

Relationships between *Staphylococcus aureus* Genetic Background, Virulence Factors, *agr* Groups (Alleles), and Human Disease

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The expression of most *Staphylococcus aureus* virulence factors is controlled by the *agr* locus, which encodes a two-component signaling pathway whose activating ligand is an *agr*-encoded autoinducing peptide (AIP). A polymorphism in the amino acid sequence of the AIP and of its corresponding receptor divides *S. aureus* strains into four major groups. Within a given group, each strain produces a peptide that can activate the *agr* response in the other member strains, whereas the AIPs belonging to different groups are usually mutually inhibitory. We investigated a possible relationship between *agr* groups and human *S. aureus* disease by studying 198 *S. aureus* strains isolated from 14 asymptomatic carriers, 66 patients with suppurative infection, and 114 patients with acute toxemia. The *agr* group and the distribution of 24 toxin genes were analyzed by PCR, and the genetic background was determined by means of amplified fragment length polymorphism (AFLP) analysis. The isolates were relatively evenly distributed among the four *agr* groups, with 61 strains belonging to *agr* group I, 49 belonging to group II, 43 belonging to group III, and 45 belonging to group IV. Principal coordinate analysis performed on the AFLP distance matrix divided the 198 strains into three main phylogenetic groups, AF1 corresponding to strains of *agr* group IV, AF2 corresponding to strains of *agr* groups I and II, and AF3 corresponding to strains of *agr* group III. This indicated that the *agr* type was linked to the genetic background. A relationship between genetic background, *agr* group, and disease type was observed for several toxin-mediated diseases: for instance, *agr* group IV strains were associated with generalized exfoliative syndromes, and phylogenetic group AF1 strains with bullous impetigo. Among the suppurative infections, endocarditis strains mainly belonged to phylogenetic group AF2 and *agr* groups I and II. While these results do not show a direct role of the *agr* type in the type of human disease caused by *S. aureus*, the *agr* group may reflect an ancient evolutionary division of *S. aureus* in terms of this species' fundamental biology.

Staphylococcus aureus is both a commensal and an extremely versatile pathogen in humans, causing three basic syndromes: (i) superficial lesions such as skin abscesses and wound infections; (ii) deep-seated and systemic infections such as osteomyelitis, endocarditis, pneumonia, and bacteremia; and (iii) toxic syndromes such as toxic shock syndrome (TSS) and staphylococcal scarlet fever (both due to toxic shock syndrome toxin 1 [TSST-1] and staphylococcal enterotoxins [SEs]), staphylococcal scalded-skin syndrome (SSSS; due to exfoliatins), and staphylococcal food poisoning (due to SEs) (1, 18, 24). With the exception of toxemia, the molecular basis of *S. aureus* pathogenicity is multifactorial, depending on the expression of a large class of accessory gene products that comprise cell wall-associated and extracellular proteins (24). Expression of most virulence factors in *S. aureus* is controlled by the *agr* locus, which encodes a two-component signaling pathway whose activating ligand is a bacterial-density-sensing peptide (autoinducing peptide) also encoded by *agr* (24). A poly-

morphism in the amino acid sequence of the autoinducing peptide and of its corresponding receptor (AgrC) has been described. *S. aureus* strains can be divided into four major groups on this basis: within a given group, each strain produces a peptide that can activate the *agr* response in the other member strains, whereas the autoinducing peptides produced by the different groups are usually mutually inhibitory (14, 16). Links between a peculiar *agr* type and a specific staphylococcal syndrome have been shown for TSS and SSSS. TSST-1-producing isolates belong to *agr* specificity group III (16) and mostly belong to a single clone, as shown by multilocus enzyme electrophoresis (MLEE) (23) and pulsed-field gel electrophoresis (PFGE) (3). Most exfoliatin-producing strains responsible for SSSS belong to *agr* group IV, but the clonality of these strains has not been investigated (14). *agr* group I was prevalent in a collection of 192 *S. aureus* strains, most of which were methicillin resistant, but no clinical information was available in this study (29).

The aim of the present study was to further investigate a possible relationship between *agr* groups (alleles) and the pattern of *S. aureus* disease. We studied 198 methicillin-susceptible strains from the French National Reference Center for Staphylococcal Toxemia strain collection, in which all clinical syndromes are represented, to determine their *agr* type and the distribution of 24 toxin genes (by PCR), as well as to determine

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TABLE 1. Oligonucleotide primers and reference strains used for toxin gene detection

Toxin	Gene	GenBank accession no.	Primer(s)	Sequence (5'-3')	Size of amplified product (bp)	Control strain
SEA	<i>sea</i>	M18970	SEA-1	GAAAAAAGTCTGAATTGCAGGGAACA	560	ATCC 13566
			SEA-2	CAAATAAATCGTAATTAACCGAAGGTTTC		
SEB	<i>seb</i>	M11118	SEB-1	ATTCTATTAAGGACACTAAGTTAGGGA	404	ATCC 13566
			SEB-2	ATCCCGTTTCATAAAGGCGAGT		
SEC	<i>sec</i>	X05815	mpSEC-1	GTAAAGTTACAGGTGGCAAACTTG	297	ATCC 19095
			mpSEC-2	CATATCATACCAAAAAGTATTGCCGT		
SED	<i>sed</i>	M28521	SED-1	GAATTAAGTAGTACCGCGCTAAATAATATG	492	FRI-1151m
			SED-2	GCTGTATTTTTCTCCGAGAGT		
SEE	<i>see</i>	M21319	SEE-1	CAAAGAAATGCTTTAAGCAATCTTAGGC	482	ATCC 27664
			SEE-2	CACCTTACCGCCAAAGCTG		
SEG	<i>seg</i>	AF064773	SEG-1	AATTATGTGAATGCTCAACCCGATC	642	A900322
			SEG-2	AACTTATATGGAACAAAAGGTACTAGTTC		
SEH	<i>seh</i>	U11702	SEH-1	CAATCACATCATATGCGAAAGCAG	376	ATCC 51811
			SEH-2	CATCTACCAACATTAGCACC		
SEI	<i>sei</i>	AF064774	SEI-1	CTCAAGGTGATTTGGTGTAGG	576	A900322
			SEI-2	AAAAAACTTACAGGCAGTCCATCTC		
SEJ	<i>sej</i>	AF053140	mpSEJ-1	TAACCTCAGACATATATACTTCTTTAACG	300	FRI-1151m
			mpSEJ-2	AGTATCATAAAGTTGATTGTTTTTCATGCAG		
SEN	<i>sen</i>	AF285760	mpSEN-1	ATGAGATTGTTCTACATAGCTGCAAT	680	A900322
			mpSEN-2	AACTCTGCTCCCACTGAAC		
SEO	<i>seo</i>	AF285760	mpSEO-1	AGTTTGTGTAAGAAGTCAAGTGTAGA	180	A900322
			mpSEO-2	ATCTTTAAATTCAGCAGATATTCATCTAAC		
SEM	<i>sem</i>	AF285760	mpSEM-1	CTATTAATCTTTGGGTTAATGGAGAAC	300	A900322
			mpSEM-2	TTCAGTTTCGACAGTTTTGTTGTCAT		
TSST-1	<i>tst</i>	J02615	TST-1	TTCACTATTTGTAAGAGTGCAGACCCACT	180	FRI-1169
			TST-2	TACTAATGAATTTTTTATCGTAAGCCCTT		
ETA	<i>eta</i>	M17347	mpETA-1	ACTGTAGGAGCTAGTGCATTGT	190	TC-7
			mpETA-3	TGGATACTTTTGTCTATCTTTTTTCATCAAC		
ETB	<i>etb</i>	M17348	mpETB-1	CAGATAAAGAGCTTTATACACACATTAC	612	TC-146
			mpETB-2	AGTGAACCTATCTTTCTATTGAAAAACACTC		
PVL components S and F	<i>lukS-PV-lukF-PV</i>	AB006796	PVL-1	ATCATTAGGTAAAATGTCTGGACATGATCCA	433	ATCC 49775
LukE-LukD	<i>lukE-lukD</i>	Y13225	NPVL-2	GCATCAASTGTATTGGATAGCAAAAAGC	269	FRI-913
LukM	<i>lukM</i>	D42144	LUKDE-1	TGAAAAAGGTTCAAAGTTGATACGAG	780	ATCC 31890
			LUKDE-2	TGTATTTCGATAGCAAAAAGCAGTGCA		
Alpha-hemolysin	<i>hla</i>	M90536	LUKM-1	TGGATGTTACCTATGCAACCTAC	780	ATCC 31890
			LUKM-2	GTTTCGTTTTCCATATAATGAATCACTAC		
Beta-hemolysin	<i>hly</i>	S72497	HLA-1	CTGATTACTATCCAAGAAATTCGATTG	209	FRI-913
			HLA-2	CTTTCCAGCCTACTTTTTTATCAGT		
Beta-hemolysin	<i>hly</i>	S72497	HLB-1	GTGCACCTACTGACAATAGTGC	309	NCTC 7428
			HLB-2-2	GTTGATGAGTAGTACCTTCAGT		
Delta-hemolysin	<i>hly</i>	AF288215	HLD-1	AAGAAATTTTTATCTTAATTAAGGAAGGAGTG	111	NCTC 9393
			HLD-2	TGATGAATTTGTTCACTGTGTGCGA		
Gamma-hemolysin components A, B, and C	<i>hly</i>	L01055	mpHLG-1	GTCAYAGAGTCCATAATGCATTTAA	535	ATCC 49775
			mpHLG-2	CACCAAAATGTTAGCTTAAAGTG		
Gamma-hemolysin variant	<i>hly-2</i>	D42143	mpHLG2-1	GACATAGAGTCCATAATGCATTYGT	390	RIMD 31092
			mpHLG2-2	ATAGTCATTAGGATTAGGTTTCACAAAAG		
EDIN	<i>edin</i>	M63917	EDIN-1	GAAGTATCTAATACTTCTTTAGCAGC	625	E-1
			EDIN-2	TCATTTGACAATTCTACACTTCCAAC		

their genetic background (by amplified fragment length polymorphism [AFLP] analysis). We then sought to determine the relationships between these characteristics and the type of clinical disease syndrome.

MATERIALS AND METHODS

Staphylococcal strains and corresponding disease syndromes. The French National Reference Center for Staphylococcal Toxemia (Lyon, France) collects more than 800 strains yearly from patients with toxic and nontoxic staphylococcal diseases throughout France. For this study we selected a subset of 198 *S. aureus* strains isolated between January 1985 and December 1999. They were isolated from nose swabs ($n = 3$) and vaginal swabs ($n = 11$) of 14 asymptomatic

carriers and from clinical specimens of 66 patients with *S. aureus* suppurative infections (necrotizing pneumonia caused by Pantone-Valentine leukocidin-producing strains [$n = 11$], furunculosis [$n = 11$], native valve endocarditis [$n = 19$], finger pulp infections [$n = 9$], osteitis [$n = 8$], cellulitis and/or myositis [$n = 4$], and arthritis [$n = 4$]), 4 patients with enterocolitis, and 114 patients with acute toxemia, including 35 cases of TSS, 33 cases of staphylococcal scarlet fever, and 46 cases of SSSS (20 cases of generalized exfoliative syndrome and 26 cases of bullous impetigo). The types of infection were defined according to published criteria (8, 18, 19). All infections were community acquired. All of the strains were collected from hospitals located throughout France and were identified as *S. aureus* by their ability to coagulate citrated rabbit plasma (bioMérieux, Marcy l'Etoile, France) and to produce a clumping factor (StaphySlide Test; bioMérieux).

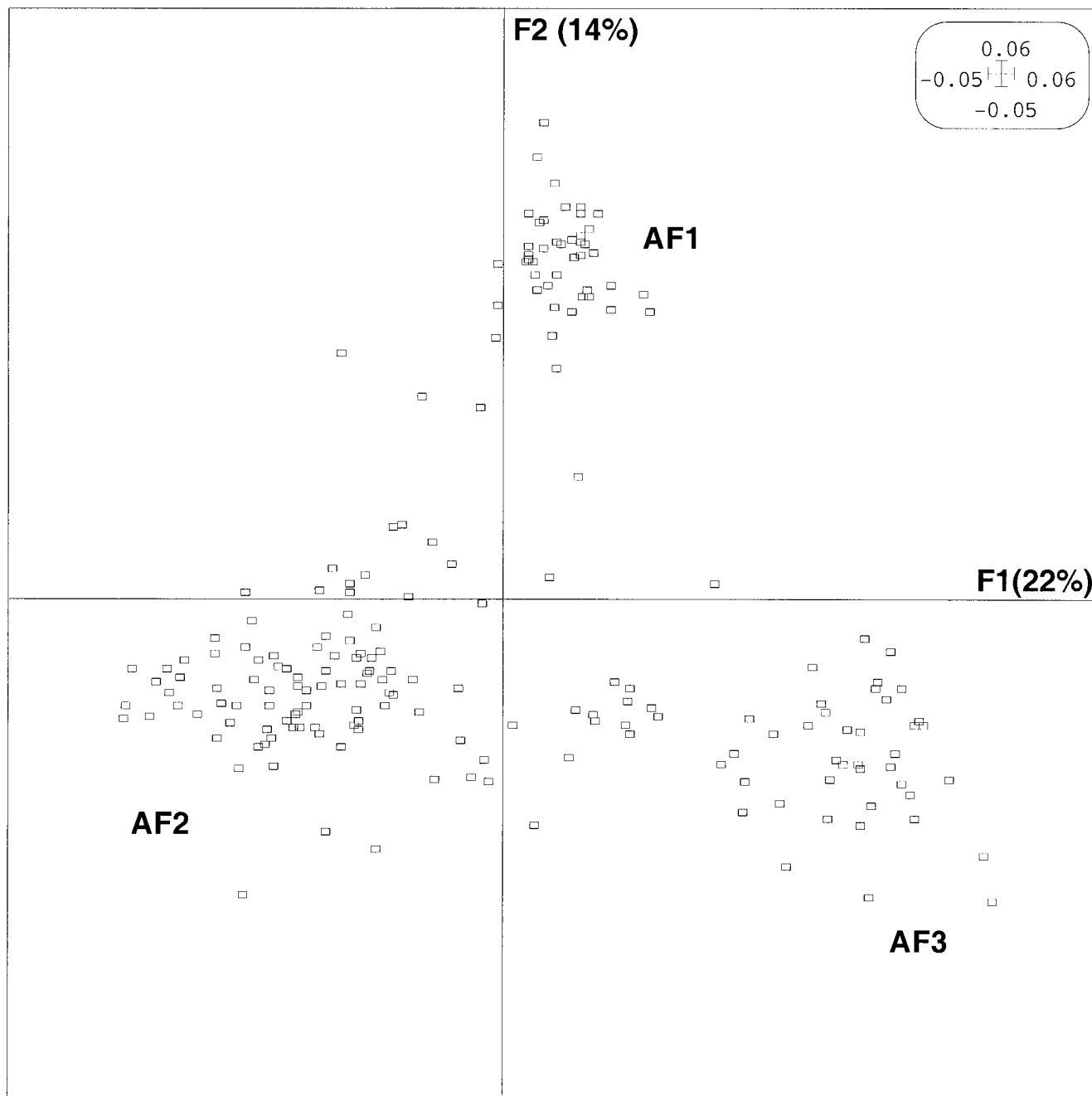


FIG. 1. PCO factor map (F1 × F2) of the 198 strains of *S. aureus* based on the AFLP data. The 198 strains are projected in the F1/F2 plane. This plane, obtained by computation, is defined by the two principal axes of the analysis: F1 explains most of the variance, and the second axis, F2 (orthogonal to F1), explains most of the remaining variance. Phylogenetic groups AF1, AF2, and AF3 are indicated. For clarity, when several strains are projected on the same point, only one is represented.

S. aureus strains RN6390 (*agr* group I), RN6923 (*agr* group II), RN8462 (*agr* group III), A980740 (*agr* group IV), and RN6911 (*agr* null) were used as controls for *agr* group identification (14, 16). Control strains used for toxin gene detection are listed in Table 1.

Culture and DNA extraction. Strains were grown on brain heart infusion agar or in the same broth at 37°C overnight. Genomic DNA used as target for PCR and AFLP assays was extracted by using a standard phenol-chloroform procedure (26), and the concentration of DNA was estimated spectrophotometrically (26).

Identification of *agr* alleles. The primers Pan-1 (5'-ATG CAC ATG GTG CAC ATG CA-3') and Pan-2 (5'-CAT AAT CAT GAC GGA ACT TGC TGC GCA-3') (Eurogentec, Seraing, Belgium) were designed from *agr* group I to IV sequences (GenBank accession numbers M21854, AF001782, AF001783, and AF288215, respectively) to amplify a 1,234-bp *agr* fragment encompassing the 3' end of *agrB*, all of *agrD*, and the 5' end of *agrC*. Amplification was carried out on a PE-9600 thermocycler (Perkin-Elmer Corp., Norwalk, Conn.) under the following conditions: an initial 5-min denaturation step at 95°C; followed by 30 cycles of 1 min of denaturation at 94°C, 1 min of annealing at 55°C, and 1 min

TABLE 2. *agr* group and AFLP group distribution according to the disease type

Disease type	Letter designation	No. of strains							
		Total (n)	<i>agr</i> group (n)				AFLP group (n)		
			I	II	III	IV	AF1	AF2	AF3
Exfoliative toxin-mediated disease									
Generalized exfoliative syndrome	A	20	0	3	1	16	15	5	0
Bullous impetigo	B	26	3	10	0	13	12	14	0
Subtotal		46	3	13	1	29	27	19	0
Enterotoxin-mediated disease									
TSS	C	24	16	7	0	1	1	20	2
Scarlet fever		16	10	5	0	1	2	15	0
Subtotal		40	26	12	0	2	3	35	2
TSST-1-mediated disease									
Menstrual TSS	D	5	0	0	5	0	1	0	4
Nonmenstrual TSS		6	1	0	5	0	0	1	5
Scarlet fever		17	1	4	12	0	1	4	12
Subtotal		28	2	4	22	0	2	5	21
Suppurative infections									
Endocarditis	E	19	8	9	2	0	0	16	3
Necrotizing pneumonia	F	11	3	0	4	4	5	3	3
Cellulitis and/or myositis	G	4	1	0	0	3	3	1	0
Furunculosis	H	11	4	0	2	5	5	4	2
Osteitis and/or osteomyelitis	I	8	1	4	3	0	0	5	3
Finger pulp infections	J	9	2	3	3	1	1	5	3
Arthritis	K	4	0	1	3	0	0	3	1
Subtotal		66	19	16	18	13	14	37	15
Enterocolitis	L	4	3	1	0	0	0	3	1
Colonization									
Nose	M	3	0	1	2	0	0	1	2
Vagina		11	8	1	1	1	1	9	1
Subtotal		14	8	2	3	1	1	10	3
Overall total		198	61	49	43	45	47	109	42

of extension at 72°C; and a final extension step at 72°C for 10 min. The PCR products were purified by using the High Pure kit (Boehringer Mannheim) and sequenced with the primers used for PCR (Genome Express, Grenoble, France). The 198 strains were assigned to one of the four *agr* groups by comparing the predicted product of *AgrD* and the N-terminal half of *AgrC* with those of the four control strains (see above).

Toxin gene detection. Sequences specific for *sea-e*, *seg-j*, *sem-o*, *tst*, *eta*, *etb*, *lukS-PV-lukF-PV*, *lukE-lukD*, *lukM*, *hla*, *hfb*, *hld*, *hlg*, *hlg-2*, and *edin*, encoding SEA-E; SEG-J; SEM-O; TSST-1; ETA; ETB; PVL components S and F; LUKE-LUKD; LUKM; the alpha-, beta-, delta-, gamma-, and gamma variant hemolysins; and EDIN, respectively, were detected by PCR on a PE-9600 thermocycler (Perkin-Elmer) as previously described (15, 19) with the primers shown in Table 1 (Eurogentec). Amplification of *gyrA* was used to confirm the quality of each DNA extract and the absence of PCR inhibitors (5). All PCR products were analyzed by electrophoresis through 1% agarose gels (Sigma, Saint Quentin Fallavier, France). The distribution of the 24 toxin genes among the 198 strains is available on-line (<ftp://pbil.univ-lyon1.fr/pub/datasets/statox.txt>).

AFLP. The Perkin-Elmer AFLP Microbial Fingerprinting Kit was used according to the manufacturer's recommendations, except that the *TaqI* primers and adapter were as described by Vos et al. (30). The following two AFLP conditions were used, with one selective nucleotide added at the 3' extremity of each primer: *EcoRI* + *A/TaqI* + C (condition A/C) and *EcoRI* + *T/TaqI* + G (condition T/G). Touchdown PCR cycling was done as recommended by the manufacturer, in a PE-9600 thermocycler (Perkin-Elmer). Processed DNA samples were loaded in pools of two with fluorescent dyes in 6% (wt/vol) denaturing polyacrylamide gels for electrophoresis (ABI Prism 373 Sequencer; Perkin-Elmer).

AFLP data processing. Perkin-Elmer GeneScan analysis software was used to extract data from electropherograms. Assignment of fragments to discrete categories, transformation of data into tabular binary matrices (LecPCR program), and calculation of genomic distance (DistAFLP program), with or without boot-

strap resamplings, were done as previously described (22). The LecPCR and DistAFLP programs are freely available on the ADE-4 web server (<http://pbil.univ-lyon1.fr/ADE-4/microb>). DistAFLP provides output files in the ADE-4 binary format suitable for multivariate analysis (28). Genomic distances are estimated rates of nucleotide substitution over whole genomes by using the Dice similarity index and Jukes-Cantor correction (22).

Statistical analysis. The table of toxin gene detection in the 198 isolates was analyzed by using classical principal component analysis (PCA). Only 19 toxin genes were used, since four were either always present (*hla* and *hld*) or always absent (*edin* and *lukM*). The AFLP results (distance matrix between the 198 strains) were analyzed by using principal coordinate analysis (PCO) (11). This method provides a description of the main structures of distance matrices in the form of factor maps, in the same way as PCA. PCA of the toxin genes and PCO of the AFLP results gave independent results, and the corresponding graphics cannot be compared. However, these two analyses can be linked by co-inertia analysis (CIA) (7), so that the results can be compared and the factor maps can be superimposed. CIA gives co-inertia axes that have the maximum possible covariance with the variables in each of the two data sets. By using the covariance instead of the correlation (as in canonical correlation analysis), CIA maximizes the product of the correlation by the projected variances, ensuring that co-inertia axes will have both a good correlation with the initial variables and real meaning for each of the two data sets (7).

RESULTS

Phylogenetic distribution of the clinical *S. aureus* strains.

Since any relationship between the *agr* group and disease type would have to be interpreted according to the strain's genetic background, we first conducted AFLP analysis of the 198

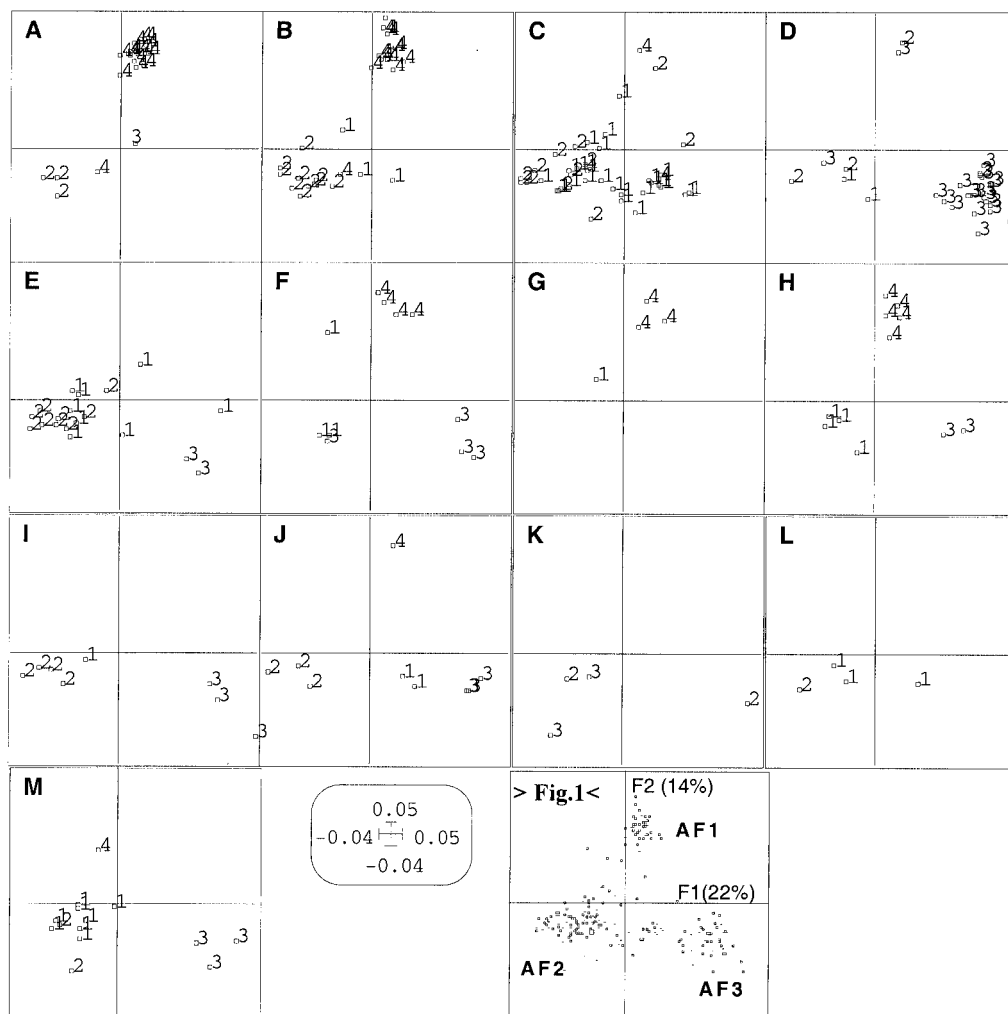


FIG. 2. AFLP PCO factor map (F1 × F2) of the 198 strains, split according to the 13 disease types designated A to M: A, generalized exfoliative syndrome; B, bullous impetigo; C, enterotoxin-mediated diseases; D, TSST-1-mediated diseases; E, endocarditis; F, necrotizing pneumonia; G, cellulitis and/or myositis; H, furunculosis; I, osteitis and/or osteomyelitis; J, finger pulp infections; K, arthritis; L, enterocolitis; M, colonization. Numbers 1 to 4 in panels A to M indicate the *agr* group of each strain. Panel “>Fig. 1<” is a reduction of Fig. 1 showing the axes (F1 and F2) and the positions of the phylogenetic groups (AF1, AF2, and AF3).

strains. PCO was performed on the distance matrix of the 198 isolates obtained by using the AFLP results to calculate genome divergence (i.e., the rate of nucleotide substitutions over whole genomes) between pairs of isolates. Figure 1 shows the factor map of the AFLP PCO. Since the first axis, F1, accounted for 22% of the variance, and the second axis, F2 (orthogonal to F1), accounted for the largest part (14%) of the variance not accounted for by F1 (data not shown), the results are expressed as the projections of each strain on a plane defined by these two axes, which were conserved for further analysis. This analysis divided the strains into three main phylogenetic groups, namely, AF1, a group with positive values on the F2 axis; AF2, a group with negative values on both the F1 and F2 axes; and AF3, a group distinguished from AF1 by negative values on the F2 axis.

To evaluate the possible relationship between the disease type and the bacterial genetic background, the same AFLP PCO factor map was split according to the different diseases

(classified in 13 disease types designated A to M, Table 2) for a better understanding of this relationship (Fig. 2). This analysis shows that the strains involved in scalded skin syndrome (disease type A) and bullous impetigo (disease type B) mainly belonged to phylogenetic group AF1 (27 of 46 strains). Enterotoxin-producing strains associated with scarlet fever and TSS (disease type C) mainly belonged to phylogenetic group AF2 (35 of 40 strains). The strains involved in TSST-1-associated disease (scarlet fever and menstrual and nonmenstrual TSS) (disease type D) belonged principally to phylogenetic group AF3 (21 of 28 strains). In contrast to toxin-mediated diseases, strains associated with suppurative infections (group E to M) did not seem to be specifically related to a particular AFLP cluster. Only endocarditis strains (disease type E) were mainly related to phylogenetic group AF2 (16 of 19 strains). The strains associated with diseases I, J, K, and L rarely belonged to group AF1. Interestingly, the strains associated with necrotizing pneumonia (disease F) and furuncles (disease H), which

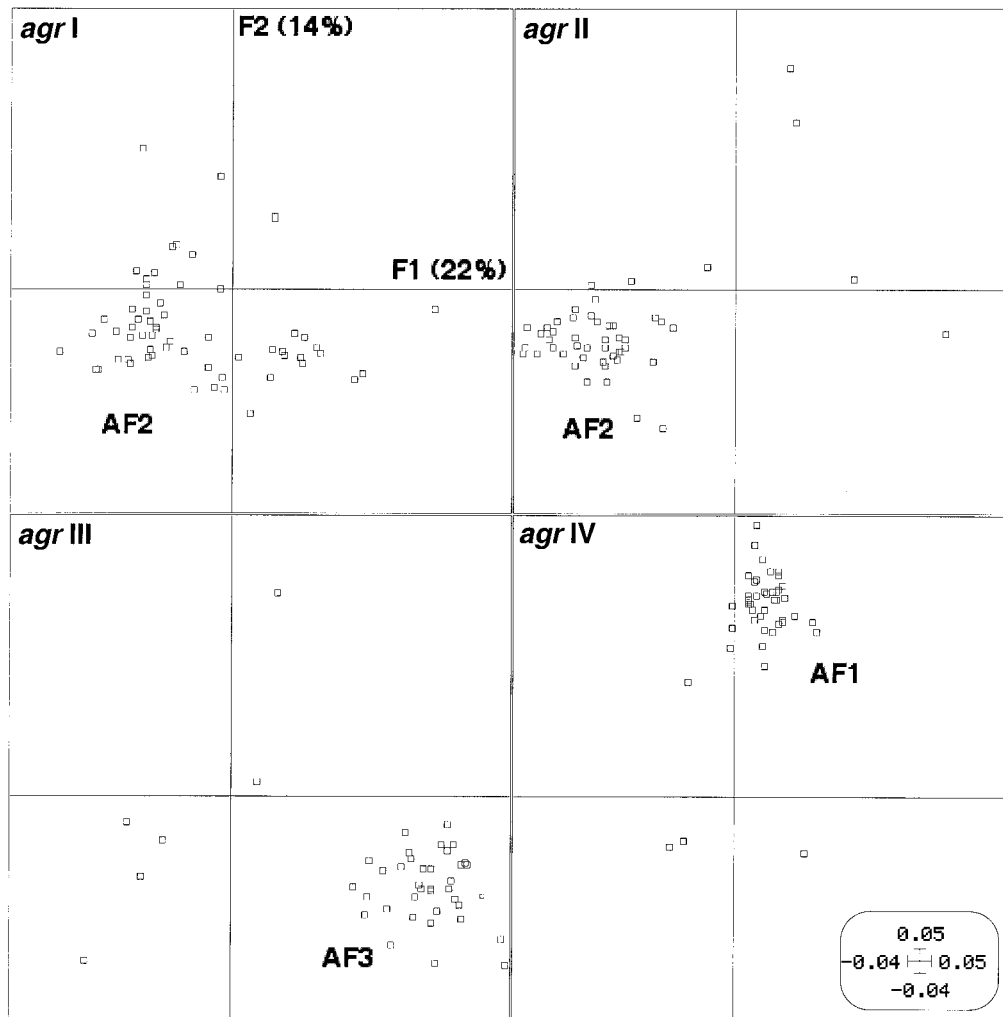


FIG. 3. AFLP PCO factor map (F1 \times F2) of the 198 strains of *S. aureus* as in Fig. 1 split according to the *agr* groups (*agr* I to IV). Phylogenetic groups AF1, AF2, and AF3 are indicated.

are both caused by Panton-Valentine leukocidin-producing strains (19), belonged to three phylogenetic groups, and the two graphics were highly superimposable.

Relationships between *agr* groups, genetic background, and disease type. The *agr* group of the 198 *S. aureus* isolates was analyzed by amplification and sequencing. All isolates were classified as part of one of the four *agr* groups, and the distribution was relatively even, with 61 strains belonging to *agr* I, 49 belonging to *agr* II, 43 belonging to *agr* III, and 45 belonging to *agr* IV (Table 2). The AFLP PCO factor map for the 198 isolates was then split according to the four *agr* groups (Fig. 3). This clearly individualized strains from groups III and IV, while strains from groups I and II were partly superimposed. Thus, there was a very strong relationship between the AFLP cluster and the *agr* group in spite of the fact that a few isolates appeared to be outliers from the clusters of isolates belonging to the same *agr* group (four strains of *agr* group I, four strains of *agr* group II, six strains of *agr* group III, and three strains of *agr* group IV). This finding clearly indicated that *agr* types are associated with genetic backgrounds of strains. *agr* group la-

belonging of each isolate on the AFLP PCO factor map showed a clear relationship between the genetic background, the *agr* group and the disease type, particularly for toxin-mediated diseases (Fig. 2, panels A to D). For instance, *agr* group IV strains involved in generalized exfoliative syndrome and bul- lous impetigo (disease groups A and B) were particularly associated with phylogenetic group AF1. Likewise, *agr* group III strains involved in TSST-1-mediated diseases (disease group D) were particularly associated with phylogenetic group AF3, and strains causing SE-mediated diseases (disease group C) belonged to *agr* group I or II and phylogenetic group AF2.

Relationships between toxin genes and genetic background. To determine whether the observed link between toxin-mediated diseases and genetic background was also found at the level of toxin genes, we analyzed the relationship between the distribution of 24 toxin genes and the genetic background. We first determined whether preferential combinations of toxin genes occurred among the 198 clinical strains, by means of PCA. Since the first axis, F1, and the second axis, F2 (orthogonal to F1), accounted for the largest part of the variance in

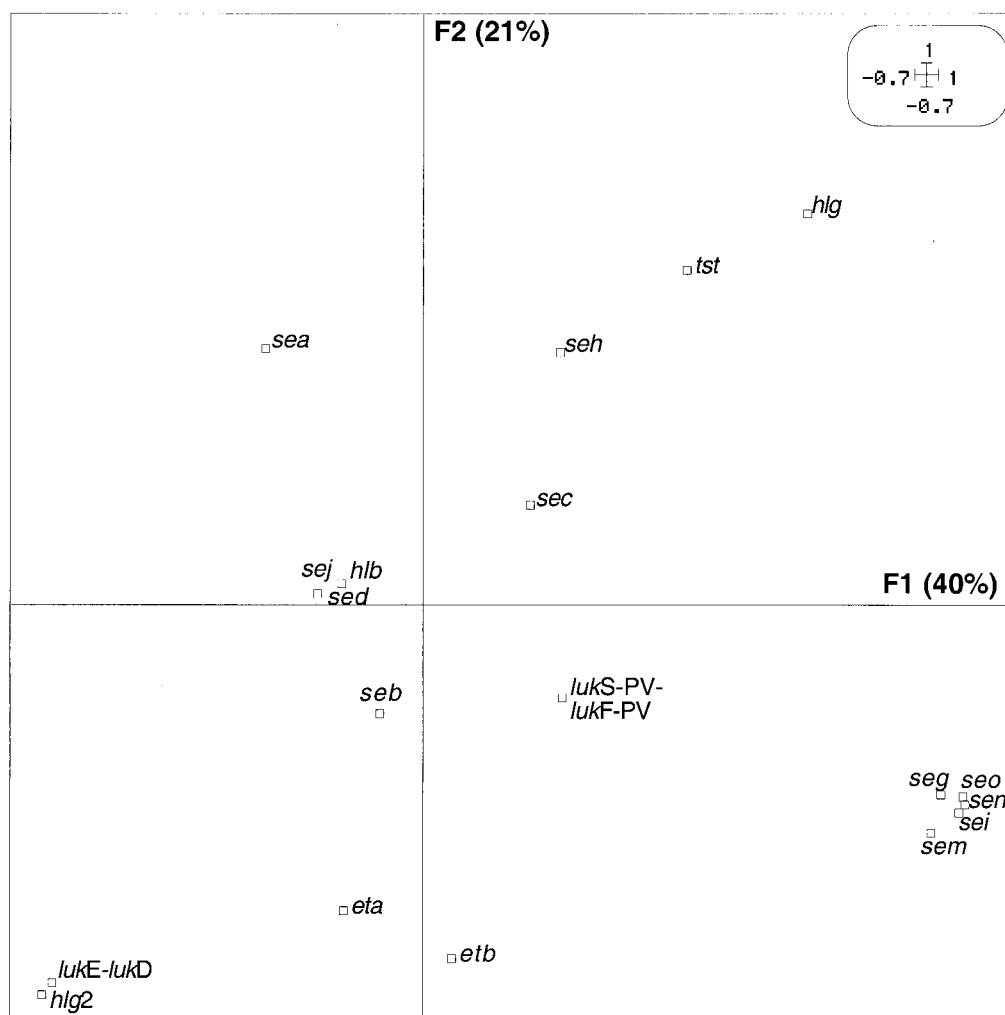


FIG. 4. PCA of 19 toxin genes in the 198 *S. aureus* strains. The variables are projected in the F1/F2 plane (defined as in Fig. 1). In each pair of axes, the variables located in a given direction relative to the origin can be considered positively associated, whereas the variables located in opposite directions can be considered antagonistic. Variables plotted near the origin cannot be interpreted. For example, the five gene toxins *seg*, *sei*, *sen*, *sem*, and *seo* can be considered associated with one another and negatively associated with *sea*. The toxin gene codes are given in Table 1.

PCA (40 and 21%, respectively), the results were expressed as the projections of each toxin gene on a plane defined by these two components (F1 and F2) (Fig. 4). On the first component, a group of five toxin genes (*seg*, *sei*, *sen*, *seo*, and *sem*, all encoded by the enterotoxin gene cluster) (15) was clearly individualized and slightly opposed to *sea*. On the second component there was an opposition between *hlg-2* (together with *lukE-lukD* and, less strongly, *eta* and *etb*) and *hlg* (and, to a lesser extent, *tst* and *seh*). The other toxin genes (*seb*, *sec*, *sed*, *sej*, *hlb*, and *lukS-PV-lukF-PV*) were too near to the origin to be interpreted. The preferential combinations or exclusions of toxin genes suggested a nonrandom distribution of toxin genes in these strains.

To confirm the link between the toxin gene distribution and the genetic background of the strains, we coupled the AFLP clusters and the toxin gene analysis by using CIA (7). The PCA of the toxin gene table and the PCO of the AFLP results were coupled. Figure 5 shows the CIA factor map for the toxin

genes, while Fig. 6 shows the CIA factor map for the AFLP patterns of the 198 isolates. These factor maps were very similar to the factor maps of the separate analyses (Fig. 3 and 4). The correlation coefficients between the co-inertia axes were 0.813 and 0.783 for the first and second axes, respectively. The percentages of variance derived from the co-inertia axes were 92 and 86% for the first axis in the toxin gene space and in the AFLP space, respectively. For the first two axes, the corresponding percentages were 98 and 83%. These very strong correlations and percentages of explained variance reflected a strong relationship between the toxin gene distribution and AFLP clusters. Hence, *tst* and *hlg* were associated with phylogenetic group AF3 and opposed to groups AF1 and AF2. The five genes belonging to the enterotoxin gene cluster (*seg*, *sei*, *sem*, *sen*, and *seo*) were associated with group AF1 and opposed to group AF2. *lukD-lukE* and *hlg-2* were opposed to phylogenetic group AF3 and associated with groups AF1 and AF2. *eta* and *etb* were associated with phylogenetic group AF1

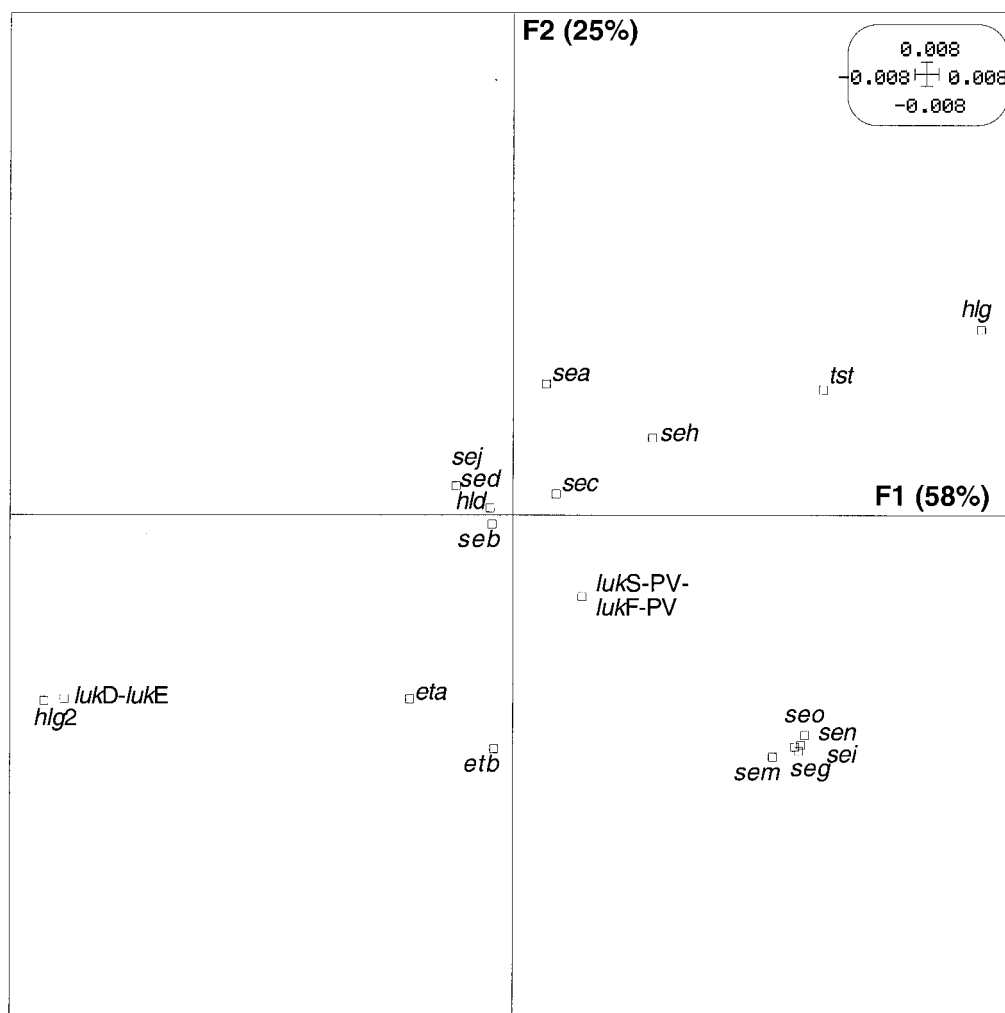


FIG. 5. First factor map ($F1 \times F2$) of the CIA of the 19 toxin genes. This factor map is very similar to the PCA factor map (Fig. 4), with a few exceptions (*sea* and *seb*, for example). Genes with similar positions on this map have a similar profile of presence or absence among the 198 strains (as in Fig. 4), but they also belong to strains that have a similar AFLP profile (see Fig. 6).

(Fig. 5 and 6). The distribution of the 24 toxin genes, the *agr* group, and the AFLP group among the 198 strains is available on-line (<ftp://pbil.univ-lyon1.fr/pub/datasets/statox.txt>).

DISCUSSION

We investigated a possible relationship between *agr* groups and the pattern of *S. aureus* disease and found a strong association between the *agr* types and certain diseases. However, in most cases the association reflected the link between the disease types, the pattern of toxin genes, and the genetic background of the strains. For instance, the strains causing SE-mediated diseases (disease group C) belonged to *agr* group I or II and phylogenetic group AF2. We thus concluded that, in most of the disease types considered (mainly toxin-mediated diseases), the *agr* alleles and toxin genes evolved contemporaneously with their parent strains and that horizontal transfer played only a marginal role. These findings confirm that specific bacterial pathogenicities are each essentially associated with a specific clone or group of clones.

The association of virulence factors with specific backgrounds has been discussed recently for several bacterial species such as *Escherichia coli*, in which a clonal distribution of virulence genes has been reported among clinical strains isolated from bloodstream infections (12, 20), neonatal meningitis (2), and extraintestinal infections (25). Regarding *S. aureus*, multilocus sequence typing comparison of isolates recovered from asymptomatic nasal carriers and from patients with severe diseases revealed that invasive diseases were primarily caused by a subset of genotypes unrepresentative of the carriage population as a whole (4). Another study of the genetic structure of *S. aureus*, involving MLEE, revealed that a single clone (designated ET41) of *S. aureus* producing TSST-1 causes most epidemiologically unrelated cases of urogenital TSS (23). The observation of a large proportion of asymptomatic female genital-tract carriers of this clone suggested that ET41 was highly adapted to the cervicovaginal tract. More recently, Booth et al. (3) used PFGE to analyze 405 clinical isolates of *S. aureus* and found that five phylogenetic lineages were highly

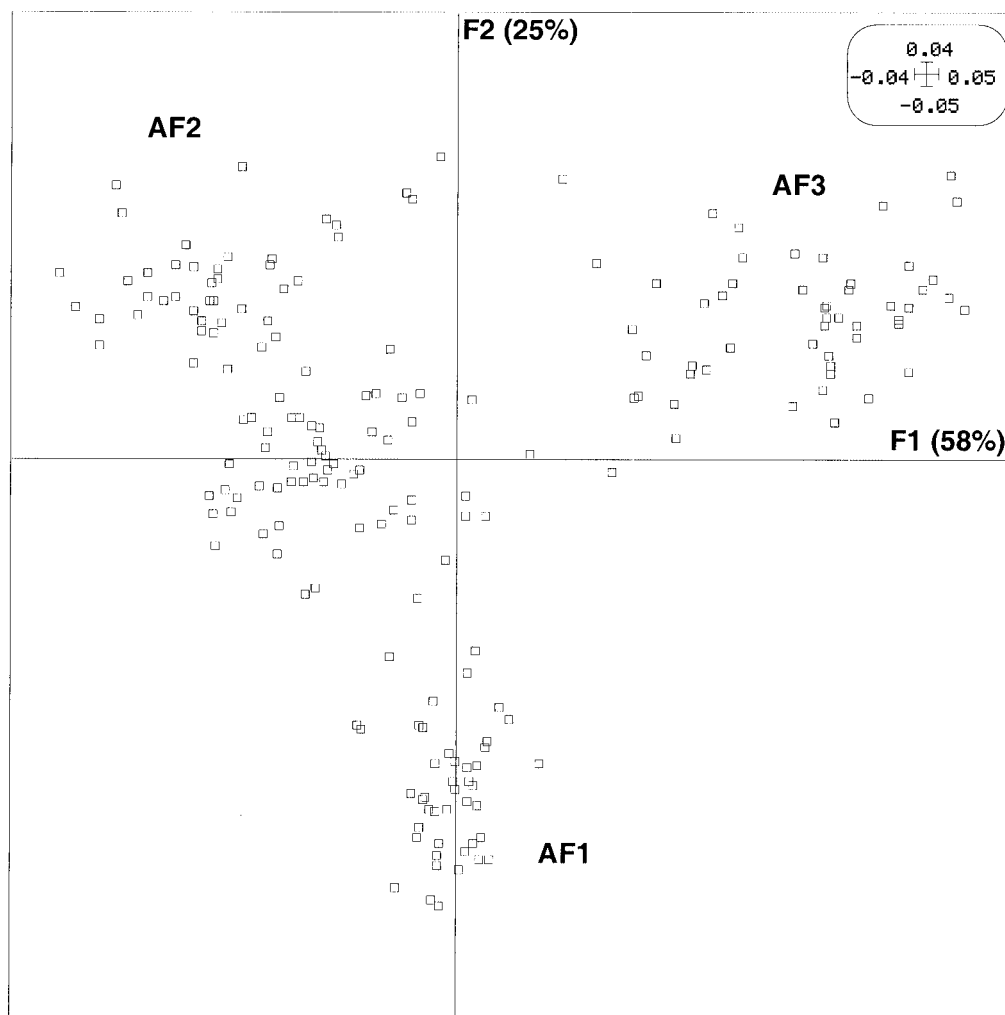


FIG. 6. First factor map ($F1 \times F2$) of the CIA of the 198 *S. aureus* strains. This factor map is very similar to the PCO factor map (Fig. 1), except that the second (vertical) axis is inverted. Phylogenetic group AF1 is in the lower part of the graph, while groups AF2 and AF3 are in the upper part. Strains with similar positions on this map have genes present in the same direction on the map in Fig. 5.

prevalent and widely distributed, in contrast to 85 other lineages which occurred with frequencies of $<2.5\%$. One of the five prevalent lineages (SAL1) comprised most TSST-1-producing strains, confirming the observation by Musser et al. (23). Booth et al. also found that SAL1 was enriched among the normal flora of the anterior nares and that lineage SAL4, which comprised 90% of methicillin-resistant *S. aureus* (MRSA) strains, was significantly associated with respiratory tract infections (3).

We studied a different population of methicillin-susceptible *S. aureus* clinical strains causing a broad spectrum of community-acquired infections. In agreement with the findings of Musser et al. (23) and Booth et al. (3), we confirmed the clonality of TSST-1-producing strains. In addition, we found that a number of diseases, such as SSSS, bullous impetigo, scarlet fever, TSS and, to a lesser extent, infective endocarditis, were preferentially associated with one of the three phylogenetic lineages that structured our strain population. In this respect, we fully agree with the concept developed by Falkow

et al. that “the basic unit of bacterial pathogenicity is the clone or lineage that expands due to the possession of unique combinations of virulence genes” (10). With the exception of infective endocarditis, most of the diseases listed above are caused by specific toxins whose genetic determinants are frequently carried by potentially mobile elements such as plasmids (*etb*, *seb*, etc.), phages, or pathogenicity islands (*tst*, *egc*, *lukFD-lukSE*, etc.) (17). CIA, used to couple PCA of the toxin gene table (Fig. 4) and PCO of the AFLP results (Fig. 1), clearly indicated that the distribution of toxin genes was closely linked to the strain’s genetic background (Fig. 5 and 6). This suggested that the virulence determinants did not spread homogeneously among various genetic backgrounds or, at least, that the efficiency of such genetic exchanges between the three major lineages was low. Similarly, Booth et al. found that *tst*, *cna* (the collagen-binding protein gene), and *hly* were associated with certain PFGE lineages and not with others, suggesting limited horizontal transfer among lineages (3). This is in accordance with the observation that, in *E. coli*, horizontal

transfer generally does not disrupt the clonal structure of the species (6, 21, 27). A stable link between virulence and phylogeny could correspond to the necessity of having virulence determinants move into a particular genetic background for the emergence of a "virulent clone" (9, 10, 25). In contrast, PVL-associated diseases (necrotizing pneumonia and furunculosis) appeared to be caused by strains of the three lineages (Fig. 2). Since PVL is encoded by a bacteriophage, it is likely that in this case the bacteriophage has spread easily among the different backgrounds. With other suppurative diseases (groups I, J, K, and L [i.e., osteitis, finger pulp infections, arthritis, and enterocolitis, respectively]), the rarity of associations with phylogenetic group AF1 (Fig. 2) cannot yet be interpreted, since we do not know the virulence factors specifically involved in these infections. Microbial surface component-recognizing adhesive matrix molecules are likely play a role in these settings (13).

Considering the possible relation between the *agr* group and the disease type, we first postulated a relation between the *agr* group and the capacity to induce a specific disease. Ji et al. observed that the vast majority of menstrual toxic shock strains belonged to *agr* group III but that strains belonging to the other two groups had no apparent clinical specificity (16). We have previously shown that most ET-producing strains belong to *agr* group IV (14). In another study, *agr* group I was the most prevalent among 192 carrier and disease isolates, but 71% of the isolates were MRSA strains, which are known to be highly clonal (29). In our carefully selected collection of *S. aureus* clinical isolates (mainly causing community-acquired infections), the four *agr* types were relatively evenly distributed (Table 2). The *agr* group distribution correlated strongly with the genetic background of the strains and thus, indirectly, with certain disease profiles. The observed link between the *agr* group and genetic background was also found among coagulase-negative staphylococci in our laboratory: amplification-sequencing yielded 25 distinct *agr* variants among 13 staphylococcal species; the overall topology of the phylogenetic tree constructed from the DNA sequences of the 25 *agr* alleles was remarkably similar to that constructed from 16S rRNA loci of the 13 staphylococcal species, arguing against significant horizontal transfer of *agr* genes between populations of staphylococci (D. Dufour et al., unpublished data). As proposed by Novick, this *agr* grouping may represent the first subdivision of *S. aureus* based on the fundamental biology of the organism (24). Finally, though we cannot attribute a direct responsibility of the *agr* type in disease initiation, we can speculate that the preferential association between certain *agr* alleles, certain toxin genes, and a particular genetic background may make the activation of virulence factors more efficient. To paraphrase Falkow (10), we propose that the basic unit of bacterial pathogenicity would be the clone or lineage, which expands because it possesses particular combinations of virulence and regulatory genes in the appropriate genetic background.

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