated and stored as first samples in the hospital collection were identified in subsequent samples (Fig. 2). From patient B6, however, we isolated more than one genotype (Fig. 2). Taken together, these data illustrate that recolonization can be caused either by new strains from the environment or by the same clone, which may have remained transiently undetectable by culture of sputum or have appeared by reinfection from a specific host or from an environmental source in close association with the patient.

High P. aeruginosa genotype diversity among recently diagnosed chronically infected patients. During the course of our studies of intermittent colonizations, two young CF patients (B3 and B19) were excluded from this group as they became chronically infected with P. aeruginosa and exhibited an elevated level of precipitins (Table 1). However, this gave us the opportunity to study the initial phase of chronicity. Genotyping of eight sequential isolates from patient B3 over a period of 24 months demonstrated a clonal colonization of initially nonmucoid P. aeruginosa followed by the appearance of mucoid variants of the same genotype (Fig. 3A). This genotype is identical to that found in a newly colonized child (B23). From B19 we have so far obtained only one isolate (unique genotype).

Low P. aeruginosa genotype diversity among CF patients with long-term chronic infection. We next determined the genotypes of a set of 45 P. aeruginosa strains isolated during 2005 from seven Danish CF patients with long-term chronic P. aeruginosa infections. The group of patients studied here has been diagnosed as being chronically infected with P. aeruginosa for more than 12 years and are cohort isolated from other patients at the CF Center (Table 1). Our genotype data based on SNP analysis show that the investigated patients were colonized mainly by one clone or very few coexisting clones (Fig. 3B). Interestingly, only five different P. aeruginosa genotypes were found among the 45 isolates tested, and considerable overlap in clones among the patients was observed (Fig. 3B). Two genotypes (marked “r” and “b” in Fig. 3B) were particularly dominant and were found in four and two patients, respectively, as well as coexisting in one patient. This finding of relatedness was confirmed by PFGE analysis of selected strains. As shown in Fig. 4, the PFGE data correlate well with the AT biochip results from representative isolates. In a reverse experiment, we used the PFGE pattern for the b geno-
type (Fig. 4) to search our PFGE database. We chose nine isolates from eight different patients with a PFGE pattern that suggested they were of the b genotype. This was in all cases confirmed by SNP typing (data not shown).

These data show that, in contrast to our findings for CF patients with intermittent colonization or recent chronic infections, an overall low level of \textit{P. aeruginosa} genotype diversity and the existence of two dominant clones characterize the Copenhagen CF patients with long-term chronic infections.

**Genetic variations among clonal isolates.** To further characterize the two dominant clones, we examined the presence and absence of variable genomic regions monitored on the AT biochips (Fig. 3B). This revealed some differences, both within isolates from a particular patient and between clones isolated from different patients. For example, among 22 isolates of the r genotype from five of the patients we found differences in 6 of the 45 investigated variable regions on the AT biochip, on the basis of which the 22 clones could be divided into 10 groups.
(presence or absence of these 6 regions) (Fig. 5A). Likewise, the 14 b clones isolated from different patients were divided into three groups based on the presence or absence of variable regions (Fig. 5B).

This shows that there are genetic variations among clonal isolates from both individual patients and different patients. This diversity probably indicates that adaptations to specific niches occur within the complex CF patient lung environment or that adaptation of *P. aeruginosa* to life in the CF lung is achieved by multiple evolutionary paths. The adaptive significance of the observed diversity will require further characterizations.

The two dominant clones are transmissible. Sequential isolates collected from six of the seven chronically infected patients for periods of 12 to 28 years (Fig. 6) were genotyped in order to investigate the *P. aeruginosa* population dynamics within the patient group. Interestingly, among the 77 archival strains tested, only 10 additional genotypes, not seen in 2005, were found and only 2 of these genotypes were found more than once (genotypes 5 and 6 in patient 2). Again, this overall low level of genotype diversity among the study patients was due to a substantial overrepresentation of the r and b genotypes colonizing the patients. In fact, all six patients have or have had at some point the b clone, while the r clone was present in four patients (Fig. 6).

Our longitudinal study also showed that, with the exception of the 14 years of clonal colonization by the b strain in patient 7, chronic infection appears to be dynamic, resulting in periodic replacement of genotypes with other genotypes (Fig. 6). For example, in patient 2 several strain replacement episodes occurred between 1978 and 2005, resulting in a temporal pattern of establishment and elimination of the 5, 6, and b clones and leaving the r clone as the dominant clone since 1994. A similar pattern was observed in patients 6 and 10, in which the initial b clone was replaced by the r clone. In patients 11 and 16 we observed the establishment of coinfections. In patient 11, the 4 clone was repeatedly found between 1991 and 2003. From that time and onward the b clone was consistently iso-
lated together with the 4 clone. In patient 16 the b clone has been isolated since 1985 and has been isolated since 2002 together with the r clone.

These results clearly show that the two dominant clones are transmissible between patients. We observed a total of 10 infection events among these patients involving either the r or b clone among the six patients with long-term infection (Fig. 6). These findings suggest that these particular clones must have strong selective advantages in colonizing CF patient airways.

Transmissibility of the dominant clones is not linked to mucoidy. During the course of CF patient lung infection *P. aeruginosa* usually undergoes a transition from nonmucoid to a mucoid, alginate-overproducing phenotype. We found both nonmucoid and mucoid strains in four of the seven chronically infected patients (Fig. 3 and 6). In patients 2 and 8 nonmucoid and mucoid isolates were of the same genotype whereas in patients 11 and 16 they were of different genotypes (Fig. 6).

It has been shown that overproduction of alginate confers a selective survival advantage for the mucoid *P. aeruginosa* strains relative to nonmucoid strains in mouse lungs (4, 12, 41, 45). Although we observed both nonmucoid and mucoid variants of both the r and b dominant genotypes among our seven study patients (Fig. 3 and 6), we nevertheless consistently found only nonmucoid clones of the b and r genotypes in patients 11 and 16 they were of different genotypes (Fig. 6).

The majority of nonmucoid isolates are mucoid revertants. Mutations in the *mucA* gene are known to cause the mucoid phenotype (4, 24). To further understand the mucoid/nonmucoid dynamics of the two dominant clones, we determined the DNA sequences of the *mucA* and *algT* genes for two isolates from the patients. One candidate

![FIG. 6. Genotyping of sequential *P. aeruginosa* isolates from patients with long-term chronic infection. Boxes below the names of the isolates indicate specific genotype classes. White boxes specify unique genotypes found only once in the data set. Labeled boxes specify that a genotype was found more than once. Asterisks indicate mucoid isolates. The results of sequence analysis of *mucA* and *algT* from selected isolates are shown in black and white text, respectively.](http://iai.asm.org/)

years and patient 10 has carried either the b or the r nonmucoid clone for a period of 20 years. Nevertheless, close inspection of the bacteriology records for these two patients showed that the mucoid variant has been found (but not stored) at a low frequency in the histories of the patients. To rule out the possibility of an existing small subpopulation of mucoid variants in these two patients, we inspected multiple sputum samples from patient 7 (9 samples) and patient 10 (15 samples) during 2005 for the presence of mucoid strains. No mucoid variants were found in the samples, and it thus appears that the clear majority of *P. aeruginosa* strains in these two patients are nonmucoid. For the r and b nonmucoid clones in these two patients it appears that adaptive features other than the transition to the mucoid phenotype enable them to persist and establish chronic infections in CF patient lungs.

The majority of nonmucoid isolates are mucoid revertants. Mutations in the *mucA* gene are known to cause the mucoid phenotype (4, 24). To further understand the mucoid/nonmucoid dynamics of the two dominant clones, we determined the DNA sequences of the *mucA* gene for selected isolates. For the b clone, we sequenced the *mucA* gene from eight nonmucoid isolates from patients 2, 7, 10, 11, and 16 and one mucoid isolate from patient 16 (Fig. 6). Interestingly, all nine isolates had the same 430ΔG (*mucA22*) mutation, suggesting that the nonmucoid isolates are in fact revertants that carry a mutation(s) that suppresses the *mucA* mutations. One candidate
gene for such a second-site reverting mutation is the algT gene (9). We found that eight of the nine isolates also carried a G55A mutation in algT, resulting in a Glu-to-Lys substitution (Fig. 6). However, as the mucoid isolate 7329B/02 from patient 16 also carried this mutation in algT, it is unlikely to be the only cause of suppression of the mucA mutation.

Sequencing of the mucA gene from five nonmucoid and two mucoid isolates of the r clone from patients 2, 10, and 16 showed a result similar to that obtained for the b clone (Fig. 6). Six of the seven isolates carried the same C349T mucA mutation, whereas one nonmucoid isolate from 2005 had the wild-type mucA gene. Of the five nonmucoid isolates, four carried a C499T mutation in algT, resulting in a Pro-to-Ser substitution, and one carried a GC insertion at position 336, which introduced a stop codon at position 340 (Fig. 6). The two mucoid isolates analyzed had no mutations in algT. Again, these data suggest that nonmucoid isolates are mucA revertants. In contrast to the findings for the b clone, we cannot exclude mutations in algT as the only cause of suppression of mutations in mucA.

Finally, we sequenced the mucA and algT genes from three nonmucoid isolates and one mucoid isolate of the 4 genotype from patient 11 (Fig. 6). The two nonmucoid isolates from 1991 and 1994 had wild-type mucA and algT sequences (Fig. 6). A nonmucoid isolate from 1995 carried a T367C mucA mutation as well as a 209AC deletion in the algT sequence, which most likely explains the nonmucoid phenotype. Again, this particular isolate appears to be a mucA revertant. A mucoid isolate from 2003 carried the same T367C mucA mutation and a wild-type algT gene.

Taken together, our sequencing results for the mucA gene show that a clear majority of nonmucoid isolates (13 out of 16) contain mutations in mucA and that they consequently are revertants originating from mucoid parent strains. Importantly, none of the 10 infection events involving either the r or the b clones among the study patients involved mucoid variants (Fig. 6).

The transmissible clones retain their potential to infect CF patient airways despite reduced virulence. It has previously been shown that transmissible and highly infectious clones such as the Liverpool epidemic strain (LES) exhibit enhanced virulence (37). To further characterize the two transmissible Copenhagen clones, we measured virulence of selected isolates using the C. elegans virulence model (Materials and Methods). As shown in Fig. 7, both an r clone isolated from patient 2 in 2005 (isolate 19945b/05) and a b clone isolated from patient 7 in 2004 (isolate 7331A/04) were completely nonviralulent in this assay, whereas an unrelated strain from the newly chronically infected patient B3 (isolate s0/04) killed the worms as readily as the virulent control strain. This finding is consistent with previous observations that virulence and virulence factors of P. aeruginosa are lost due to mutations during long-term colonization of the CF patient airways (20, 23, 40). Importantly, we also found a b clone cross-infecting patient 11 in 2003 (isolate 1299d/03) to be completely nonviralulent in the worm assay (Fig. 7). These results suggest that, although virulence has been reduced for both the b and r clones during colonization of Danish CF patients over a period of 20 years, the clones have preserved their capacity to spread among the patients.

**DISCUSSION**

We have examined the initial P. aeruginosa colonization and chronic infection dynamics in a group of CF patients at the Danish CF Center in Copenhagen. We have shown that an overall high level of P. aeruginosa genotype diversity characterizes CF patients with intermittent colonization or recently acquired chronic infections and that this reflects the genotype diversity found in other clinical settings and the environment (Fig. 1 and 3). In agreement with previous findings, this suggests that CF patients acquire unique P. aeruginosa strains independently, presumably from different environmental reservoirs (5, 16, 35, 36). In general, these isolates are nonmucoid, virulent, and susceptible to antibiotics. For intermittently colonized patients we furthermore observed that, following initial eradication by antibiotic treatment, recolonization by P. aeruginosa occurred, with either a different genotype, suggesting a new environmental source of infection, or with the same genotype, suggesting either undetected colonization (e.g., in the paranasal sinuses) or a persistent environmental source. Longitudinal genotyping of isolates from recently chronically infected patients showed clonal infections and development of the mucoid phenotype, illustrating what is generally thought to be a characteristic pattern of infection of the CF patient airways.

In contrast, genotyping of longitudinal isolates from patients with long-term chronic infection revealed a completely different pattern. We observed a clear overrepresentation of two distinct clones (the r and b clones in Fig. 6). All six patients with long-term chronic infection studied in detail have at some point in time been infected by either the r or the b clone or both, and in total for these patients we observed 10 infection events involving these clones (Fig. 6). These findings strongly suggest that the clones are transmissible and have the ability to spread among CF patients. We do not know whether the transmission route is direct (person to person) or indirect (i.e., bacteria from a patient are transiently located at the CF Center and inhaled by another patient). The first observation of the
transmissible clones was in 1987 for the r clone (isolate 7505/87 from patient 2) and 1984 for the b clone (isolate 17368/84 from patient 6). Although the two transmissible clones have been present in patients at the Copenhagen CF Center for more than 20 years, they are still transmitted from patient to patient. For example, the r clones 81052B1/01 and 7329a/02 colonized patient 10 and 16 in 2001 and 2002, respectively, and the b clone 1299b/03 colonized patient 11 in 2003 (Fig. 6). So far we have not observed the two dominant clones among other groups of CF patients (intermittently colonized and recently chronically infected), suggesting that the cohort isolation procedures employed at the Danish CF Center in Copenhagen appear to have contained these clones to the older group of patients (long-term chronically infected) and prevented further spreading.

As an alternative explanation to the recent occurrences of either r or b clones in patients 10, 11, and 16 (which we interpret as recent transmission events), it might be argued that transmissibility was an original but no longer present trait of the clones, which allowed spreading of the clones among patients some time ago. If so, their sudden and recent appearance in different patients reflects the presence of undetected, indigenous subpopulations of either of the clones, which due to unknown factors (such as changes in treatment strategies, altered airway pathology, etc.) suddenly gained advantages over the present dominant population of P. aeruginosa. To disclose the presence of such already-present subpopulations, we specifically searched for small subpopulations of the r clone in patient 7 and for subpopulations of the b clone in patient 10. Phenotype screening (colony typing and antibiotic resistance profiling) coupled to genotyping by AT biochips of many P. aeruginosa isolates from multiple sputum samples taken during 2005 from these two patients failed to provide evidence for the presence of such residual subpopulations (data not shown). We therefore suggest that the two dominant clones have preserved their transmissibility during their presence in the airways of Danish CF patients for more than 20 years and that these clones have strong selective advantages for colonizing CF patient lungs.

Our data further suggest that the virulence potential of both the r and the b clones plays little if any role in their ability to persist in CF airways and to spread among patients. In fact, virulence as measured both in the C. elegans virulence model (Fig. 7) and by the Bacillus subtilis killing system (32) (data not shown) was severely reduced for both genotypes recently isolated after long-term colonization of specific patients, or after a recent transmission to a different patient. In support of the latter finding, it was recently shown that recolonization of the airways in a CF patient after lung transplantation was by an r clone that was found to be severely reduced in virulence factor production (20). Our findings for the two dominant clones at the Copenhagen CF Center are different from reports concerning other transmissible clones, such as the Liverpool epidemic strain, which display enhanced virulence that could contribute to their spread throughout CF patient populations (37).

We found that the P. aeruginosa population structure within the CF patient airways is highly dynamic, even during long-term chronic infection. Although in most patients only one clone or very few clones are present at any given time, we frequently observed replacements of infecting clones (Fig. 6). For example, in patients 2, 6, and 10 we observed several periods of establishment of infections followed by elimination of specific clones. Equally important, we also observed several infecting clones that failed to establish in the lungs of the patients (Fig. 6). The latter clones could be environmental or clinical isolates, which transiently established in the lungs but were competed out by the resident population or eliminated by antibiotics and/or the immune system. Clearly, infection of an already chronically infected CF patient lung requires special properties. The most noticeable exception from this pattern was patient 7, who showed a stable infection by a nonmucoid b clone for a period of 14 years.

The P. aeruginosa population dynamics in patient 10 is of special interest. In this patient, we observed a complete replacement of the nonmucoid b clone by the nonmucoid r clone (Fig. 6). From 1986 to 2002 only the b clone was found to colonize the airways of this patient. From 2002 onward only the r clone was found. A similar pattern of the b clone replaced by the r clone was also observed in patients 2 and 6 (Fig. 6). Changing patient treatment protocols cannot readily explain this eradication of the b clone, as this particular clone was not eradicated in patients 7, 11, and 16. From an ecological perspective these data suggest that the r and b clones compete in the same specific niches of the CF patient lung and that in the cases of patients 2, 6, and 10 better fitness of the r clone than of the residing b clones results in niche exclusion and eradication of the b clone.

Based on these results, we therefore propose that competition among different P. aeruginosa genotypes plays a significant role in determining the P. aeruginosa population structure within the CF patient airways. Competition and the presence of two highly adapted and transmissible clones together with the aggressive antibiotic therapy employed at the CF clinic in Copenhagen result in the specific P. aeruginosa population structure reported here, in which new incoming clones are unable to establish themselves in the CF patients and already-present clones that cannot be eradicated by conventional therapy in fact can be displaced completely (eradicated) by interclonal competition. Although the specific mechanism of competition is unknown, it is obvious that the nutritional conditions, tolerance to antibiotics and the immune system, and responses to other stress factors could constitute significant competitive factors.

During the course of CF patient lung infections, P. aeruginosa usually undergoes a transition from a nonmucoid to a mucoid phenotype. Significant attention has been given to the emergence of the mucoid phenotype as the presence of mucoid variants is thought to mark the transition to the fatal, chronic stage of the infection (14). The mucoid variants are indeed better protected than nonmucoid bacteria against the inflammatory defense mechanisms of the host and show improved persistence in mouse lungs (4, 6, 12, 41, 45). Clinically, the presence of mucoid variants is associated with poor prognosis, deteriorating lung function, and increased tissue damage (33). In agreement with these observations, our longitudinal studies of P. aeruginosa isolates from different patients showed the development of the mucoid phenotype during the course of infection. For example, clone 3 in patient B3, clone 4 in patient 11, the r clone in patient 2, and the b clone in patient 16 all...
developed the mucoid phenotype starting from an initial nonmucoid population.

Nevertheless, in patients 7 and 10 we observed long-term colonization mainly by nonmucoid variants. For the r and b nonmucoid clones in these two patients, mucoidy must be replaced by other adaptive features that provide the bacteria with selective advantages for living in the two specific CF patient lungs. From an epidemiological point of view, the nonmucoid variants of the r and b genotypes are also of special interest, since all 10 infection events by the r or b clone evident in our data set were by nonmucoid variants. Our data thus emphasize that persistence and transmissibility mechanisms of nonmucoid strains associated with the chronic infection process should be further investigated.

From an ecological and evolutionary perspective loss of the transcriptional regulator mucA by mutation and the resulting conversion to the mucoid phenotype are an example of adaptive evolution that results in ecological specialization by providing the bacteria with adaptive advantages for establishment and survival specifically in the unique CF patient lung niche. Indeed, mucoid variants are rarely if ever isolated as free-living bacteria. In that sense mucoidy can be considered an important virulence factor in case of CF infections. Development of the mucoid phenotype thus resembles the concept of within-host, shortsighted evolution of virulence as proposed by Levin and Bull (21). The model is based on three conditions (21), which are all fulfilled in the case of emergence of mucoid variants during infection of the CF patient lung (1). The bacteria responsible for increased morbidity and mortality of a given host are a result of niche expansion by a genetically distinct subpopulation that arises by mutations within the host during the course of infection (2). The subpopulation becomes established because of local advantages over the ancestral population that they have within the host (3). The selective advantage of the virulent subpopulation is shortsighted because it is uniquely local, that is, within the host. Its members have a disadvantage in their capacity for, or likelihood of, infectious transmission to new hosts. In the extreme case, they are evolutionary dead-ends.

The protection against the immune system provided by the alginate encasement of the mucoid variants probably results in niche expansion, maybe causing infections of the alveoli in the aerobic compartments of the lung as suggested by Hoiby (13). The alginate-overproducing mutants, however, are characterized by a reduced fitness in other environmental contexts due to the severe cost of producing the polysaccharide in large amounts, and in this respect they could be viewed as evolutionary dead-ends.

One prediction from the described concept of shortsighted evolution is that evolution of an alginate-overproducing phenotype may open up an alternative niche that is not occupied by the parent cells. This would allow the two different cell lines to coexist stably in the CF patient lung, in agreement with the observations that in patients 2, 11, and 16 we consistently found both mucoid and nonmucoid clones in the sputum samples—in patient 2 derived from the same ancestral clone and in patients 11 and 16 derived from different clones. However, in patients 7 and 10, harboring nonmucoid clones, no stable coexistence of different genotypes was observed. These results indicate that the CF patient lung constitutes more than one ecological niche (the aerobic respiratory zone and the conductive zone containing anaerobic sputum, as suggested by Hoiby (13) and that competition between cells does take place within a niche but not between cells in separate niches.

Sequencing of the mucA gene in several isolates of different genotypes showed that most of the nonmucoid isolates were in fact revertants from mucoid parent strains (13 out of 16 nonmucoid isolates carried mutations in mucA). For some of the nonmucoid revertants we found mutations in algT that could explain the suppression of the mucA mutation (9), but most isolates seem to harbor suppressor mutations in so-far-unidentified genes. Whatever the genetic mechanisms responsible for suppression of the mucA mutation, we suggest that this is an example of further within-host evolution that consequently results in escape from the potential evolutionary dead-end of being mucoid. The nonmucoid revertants (containing mucA mutations as well as suppressor mutations) must have evolved alternative survival strategies that can substitute for mucoidy. We suggest that these adaptive properties, which could override the significance of mucoidy, might include increased antibiotic resistance, improved utilization of the lung environment, and increased tolerance of the immune response and stress in general. By using these alternative strategies for survival they have retained their ability to spread to new hosts.

In conclusion, this epidemiological study of P. aeruginosa isolates from a group of CF patients associated with the Copenhagen CF Clinic in Denmark has documented the dominance of two distinct genotypes infecting the patients and pointed out cross-infection among the patients as a major cause of the occurrence of these particular clones in connection with chronic lung infections. In most patients these two clones have competed with each other and with other strains, resulting in clonal substitutions at certain time points during the course of the chronic infection. Many patients harbor alginate-overproducing mucoid variants (of either genotype) in coexistence with nonmucoid cells of the same or different genotypes. The data support the theory that these mucoid variant cells occupy a specific niche in the CF patient lung which is separate from that harboring nonmucoid P. aeruginosa (13). Further investigations are required to directly document this type of niche specialization.

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