



First Genome-Based Characterisation and Staphylococcal Enterotoxin Production Ability of Methicillin-Susceptible and Methicillin-Resistant *Staphylococcus aureus* Strains Isolated from Ready-to-Eat Foods in Algiers (Algeria)

Francesca Fanelli ^{1,+}, Daniele Chieffi ^{1,+}, Gyu-Sung Cho ², Justyna Schubert ³, Omar Amine Mekhloufi ⁴, Jacek Bania ⁵, Charles M. A. P. Franz ² and Vincenzina Fusco ^{1,*}

- ¹ National Research Council of Italy, Institute of Sciences of Food Production (CNR-ISPA), 70126 Bari, Italy
- ² Department of Microbiology and Biotechnology, Max Rubner-Institut, Hermann-Weigmann-Straße 1, 24103 Kiel, Germany
- ³ Research and Development, Silantes GmbH, Gollierstrasse 70C, 80339 Munich, Germany
- ⁴ Laboratory of Hygiene and Animal Pathology, Institute of Veterinary Sciences, University of Tiaret, 14000 Tiaret, Algeria
- Department of Food Hygiene and Consumer Health Protection, Wrocław University of Environmental and Life Sciences, 50-375 Wrocław, Poland
- + These authors contributed equally to this work and thus should be designated both as first authors.
- Correspondence: vincenzina.fusco@ispa.cnr.it

Abstract: Staphylococcus aureus is a pathogenic microorganism of humans and animals, able to cause foodborne intoxication due to the production of staphylococcal enterotoxins (SEs) and to resist antibiotic treatment as in the case of methicillin-resistant S. aureus (MRSA). In this study, we performed a genomic characterisation of 12 genetically diverse S. aureus strains isolated from ready-to-eat foods in Algiers (Algeria). Moreover, their ability to produce some classical and new staphylococcal enterotoxins (SEs) was investigated. The 12 S. aureus strains resulted to belong to nine known sequence types (STs) and to the novel ST7199 and ST7200. Furthermore, S. aureus SA46 was assigned to the European clone MRSA-ST80-SCCmec-IV. The 12 strains showed a wide endowment of se and sel (staphylococcal enterotoxin-like toxin) genes (sea, seb, sed, seg, seh, sei, selj, sek, sem, sen, seo, seq, ser, selu2, selw, selx, sey, sel30; ψent1-ψent2), including variants and pseudogenes, and harboured the enterotoxin gene cluster (egc) types 1 and 5. Additionally, they produced various amounts of SEA (64.54-345.02 ng/mL), SEB (2871.28-14739.17 ng/mL), SED (322.70-398.94 ng/mL), SEH (not detectable-239.48 ng/mL), and SER (36720.10-63176.06 ng/mL) depending on their genotypes. The genetic determinants related to their phenotypic resistance to β-lactams (blaZ, mecA), ofloxacin (gyrA-S84L), erythromycin (ermB), lincomycin (lmrS), kanamycin (aph(3')-III, ant(6)-I), and tetracyclin (tet(L), tet(38)) were also detected. A plethora of virulence-related genes, including major virulence genes such as the tst gene, determinant for the toxic shock syndrome toxin-1, and the lukF-PV and lukS-PV genes, encoding the panton-valentine leukocidin (PVL), were present in the S. aureus strains, highlighting their pathogenic potential. Furthermore, a phylogenomic reconstruction including worldwide foodborne S. aureus showed a clear clustering based on ST and geographical origin rather than the source of isolation.

Keywords: genomics; staphylococcal enterotoxins; *Staphylococcus aureus*; ready-to-eat food; virulence; antibiotic resistance; food safety; methicillin-resistant *Staphylococcus aureus* (MRSA); panton-valentine leukocidin (PVL); *luk*F-PV and *luk*S-PV genes

Key Contribution: This study reported for the first time a genome-based characterisation and SEs production of foodborne *S. aureus* in Algeria, and helped to enlarge the knowledge on the enterotoxigenicity, overall virulence, antimicrobial resistance, as well as molecular-based epidemiology of this microorganism in the African continent.

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1. Introduction

Despite the increased awareness of foodborne diseases and huge progress in food safety diagnostics, enterotoxigenic *Staphylococcus* (*S.*) *aureus* continues to cause staphylococcal food-poisoning (SFP) cases worldwide. Following the discovery of the classical enterotoxins (SEA-SEE), 33 staphylococcal enterotoxin (*se*) and staphylococcal enterotoxin-like toxin (*sel*) genes are known to date [1], highlighting the great enterotoxigenic potential of *S. aureus*. Moreover, this pathogenic bacterium may show resistance to several antibiotics [2] that may affect the antibiotic treatment when *S. aureus* infections occur. In particular, community-, livestock-, and healthcare-associated methicillin-resistant *S. aureus* (MRSA), even enterotoxigenic strains, are emerging in food products, animals used for foods, animal husbandry, and food production or processing workers [3].

Recently, we investigated the prevalence, enterotoxigenic potential, and antimicrobial resistance of 48 methicillin-susceptible S. aureus (MSSA) and MRSA isolated from 207 ready-to-eat foods randomly sampled from hotels, restaurants, fast foods, and pizzerias in Algiers, the capital of Algeria, during 2018 and 2019 [4]. The resulting S. aureus prevalence was 23.2% (48/207) [4]. For strain typing, intergenic spacer region (ISR) typing showed a higher discriminatory power than the staphylococcal enterotoxin gene (SEg) typing. Indeed, the same SEg type was found in S. aureus belonging to different ISR types. In particular, the 48 S. aureus strains were grouped in seven SEg types, while 12 ISR types were detected. Moreover, 2 out of the 48 S. aureus isolates, belonging to the same ISR type, were found to harbour the mecA gene that encodes for the penicillin-binding protein 2a (PBP2a), a transpeptidase protein with low affinity for most β -lactam antibiotics, including methicillin as well as oxacillin and cefoxitin [4,5]. Twelve strains, each belonging to one of the twelve detected ISR types, were subjected to antibiotic susceptibility testing. Some strains were resistant towards antimicrobials including benzylpenicillin, ofloxacin, erythromycin, lincomycin, tetracycline, kanamycin, oxacillin, and cefoxitin; the mecA-positive strain, being resistant to oxacillin and cefoxitin, was confirmed as MRSA, and two other tested strains showed multidrug resistance [4]. However, the performed molecular detection of se and sel genes did not permit to assess the presence of enterotoxin gene variants or pseudogenes, which may affect the enterotoxigenic activity and the virulence of the S. aureus strains, nor to distinguish among the ψ *ent1*/ ψ *ent2* pseudogenes and the *selu2* gene in the enterotoxin gene cluster (*egc*) found in the S. aureus strains [4]. This drawback did not allow determining the egc type they harboured. This cluster plays a crucial role in the enterotoxigenicity of S. aureus since phylogenetic analyses indicated that potentially all the se and sel genes may derive from the egc [6], and, to date, at least eight egc types have been described in S. aureus [1,7]. Moreover, the presence of genetic determinants of antibiotic resistance, other than mecA, was not investigated in the 12 analysed strains [4]. Although some efforts are being made to characterise foodborne S. aureus in Africa, including the Algerian Country [8-11], specific data on S. aureus from ready-to-eat foods are scarce. Additionally, data on S. aureus molecular epidemiology are fairly increasing in the African continent, but studies are mainly performed on humans and animals rather than food isolates and MRSA rather than MSSA [12,13], underlying the need of collecting data on foodborne African isolates, including the Algerian Country. For all these reasons, the 12 Algerian MSSA and MRSA strains, representative of the different ISR types, were subjected to whole-genome sequencing. Herein, we reported a comprehensive genomic characterisation of these 12 genetically diverse S. aureus, exploring their genomic relatedness, investigating the genetic features in relation to their virulence and antibiotic resistance, and performing a phylogenomic analysis including worldwide foodborne S. aureus isolates. We also assessed the SEs production and detected all the se and sel genes known to date in order to improve the characterisation of foodborne S. aureus in the African continent.

2. Results

2.1. Genomic Statistics

The genomic features of *S. aureus* strains are reported in Supplementary Table S1. Genomes resulted in a median length of 2.7 Mbp and an average N50 value of 270 kbp. The average mol% GC content was 32.59%, with a median number of coding genes of 2606 and a coding density of 83.95%. The quality of the assemblies was excellent for all strains, with the exception of SA02, which resulted in only 95.5% completeness.

2.2. Multilocus Sequence and spa Typing

The sequence types (STs) of the 12 sequenced *S. aureus* strains are shown in Table 1, whereas the multilocus sequence type (MLST) allelic profiles are detailed in Supplementary Table S2. All strains belong to different STs, with the exception of strains SA02 and SA51, which both belong to ST5. Two novel STs have been deposited in PubMLST as ST7199 and ST7200: ST7199 differs from the known ST15 only for the novel *aro*E-1025 allele, whereas ST7200 differs from the known ST72 for the novel *yqi*L-954 allele. Additionally, *spa* types of the 12 sequenced *S. aureus* strains are reported in Supplementary Table S3.

Table 1. Main genetic features of the sequenced *S. aureus* strains isolated from Algerian ready-toeat foods.

Strain	MSSA or MRSA	Origin	ST	se and sel genes	mecA (SCCmec-Type)	tst	<i>luk</i> F-PV and <i>luk</i> S- PV
SA01	MSSA	Potato in sauce	15	selw ^a , selx ^h	-	-	-
SA02	MSSA	Cooked meat	5	egc1 (seg, sei, sem, sen, seo, ψent1- ψent2 *), selw *,ʰ, selx1	-	-	-
SA04	MSSA	Cooked meat	1	sea1, seh, sek, seq, selw ª, selx5	-	-	-
SA07	MSSA	Cooked meat	7199 ^k	selw ^a , selx ^h	-	-	-
SA08	MSSA	Cooked meat	25	egc5 (seg, sei, sem, sen, seo, selu2), selw ^d , selx9 ⁱ	-	-	-
SA10	MSSA	Cooked meat	101	selw ^e , selx ^h	-	-	-
SA18	MSSA	Pizza	97	<i>seb</i> v1 <i>, selw</i> *, ^f <i>, selx</i> bov2	-	-	-
SA20	MSSA	Cooked meat	8	sed*,j, selj, ser, selw ^c , selx2	-	-	-
SA24	MSSA	Lentil soup	7200 ^k	egc5 (seg, sei, sem, sen, seo, selu2), selw ^{*,t} , selx5	-	-	-
SA46	MRSA	Braised beef	80	seh *, selw ^g , selx ^h , sey	+ (SCC <i>mec</i> -IV)	-	+
SA51	MSSA	Fermented milk	5	sea2 ⁱ , egc1 (seg, sei, sem, sen, seo, ψent1-ψent2 *), selw *,ʰ, selx1, sel30	-	-	-
SA82	MSSA	Sautéed beef with potato	22	egc5 (seg, sei, sem, sen, seo, selu2), selw ^d , selx6	-	+	-

* pseudogene; + present; - absent; ^a *selw* belonging to group 3 [14]; ^b Nonfunctional *selw* belonging to group 4 [14]; ^c *selw* belonging to group 2 [14]; ^d 98% similarity to *selw* genes belonging to group 1 [14]; ^e 98% similarity to *selw* gene belonging to group 2 [14]; ^f 98% similarity to nonfunctional *selw* gene belonging to group 4 [14]; ^g 99% similar to *selw* gene belonging to group 6 [14]; ^h Different allelic variant than those previously described for *selx* [7,15,16]; ⁱ 1 nucleotide substitution compared with the described allelic variant (not leading to amino acid sequence variation in the predicted protein) [15,17]; ^j Different allelic variant than the truncated *sedv*3 described by Johler et al. [18]; ^k Novel ST.

2.3. Phylogenetic analysis

A RAxML phylogenetic tree is reported in Figure 1. It also shows the ST, source of isolation (S), and geographical origin (O) of the analysed strains.

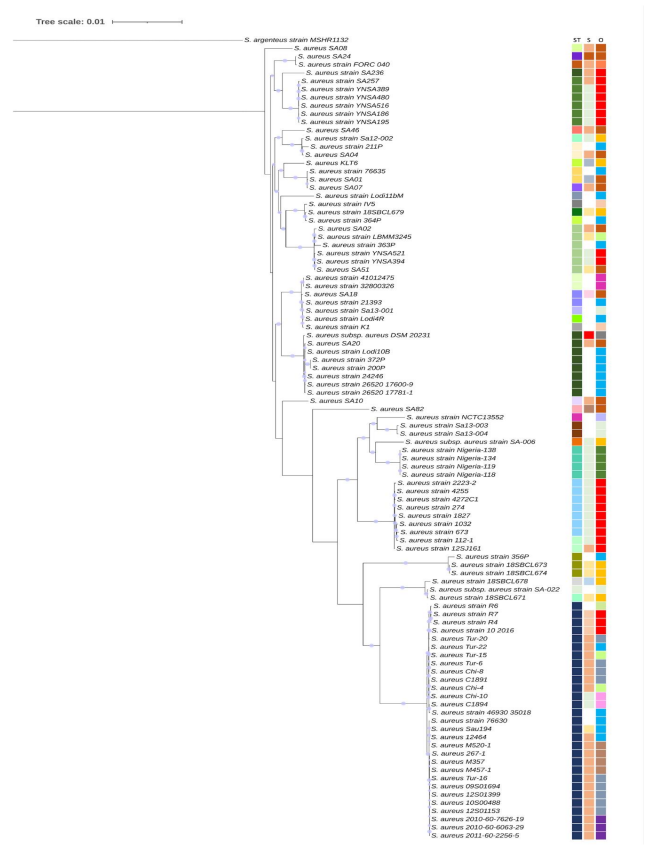


Figure 1. Phylogeny of *S. aureus* strains. Genome-based phylogenetic tree inferred by using maximum likelihood method RAxML with progressive refinement. *S. argenteus* MSHR1132 was used as outgroup. Support values are represented by scaled circles at each node. Tree is annotated with sequence type (ST), source of isolation (S), and geographical origin (O). See Supplementary Table S4 for detailed information.

Globally, *S. aureus* strains are grouped in the tree mainly depending on the ST and geographical origin rather than the source of isolation, although a certain association appears for meat, milk, and dairy products. The grouping is clear for *S. aureus* isolates from China (red squares in the third column origin (O)), Nigeria (dark green squares in the third column O), and Italy (light blue in the third column O).

S. aureus strains sequenced in this study clustered on different branches of the tree. Those belonging to the same ST, namely SA02 and SA51, and those belonging to STs that differ by only a single allele, namely SA01 and SA07, closely grouped in the tree, whereas SA08, SA10, and SA82, belonging to STs that differ in numerous alleles, were individually located in the tree. The genetic divergence of SA82 from the other sequenced strains was also confirmed by ANI analysis, which reported average values of 98.3% between SA82 and the other sequenced strains, which, among them, share >99% of ANI (Supplementary Table S5).

2.4. Pan-Genome Analysis

The pan-genome of all the *S. aureus* isolates analysed in this study, which includes orthologues and unique genes, comprised a set of 7537 genes. The core genome (shared by >99% of *S. aureus* isolates), on the other hand, consisted of 1596 genes. The accessory genome (total core: genes in >2 isolates but not in all) consisted of 4059 genes, whereas the unique genome was composed of 1882 genes. The soft core, shell, and cloud genomes contained 261, 1323, and 4357 genes, respectively (Supplementary Table S6). The output of the Roary analysis is shown in Supplementary Figure S1, which confirms the clustering obtained by the phylogenetic analysis.

2.5. Enterotoxin Genes Presence and Localisation

The *se* and *sel* genes harboured by the 12 sequenced *S. aureus* strains, including variants and pseudogenes, are shown in Table 1. The nucleotide and amino acid sequences of the detected se and sel genes and predicted proteins, as well as the percentage of identity with previously reported reference sequences, are shown in Supplementary Table S7 [1,6,7,14,15,17-31].

selw is located on the chromosome. In SA02, SA51, SA18, and SA24, the *selw* gene was predicted as a pseudogene due to a frameshift mutation (SA02 and SA51) or an internal stop codon (SA18 and SA24). It should be pointed out that for SA82, SA08, and SA10 *selw*, the NCBI Prokaryotic Genome Annotation Pipeline (PGAP) annotated the predicted protein as *se* type 26 [1]. In this study, however, we retained the *selw* nomenclature as proposed since its discovery by Okumura et al. [32].

selx is also located on the chromosome in all the analysed strains.

In SA02, the enterotoxin gene cluster (*egc*) is located in a *S. aureus* pathogenicity island (SAPI) of vSa β type I (SaPim3/n3), with *seg*, *sen*, ψ *ent*1- ψ *ent*2, *sei*, *sem*, and *seo* (J3P54_11775-11800) (*egc*1; [6]) (Supplementary Figure S2). This pathogenicity island harbours genes for the bi-component leukocidin LukED, serine protease genes *spl*ABCDEF with *spl*A being a pseudogene, genes for the restriction endonucleases M and S, and several transposase coding genes, some of which predicted as pseudogenes.

SA08 harbours the *egc5* (*seg*, *sen*, *selu2*, *sei*, *sem*, *seo*: KQP42_01815-01790) [7] in a vSaβ of type IV, with *luk*ED, serine protease genes, and a lantibiotic encoding gene cluster (KQP42_01860-01895) [33] (Supplementary Figure S2). SA24 harbours the *egc5* (*seg*, *sen*, *selu2*, *sei*, *sem*, *seo*: J3135_04230-04205) located in a SAPI of vSaβ type I. We hypothesised a vSaβ of type III in SA82, but the contig is interrupted downstream the *egc5* (*seg*, *sen*, *sei*, *sem*, *seo*: KQP39_04565-04540) (Supplementary Figure S2). In strain SA51, the *egc1* (*seg*, *sen*, *ψent1-ψent2*, *sei*, *sem*, and *seo*: J3T04_07830-07855) is located in a vSaβ type I (SaPim3/n3) (Supplementary Figure S2). This genome also harbours *sea* (J3T04_10205) in a prophage region (Φ Mu3A) (Figure 2A), harbouring the *sa*K gene and sphingomyelin phosphodiesterase pseudogene as a result of a frameshift mutation. In the genome of SA51, we also identified the *sel30* gene (J3T04_08740). The plasmid location of this gene

[1] appears to be confirmed by the nearby presence of *rep* pseudogene and the high homology (>99%) retrieved between a 17 kb flanking region comprising *sel30* in *S. aureus* NCTC6135 and SA51 genome.

The *sea* gene is also present in the genome of SA04 in a prophage region of φ SA3 type (Figure 2A). This large region includes 40 kb upstream *sea*, *seq*, and *sek* (Figure 2A), close to a site-specific integrase, a pseudogene for a sphingomyelin phosphodiesterase, a MBL fold metallo-hydrolase, the bi-component leukocidin LukGH, and the chaperonin GroEL.

In SA04, the genomic locus including the *seh* gene (J3T03_06310) is shown in Figure 2B and it is typical of a phage of φ Sa3ms type. Close to the *seh* gene, a pseudogene for the exotoxin *seo*, and the *cst* operon, which protects *S. aureus* from sulfide toxicity [34], can be identified.

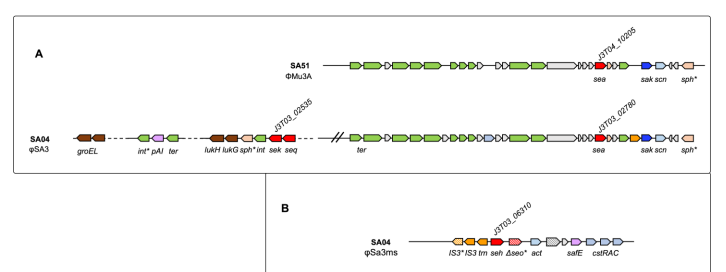


Figure 2. *sea* and *seh* prophagic regions in *S. aureus* SA04 and SA51 genomes. (**A**) Φ Mu3A region comprising *sea* gene in SA51 genome and φ SA3 region comprising *sea*, *sek*, and *seq* genes in SA04 genome. *se* genes are shown in red; *sak*: staphylokinase; *scn*: complement inhibitor SCIN-A; *sp*H: sphingomyelin phosphodiesterase; *int*: integrase; *p*AI: pathogenicity island family protein; *ter*: terminase; *luk*GH: bi-component leukocidin LukGH; *gro*EL: chaperonin GroEL; * indicates pseudogenes. (**B**) φ Sa3ms region comprising *seh* gene in SA04 genome. *act*: acetyltransferase; *saf*E: SafE family protein; *cst*R: persulfide-sensing transcriptional repressor CstR; *cst*A: persulfide response sulfurtransferase CstA; *cst*B: persulfide dioxygenase-sulfurtransferase CstB; * indicates pseudogenes.

In the SA18 genome, the *seb* gene is located in a pathogenicity island with 99.4% similarity with the SaPlivm60 described by Sato'o et al. [35].

In the genome of strain SA20, the pseudogene *sed* (J3T02_03365) and the genes *ser* (J3T02_03360) and *selj* (J3T02_03355) are one by one located at the beginning of NODE_17, immediately followed by several *rep* protein genes, indicating the presence of a putative multiresistance pIB485-like plasmid [36] (Figure 3). This plasmid contains *blaI*, *bla*R1, and *blaZ* genes and shows a 100% nucleotide homology with the SAP048A plasmid (Accession: NC_019007.1), with an inversion due to the role of the two recombinase/invertase present in this locus (*bin* gene) [37,38].

In SA46, the *sey* gene (J3T00_13960) is located on the chromosome [7]. The genomic locus comprising *seh* (J3T00_044075), annotated as a pseudogene, also includes the staphyloferrin operon and three tandem-type lipoproteins.

Lastly, the sequences of allelic variants of *sea* [17], *sed* [18] and *selx* [7,15,16], different than those previously described, have been deposited in DDBJ/ENA/GenBank under the following accessions: seA-SA51 (ON205830.1); seD-SA20 (ON205834.1); selX-SA01 (ON205831.1); selX-SA08 (ON205832.1); selX-SA10 (ON205833.1); selX-SA46 (ON205835.1).

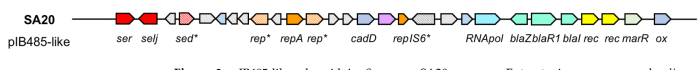


Figure 3. pIB485-like plasmid in *S. aureus* SA20 genome. Enterotoxin genes *ser* and *selj* are represented in red. *rep*: replication protein; *cad*D: cadmium resistance transporter CadD; RNA*pol*: RNA-directed DNA polymerase; *rec*: recombinase; IS6: IS6 family transposase; *blaZ*: beta-lactamase; *bla*R1: regulatory sensor/signal transducer BlaR1; *bla*I: beta-lactamase repressor BlaI; *mar*R: MarR family transcriptional regulator; *ox*: oxidoreductase; * indicates pseudogenes.

2.6. Antibiotic Resistance Genetic Determinants

Table 2 shows the antibiotic resistance profile, as assessed by Mekhloufi et al. [4], and the related genetic determinants identified in the *S. aureus* genomes analysed in the present study.

Table 2. Antimicrobial resistance profiles and related genetic determinants of *S. aureus* strains isolated from Algerian ready-to-eat foods.

Strain	Resistance Profile ^a	Genetic Determinants
SA01	Р	blaZ
SA02	OF, ERY, L, TE	gyrA-S84L, ermB, lmrS, tet(L), tet(38)
SA04	Р	blaZ
SA07	Р	blaZ
SA08	Р	blaZ
SA10	Р	blaZ
SA18	P, KAN, TE	blaZ, aph(3')-IIIa, ant(6)-I, tet(L), tet(38)
SA20	Р	blaZ
SA24	Р	blaZ
SA46	P, OXA, FOX, KAN	mecA, aph(3')-IIIa, ant(6)-I
SA51	_b	blaZ
SA82	Р	blaZ

P: Benzylpenicillin; OF: ofloxacin; ERY: erythromycin; L: lincomycin; KAN: kanamycin; TE: tetracycline; OXA: oxacillin; FOX: cefoxitin; ^a As reported by Mekhloufi et al. [4]; ^b Sensitive to all tested antibiotics (benzylpenicillin, oxacillin, cefoxitin, gentamicin, kanamycin, tobramycin, ofloxacin, erythromycin, lincomycin, clindamycin, pristinamycin, linezolid, teicoplanin, vancomycin, tetracycline, fosfomycin, nitrofurantoin, fusidic acid, rifampicin, and co-trimoxazole) [4].

With the exception of SA02 and SA46, in all the other strains we identified *blaZ*, *bla*R1, and *bla*I genes, which encode for the penicillin-hydrolysing class A beta-lactamase BlaZ, the beta-lactam sensor/signal transducer BlaR1, and the penicillinase repressor BlaI, respectively. In *S. aureus*, the *blaZ* gene is carried by transposons [39], either located on a large plasmid or integrated into the bacterial chromosome. In strain SA24, we did not find any *rep* gene, which would suggest the presence of a plasmid. In contrast, such a *rep* gene was found in all the other *blaZ*-carrying strains. Moreover, the *blaZ* genomic context in SA24 does not appear to be similar to the other strains; indeed, it harbours few transposases, genes coding for YoID-like family proteins, integrases and recombinases, and the operon *pmt*ABCD, coding for the transporter responsible for the export of phenolsoluble modulins (PSMs), which modulate both biofilm formation and virulence in *S. aureus* [40]. In strain SA20, besides the presence of beta-lactam resistance genes, this region also harbours the pseudogene *sed*, as well as *selj* and *ser* genes, as described above (Figure 3).

In the *S. aureus* strain SA02 (resistant to ofloxacin, erythromycin, lincomycin, and tetracycline), we found an S84L substitution as a result of a mutation in the *gyrA* gene, which was reported to confer resistance to quinolones and fluoroquinolones [41]. The

SA02 genome also harbours the *erm*B gene encoding a 23S rRNA (adenine(2058)-N(6))dimethyltransferase associated with erythromycin resistance, a gene coding for the multidrug efflux MFS transporter LmrS involved in lincomycin resistance, as well as tet(L)and tet(38) genes, which are related to tetracycline resistance. tet(L) and tet(38) were closely located on a putative plasmid.

In the *S. aureus* strain SA18, we identified the aph(3')-IIIa and ant(6)-I genes, which encode the aminoglycoside O-phosphotransferase APH(3')-IIIa and the aminoglycoside nucleotidyltransferase ANT(6)-Ia, respectively, involved in the aminoglycoside resistance. These genes are located near *blaZ*, *bla*R1, and *bla*I on a mobile element. Furthermore, the genome of the strain SA18 also harbours the *tet*(L) and *tet*(38) genes involved in tetracycline resistance.

In addition, in the genome of the *S. aureus* strain SA46, which is resistant to benzylpenicillin, oxacillin, cefoxitin, and kanamycin, we identified the *aph*(3')-IIIa and *ant*(6)-I genes. The *blaZ* gene was not present, whereas the genome harbours the *mecA* gene coding for the low-affinity penicillin binding protein (PBP) 2a responsible for the β -lactam resistance [42].

The strain SA51 was susceptible to the benzylpenicillin, as well as to the other tested antimicrobials (oxacillin, cefoxitin, gentamicin, kanamycin, tobramycin, ofloxacin, erythromycin, lincomycin, clindamycin, pristinamycin, linezolid, teicoplanin, vancomycin, tetracycline, fosfomycin, nitrofurantoin, fusidic acid, rifampicin, and co-trimoxazole; [4]), even though the *blaZ* gene was shown to be present and intact in the genome of this strain.

2.7. Staphylococcal Chromosomal Cassette mec

In the genome of SA46, we identified a locus that has a homology of 88.74% with the staphylococcal chromosomal cassette mec (SCCmec) subtype-IVc(2B) and is identical to the clinical isolate GR2 (CP010402.1). The sequencing assembly has provided the dislocation different contigs: *mecA* of the SCC*mec* elements on two (J3T00_03665, NODE_6:211567..213573), (J3T00_03670, NODE_6:213673..214659), mecR1 IS1272 (J3T00_03680*, predicted as incomplete, NODE_6:214648..216490), ccrA2 (J3T00_04455, NODE_7:179619..180968), *ccr*B2 (NODE_7:180969..182618), and subtype-IVc(2B) (J3T00_04435, NODE_7:174448..175602). Although the prediction of SCCmecFinder classifies the SA46 mec complex as class B, the PAG pipeline predicted mecR1 as intact, with a sequence 100% identical to WP_001549960.1, whereas IS1272 was predicted as incomplete in the middle of the contig.

In SA04, genes with 92.2% of homology with *ccr*B1 (NODE_2:290655..292279) and 94.37% with *ccr*A1 (NODE_2:292301..293650) were predicted, but no whole SCC*mec* cassette was identified. In SA20, genes with 98.7%, 94.0%, and 99.8% of homology, respectively, were predicted with *ccr*A2 (NODE_20:3675..5024), *ccr*B2 (NODE_20:5046..6653), and subtype-IVa(2B): (NODE_10:93114..94604); moreover, in this case, no complete SCC*mec* cassette can be detected.

2.8. Virulence Determinants

Comparative analysis of virulence determinants is described by the heatmap shown in Figure 4, whereas the list of genetic determinants is reported in Supplementary Table S8. All strains harbour a repertoire of genes coding for a) proteins involved in the adherence, such as the autolysin gene, clumping factors *clfA* and *clfB* (this latter is a pseudogene in SA08), and elastin binding protein gene; b) exoenzymes such as the hyaluronate lyase gene *hysA*, serine proteases, lipases, and the coagulase gene *vWbp* (von Willebrand factor-binding protein); c) proteins involved in the immune modulation and in the heme uptake system; and d) several exotoxins (Figure 4; Supplementary Table S8).



Figure 4. Virulence determinants in *S. aureus* genomes. Heatmap of virulence determinants in *S. aureus* genomes. Presence and numbers of genes are indicated by green colour intensity; absence of gene was indicated by a white square; pseudogenes are indicated as red-striped squares. Heatmap was visualised by using heatmapper web server [43] with average linkage as clustering method and Euclidean distance measurement method.

The staphylococcal superantigen-like (*ssl*) genes are arranged as tandem repeats in the genomic island α (*ssl*1-11) and in the immune evasion cluster 2 (IEC2, *ssl*12-14) on a vSa γ island on the bacterial chromosome [44]. This last island also comprises the *hyl* gene, which codes for the α -haemolysin, one fibrin-binding protein, and the *scn* gene coding for the staphylococcal complement inhibitor [45]. Only in *S. aureus* SA02 the contig is interrupted downstream *hyl*.

S. aureus SA82 is the only strain that harbours the *tst* gene coding for the toxic shock syndrome toxin (TSST-1), whereas the *luk*F-PV and *luk*S-PV genes coding for the Panton–Valentine leukocidin (PVL) are present on a bacteriophage φ Sa2 only in the genome of *S. aureus* SA46.

In the heatmap built on the presence or absence of genes or pseudogenes coding for virulence determinants, the strains are clustered into two groups, with SA82 in an individual clade, as already shown by the phylogenetic analysis (Figure 1). The SA82 genome, although harbouring the *tst* gene, lacked genes coding for a) four out of five secreted virulence factors (EsxB, EsxC, EscD, and EsaD); b) the soluble cytosolic protein EsaE of the type VII secretion system; and c) the bi-component staphylococcal leukotoxins family LukDE [46]. It also missed d) the *efb* gene coding for a fibrinogen binding protein; e) the *fnb*B gene, encoding a fibronectin binding protein; and f) all the genes coding for serine proteases. The *hlg*A gene, coding for the gamma haemolysin, is predicted to be a pseudogene. In contrast to all the other strains, the SA82 genome harbours two *cna* genes, encoding the collagen binding protein, which mediates bacterial adherence to collagen substrates and collagenous tissues, and is strongly associated with the pathogenesis of osteomyelitis and septic arthritis [47].

The *hlb* gene is interrupted due to a prophage integration in all strains with the exception of SA20.

2.9. Growth of S. aureus Strains in BHI + YE Broth and Production of Staphylococcal Enterotoxins

The concentrations of *S. aureus* strains SA04, SA18, SA20, SA46, and SA51 (harbouring *sea, seb, sed, seh,* and *ser* genes) at 0, 24, and 48 h of incubation in brain heart infusion broth supplemented with yeast extract (BHI + YE) are shown in Table 3. The pH values of the corresponding broth cultures are also reported (Table 3). At 0 h (immediately after the inoculum) the initial counts of all strains were similar (ranging between 6.38 ± 0.28 and 6.91 ± 0.29 Log cfu/mL). Major changes were observed after 24 h of incubation, when all bacterial numbers increased by more than 2 Log cfu/mL (ranging between 8.97 ± 0.09 and 9.34 ± 0.23 Log cfu/mL). Concentrations after 48 h of incubation were found at similar levels to those found after 24 h of incubation (ranging between 9.08 ± 0.13 and 9.44 ± 0.28).

Chusin	0 h		24 h		48 h		
Strain	Log cfu/ml	pН	Log cfu/ml	pН	Log cfu/ml	pН	
SA04	6.77 ± 0.58	7.61	9.22 ± 0.03	6.86	9.22 ± 0.12	7.53	
SA18	6.91 ± 0.07	7.61	9.34 ± 0.23	6.97	9.44 ± 0.28	7.57	
SA20	6.91 ± 0.29	7.61	8.97 ± 0.09	6.71	9.08 ± 0.13	7.29	
SA46	6.39 ± 0.26	7.61	9.29 ± 0.08	7.05	9.26 ± 0.08	7.61	
SA51	6.38 ± 0.28	7.61	9.15 ± 0.11	6.81	9.16 ± 0.07	7.23	

Table 3. Growth and pH values of *S. aureus* cultures in BHI + YE broth.

With the exception of SA46, for which the SEH production was 1.76 ± 2.26 ng/mL at 24 h and not detectable at 48 h, all the other strains produced higher amounts of SEA, SEB, SED, SEH and SER at both 24 h and 48 h of incubation in the BHI + YE broth, as shown in Table 4.

Table 4. Production of SEA, SEB, SED, SEH, and SER (ng/mL) by S. aureus strains in BHI + YE broth.

Strain	SEA		SEB		SED		SEH		SER	
	24 h	48 h	24 h	48 h	24 h	48 h	24 h	48 h	24 h	48 h
SA04	$156.04 \pm$	$345.02 \pm$	-	-	-	-	221.76 ±	$239.48 \pm$	-	-
	36.58	62.67					38.15	84.83		
SA18	-	-	$2871.28 \pm$	14739.17±			-	-	-	-
			811.09	5077.70	-	-				
SA20	-			-	$322.70 \pm$	$398.94 \pm$	-		$36720.10 \pm$	$63176.06 \pm$
		-			41.17	64.79			6272.32	14422.93
SA46	-	-	-	-	-	-	1.76 ± 2.26	N.D.	-	-
SA51	$64.54 \pm$	74.91 ±	-	-	-		-	-		
	9.29	7.36				-			-	-

N.D.: not detectable.

2.10. Expression of the seh Gene

The relative expression of the *seh* gene was evaluated in SA46 as well as in the other *seh*-positive strain SA04, at 5 and 24 h of incubation in the BHI + YE broth. As shown in Figure 5, in SA46 no transcription of the *seh* gene was observed at 5 h of incubation, whereas a very low transcription level compared to SA04 (0.13 fold) was detected at 24 h of incubation. In SA04 *seh* expression decreased at 24 h compared to that at 5 h (0.39-fold).

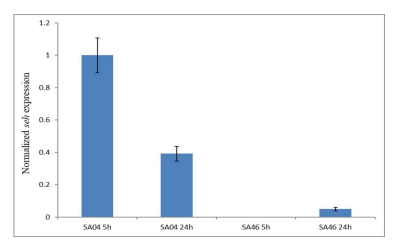


Figure 5. *seh* relative gene expression. *seh* relative gene expression in *S. aureus* SA04 and SA46 at 5 h and 24 h of incubation in BHI + YE broth.

3. Discussion

In this study, we performed a comprehensive genomic analysis of 12 *S. aureus* strains isolated from 207 Algerian ready-to-eat foods to explore their genetic relatedness, gain further data on *S. aureus* molecular epidemiology, and investigate their virulence and antibiotic resistance genetic determinants that may help to better understand the features of foodborne *S. aureus* in Algeria as well as in the African continent.

Two novel alleles, aroE-1025 and yqiL-954, and two novel STs, 7199 and 7200, were identified and deposited in the PubMLST public database, thus contributing as new suitable resources for monitoring the global and local epidemiology of *S. aureus*. Among the other nine detected STs, six have been previously reported in Algeria, i.e., ST1-, ST8-, ST15-, ST22-, and ST97-MSSA clones as well as the ST80-MRSA clone [48,49], whereas ST5-, ST25-, and ST101-MSSA clones have been detected in other African countries [12,50,51]. In our study, the detection of a strain that is a member of the ST80-MRSA-SCCmec-IV lineage, also referred to as the European clone [52], is of particular interest since it has been described as having an alarming dissemination in Algeria [49]. It is worth mentioning that the clones herein found in Algerian ready-to-eat foods have been also found in humans (ST1-, ST5-, ST22-, ST25-, ST97-, and ST101-MSSA and ST80-MRSA-SCCmec-IV; [12,49-51]) and animals (ST1-, ST8-, ST15-, ST22-, and ST97-MSSA and ST80-MRSA-SCCmec-IV; [48,49]). This confirms the complexity of S. aureus epidemiology and highlights the fact that ready-to-eat foods, besides posing a risk to consumers for the onset of SFP, may also serve as means for S. aureus dissemination via food ingestion or handling, as pointed out also by other authors [3,53,54].

Considering the role of *S. aureus* in foodborne diseases due to the production of SEs [3], understanding the enterotoxigenic capabilities of circulating strains is a pivotal issue. Following the discovery of the classical SEs (SEA-SEE) [55,56], the number of the newly discovered SEs and SEls is constantly increasing. In particular, a total of 33 se and sel genes (sea-see; seg-selu; selu2; selv-selz; sel27-sel33) are known to date, including the last five sels recently reported (sel29 pseudogene; sel30-sel33) [1], thus indicating the wide enterotoxigenic potential of *S. aureus*. To the best of our knowledge, this is the first study searching for all the se and sel genes known to date and reporting the sel30 gene in Algerian S. aureus isolates from food after its recent discovery [1]. The gene, detected in SA51, shares 100% homology with the sequence reported by Dicks et al. [1]. Recently, some authors made efforts to gain data on the enterotoxigenic potential of foodborne S. aureus in Algeria, searching for some of the known se and sel genes (sea-see; seg-sek; selp and ser) [8,10,11], as well as the presence of the enterotoxin gene cluster (egc) [8]. These authors found the sea, seb, sec, see, seg, seh, sei, selj, sek, selp, ser, and the egc, although, in the latter case, no information on the actual egc gene content and the egc-type was provided [8,10,11]. We should remark that for the assessment of S. aureus enterotoxigenicity, not only the presence of *se* and *sel* genes but also the evaluation of genetic variants and the presence of pseudogenes, as herein performed, should be considered, since they may directly affect the enterotoxin production and virulence of the hosting *S. aureus* strains.

According to the results of our genome analysis, *selw* and *selx* genes were identified in all the sequenced strains, both located in the chromosome, in line with other studies [15,32]. *selw* was predicted as a pseudogene in 4 of 12 *S. aureus* strains, confirming that this gene, likely due to its chromosomal localisation, has a high rate of pseudogenisation [1]. Otherwise, no pseudogenes for *selx* were herein detected, corroborating previous reports in which an overall low prevalence of *selx* pseudogenes was observed, probably linked only to specific *S. aureus* lineages [1,16].

Most of the SFP diseases are associated with the classical SEs (SEA-SEE) [57]; furthermore, SEH has been found as a causative agent of SFP outbreaks [25,58]. Moreover, other newly discovered SEs such as SER and those belonging to the enterotoxin gene cluster (*egc*) are being reported with increasing evidence to have a possible role in foodborne intoxications [59-61].

The sea gene belonging to allele class 1 (sea1) [17] was identified in SA04, whereas SA51 harbours the *sea*2, a single-nucleotide sequence variant of the *sea*2 reported by Borst and Betley [17], that leads to no amino acid change in the predicted protein sequence (Table 1; Supplementary Table S7). SA04 produced higher amounts of SEA (156.04 ± 36.58- 345.02 ± 62.67 ng/mL) compared with those produced by SA51 ($64.54 \pm 9.29-74.91 \pm 7.36$ ng/mL), being 2.4- and 4.6-folds higher at 24 h and 48 h, respectively. SEA, responsible for the majority of the SFP cases (about 80%) [57], is reported to cause human intoxication symptoms at a low dose of about 20–200 ng [62,63], and it is frequently detected in small amounts (ranging from 0.015 ng to >6 ng per gram or millilitre of food) in foods involved in SFP outbreaks [62-65]. These findings highlight the potential pathogenicity that our SEA-producing *S. aureus* strains may have in the contest of SFP. As regard the difference in SEA production observed in SA04 and SA51, it has been associated, by Borst and Betley [17], to the region upstream the translational start sites of *sea*. This region in SA04 is identical to that of the high-SEA-producing S. aureus FRI100 (harbouring sea allele class 1), whereas that in SA51 is identical to that of the low-SEA-producing S. aureus FRI281A (harbouring sea allele class 2) (Supplementary Table S7). However, the dissimilarities in the SEA amounts registered for SA04 and SA51 were not so pronounced as for S. aureus FRI100 and S. aureus FRI281A, since also other loci may affect the phage-mediated mechanism responsible for SEA expression [17].

*sebv*1 was identified in *S. aureus* SA18. This strain was able to produce SEB (2871.28 \pm 811.09-14739.17 \pm 5077.70 ng/mL) corroborating the findings of Johler et al. [18] who reported that all the described *seb* variants (*sebv*1-v4) are expressed, leading to the SEB production. In the literature, SEB-related SFPs are poorly documented; nevertheless, in a described outbreak involving boiled eggs, a high SEB concentration, greater than 1000 ng/g [66] and thus similar to that produced in the BHI + YE broth by our *S. aureus* strain, was reported. Unlike *sea*, differences in the *seb* upstream region were reported as not directly correlated with the SEB production level, which, on the other hand, may be linked to the type of the *seb*-harbouring SAPIs. Interestingly our *seb*-positive strain harbours a SAPI that has 99.4% similarity with SaPIivm60, which has been associated with strong SEB production by Sato'o et al. [35].

In SA20, sed was predicted as a pseudogene, which differs for a nucleotide substitution (leading to 1 amino-acid substitution) from the truncated sed described by Johler et al. [18] (Table 1; Supplementary Table S7). SA20 was able to produce SED (322.70 \pm 41.17–398.94 \pm 64.79 ng/mL), predicted as shorter than the intact protein (179 compared with the 258 amino acids of the intact SED) [18]. We should highlight that the prediction of the toxic properties based on the changes in the length and/or amino acid sequence of the SEs may be difficult, considering that the mechanism of action of SEs is not fully elucidated [18]. However, it should be highlighted that the truncated SED may have an impaired toxic function, resulting in a reduced virulence of the producing *S. aureus* strain. Low (0.052 ng/g; [67]) to high (>200 ng/g; [63]) concentrations of SED have been detected in foods involved in SFP outbreaks, although, to the best of our knowledge, no estimation of the oral dose capable of causing human symptoms has been reported. Some S. aureus strains harbouring the truncated *sed* gene were found either to not produce the relevant protein at detectable levels or even to not transcribe this pseudogene [18,68]. Nevertheless, Johler et al. [18] reported that certain S. aureus harbouring the truncated sed were able to produce the SED protein, being in line with our finding, although they found that the amounts were far lower than those produced by strains harbouring the intact sed gene.

selj and *ser* genes were also present in SA20 and located along with *sed* on the same putative pIB485-like plasmid as previously described [36]. Although also *ser* pseudogenes have been recently reported [7,69], SA20 harbours an intact *ser* gene and was able to produce SER ($36720.10 \pm 6272.32-63176.06 \pm 14422.93$ ng/mL) in amounts more than 100-folds higher than SED. Interestingly, *S. aureus* strains harbouring the plasmid-related *sed*, *selj*, and *ser* genes have been isolated from seven SFP outbreaks in Japan [60], and their

ability to produce SER in BHI + YE (523–2925 ng/mL) has been demonstrated, suggesting that SER may have played a possible role in the onset of the described SFPs [60].

The *seh* gene was detected as intact or as a pseudogene in two of our sequenced strains. SA04, harbouring the intact seh, was able to produce the enterotoxin SEH (221.76 \pm 38.15-239.48 \pm 84.83 ng/mL) (along with SEA, as discussed above), and, notably, the nucleotide sequence was identical to that of the *seh* gene found in *S. aureus* strains that caused a SEH-mediated SFP outbreak ([25]; Supplementary Table S7), indicating, therefore, the potential virulence of this strain. In particular, in the reported outbreak, a concentration of SEH equal to 55.5 ng/g was estimated in the consumed mashed potato made with raw milk, and the tested S. aureus strains, isolated from these leftovers and the farm bulk raw milk, were able to produce SEH in concentrations similar (96–108 ng/mL) to that produced by our SA04 strain, when grown in a microbial broth [25]. The seh pseudogene was instead detected in the SA46 strain belonging to the European MRSA-ST80-SCCmec-IV clone, corroborating the study of Dicks et al. [1] who recently reported this pseudogene as strongly associated with the MRSA-ST80-SCCmec-IV clone. SA46 also harboured the sey gene, whose sequence was previously detected in a ST80 strain [24]. SA46 produced SEH at almost undetectable levels. To investigate if the difference in the SEH amounts produced by SA04 and SA46 may depend on differences in seh expression, we assessed it taking into account that, in experimental conditions similar to those herein applied, the *seh* gene expression occurs during the exponential phase of growth [70]. The expression analysis detected no transcripts at 5 h for SA46, whereas at 24 h, their levels were much lower than those of the intact seh gene, harboured by SA04, confirming that, in this case, the failure of the SEH production in SA46 depends on transcriptional regulation. This is interesting since certain se pseudogenes may be transcribed at levels similar to the intact se genes, as found by Lis et al. [68], suggesting that the protein synthesis failure may occur later, at the translational level.

Although the involvement of MRSA in SFP has been reported [71] based on these findings and considering the relative little assortment of *se* and *sel* genes (*seh* pseudogene, *sey*, *selx*, and *selw*), it seems that our MRSA strain (SA46) may play a minor role in the onset of SFP compared with the other enterotoxin producing MSSA strains herein detected. A previous study found one *S. aureus* strain belonging to the European clone carrying the *egc*-related genes *seg*, *sei*, *sen*, *seo*, and *selu* [72], whereas *seb* and *sek* were harboured by atypical MRSA-ST80-SCC*mec*-IV *pvl*-negative isolates [73]. Therefore, our findings may help to enrich the knowledge regarding the endowment of *se* and *sel* genes within the European clone in order to improve the understanding of the enterotoxigenic potential of this lineage.

The *egc* was found in 5 out of the 12 sequenced *S. aureus* strains, corroborating previous investigations that reported that *egc*-related genes are among the most prevalent genes in foodborne *S. aureus* isolates [74]. Beyond its correlation to SFP, the *egc* presence seems to be relevant also in other clinical conditions, such as chronic infections in cystic fibrosis patients [75]. This cluster may harbour several *se* and *sel* genes (i.e., *seo, sem, sei, selu, sen, seg, selu2, selv,* and *sel33*) and pseudogenes (ψ *ent1*- ψ *ent2*) in different combinations, and, to date, at least eight *egc* types have been described in *S. aureus* [1,7]. Two of our strains harbour the *egc1* (*seg, sei, sen, seo, and \psient1-\psient2*) and three strains harbour the *egc5* (*seg, sei, sen, seo, and selu2*). Both these types have been detected in foodborne *S. aureus* isolates [7], but there is still scarce knowledge on their actual prevalence mostly due to the fact that *egc5* was only recently described [7].

The emergence of antimicrobial resistance in *S. aureus* is a clinical and public health challenge due to the reduction of the antimicrobial assortment that is effective when *S. aureus* infections occur. In addition to MRSA, whose presence has been globally reported in various ecological niches including African food, animals, and humans [13,76], also multidrug-resistant MSSA, typically showing resistance to three or more antimicrobials of different classes, as herein reported for SA02 and SA18, are being detected in various sources comprising food products (especially animal-derived unprocessed products) in

Africa [23,77-79] and worldwide [54,80-82]. Genome analysis allowed us to detect the antimicrobial resistance genes related to the phenotypic resistance profile shown by our sequenced *S. aureus* strains. In particular, the *blaZ* gene and regulatory system composed of *bla*R1 and *bla*I that we found in 10 out of the 12 sequenced strains represent the genetic mechanism conferring resistance to benzylpenicillin and other penicillinase-labile penicillins. We should emphasise that in *S. aureus* SA51 we found the *bla* locus (*blaZ*, *bla*R1, and *bla*I), although it was reported as benzylpenicillin-susceptible. Since 2012, CLSI acknowledged that penicillin-susceptible results should be further investigated before being confirmed, performing additional phenotypic tests to detect β -lactamase production (i.e., nitrocefin-based tests and/or penicillin zone edge) [83]. However, these phenotypic methods were described as less sensitive compared with the molecular detection of the *blaZ* gene, and, therefore, different authors considered the latter the gold standard method for β -lactamase detection in *S. aureus* [84,85].

blaZ is also frequently found in MRSA [86,87], including the ST80-MRSA-SCC*mec*-IV lineage as previously reported for most of the analysed isolates by Monecke et al. [86]. Our ST80-MRSA-SCC*mec*-IV strain, SA46, although harbouring the *mec*A gene conferring resistance to β -lactam antimicrobials [87] such as benzylpenicillin, oxacillin, and cefoxitin, does not harbour the *blaZ* gene, and, in Africa (in Cape Verde), a similar feature was recently reported for other MRSA isolates belonging to the ST5-SCC*mec*-VI lineage [88]. Although the *bla* system is involved in the regulation of the *mec*A expression [89], it has been elucidated that the loss of the *bla* system may allow the constitutive expression of the *mec*A gene, conferring β -lactam resistance in SCC*mec*-IV harbouring MRSA [89].

It has also been reported that 91.6% of ST80-MRSA isolates are resistant to kanamycin, and the kanamycin-resistance gene aph(3')-III has been detected in the ST80-MRSA clone [90], which is consistent with our finding. Beyond aph(3')-IIIa, also the other detected genes conferring resistance to aminoglycosides (ant(6)-I), lincosamides (lmrS), tetracyclines (tet(L), tet(38)), macrolides (ermB), quinolones, and fluroquinolones (gyrA-S84L) have been previously reported in *S. aureus* isolates [91-94]. Nevertheless, a paucity of studies detected a range of antimicrobial resistance determinants in foodborne *S. aureus* in Africa, and, in particular, only few of the aforementioned genes were previously reported in Tunisia (aph(3')-IIIa, tet(L); [95]), Kenya (aph(3')-IIIa; [96]), and South Africa (aph(3')-1-IIIa, gyrA-S84L; [77,97]). Therefore, the present study helps to further expand the knowledge on the genetic determinants harboured by antimicrobial-resistant *S. aureus* isolated from food in the African continent.

The results of the genomic analysis highlighted a plethora of different virulence determinants in all the sequenced strains, involved in adhesion, colonisation, spreading, and immune modulation. Adherence is the first process involved in the pathogenicity of S. aureus. It is determined by the expression of several genes such as that coding for the autolysin, which mediates adherence to immobilised fibrinogen and fibronectin [98], the clumping factors *clfA* and *clfB* that bind to different sites in fibrinogen, and fibronectin binding proteins A and B, which facilitate the attachment of S. aureus to host cells [99].These genes are present in all the sequenced strains. Exoenzymes such as the hyaluronate lyase coded by *hys*A are required for the degradation of hyaluronic acid, which contributes to the local dissolution of the extracellular matrix, whereas vWbp enables the bacteria to disseminate and resist to opsonophagocytic clearance by host immune cells [100]. Among the exotoxins, haemolysin genes were retrieved in all the strains. The *hlb* gene, which codes for a sphingomyelinase that cleaves sphingomyelin into phosphocholine and ceramide [101], was a pseudogene in all strains, with the exception of SA20. The interruption of this gene is caused by the integration of a prophage of Sa3int type. Loss and reintegration of this phage are correlated with the transmission of S. aureus from humans to livestock and vice versa [102]. The restoration of the hlb gene, which occurred in SA20, might be relevant for important specific infections [103,104].

Panton-Valentine leukocidin (PVL) is a major virulence factor of *S. aureus*, although it presents in less than 5% of *S. aureus* strains. It is a two-component toxin that induces

pore formation in the leukocyte cell membrane complement receptors [105]. The results of our genomic analysis detected *pvl* genes only in *S. aureus* SA46, harbouring SCC*mec* of type IV. PVL has been proposed as a marker of CA-MRSA [106], although PVL-negative CA-MRSA clones have been reported [52].

According to the international working group on the Classification of Staphylococcal Cassette Chromosome [107], the genomic sequence of SA46 meets the criteria for defining an SCC*mec*: i) carriage of *mecA* in a *mec* gene complex; (ii) presence of *ccr* gene(s) (*ccrAB* and/or *ccrC*) in the *ccr* gene complex; (iii) integration at a specific site in the staphylococcal chromosome, designated as the integration site sequence (ISS) for SCC, which serves as a target for *ccr*-mediated recombination; and iv) presence of flanking direct repeat sequences containing the ISS. SCC*mec* of SA46 was predicted as SCC*mec* subtype-IVc(2B), although the B class of the *mec* complex is characterised by the presence of *mecA* and a truncated *mec*R1 resulting from the insertion of IS1272 upstream *mecA* [108]. On the contrary, the annotation of this genome returned an intact *mec*R1 and an incomplete IS1272, in the middle of a contig. Indeed, the whole cassette was dislocated in two separate nodes. This failure in reconstructing the entire SCC*mec* as the result of short-read sequencing technology is due to the multiple insertion sequences, which cause difficulties in the assembly.

To date, only two studies performed a genomic analysis of Algerian *S. aureus* strains: one by Mairi et al. [109], who reported the first occurrence of the "Maltese clone" (CC5-MRSA-IV-SCC*fus*) in Bat Guano, and a second by Aouati et al., [110], who uncovered the emergence of MRSA SCC*mec*-III Mercury in clinical isolates in Eastern Algeria. Other sequence read archives (SRAs) obtained by sequencing *S. aureus* isolated from Algerian dairy products were deposited in NCBI (Accession ERS4338951-ERS4338960).

This is the first genomic study performed on Algerian foodborne *S. aureus* strains and, to the best of our knowledge, the first comparative and phylogenetic reconstruction of foodborne *S. aureus* of worldwide origin. According to our data, there is a clear phylogenetic association of *S. aureus*, based on the ST or the geographical origin, rather than the source of isolation. This reflects the dynamics and evolution of the *S. aureus* population worldwide. The arising grouping based on the food source might be an effect of the sample collection rather than a clustering based on the use of a specific ecological niche.

The Algerian isolates are dispersed in the phylogenetic tree with the exception of SA51 and SA02, which both belong to the ST5, and SA01 and SA07, which differ for the *aroE* allele in the MLST. From an evolutionary point of view, SA08 is the less recent strain, whereas SA82, which is individually located in all the cluster analyses we have performed, is the most recent. However, such phylogenetic reconstructions are affected by the fact that most strains, although sharing the same geographical origin, have been isolated from different food products, some generically indicated as food, and no information is available on the manufacturing process nor on the origin of raw material used, thus hindering an exhaustive reconstruction of phylogenetic and epidemiological maps.

This underlines the urgent need for genomic studies on foodborne *S. aureus* in Algeria to have more detailed insights regarding the structure of the *S. aureus* population as well as the harboured genetic determinants of virulence and antibiotic resistance, in order to be able to estimate the public health burden related to the spread of this pathogen in food.

4. Conclusions

A comprehensive genomic characterisation and the production of some classical and newly described staphylococcal enterotoxins of 12 Algerian *S. aureus* strains, previously isolated from 207 ready-to-eat foods, were herein performed. Two novel STs were identified, and a strain belonging to the alarming MRSA-ST80-SCC*mec*-IV European clone was detected. An up-to-date assessment of *se* and *sel* genes in our foodborne *S. aureus* strains was carried out, and we found that MSSA strains produced higher amounts of enterotoxins than the MRSA-ST80-SCC*mec*-IV strain herein sequenced. The detection of antimicrobial resistance genetic determinants in the 12 *S. aureus* strains supported the results of the antibiotic susceptibility testing previously performed by Mekhoulfi et al. [4]. Furthermore, a plethora of virulence genes were found in the 12 *S. aureus* genomes, although SA82 lacked several determinants involved in adhesion and colonisation, confirming its outgrouping position in the taxonomic clustering.

The results of the genomic analysis, together with the production of enterotoxins investigated in this study, confirm the risk associated with the spread of this pathogen in food, which is worsened by the observation that some of these *S. aureus* strains are multidrug resistant, and by the detection of the *tst* gene, determinant for the toxic shock syndrome toxin-1, in *S. aureus* SA82, and the PVL-encoding genes in the methicillin-resistant SA46 strain.

5. Materials and Methods

5.1. S. aureus Strains Used in the Study

The 12 *S. aureus* strains sequenced in this study are included in Supplementary Table S4. These strains were cultured on Baird–Parker egg yolk (BPEY, Oxoid, France), and their purity was checked by streaking on the same agar medium. The pure cultures were stored as stock cultures at -80° C in brain heart infusion broth (BHI; Conda. Pronadisa, Spain) supplemented with 0.6% yeast extract (Biolife Italiana, Milan, Italy) and 20% glycerol. For DNA extraction, the working culture was prepared as described by Fusco et al. [111].

5.2. Whole-Genome Sequencing and Analysis

Two millilitres of fresh working cultures of each of the *S. aureus* strains were used for the DNA extraction following the protocol described by Chieffi et al. [7]. The integrity, purity, and quantity of DNA were assessed by agarose gel electrophoresis, a Nanodrop photometer (Peqlab-VWR International Srl, Darmstadt, Germany), and a Qubit 3.0 fluorimeter (Thermo Fisher Scientific, Waltham, Massachusetts, USA). DNA was subjected to whole-genome shotgun sequencing by using an Illumina mate pair library prep kit (Illumina, San Diego, California, USA), according to the manufacturer's instructions, and then sequenced on an Illumina MiSeq platform with the 2×250 mate pair procedure. Reads were then trimmed by NxTrim (V2) [112], and *de novo* assembly was performed using SPAdes version 3.10.1 (©St. Petersburg State University, St. Petersburg, Russia; [113]) with the following parameters: Kmer 21, 33, 55, 77 and "careful".

The overall contiguity of the assembly and genome statistics were determined with MIGA [114]. The completeness of the *de novo* assemblies was measured by the presence of 106 single-copy genes that are observed across almost all prokaryotic genomes by using MiGA [114]. Contamination was calculated based on the copy number of essential genes (in %) present in the genome. Quality scores were then calculated as completeness percentage minus five times contamination percentage.

The whole-genome shotgun projects have been deposited at DDBJ/ENA/GenBank under the accessions reported in Supplementary Table S4. The versions described in this paper are JAHLTR010000000 for S. aureus SA01, JAFNJM010000000 for S. aureus SA02, JAFNJN010000000 for S. aureus SA04, JAHLTS010000000 for S. aureus SA07, JAHLTT010000000 for S. aureus SA08, JAHLTU010000000 for S. aureus SA10, JAFNJO01000000 for aureus SA18, JAFNJP01000000 S. aureus S. SA20, for S. aureus SA24, JAFNJQ010000000 for S. aureus JAFRED01000000 SA46, JAFNJR010000000 for S. aureus SA51, and JAHLTV010000000 for S. aureus SA82.

5.3. Bioinformatic Methods

5.3.1. Gene prediction, ANI, and Phylogenomic Analysis

Genes were predicted and annotated using the PROKKA pipeline implemented in the Galaxy platform (Galaxy Tool Version 1.14.5; [115]) and by NCBI (National Center for Biotechnology Information) Prokaryotic Genome Annotation Pipeline (PGAP; [116]). Protein ID used in the manuscript indicated those obtained by PGAP.

All the protein sequences used in this study were retrieved from GenBank (NCBI). The homology-based relationship of *S. aureus* genes and predicted proteins towards reference sequences was determined by the BLASTN and BLASTP algorithms on the NCBI site [117]. Gene models were manually determined, and clustering and orientation were subsequently deduced for the closely linked genes.

Strains used to perform pan-genome and phylogenetic analyses are listed in Supplementary Table S4. Genomic sequences were downloaded from the NCBI and submitted to the PROKKA pipeline. The obtained .gff3 files were used as input for Roary (Galaxy Version 3.13.0+galaxy1), the pangenome pipeline [118], to generate a core gene alignment with 95% as the minimum percentage identity for blastp analysis and 99% as the total percentage of the isolates in which the gene needs to be present for it to be considered a core gene. The pan-genome was represented as the core genome (shared by >99% of strains), accessory genome (genes present in >2 strains but not in all), and unique genome (genes unique to individual strains). The total pan-genome was also shown as core (99% \leq strains \leq 100%), soft core (95% \leq strains < 99%), shell (15% \leq strains < 95%), and cloud (0% \leq strains < 15%). Visualisation of the output was achieved by using Phandango (version 1.3.0; [119]).

The genetic divergence among *S. aureus* species was calculated using the ANI calculator [120,121], which estimates the average nucleotide identity (ANI) using both best hits (one-way ANI) and reciprocal best hits (two-way ANI) between genomic datasets.

Genome-based phylogeny using 1000 single copy genes was reconstructed employing the Phylogenetic Tree Building Service implemented in the BV-BRC platform [122], with the maximum likelihood method RAxML (version 8.2.11) and progressive refinement [123]. *S. argenteus* MSHR1132 was used as an outgroup, and visualisation of the phylogenetic trees was performed by using iTOL (version 6.3.2; [124]).

5.3.2. MLST and spa typing

Multilocus sequence typing (MLST) was performed by MLST 2.0 (software version: 2.0.4 (2019-05-08); database version: 2.0.0 (2021-09-13)) of the Center for Genomic Epidemiology [125,126]. *spa* typing was performed by spaTyper (Software version 1.0 Database version: 2022-08-22) of the Center for Genomic Epidemiology [127,128].

5.3.3. Antibiotic Resistance and Virulence Determinants Analyses

Antibiotic resistance determinants were computationally predicted within the BV-BRC platform [129] by a k-mer-based detection method and BLAST analysis, and then manually curated. Virulence determinants were screened by using the comparative pathogenomics-based VF analysis pipeline VFanalyzer [130] implemented in the virulence factor database (VFDB) [131], and then manually curated. A heatmap was manually constructed and visualised by using the heatmapper web server [43,132] with average linkage as the clustering method and Euclidean distance measurement method.

Enterotoxin genes were determined by homology search using previously reported reference sequences ([1]; Supplementary Table S7) using an E-value cut-off of 0.001, a minimum percentage of alignment identity of 80%, and a relative coverage threshold of >80% [41,133].

Prediction of SCC*mec* elements in sequenced *S. aureus* isolates was performed by using the SCC*mec*Finder v1.2 [134,135] with default parameters, then manually curated.

5.4. Cultivation of S. aureus for Staphylococcal Enterotoxin Production Assessment

S. aureus strains SA04, SA18, SA20, SA46, and SA51, harbouring the classical and some newly described *se* genes (namely *sea, seb, sed, seh,* and *ser*), were chosen to assess the ability to produce SEs. Bacterial cultures were prepared as described by Schubert et al. [136]. Briefly, 100 mL of brain heart infusion supplemented with 1% yeast extract (BHI + YE, Biocorp, Warsaw, Poland) were inoculated with pre-culture to reach an optical density of 0.02 at 600 nm (OD₆₀₀). Prior to inoculation, the pre-cultures were washed twice with phosphate-buffered saline (PBS) to remove residual BHI broth and enterotoxins (repeated centrifugation at 12,000× g for 5 min and resuspension in PBS). Cultures were incubated at 37 °C with constant agitation at 230 rpm. The bacterial cell concentration of the broth cultures at 0, 24, and 48 h of incubation was determined by plating serial tenfold dilutions, prepared in quarter-strength Ringer's solution (Oxoid, Basingstoke, United Kingdom), onto BHI agar. The pH of the broth cultures was measured using a FE20-FiveEasyTM pH-meter (Mettler-Toledo, Greifensee, Switzerland).

5.5. Sandwich ELISA for SEA, SEB, SED, SEH, and SER Detection

Recombinant staphylococcal enterotoxins (rSED, rSEH, and rSER) were obtained as previously described by Schubert et al. [136,137]. Rabbit polyclonal anti-A, anti-B, anti-SED, and anti-SER antibodies were purchased from OriGene Technologies GmbH (Herford, Germany), whereas rabbit polyclonal anti-H and anti-H(HRP) antibodies were purchased from Abcam (Cambridge, UK). Samples for staphylococcal enterotoxins (SEs) detection were collected after 24 and 48 h of growth and stored at -20 °C until analysed. Supernatants were pre-incubated with a 20% normal rabbit serum to bind protein A and then diluted in PBS containing 0.1% Tween-20 (Merck, Darmstadt, Germany). The enzyme linked immunosorbent assay (ELISA) was also performed as described by Schubert et al. [136,137]. The concentration of the SEs in samples was measured with SEA, SEB (Merck KGaA, Darmstadt, Germany), rSED, rSEH, and rSER as standards, using a 4-parameter logistic curve fit. The ELISA assays for each SE were run in two biological repeats with two replicates. Data analysis was carried out using GraphPad Prism software 8.0.1 (GraphPad Software Inc., La Jolla, CA, USA).

5.6. RNA Extraction and Reverse Transcription–Quantitative PCR to Assess seh Gene Expression

Expression of the *seh* gene was assessed in the *seh*-positive *S. aureus* SA04 and SA46 strains. Samples for RNA isolation were collected from the BHI+YE broth after 5 and 24 h of growth. RNA extraction, purification, cDNA synthesis, and reverse transcription–quantitative polymerase chain reaction (RT-qPCR) were performed as described by Schubert et al. [136]. The *rpo*B housekeeping gene was used for normalisation [138]. Transcript levels of *seh* relative to *rpo*B were calculated according to Pfaffl [139]. Each RT-qPCR assay consisted of two biological repeats with three replicates. Data analysis was carried out using Bio-Rad CFX Manager software.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/toxins14110731/s1, Supplementary Figure S1. Pan-genome analysis of *S. aureus* strains. Visualisation of pan-genome analysis of *S. aureus* strains was performed by using Phandango (version 1.3.0). Supplementary Figure S2. Enterotoxin gene cluster (*egc*) localisation in *S. aureus* strains. Protein IDs are indicated in the figure. Red: enterotoxins; yellow: serine proteases; brown: leukocidins; grey: hypothetical proteins; light blue: restriction endonucleases; pink: transposases; turquoise: lantibiotic cluster; *cap*B: calcium binding protein; *pep*A1: glutamyl aminopeptidase; * indicates a pseudogene. Supplementary Table S1. Genomic features of *S. aureus* strains. Supplementary Table S3. *spa* typing of *S. aureus* strains. Supplementary Table S4. List of *S. aureus* strains used in this study. Supplementary Table S5. ANI values of *S. aureus* strains. Supplementary Table S6. Roary statistics. Supplementary Table S7. *se* and

sel genes and predicted protein sequences in *S. aureus* strains. Supplementary Table S8. Virulence determinants in *S. aureus* strains.

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