In 2010, a new methicillin-resistant *Staphylococcus aureus* clone, belonging to the clonal complex (CC) 913, Panton-Valentine leukocidin (PVL)-negative, staphylococcal cassette chromosome mec type IV, was isolated from Bedouin children in Israel (1). In 2012, isolates of CC913 were further analyzed, and their *spa* type was revealed as t991. Four t991 isolates were identified in hospitals across Israel, indicating the spread of the clone to communities beyond the Bedouin population in southern Israel (2). In 2015, a total of 12 t991 isolates were obtained from 280 patients (3), and in 2019, a total of 6 t991 isolates were obtained from 112 patients (4), mainly from children. Since 2015, MRSA isolates of *spa* type t911 have emerged to become one of the main lineages in hospitals and health maintenance organizations in Israel. However, despite its significance, comprehensive characterization of *spa* type t911 clone is lacking. We explore its genomic context and antibiotic profile in this study.

The Study
During 2012–2020, the *S. aureus* national reference laboratory of Israel received a total of 4,646 MRSA isolates, obtained from skin and soft tissue infections (SSTIs), that were classified into 284 different *spa* types. Types t002, t008, and t032 were the most prevalent; t002 comprised 25% of total MRSA SSTI isolates, t008 comprised 15%, and t032, 5%. During that period, the proportion of t991 MRSA gradually increased to 13% of total MRSA SSTI isolates, whereas the leading *spa* types in MRSA SSTIs (t002 and t008) remained stable (Appendix Figure 1, https://wwwnc.cdc.gov/EID/article/30/8/23-0981-App1.pdf). During that period, 689 *S. aureus* samples of *spa* type t991, *mecA*-positive, PVL-negative, were received at the *S. aureus* national reference laboratory (Appendix). Most of the samples (406, 70%) were isolated from SSTIs; 66% isolates were from patients <5 years of age (p = 0.0001), whereas 4% were isolated from patients >60 years of age. In addition, most patients resided in localities associated with Arab and Orthodox Jewish populations (5). The number of t991 MRSA isolates from SSTI and blood increased dramatically, from 5 in 2012 to 180 in 2019 and 146 in 2020 (https://microreact.org/project/r4dFwGXuqh3gyWtElf87-t991final) (Figure 1).

We conducted whole-genome sequencing on 20 t991 MRSA isolates that were selected (Appendix Table), along with 3 t991 MRSA isolates from Germany (6,7). The isolates clustered into 4 separate clades (Figure 2). Clade A consisted of the 3 t991 isolates from Germany and is 130 whole-genome multilocus sequence typing (wgMLST) alleles distant from the first isolate of clade B, which consisted of 7 isolates from patients who lived in the Negev and were admitted to the same hospital. Clade C consisted of 5 isolates from patients residing in the Jerusalem district. Clade D consisted of 8 isolates, 7 of which were from Orthodox Jewish patients.

We tested 116 t991 MRSA isolates for phenotypic susceptibility by using the broth microdilution method (Figure 3). Isolates from patients living in Arab localities were more resistant to erythromycin and chloramphenicol, whereas those isolated from patients living in Jewish localities showed higher resistance to gentamicin, ciprofloxacin, levofloxacin, and...
moxifloxacin (Appendix Figure 2). That tendency was statistically significant for chloramphenicol ($p = 0.01$) and gentamicin ($p = 0.01$).

We found 9 antimicrobial resistance (AMR) determinants, 8 AMR genes and 1 point mutation, among the 20 t991 MRSA sequences (Figure 4). Overall,
correlation between genotype prediction based on WGS and phenotypic AMR was 99% with a sensitivity of 94% and specificity of 100%. All discrepancies were associated with an absence of resistance determinant among phenotypically resistant isolates. No data for quinolone resistance genes or mutational resistance were predicted by the BioNumerics (https://www.bionumerics.com) or AMRFinder (https://github.com/ncbi/amr/releases/tag/amrfinder_v3.10.21) algorithms. In addition, we did not test phenotypic resistance against mupirocin.

We compared virulence profiles of 20 WGS t991 isolates with 3 t991 isolates from Germany, community-acquired MRSA USA300, and USA400 (8) as a reference using the functional genotyping tool of BioNumerics version 8.0. The main difference in virulence gene profile is reflected in the group of genes associated with adherence (Appendix Figure 3). In addition, virulence profiles can be grouped into 6 patterns on the basis of the presence of specific adherence factor genes in the genomes (Appendix Figure 4). We found no correlation between virulence profile and AMR, age, sex, residence location, or association with 1 of the genomic clades.

Next, to assess the genetic relationship of t991 isolates to other MRSA strains circulating worldwide, we created GrapeTree on the pubMLST site (https://pubmlst.org/organisms/staphylococcus-aureus) based on wgMLST data of representative local MRSA t991 strain (SA14675) along with 37,883 S. aureus global isolates (Appendix Figure 5) (9). The closest node to strain SA14675 is at a distance of 74 wgMLST alleles and is composed of 7 isolates. The next closest node is at a distance of 1,486 wgMLST alleles and composed of 7 isolates. The next closest node is at a distance of 1,486 wgMLST alleles and composed of isolates that belong mainly to CC1.

Conclusions
Most t991 cases were isolated from young patients who live in strictly Orthodox and Arab localities. A possible explanation for this phenomenon is a similar lifestyle of the 2 sectors, characterized by overcrowding and large families. Regarding the evolution of this clone and its spread into the population, this strain appears to have evolved by multiple different genetic events. This assumption is supported by several findings. First, t991 MRSA isolates demonstrated classification into 4 distinct clades on the basis of geographic location and sectoral association; we noted genetic variation and weak clonality evident from the considerable distances between nodes, even within the same clade. Second, antibiotic resistance patterns vary between isolates obtained from patients who live in Jewish and Arab localities (Appendix Figure 2). Finally, spa type t991 composition is very short; it consists of 3 repeats (07-33-23) and can be formed as a result of genetic rearrangement of numerous MRSA strains harboring longer spa type repeats in which the repeats of spa type t991 from a part of their repeat succession. Worldwide phylogenetic analysis indicates that t991 MRSA stands out as a distinct emerging lineage because it appears considerably distant from most strains included in the GrapeTree (Appendix Figure 5).

Phenotypic AMR data for global isolates were available for 2 isolates (7,10). One isolate obtained in Kuwait was resistant to erythromycin, clindamycin, trimethoprim, and fusidic acid (10), and the

![Figure 3. Percentage of resistant isolates to antibacterial agents among 116 t991 MRSA isolates from Israel tested for antimicrobial susceptibility using the broth microdilution method. MRSA, methicillin-resistant Staphylococcus aureus.](image-url)
ST913-IVa-t991 MRSA, Israel

other was isolated in Germany from a refugee from Syria (7) and was resistant to erythromycin, clindamycin, and tetracycline. Out of the 116 tested t991 isolates, none showed resistance to tetracycline by antimicrobial susceptibility. For 2 isolates, we observed phenotypic resistance for erythromycin and chloramphenicol without prediction of AMR determinant. Close inspection of those isolates revealed they were actually positive for \textit{ermC} and \textit{cat}, and their sequences were fragmented into multiple contigs. Consistent with previous publications (10–12), all isolates tested, except for the strain from the Syria refugee (7), were positive for \textit{eta}, a toxin responsible for skin infections seen mainly among young patients (3,4,7). Those findings are in accordance with the observation that t991 MRSA is predominantly isolated from children.

In conclusion, our study shows the emergence of t991 MRSA in Israel. These strains affect mainly pediatric populations, and a geographic distribution is limited mainly to the Middle East. The epidemiologic and genomic information our research provides will assist further investigation on the origin and dissemination of this clone.

About the Author

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References

Seeing a several-centimeters-long worm traversing the conjunctiva of an eye is often the moment when many people realize they are infected with *Loa loa*, commonly called the African eye-worm, a parasitic nematode that migrates throughout the subcutaneous and connective tissues of infected persons. Infection with this worm is called loiasis and is typically diagnosed either by the worm’s appearance in the eye or by a history of localized Calabar swellings, named for the coastal Nigerian town where that symptom was initially observed among infected persons. Endemic to a large region of the western and central African rainforests, the *Loa loa* microfilariae are passed to humans primarily from bites by flies from two species of the genus *Chrysops*, *C. silacea* and *C. dimidiate*. The more than 29 million people who live in affected areas of Central and West Africa are potentially at risk of loiasis.

Ben Taylor, cover artist for the August 2018 issue of EID, discusses how his personal experience with the *Loa loa* parasite influenced this painting.