

Multilocus Sequence Typing of *Streptococcus pyogenes* and the Relationships between *emm* Type and Clone

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Received 21 November 2000/Returned for modification 4 January 2001/Accepted 24 January 2001

Multilocus sequence typing (MLST) is a tool that can be used to study the molecular epidemiology and population genetic structure of microorganisms. A MLST scheme was developed for *Streptococcus pyogenes* and the nucleotide sequences of internal fragments of seven selected housekeeping loci were obtained for 212 isolates. A total of 100 unique combinations of housekeeping alleles (allelic profiles) were identified. The MLST scheme was highly concordant with several other typing methods. The *emm* type, corresponding to a locus that is subject to host immune selection, was determined for each isolate; of the >150 distinct *emm* types identified to date, 78 are represented in this report. For a given *emm* type, the majority of isolates shared five or more of the seven housekeeping alleles. Stable associations between *emm* type and MLST were documented by comparing isolates obtained decades apart and/or from different continents. For the 33 *emm* types for which more than one isolate was examined, only five *emm* types were present on widely divergent backgrounds, differing at four or more of the housekeeping loci. The findings indicate that the majority of *emm* types examined define clones or clonal complexes. In addition, an MLST database is made accessible to investigators who seek to characterize other isolates of this species via the internet (<http://www.mlst.net>).

Group A streptococci (GAS; *Streptococcus pyogenes*) are highly prevalent bacterial pathogens, having a worldwide distribution, whereby humans serve as their primary biological host. Most often, GAS infect superficial tissue sites, involving the mucosal epithelium of the upper respiratory tract (URT) or the epidermal layer of the skin, leading to pharyngitis or impetigo, respectively. On rare occasions, a GAS infection can lead to invasive disease that includes cellulitis, bacteremia, necrotizing fasciitis, and toxic shock syndrome, which can be life-threatening conditions. In addition, GAS contribute to morbidity through delayed nonsuppurative sequelae, such as rheumatic fever and acute glomerulonephritis.

The M and M-like proteins of GAS form surface fibrils that provide the basis for a widely used serological typing scheme. For many molecules studied in detail, the M serotype (M type) is usually defined by antigenic target sites contained within the distal, amino-terminal ends of these fibrillar proteins, and >80 distinct M types have been identified. M and M-like proteins are also key virulence factors, and protective immunity against GAS infection is type specific (8, 23). More recently, a genotypic typing scheme based on the *emm* genes that encode M and M-like proteins has become widely used and >150 different *emm* types have been characterized (15; <http://www.cdc.gov/ncidod/biotech/strep/strains.html>). The antigenic heterogeneity exhibited by this family of proteins reflects the strong im-

pact of host immunity on the generation of diversity within this bacterial species.

Numerous other genotypic methods have been developed for the typing of GAS isolates. Vir-typing measures restriction fragment length polymorphisms within the *emm* chromosomal region (18). Pulsed-field gel electrophoresis and arbitrary-primed PCR can provide high levels of resolution between strains by measuring multiple loci for differences that are not necessarily under selection (10, 17, 18, 33). Another important tool for discrimination among strains of GAS is multilocus enzyme electrophoresis (MLEE), which indexes differences in the net charge of housekeeping enzymes resulting from certain mutations (29, 32).

Multilocus sequence typing (MLST) is a nucleotide sequence-based method that is well suited towards characterizing the genetic relationships between the organisms of a bacterial species (12–14, 26). Because it is based on nucleotide sequence, it provides unambiguous results and is easily portable from lab to lab. Housekeeping loci are chosen for analysis because they are present in every organism (i.e., their products serve a vital function), and mutations within them are largely assumed to be selectively neutral (32). Clones, defined as isolates that are descendants of a recent common ancestor, can be identified as having shared alleles at each of the housekeeping loci. In this report, an MLST scheme using seven housekeeping loci was used to evaluate >200 GAS isolates that were derived from several continents, spanning a time period of >50 years and representing 78 distinct *emm* types.

MATERIALS AND METHODS

Bacterial strains. The GAS isolates of the MGAS series were kindly provided by Susan Hollingshead (University of Alabama at Birmingham), who had received them from James Musser, and the isolates have been previously described in detail (29, 30). The GAS isolates of the CT98 series were kindly provided by

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TABLE 1. Housekeeping loci under study^a

Locus	Putative function of gene product	Size of sequenced fragment (bp)	No. of alleles identified	No. of polymorphic nucleotide sites	No. of polymorphic amino acid sites	% Nucleotide divergence between pairs of alleles		Chromosomal location (no. of kb from <i>ori</i>) ^b	GenBank no.
						Maximum	Average		
<i>gki</i>	Glucose kinase	498	34	36	12	2.6	1.3	1259	AJ302470–AJ302503
<i>gtr</i>	Glutamine transport ATP-binding protein	450	32	30	9	2.6	1	1237	AJ302504–AJ302535
<i>murI</i>	Glutamate racemase	438	24	21	6	1.4	0.8	313	AJ406970–AJ406993
<i>mutS</i>	DNA mismatch repair (HexA)	405	21	23	9	2.2	0.9	1788	AJ406994–AJ407014
<i>recP</i>	Transketolase	459	35	59	11	6.1	1.9	1392	AJ407015–AJ407049
<i>xpt</i>	Xanthine phosphoribosyltransferase	450	29	31	10	2.9	0.8	931	AJ291638–AJ291666
<i>yqiL</i>	Acetoacetyl-CoA thiolase (AtoB)	434	22	22	12	1.4	0.7	129	AJ291616–AJ291637

^a Alleles of the seven housekeeping loci can be obtained (<http://www.mlst.net>).

^b The *emm* gene is located at kb 1685, relative to the origin, based on the genome of GAS strain 700294 (www.genome.ou.edu; 10 February 2001). The size of the strain 700294 genome is 1,852 kb.

James Hadler and Nancy Barrett (State of Connecticut Department of Public Health, Hartford). Strain 700294 was purchased from the American Tissue Culture Collection (Manassas, Va.). All other GAS isolates have been previously described (6, 17).

Multilocus sequence typing. Chromosomal DNA was prepared from freshly grown GAS by previously described methods (6). Internal fragments of the glucose kinase (*gki*), glutamine transporter protein (*gtr*), glutamate racemase (*murI*), DNA mismatch repair protein (*mutS*), transketolase (*recP*), xanthine phosphoribosyl transferase (*xpt*), and acetyl coenzyme A (acetyl-CoA) acetyltransferase (*yqiL*) genes were amplified by PCR using the following primer pairs: *gki*-up, 5'-GGC ATT GGA ATG GGA TCA CC-3', and *gki*-dn, 5'-TCT CCT GCT GCT GAC AC-3'; *gtr*-up, 5'-GAG GTT GTG GTG ATT ATT GG-3', and *gtr*-dn, 5'-GCA AAG CCC ATT TCA TGA GTC-3'; *murI*-up, 5'-TGC TGA CTC AAA ATG TTA AAA TGA TTG-3', and *murI*-dn, 5'-GAT GAT AAT TCA CCG TTA ATG TCA AAA TAG-3'; *mutS*-up, 5'-GAA GAG TCA TCT AGT TTA GAA TAC GAT-3', and *mutS*-dn, 5'-AGA GAG TTG TCA CTT GCG CGT TTG ATT GCT-3'; *recP*-up, 5'-GCA AAT TCT GGA CCA CCA GG-3', and *recP*-dn, 5'-CTT TCA CAA GGA TAT GTT GCC-3'; *xpt*-up, 5'-TTA CTT GAA GAA CGC ATC TTA-3', and *xpt*-dn, 5'-ATG AGG TCA CTT CAA TGC CC -3'; *yqiL*-up, 5'-TGC AAC AGT ATG GAC TGA CCA GAG AAC AAG ATG C-3', and *yqiL*-dn, 5'-CAA GGT CTC GTG AAA CCG CTA AAG CCT GAG-3'. The PCRs were performed in volumes of 50 μ l, with an initial denaturation at 95°C for 4 to 5 min, followed by 28 cycles of 95°C for 1 min, 55°C for 1 min, and 72°C for 1 min. The amplified DNA fragments were purified either by precipitation with polyethylene glycol or using a PCR purification kit (Qiagen, Valencia, Calif.). The sequence of each fragment was obtained on both strands by using the same primers as those in the initial PCR amplifications and an ABI377 or ABI3700 DNA sequencer (Perkin-Elmer Applied Biosystems, Foster City, Calif.).

For each locus, every different sequence was assigned a distinct allele number, and each isolate was defined by a series of seven integers (the allelic profile) corresponding to the alleles at the seven loci, in the order (alphabetical) of *gki*, *gtr*, *murI*, *mutS*, *recP*, *xpt*, and *yqiL*. Isolates with an identical allelic profile were assigned to the same sequence type (ST).

***emm* sequence typing.** *emm* sequence typing is based on the 5' end of the central *emm* gene within the *emm* chromosomal region (for map, see references 5 and 6). A unique *emm* type is defined as having <95% sequence identity to any other known *emm* type over 160 bp near the 5' end, as specified (<http://www.cdc.gov/ncidod/biotech/strep/strains.html>). There is a very strong correspondence between M type, as determined by serology, and the *emm* type that meets the stated definition (3, 15). In addition to a sequence identity of $\geq 95\%$, indels of four or fewer codons and/or frameshift mutations relative to the reference *emm* typing strain are allowed for classification as an established *emm* type. Until validation is complete, new *emm* types are assigned the nomenclature "emmst," which stands for *emm* sequence type (15) and is not to be confused with "ST," which refers to the MLST allelic profile.

Computations. A matrix of pair-wise differences in allelic profiles was constructed, and the similarities between the allelic profiles of the isolates were assessed by cluster analysis using the unweighted pair-group method with arithmetic averages (UPGMA) and the percent disagreement distance measure (Statistica version 5.5; StatSoft, Tulsa, Okla.). The maximum percent nucleotide divergence and average percent nucleotide divergence between pairs of alleles at a given locus were calculated using Mega version 2.0 (<http://www.megasoftware.net>). The Index of Association (27) was used to test for linkage disequilibrium

between alleles at the seven housekeeping loci. The observed variance in the distribution of allelic mismatches in all pair-wise comparisons of the allelic profiles was compared to that expected in a freely recombining population (linkage equilibrium). The significance of the difference in the observed and expected variance was evaluated by computing the maximum variance in the distribution of allelic mismatches obtained using 100 randomizations of the data set. Significant linkage disequilibrium was established if the observed variance obtained with the actual data was greater than that found with any of the 100 randomized data sets; otherwise, there was no evidence of a departure from linkage equilibrium.

RESULTS

Housekeeping loci used for MLST. Seven housekeeping loci were chosen for the characterization of GAS isolates by MLST and for determining their population genetic structure (Table 1). The nucleotide sequence was determined for an internal portion of about 400 to 500 bp of each gene. The loci that were chosen had been used successfully for pneumococci (14) or were selected with guidance by data from the University of Oklahoma GAS genome sequencing project that is available on the World Wide Web. Large contigs from the database (www.genome.ou.edu) were used in BLASTX searches against the GenBank database. Housekeeping loci were identified based on their putative function. Loci selected for this study were devoid of flanking regions containing genes that are likely to be under selection for variation (e.g., genes encoding cell surface proteins that may be under diversifying selection from the host immune response). The only possible exception was *recP*, positioned ~9 kb from a putative penicillin-binding protein gene (*pbp2x* homologue). However, analysis of a set of 14 isolates showed nucleotide sequence divergence of <1.0% for an internal portion of *pbp2x* and a lack of evidence for inter-species recombinational events, as has been observed for pneumococcal and meningococcal *pbp* genes (11) (data not shown). Furthermore, GAS isolates that are resistant to penicillin have not been described as occurring in nature. Ten housekeeping loci were initially examined in a small subset of strains and the least and most polymorphic ones were discarded. The chromosomal distance between any two loci, calculated on the basis of the tentative genome map of strain 700294, ranges from 20 to 600 kb (www.genome.ou.edu); it is possible that for other strains, the genomic location of the loci under study may differ.

The number of unique alleles identified for each of the seven housekeeping loci ranged from 21 (for *mutS*) to 35 (for *recP*) (Table 1). The maximum percent nucleotide sequence diver-

gence between the alleles of a given locus ranged from 1.4% (for *yqiL* and *murI*) to 6.1% (for *recP*). For one housekeeping locus, *recP*, there were four widely divergent alleles (*recP7*, *recP15*, *recP21*, *recP29*) which may have arisen by importation of homologous regions from closely related species. As noted above, the *recP* gene is ~9 kb from a *pbp2x* gene; however, *pbp2x* alleles display low levels of polymorphism, and there were no obvious differences between the *pbp2x* alleles of isolates recovered in the pre-antibiotic era (early 1940s) and those obtained in recent decades (data not shown). The sequence was determined for part of the *pbp2x* gene of an isolate containing one of the diverged *recP* alleles (*recP7*); this strain (C135) possessed the most prevalent *pbp2x* allele, and there is no evidence that the increased divergence of some *recP* alleles is due to hitchhiking driven by selection for interspecies recombination at the *pbp2x* locus. A more complete analysis of the housekeeping alleles is presented elsewhere (16; A. Kalia, M. C. Enright, B. G. Spratt, and D. E. Bessen, submitted for publication).

MLST of the GAS population. The collection of 212 GAS isolates (Table 2) was assembled with several goals in mind. First, a genetically diverse group of GAS strains was desired. As will be shown in this report, *emm* type is a sensitive measure of genetic diversity. Of the >150 *emm* types characterized to date (<http://www.cdc.gov/ncidod/biotech/strep/strains.html>), isolates representing 78 *emm* types were included in the MLST analysis. Secondly, it was of interest to evaluate GAS with large temporal and/or spatial distances between their isolation from human tissue, in order to assess the stability of clones. In addition, the selected GAS isolates were recovered in association with a variety of host tissues and diseases, including deep soft tissue infections. Finally, several GAS that had been previously analyzed using different molecular typing schemes were chosen for comparison to MLST, in order to provide validation of the new method.

The sequences of the seven loci were determined for each of the 212 GAS isolates, and their allelic profiles were assigned. One hundred different allelic profiles were found, corresponding to ST1 through ST100. Sixty-six of the 100 STs were represented by only a single isolate; the number of isolates assigned to the other STs ranged from 2 to 16.

The average number of alleles per locus was 28.1, and therefore, the GAS MLST scheme is able to distinguish >13 billion different allelic profiles. An isolate with the most common allele at each of the seven loci is expected to occur, by chance, at a frequency of 7.5×10^{-5} (no isolates with this allelic profile were found among the 212 strains); most allelic profiles will occur by chance at much lower frequencies. Thus, it is extremely unlikely that two unrelated GAS isolates will have the same allelic profile.

Relationships between *emm* type and MLST. A matrix of pair-wise differences in allelic profiles was determined, and a dendrogram displaying the genetic linkage distance between the 212 isolates was constructed by cluster analysis using UPGMA (Fig. 1). In the dendrogram presented in Fig. 1, the 15 STs that are represented by four or more isolates are depicted. In 13 of these STs, all isolates are of a singular *emm* type. It was of interest to further ascertain the strength of the associations between *emm* types and ST among GAS. Or, in other words, how well does *emm* type equate to clone?

For analysis of the relationships between *emm* type and MLST, selection criteria for GAS isolates were set to minimize the inclusion of epidemiologically related clones. Therefore, our analysis was specifically intended to provide a conservative estimate of the strength of the association between *emm* type and allelic profile. Multiple isolates of the same *emm* type and ST combination were included in the analysis only if they were recovered from subjects located on different continents or isolated >1 year apart within the same continent. Also, at least one representative of all unique *emm* type-ST combinations were included. *emm* types represented by four or more isolates satisfying the above-stated epidemiologic criteria ($n = 15$ *emm* types and 81 isolates in total) were assessed for the genetic distances between all possible pair-wise comparisons of alleles of the seven housekeeping loci (Table 3). This provides a measure of the genetic diversity at multiple loci within a set of epidemiologically unrelated organisms that share an *emm* type.

For six of the 15 *emm* types assessed (*emm2*, *emm5*, *emm6*, *emm12*, *emm18*, *emm33*), representing a total of 30 isolates, all isolates within an *emm* type displayed identical allelic profiles and can be regarded as clones (Table 3). Identical allelic profiles were observed for some organisms isolated >50 years apart (Table 2), indicating that GAS clones can be stable over this prolonged time period. One *emm* type (*emm19*) had isolates differing at one locus only, whereas two *emm* types had isolates differing at two loci (*emm3*, *emm89*). Isolates differing at two or fewer housekeeping loci (out of seven) can be regarded as clones or clonal complexes (16).

For epidemiologically distant organisms, as defined above, that were represented by only two or three isolates of the same *emm* type ($n = 18$ *emm* types), 11 *emm* types had identical allelic profiles, whereas five *emm* types differed at only one or two of the seven loci (Table 2). Although in some instances the sample size was small, *emm* type appears to closely correlate with clone or clonal complex for the majority (25 out of 33, or 76%) of *emm* types studied.

For several *emm* types represented by four or more epidemiologically distant isolates, there was a higher degree of genetic diversity. For three *emm* types—*emm4*, *emm11*, and *emm49*—pair-wise comparisons showed differences among three of the seven housekeeping loci (Table 3). An additional three *emm* types displayed differences at five or more of the housekeeping loci: *emm1*, *emm44/61*, and *emm77* (also known as *emm27L/77*). Perhaps it is of biological relevance that isolates of two of the *emm* types (*emm44/61* and *emm77*) were recently reported to be found in association with more than one *sof* allele, which provides the basis for a second major serological typing scheme for GAS (4). For *emm1* isolates, pair-wise comparisons indicated that this group is the most genetically diverse (Table 3). However, of the nine epidemiologically distant isolates evaluated, eight differed from one another at three or fewer of the seven loci (Table 2); furthermore, the *emm1* isolates cluster together, and there is a single node on the dendrogram from which all but one of the 23 *emm1* isolates descend (Fig. 1). One *emm1* isolate (MGAS2110; ST91) differs from the other *emm1* isolates at six or seven of the seven housekeeping loci. In addition to the *emm1*, *emm44/61*, and *emm77* isolates, the only other examples found for a single *emm* type on widely divergent genetic backgrounds are *emm91*

TABLE 2. MLST of 212 GAS isolates^a

Strain	emm type	code ^b	ST	Assigned no. for allele							Source	Year	Origin
				<i>gki</i>	<i>gtr</i>	<i>murI</i>	<i>mutS</i>	<i>recP</i>	<i>xpt</i>	<i>yqiL</i>			
86-779	1	m1-1	28	4	3	4	4	4	2	4	URT ^c	1986	Ohio
87-214	1	m1-2	28	4	3	4	4	4	2	4	URT	1987	Chile
CT95-111	1	m1-3	28	4	3	4	4	4	2	4	Invasive	1995	Connecticut
CT95-120	1	m1-4	28	4	3	4	4	4	2	4	Invasive	1995	Connecticut
CT95-131	1	m1-5	28	4	3	4	4	4	2	4	Invasive	1995	Connecticut
CT95-132	1	m1-6	28	4	3	4	4	4	2	4	Invasive	1995	Connecticut
CT95-180	1	m1-7	28	4	3	4	4	4	2	4	Invasive	1995	Connecticut
CT98360	1	m1-8	28	4	3	4	4	4	2	4	Invasive	1998	Connecticut
CT98419	1	m1-9	28	4	3	4	4	4	2	4	Invasive	1998	Connecticut
CT98520	1	m1-10	28	4	3	4	4	4	2	4	Invasive	1998	Connecticut
700294-ATCC	1	m1-11	28	4	3	4	4	4	2	4	Invasive	ND ^d	ND
MGAS166	1	m1-12	28	4	3	4	4	4	2	4	Invasive	ND	Minnesota
MGAS2127	1	m1-13	28	4	3	4	4	4	2	4	ND	ND	India
MGAS283	1	m1-14	28	4	3	4	4	4	2	4	Invasive	ND	Colorado
MGAS302	1	m1-15	28	4	3	4	4	4	2	4	Invasive	ND	Washington
MGAS307	1	m1-16	28	4	3	4	4	4	2	4	Invasive	ND	Texas
MGAS2110	1	m1-17	91	26	2	18	5	31	3	1	ND	ND	New Zealand
MGAS2120	1	m1-18	92	27	25	4	4	32	2	1	Impetigo	ND	Australia
MGAS2125	1	m1-19	92	27	25	4	4	32	2	1	Impetigo	ND	Australia
MGAS2109	1	m1-20	93	28	25	4	4	4	2	1	URT	ND	New Zealand
MGAS2144	1	m1-21	93	28	25	4	4	4	2	1	URT	ND	New Zealand
MGAS2226	1	m1-22	93	28	25	4	4	4	2	1	URT	ND	New Zealand
MGAS2123	1	m1-23	94	28	25	4	4	4	20	1	Impetigo	ND	Australia
B344	2	m2-1	55	11	9	1	9	2	3	4	URT	1950	Massachusetts
D725	2	m2-2	55	11	9	1	9	2	3	4	URT	1973	New York
89-465	2	m2-3	55	11	9	1	9	2	3	4	URT	1989	Nebraska
CT95-197	2	m2-4	55	11	9	1	9	2	3	4	Invasive	1995	Connecticut
CT98549	2	m2-5	55	11	9	1	9	2	3	4	Invasive	1998	Connecticut
MGAS286	2	m2-6	55	11	9	1	9	2	3	4	Invasive	ND	Colorado
MGAS327	2	m2-7	55	11	9	1	9	2	3	4	Invasive	ND	Arizona
C199	3	m3-1	15	2	6	8	5	2	3	2	URT	1942	New York
1GL90	3	m3-2	15	2	6	8	5	2	3	2	URT	1946	Illinois
2GL215	3	m3-3	15	2	6	8	5	2	3	2	URT	1946	Illinois
1RP228	3	m3-4	15	2	6	8	5	2	3	2	URT	1959	New York
D425	3	m3-5	15	2	6	8	5	2	3	2	Invasive	1971	New York
CT95-117	3	m3-6	15	2	6	8	5	2	3	2	Invasive	1995	Connecticut
CT95-119	3	m3-7	15	2	6	8	5	2	3	2	Invasive	1995	Connecticut
CT95-133	3	m3-8	15	2	6	8	5	2	3	2	Invasive	1995	Connecticut
CT95-145	3	m3-9	15	2	6	8	5	2	3	2	Invasive	1995	Connecticut
CT95-186	3	m3-10	15	2	6	8	5	2	3	2	Invasive	1995	Connecticut
CT95-204	3	m3-11	15	2	6	8	5	2	3	2	Invasive	1995	Connecticut
CT98429	3	m3-12	15	2	6	8	5	2	3	2	Invasive	1998	Connecticut
MGAS157	3	m3-13	15	2	6	8	5	2	3	2	Invasive	ND	Minnesota
MGAS159	3	m3-14	15	2	6	8	5	2	3	2	Invasive	ND	Utah
MGAS268	3	m3-15	15	2	6	8	5	2	3	2	Invasive	ND	Minnesota
MGAS277	3	m3-16	15	2	6	8	5	2	3	2	Invasive	ND	Colorado
88-019	3	m3-17	16	2	6	8	16	2	3	2	URT	1988	North Carolina
CT95-200	3	m3-18	18	2	27	8	5	2	3	2	Invasive	1995	Connecticut
2RP113	4	m4-1	38	5	7	8	5	15	2	1	Invasive	1952	New York
1RP156	4	m4-2	38	5	7	8	5	15	2	1	URT	1954	New York
A837	4	m4-3	38	5	7	8	5	15	2	1	Invasive	1965	Unknown
87-231	4	m4-4	39	5	11	8	5	15	2	1	URT	1987	Ohio
CT95-195	4	m4-5	39	5	11	8	5	15	2	1	Invasive	1995	Connecticut
YL05	4	m4-6	39	5	11	8	5	15	2	1	Invasive	1997	Connecticut
CT98693	4	m4-7	39	5	11	8	5	15	2	1	Invasive	1998	Connecticut
YL08	4	m4-8	39	5	11	8	5	15	2	1	Invasive	1999	Connecticut
09C02	4	m4-9	39	5	11	8	5	15	2	1	Invasive	ND	ND
MGAS320	4	m4-10	39	5	11	8	5	29	2	1	Invasive	ND	Texas
CT98529	4	m4-11	40	5	11	8	5	29	2	4	Invasive	1998	Connecticut
1RP144	5	m5-1	99	33	30	7	5	5	26	3	URT	1953	New York
87-292	5	m5-2	99	33	30	7	5	5	26	3	URT	1987	Ohio
CT95-118	5	m5-3	99	33	30	7	5	5	26	3	Invasive	1995	Connecticut
CT98302	5	m5-4	99	33	30	7	5	5	26	3	Invasive	1998	Connecticut
Manfredo	5	m5-5	99	33	30	7	5	5	26	3	URT	1958	Illinois
MGAS254	5	m5-6	99	33	30	7	5	5	26	3	Invasive	ND	California
MGAS258	5	m5-7	99	33	30	7	5	5	26	3	Invasive	ND	California
1RP112	6	m6-1	37	5	2	5	5	5	4	3	URT	1952	New York
D471	6	m6-2	37	5	2	5	5	5	4	3	ND	1971	Egypt
87-285	6	m6-3	37	5	2	5	5	5	4	3	URT	1987	Utah
87-376	6	m6-4	37	5	2	5	5	5	4	3	URT	1987	Colorado
CT95-115	6	m6-5	37	5	2	5	5	5	4	3	Invasive	1995	Connecticut
MGAS291	6	m6-6	37	5	2	5	5	5	4	3	Invasive	ND	Colorado

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TABLE 2—Continued

Strain	emm type	code ^b	ST	Assigned no. for allele							Source	Year	Origin
				<i>gki</i>	<i>gtr</i>	<i>murI</i>	<i>mutS</i>	<i>recP</i>	<i>xpt</i>	<i>yqiL</i>			
29740	8	m8	59	13	2	8	19	1	3	4	Impetigo	1988	Czech Republic
D733	9	m9-1	73	15	14	7	7	19	3	1	ND	1973	Unknown
1RP278	9	m9-2	74	15	14	7	16	19	3	1	URT	1964	New York
MGAS280	9	m9-3	75	15	14	7	18	19	3	1	Invasive	ND	Colorado
CT95-126	11	m11-1	20	3	4	6	5	1	5	1	Invasive	1995	Connecticut
CT95-155	11	m11-2	20	3	4	6	5	1	5	1	Invasive	1995	Connecticut
CT98303	11	m11-3	20	3	4	6	5	1	5	1	Invasive	1998	Connecticut
CT98414	11	m11-4	20	3	4	6	5	1	5	1	Invasive	1998	Connecticut
CT98538	11	m11-5	21	3	4	6	5	1	5	2	Invasive	1998	Connecticut
ALAB79	11	m11-6	22	3	4	6	7	1	5	4	Impetigo	1986	Alabama
D691	11	m11-7	23	3	4	6	7	11	5	4	URT	1972	Trinidad
2RP196	12	m12-1	36	5	2	2	6	6	2	2	URT	1956	New York
A374	12	m12-2	36	5	2	2	6	6	2	2	URT	1960	Trinidad
CT95-127	12	m12-3	36	5	2	2	6	6	2	2	Invasive	1995	Connecticut
CT98386	12	m12-4	36	5	2	2	6	6	2	2	Invasive	1998	Connecticut
CT98647	12	m12-5	36	5	2	2	6	6	2	2	Invasive	1998	Connecticut
MGAS278	12	m12-6	36	5	2	2	6	6	2	2	Invasive	ND	Colorado
CT95-122	13	m13-1	89	24	2	3	5	1	3	1	Invasive	1995	Connecticut
MGAS248	13	m13-2	89	24	2	3	5	1	3	1	Invasive	ND	California
25RS84	14	m14	85	20	22	2	3	27	18	14	URT	1942	New York
1GL217	17	m17	83	19	21	1	1	5	3	3	URT	1946	Illinois
88-092	18	m18-1	41	6	5	7	1	5	3	3	URT	1988	Pennsylvania
1RP62	18	m18-2	42	6	5	7	1	5	6	3	URT	1949	New York
4RP104	18	m18-3	42	6	5	7	1	5	6	3	URT	1951	New York
1RP268	18	m18-4	42	6	5	7	1	5	6	3	URT	1963	New York
A941	18	m18-5	42	6	5	7	1	5	6	3	URT	1967	New York
87-282	18	m18-6	42	6	5	7	1	5	6	3	URT	1987	Ohio
87-373	18	m18-7	42	6	5	7	1	5	6	3	URT	1987	Utah
MGAS300	18	m18-8	42	6	5	7	1	5	6	3	Invasive	ND	Washington
1RP118	19	m19-1	65	13	7	8	1	13	9	8	URT	1952	New York
1RP232	19	m19-2	65	13	7	8	1	13	9	8	URT	1959	New York
1GL205	19	m19-3	69	13	7	8	10	13	9	8	URT	1946	Illinois
D709	19	m19-4	69	13	7	8	10	13	9	8	URT	1973	New York
B243	22	m22-1	45	9	8	1	1	1	3	3	ND	1947	United Kingdom
MGAS275	22	m22-2	46	9	8	1	1	1	3	4	Invasive	ND	Colorado
MGAS330	22	m22-3	46	9	8	1	1	1	3	4	Invasive	ND	ND
MGAS339	22	m22-4	46	9	8	1	1	1	3	4	Invasive	ND	Texas
22RS72	24	m24-1	66	13	7	8	1	13	12	8	URT	1941	New York
1RP284	24	m24-2	70	13	18	8	1	13	9	8	URT	1964	New York
D316	25	m25	54	11	7	15	7	17	3	4	Invasive	1969	New York
11RS100	26	m26	67	13	7	8	1	26	3	13	URT	1942	New York
CT95-189	28	m28-1	52	11	6	14	5	9	17	19	Invasive	1995	Connecticut
CT98220	28	m28-2	52	11	6	14	5	9	17	19	Invasive	1998	Connecticut
CT98601	28	m28-3	52	11	6	14	5	9	17	19	Invasive	1998	Connecticut
MGAS255	28	m28-4	52	11	6	14	5	9	17	19	Invasive	ND	California
MGAS325	28	m28-5	52	11	6	14	5	9	17	19	Invasive	ND	ND
3RP70	29	m29-1	65	13	7	8	1	13	9	8	URT	1949	New York
D470	29	m29-2	65	13	7	8	1	13	9	8	ND	1971	Egypt
10RS101	32	m32	1	1	1	1	1	1	1	1	URT	1942	New York
13RS60	33	m33-1	3	2	2	1	2	2	2	2	URT	1941	New York
A982	33	m33-2	3	2	2	1	2	2	2	2	Impetigo	1968	New York
6010-5	33	m33-3	3	2	2	1	2	2	2	2	Impetigo	1971	Alabama
29487	33	m33-4	3	2	2	1	2	2	2	2	Impetigo	1987	Czech Republic
MGAS306	33	m33-5	3	2	2	1	2	2	2	2	Invasive	ND	ND
C142	34	m34	64	13	7	8	1	9	22	3	ND	1942	Unknown
A457	36	m36	47	9	16	1	20	25	3	16	ND	1961	Yugoslavia
19RS14	39	m39	32	4	17	2	5	1	25	1	Invasive	1941	New York
1RS79	42	m42	80	16	15	19	7	1	2	17	Invasive	1941	New York
D407	43	m43	4	2	2	1	2	2	3	2	Invasive	1971	New York
C135	49	m49-1	29	4	6	2	7	7	7	5	URT	1942	New York
A945	49	m49-2	29	4	6	2	7	7	7	5	Impetigo	1966	Minnesota
B737	49	m49-3	30	4	6	2	7	21	7	1	Impetigo	1957	Minnesota
MGAS163	49	m49-4	34	4	29	2	7	7	7	4	Invasive	ND	Louisiana
A203	50/62	m50	2	1	2	20	5	35	3	4	Mouse	1959	New York
MGAS323	50/62	m62-1	2	1	2	20	5	35	3	4	Invasive	ND	Idaho
MGAS324	50/62	m62-2	2	1	2	20	5	35	3	4	Invasive	ND	Idaho
A291	51	m51	84	20	22	2	3	27	18	12	ND	1960	Delaware
A946	52	m52	43	7	2	8	3	8	2	2	Impetigo	1971	Trinidad
D948	53	m53-1	11	2	6	1	2	2	2	2	Impetigo	1976	Trinidad
ALAB49	53	m53-2	11	2	6	1	2	2	2	2	Impetigo	1986	Alabama
AP53	53	m53-3	11	2	6	1	2	2	2	2	Impetigo	ND	Minnesota
D488	55	m55	100	34	2	2	21	1	29	16	Impetigo	1971	Trinidad
D306	57	m57	33	4	26	23	7	1	28	3	ND	1968	Trinidad

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TABLE 2—Continued

Strain	emm type	code ^b	ST	Assigned no. for allele							Source	Year	Origin
				<i>gki</i>	<i>gtr</i>	<i>murI</i>	<i>mutS</i>	<i>recP</i>	<i>xpt</i>	<i>yqiL</i>			
29454	58	m58	19	3	2	3	3	3	3	3	Impetigo	1987	Czech Republic
4500-S	60	m60	53	11	6	22	7	9	2	17	Impetigo	1960	Alabama
A956	63	m63	88	23	24	2	3	23	19	18	ND	1967	Unknown
ALAB53	66	m66-1	44	8	7	8	8	9	3	1	Impetigo	1986	Alabama
MGAS167	66	m66-2	44	8	7	8	8	9	3	1	Invasive	ND	Texas
D795	67	m76	61	13	2	16	1	5	3	1	ND	1975	Georgia
5552-S	68	m68	86	21	31	1	3	28	3	4	Impetigo	1971	Alabama
D998	70	m70	24	3	4	24	5	4	23	6	ND	1976	Japan
CT95-121	73	m73	7	2	2	9	3	18	3	4	Invasive	1995	Connecticut
86-809	75	m75-1	49	11	2	1	3	12	3	7	URT	1986	Utah
CT95-154	75	m75-2	49	11	2	1	3	12	3	7	Invasive	1995	Connecticut
CT95-171	75	m75-3	49	11	2	1	3	12	3	7	Invasive	1995	Connecticut
MGAS264	75	m75-4	49	11	2	1	3	12	3	7	Invasive	ND	California
CT98200	76	m76-1	50	11	6	3	6	6	27	4	Invasive	1998	Connecticut
CS110	76	m76-2	51	11	6	12	6	6	2	4	ND	ND	Unknown
MGAS261	77	m77-1	35	4	31	2	13	34	3	21	Invasive	ND	California
CT95-159	77	m77-2	63	13	6	2	3	23	3	11	Invasive	1995	Connecticut
CT98594	77	m77-3	63	13	6	2	3	23	3	11	Invasive	1998	Connecticut
MGAS331	77	m77-4	63	13	6	2	3	23	3	11	Invasive	ND	New York
D812	78	m78-1	76	16	2	8	3	1	3	1	Impetigo	1975	Trinidad
CT95-157	78	m78-2	79	16	2	11	3	1	11	1	Invasive	1995	Connecticut
CT95-104	80	m80	8	2	2	9	3	18	11	2	Invasive	1995	Connecticut
29665	81	m81	57	11	23	2	7	24	3	2	Impetigo	1988	Czech Republic
2RSC3	38/40	m38/40	81	17	19	1	1	1	15	3	URT	1942	New York
3RP141	44/61	m44/61-1	25	4	2	3	11	17	3	1	URT	1953	New York
D938	44/61	m44/61-2	25	4	2	3	11	17	3	1	Impetigo	1976	Trinidad
1RP18RN	44/61	m44/61-3	31	4	10	3	6	14	10	4	URT	1946	New York
5569-S	44/61	m44/61-4	31	4	10	3	6	14	10	4	Impetigo	1971	Alabama
MGAS296	82 (pt180)	m82	26	4	2	21	16	17	3	1	Invasive	ND	Washington
ALAB55	83 (pt2110)	m83-1	5	2	2	2	3	2	3	2	Impetigo	1986	Alabama
29689	83 (pt2110)	m83-2	5	2	2	2	3	2	3	2	Impetigo	1988	Czech Republic
D943	84 (pt2233)	m84-1	58	12	2	1	3	12	8	7	Impetigo	1976	Trinidad
MGAS1914A	84 (pt2233)	m84-2	58	12	2	1	3	12	8	7	ND	ND	ND
D964	86 (pt2631)	m86	9	2	2	9	3	18	14	2	Impetigo	1976	Trinidad
10RS57	87 (pt2841)	m87	62	13	3	10	12	20	2	3	URT	1941	New York
D976	88 (pt3875)	m88	90	25	2	13	6	30	25	15	ND	1976	Trinidad
MGAS2017	89 (pt4245)	m89-1	77	16	2	8	3	1	11	1	URT	1991	United States
MGAS2018	89 (pt4245)	m89-2	77	16	2	8	3	1	11	1	URT	1992	United States
CT95-176	89 (pt4245)	m89-3	77	16	2	8	3	1	11	1	Invasive	1995	Connecticut
CT98677	89 (pt4245)	m89-4	77	16	2	8	3	1	11	1	Invasive	1998	Connecticut
D424	89 (pt4245)	m89-5	78	16	2	8	7	1	13	1	Invasive	1971	New York
89-456	90 (pt4931)	m90	56	11	9	2	5	2	5	4	URT	1989	Kuwait
C506	91 (pt5757)	m91-1	12	2	6	1	5	2	2	2	URT	1943	Unknown
D821	91 (pt5757)	m91-2	13	2	6	2	5	22	3	2	Impetigo	1975	Trinidad
4426-S	92 (st2974)	m92-1	82	18	20	7	9	2	2	15	Impetigo	1970	Alabama
MGAS270	92 (st2974)	m92-2	82	18	20	7	9	2	2	15	Invasive	ND	Minnesota
D466	93 (potter41)	m93-1	10	2	2	9	13	2	14	2	ND	1971	Egypt
D502	93 (potter41)	m93-2	17	2	13	8	5	2	3	10	Impetigo	1971	Trinidad
2GL32	st1RP31	strp31-1	65	13	7	8	1	13	9	8	URT	1946	Illinois
1RP31	st1RP31	strp31-2	68	13	7	8	1	26	9	8	URT	1947	New York
CT95-169	st2346	st2346-1	60	13	2	14	1	9	3	1	Invasive	1995	Connecticut
MGAS319	st2346	st2346-2	60	13	2	14	1	9	3	1	Invasive	ND	Idaho
29486	st2370.1	st2370.1	6	2	2	8	3	2	16	12	Impetigo	1987	Czech Republic
MGAS341	st3365	st3365	97	31	2	2	5	35	24	12	Invasive	ND	Missouri
MGAS308	st4529	st4529	72	14	28	2	17	4	5	1	Invasive	ND	Washington, D.C.
6250-S	st4935	st4935	27	4	3	2	5	2	26	1	Impetigo	1972	Alabama
D997	st4973	st4973	98	32	6	8	3	2	8	22	ND	1976	Japan
MGAS273	st64/14	m64/14-1	14	2	6	8	3	9	3	1	Invasive	ND	California
MGAS2111	st64/14	m64/14-2	14	2	6	8	3	9	3	1	ND	ND	ND
MGAS2140	st833	st833	95	29	32	2	5	33	21	4	ND	ND	ND
A756	stA207	stA207-1	87	22	10	17	14	14	3	20	Invasive	1964	New York
MGAS288	stA207	stA207-2	87	22	10	17	14	14	3	20	Invasive	ND	Colorado
D432	stD432	stD432	71	14	12	2	7	16	5	9	ND	1971	Egypt
D626 ^e	stD626	stD626	9	2	2	9	3	18	14	2	Impetigo	1972	Trinidad
D631	stD631	stD631	96	30	2	9	13	18	14	2	Impetigo	1972	Trinidad
D633	stD633	stD633	48	10	6	2	6	10	3	6	Impetigo	1972	Trinidad
D641	stNS5	stNS5	11	2	6	1	2	2	2	2	Impetigo	1972	Trinidad

^a Note: *emm13* is equivalent to *emm13W*, *emm77* is equivalent to *emm27L*, and *emm50* is equivalent to *emm62* (<http://www.cdc.gov/ncidod/biotech/strep/strains.html>).

^b Codes for strains were used in dendrogram (Fig. 1).

^c URT, upper respiratory tract.

^d ND, not determined.

^e Strain D626 was previously assigned as *emmst88/31* (6) but is now correctly considered to be a new type (*emmstD626*; GenBank accession no. AF338251).

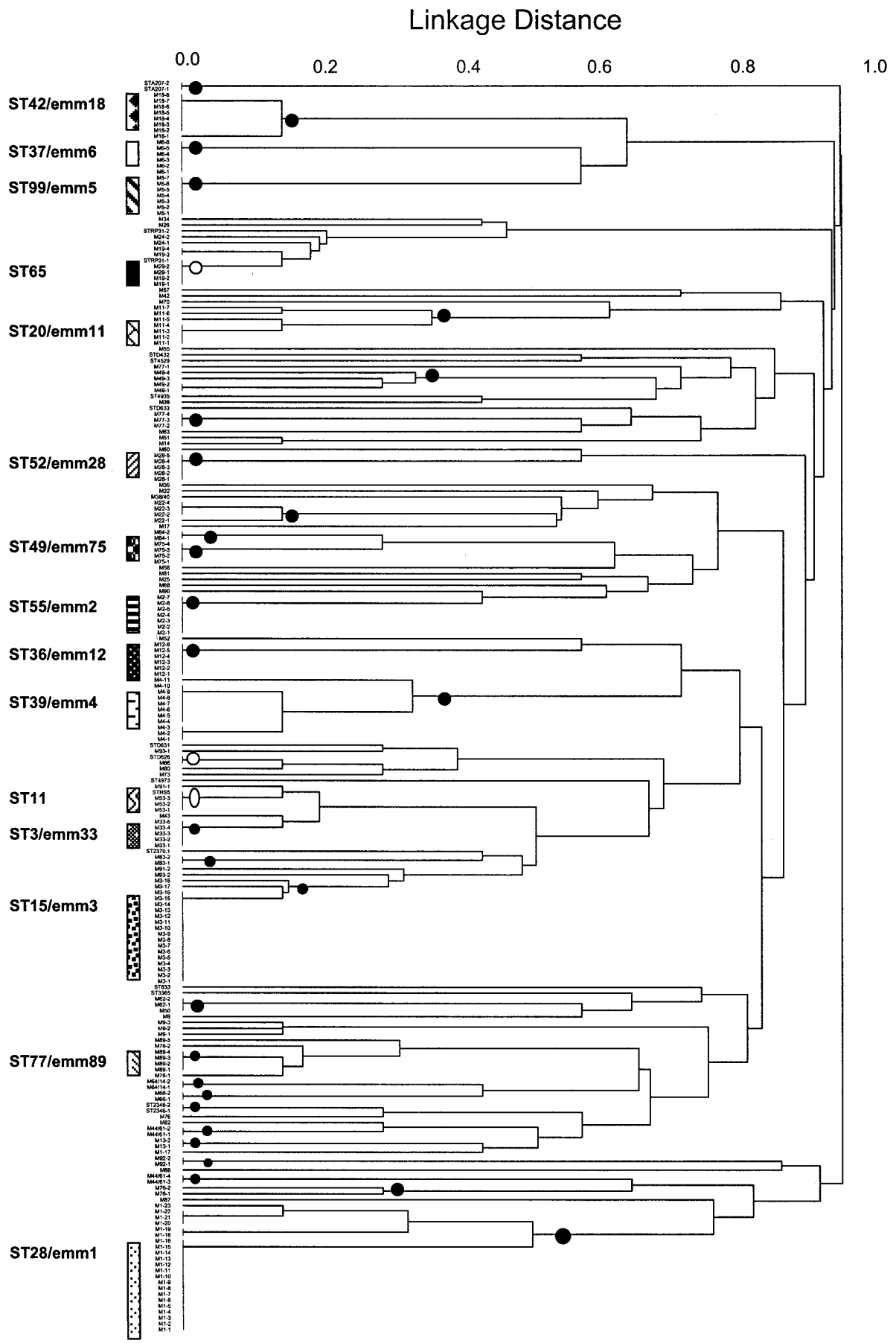


FIG. 1. Dendrogram showing UPGMA cluster analysis of 212 GAS isolates. Bars to the left show allelic profiles (STs) represented by four or more isolates. Codes for strain designations at branch tips are listed in Table 2. Filled circles ($n = 3$) mark branches in which multiple descendants are all represented by a single *emm* type. Open circles ($n = 3$) mark branches containing isolates with different *emm* types but sharing identical allelic profiles.

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TABLE 3. Pair-wise comparisons of housekeeping alleles among isolates of the same *emm* type

<i>emm</i> type ^a	No. of isolates	No. of pair-wise comparisons	% of pairs with identical loci:							
			7 of 7	6 of 7	5 of 7	4 of 7	3 of 7	2 of 7	1 of 7	0 of 7
1	9	36	28	3	3	17	28	0	8	4
2	5	10	100	0	0	0	0	0	0	0
3	9	36	58	39	3	0	0	0	0	0
4	8	28	32	43	14	11	0	0	0	0
5	4	6	100	0	0	0	0	0	0	0
6	4	6	100	0	0	0	0	0	0	0
11	5	10	10	30	30	30	0	0	0	0
12	5	10	100	0	0	0	0	0	0	0
18	7	21	100	0	0	0	0	0	0	0
19	4	6	33	67	0	0	0	0	0	0
33	5	10	100	0	0	0	0	0	0	0
44/61	4	6	33	0	0	0	0	67	0	0
49	4	6	17	0	67	17	0	0	0	0
77	4	6	50	0	0	0	0	50	0	0
89	4	6	50	0	50	0	0	0	0	0
Total	81	203								
Average			56.7	16.7	7.9	6.4	4.9	3.4	1.5	2.5
All STs										
Total	100	4,950								
Average			0	0.4	0.7	1.1	3.4	13	35	47

^a In order to avoid comparisons between epidemiologically related clones, for each *emm* type, all isolates were collected ≥ 1 year apart or from different continents, or isolates differed in allelic profile. All *emm* types listed in Table 2, of which four or more isolates meet the above-stated criteria, are included in the analysis.

and *emm93*, whereby two isolates of each type differ at three and five of the seven housekeeping loci, respectively (Table 2).

The genetic distances within an *emm* type can be compared to the genetic distance between the 100 different STs identified. By definition, none of the isolates representing each of the 100 unique STs shared alleles at all seven of the housekeeping loci. Whereas the majority of epidemiologically distant isolates within an *emm* type differed at two or fewer loci, 95% of the distinct allelic profiles (i.e., ST1 through ST100) differed from each other at five or more loci (Table 3). Furthermore, nearly half of the 4,950 possible pair-wise comparisons among the 100 STs differed at all seven housekeeping loci. Thus, comparisons between individual GAS clones most often reveal large genetic distances, contrasting sharply with the similar genotypes typically found within an *emm* type.

There were several examples of isolates with identical allelic profiles that differed in *emm* type: *emm86* and *emmstD626* (ST9), *emm53* and *emmstNS5* (ST11), and *emm19*, *emm29*, and *emmstRP31* (ST65) (Fig. 1). It is extremely unlikely that these examples of multiple *emm* types within a clone are due to a lack of discrimination of the GAS MLST scheme. For example, a single isolate with the allelic profile of ST65 was expected to occur by chance in the data set at a frequency of 2.2×10^{-8} , and the likelihood of unrelated *emm19*, *emm29*, and *emmstRP31* isolates having this allelic profile is essentially zero.

One *emm* type present on two or more genetically distant backgrounds, or multiple *emm* types present on a single genetic background, may have arisen as a consequence of the lateral movement of *emm* genes between different GAS strains. In GAS, generalized transduction by bacteriophage is the most probable mechanism for horizontal gene transfer.

Levels of linkage disequilibrium within the GAS population. The extent of recombination within the GAS population was

assessed by the Index of Association (27). Using one isolate of each of the 100 STs, there was significant linkage disequilibrium between the alleles at each of the seven housekeeping loci. However, in populations in which recombination is sufficient to randomize the alleles at different loci over a longer term, the recent expansion of clones can result in the appearance of multiple isolates with similar genotypes (27). Therefore, the Index of Association was recalculated using one isolate of each of the 72 STs obtained by truncating the dendrogram (Fig. 1) at a genetic distance of 0.3; no significant linkage disequilibrium between alleles was observed. The truncation effectively reduced each clonal complex to a single representative strain and thereby diminished any bias introduced by the oversampling of select *emm* types.

Comparison of MLST to other typing methods. The high degree of concordance between ST and *emm* type provides strong evidence that the MLST typing scheme leads to accurate identification of clones or clonal complexes. The MLST scheme can be further validated by comparison to other typing methods. Isolates that had been previously assessed by MLEE, as reported by others (22, 29, 30), were compared for *emm* type, ST, and electrophoretic type (ET) (Table 4). For organisms represented by one or more isolates of the same *emm* type-ST combination, 20 were also concordant for ET, whereas 9 were discordant with ET; however, for the discordant ETs, several were genetically close in their relationship. For organisms represented by one or more isolates of the same *emm* type-ET combination, 20 out of 21 were also concordant for ST.

Arbitrary-primed PCR, yielding random amplified polymorphic DNA (RAPD) profiles, has been previously conducted on another subset of the GAS isolates reported here (17). For organisms represented by one or more isolates of the same *emm* type-ST combination, nine also had concordant RAPD

TABLE 4. Comparison of MLST to other typing methods

Strain ^a	<i>emm</i> type	MLST-ST	ET based on MLEE	RAPD profile
MGAS166	1	28	1	
MGAS307	1	28	1	
MGAS302	1	28	15	
MGAS283	1	28	25	
MGAS2127	1	28	1D	
MGAS2110	1	91	1B	
MGAS2120	1	92	1C	
MGAS2125	1	92	1C	
MGAS2109	1	93	1F	
MGAS2144	1	93	1F	
MGAS2226	1	93	1F	
MGAS2123	1	94	1F	
MGAS327	2	55	4	
MGAS286	2	55	6	
MGAS157	3	15	2	
MGAS159	3	15	2	
MGAS268	3	15	2	
MGAS277	3	15	20	
MGAS254	5	99	28	
MGAS258	5	99	28	
MGAS300	18	42	20	
1RP232	19	65	45	
MGAS330	22	46	23	
MGAS339	22	46	23	
MGAS275	22	46	24	
1RP284	24	70	45	
D316	25	54	55	
MGAS255	28	52	10	
MGAS325	28	52	10	
D470	29	65	46	
MGAS323	50/62	2	9	
MGAS324	50/62	2	9	
MGAS264	75	49	18	
86-809	75	49	59	
MGAS261	77	35	4	
MGAS331	77	63	13	
MGAS2017	89	77	67	
MGAS2018	89	77	67	
CT95-120	1	28		1
CT95-111	1	28		1
CT95-131	1	28		1
CT95-132	1	28		1
CT95-180	1	28		1
CT95-133	3	15		2
CT95-145	3	15		3
CT95-204	3	15		4
CT95-117	3	15		5
CT95-119	3	15		5
CT95-186	3	15		8
CT95-200	3	18		5
CT95-155	11	20		19
CT95-126	11	20		19
CT95-154	75	49		21
CT95-171	75	49		22
CT95-159	77	63		15
CT95-176	89	77		15

^a Includes all isolates listed in Table 2 whereby there was more than one representative of a given *emm* type or ST and either an ET or RAPD profile based on previous reports (17, 22, 29, 30).

profiles, whereas seven displayed distinct RAPD profiles (Table 4). However, for organisms represented by one or more isolates of the same *emm* type-RAPD profile combination, 9 out of 10 were also concordant for ST.

Although the level of strain resolution differs for *emm* typ-

ing, MLEE, and RAPD analysis, each method displays high levels of concordance with the new MLST scheme.

GAS causing invasive disease. A total of 84 GAS isolates associated with invasive disease in the United States between 1986 and 1999 were included in this study. Thirty distinct *emm* types were represented by 34 unique allelic profiles (Fig. 2). Among the subset of invasive disease isolates, there was a high one-to-one correspondence between *emm* type and ST. However, for the vast majority of pair-wise comparisons between invasive disease isolates of different *emm* types, there were differences at four or more loci. Therefore, invasive disease caused by GAS can be attributed to a large number of genetically diverse strains or clones, confirming other reports (2, 17, 29, 35). However, two major clusters of isolates with identical or very similar allelic profiles were identified. These two clusters contained isolates of *emm1* and *emm3*, which are the *emm* types most commonly recovered from invasive disease in the United States during the 1990s (2, 17, 35).

DISCUSSION

A primary objective of this report is to provide the foundation for a new typing scheme for GAS that can be readily expanded upon by other investigators. In general terms, the value of molecular typing schemes lies in their ability to discriminate between the various strains within a bacterial species. However, high levels of discrimination are often achieved by indexing variation that accumulates very rapidly, making it difficult to demonstrate the relatedness of isolates that have diversified from a common ancestor that existed many decades ago. Variation within the nucleotide sequences of housekeeping genes accumulates relatively slowly, and as demonstrated in this report, isolates with the same allelic profile can be recovered many decades apart. Although the genetic variation indexed by MLST accumulates slowly, the multilocus approach allows for a vast number of distinct genotypes to be distinguished. Furthermore, MLST has high resolving power and, in many instances, it can discriminate among isolates of a single *emm* type.

The clustering of isolates achieved by MLST was in good agreement with those obtained using other typing procedures, and thus, the GAS MLST scheme provides a validated method for the unambiguous identification of GAS isolates. Since it is based on nucleotide sequence data, MLST allows different laboratories to compare their results via the internet. A website containing an initial database of the allelic profiles and molecular properties of the 212 GAS isolates and associated epidemiological data, together with interrogation and analysis software, is available (<http://www.mlst.net>).

The organisms initially selected for analysis by MLST represented a total of 78 *emm* types, and their isolation from human subjects dates back nearly 60 years. A future goal is to apply the MLST scheme to at least one isolate of every known *emm* type, collected from worldwide sources. A thorough documentation of existing GAS clones will lay the groundwork for gaining a better understanding of the epidemiological trends underlying GAS disease and aid in deciphering the molecular basis for biological diversity within this species.

emm type provides the basis for a serological typing scheme that differentiates between antigenic epitopes contained within

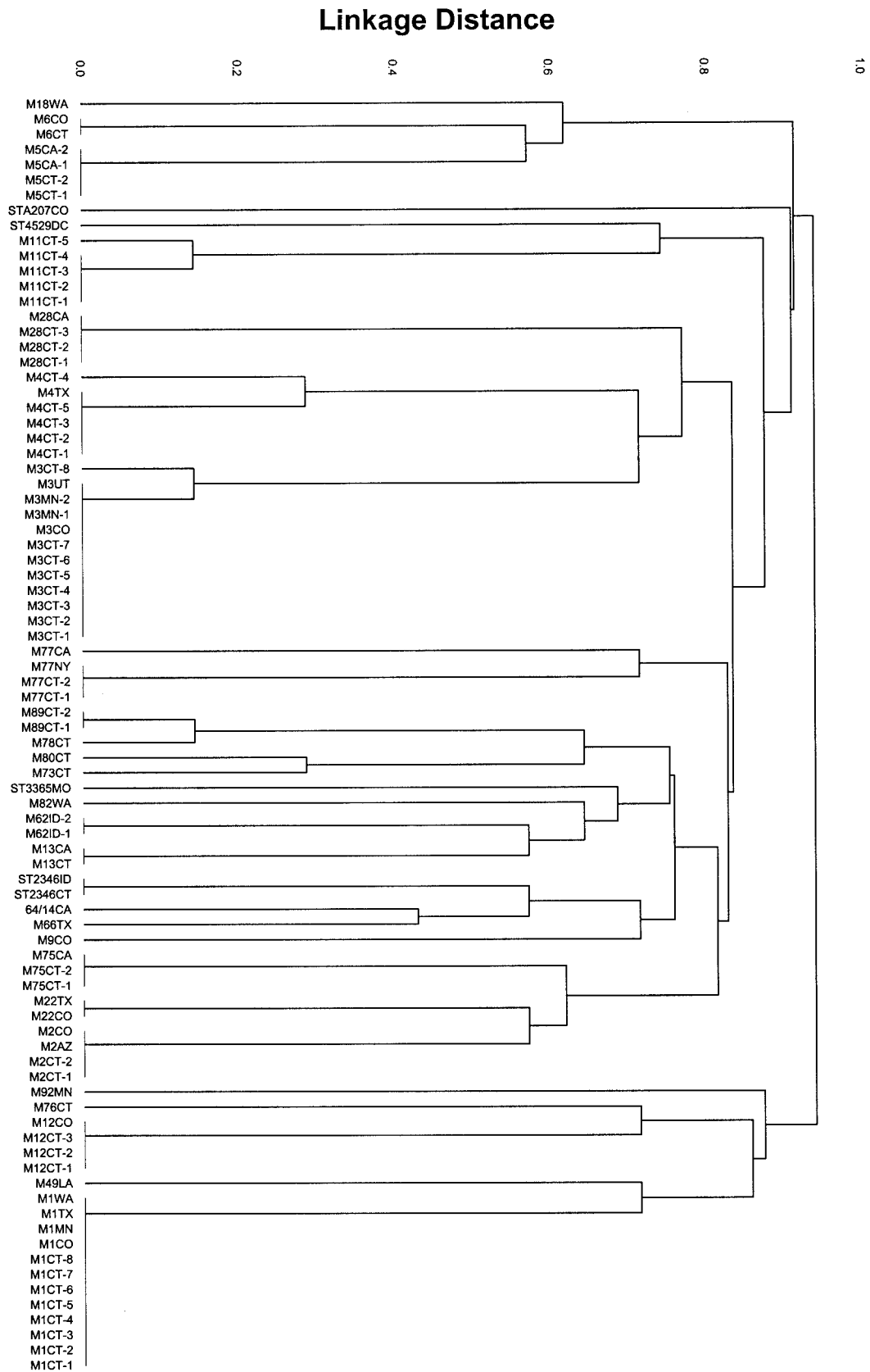


FIG. 2. Dendrogram of invasive isolates from the United States (1986 to 1999). UPGMA cluster analysis of all 84 isolates derived from normally sterile tissue sites, as listed in Table 2, is shown. The nomenclature at the branch tips indicate *emm* type, followed by the two-letter abbreviation of the state of origin and a unique isolate number (where necessary).

the amino-terminal, distal region of M-protein surface fibrils. Serum immunoglobulin G directed to M-type-specific epitopes leads to protective immunity for most strains that have been studied (1, 9, 25). Furthermore, the M proteins are key virulence factors, displaying a wide array of functional activities that act to promote disease (8). Unlike the housekeeping loci, *emm* genes are highly variable as a consequence of diversifying selection applied by the host immune response. It might therefore be expected that *emm* type would change more rapidly than alleles at housekeeping loci, resulting in variation within *emm* type among isolates of a clone or clonal complex. However, *emm* type is not defined by a unique nucleotide sequence but by $\geq 95\%$ sequence identity. Consequently, descendants of an ancestral strain may accumulate as many as eight nucleotide changes (and small indels or frameshifts) within the 160-bp sequenced region of the *emm* gene without altering the *emm* type, whereas even a single nucleotide change in the ~ 450 bp sequenced regions of any of the seven housekeeping loci results in a change in allelic profile. There are a few examples of isolates with identical allelic profiles having different *emm* types, such as ST65, which includes isolates of *emm19*, *emm29*, and *emmst1RP31*. Presumably, in these isolates, recombinational exchanges have resulted in the replacement of the region of the *emm* gene that defines *emm* type with the corresponding region from isolates of different *emm* types, since their divergence in *emm* type far exceeds 5%. Another multilocus typing method—MLEE—has also uncovered examples of isolates of the same genotype having different *emm* types (24, 30, 34).

A striking finding of this report is the degree to which multiple isolates of a given *emm* type share identical or highly similar allelic profiles (Table 3). Isolates of these *emm* types are considered to be clones or to form a clonal complex consisting of isolates with closely related allelic profiles. A much more extensive sampling of the GAS population will confirm the validity of this concept. The finding of a high one-to-one correspondence between *emm* type and clones or clonal complex suggests that GAS clones typically emerge and begin to diversify without changing their ancestral *emm* type. Recent studies using statistical tests of congruence between different housekeeping loci have indicated that recombination may be relatively common in GAS (16). This view was also supported by the lack of significant linkage disequilibrium between alleles that was observed when multiple isolates with similar genotypes were removed from the MLST data set, as measured by the Index of Association (27). Given this evidence for a major impact of recombination in the evolution of GAS populations, it is surprising that horizontal gene transfer appears to have rarely resulted in the presence of the same *emm* type in distantly related lineages. There are examples of this phenomenon, but they are uncommon. For example, among *emm1* isolates (the most intensively sampled *emm* type), 22 of the 23 isolates form a cluster of lineages that all descend from the same relatively deep node (genetic distance of 0.5), whereas the other *emm1* isolate differed from the former *emm1* isolates at six or seven of the seven loci (Fig. 1; Table 2) (30).

MLST studies of *Streptococcus pneumoniae* have also shown that isolates with identical or closely related allelic profiles almost invariably have the same serotype. However, in contrast to the findings on GAS, there are often multiple examples of distantly related clones or clonal complexes sharing the same

pneumococcal serotype. The paucity of distantly related GAS lineages sharing the same *emm* type may reflect differences in the strength of the immune response against pneumococcal capsular polysaccharides compared to that against M proteins, leading to differences in the strength of competitive exclusion between clones with the same capsular serotype or *emm* type. However, it might also be explained by the likelihood that changes in GAS serotype (i.e., *emm* type) occur by both mutation and recombination, whereas recombination involving the capsular biosynthetic genes is the only known mechanism underlying serotype changes in pneumococci (7). In the presence of strong selective immunological pressures, the diversification of *emm* genes might be further promoted by highly mutable processes, such as frameshift mutation and DNA slipped-strand mispairing (21, 28, 31). Unless recombinational exchanges that result in the presence of the same *emm* type in different lineages have occurred relatively recently, the diversifying selection applied by the host immune system is likely to result in the divergence of the *emm* types of the parental and recipient lineages. Thus, descendants of ancient horizontal genetic transfer events that distributed a particular pneumococcal capsular locus into multiple lineages may have retained the same serotype, whereas it is far less likely that the descendants of a similar ancient horizontal distribution of an *emm* gene will have retained the original *emm* type. The different extent to which the same capsular or M type is found in different lineages of pneumococci or GAS may rest more on the ease with which serotypes can change in these species, rather than differences in the rates of horizontal gene transfer.

The GAS MLST scheme provides a new and unambiguous method for characterizing GAS isolates for epidemiological purposes by using the internet. The MLST data can be used to address several epidemiological issues concerning GAS disease. Changes in epidemiological trends can be more readily ascribed to the emergence of new clones. Vaccine design strategies can be further refined, and vaccine efficacy can be measured with greater precision. The sequences of fragments of seven housekeeping genes from hundreds of GAS isolates provide data that can be used to address aspects of the population and evolutionary biology of the species. For example, the ancestral relationships and patterns of descent among closely related isolates can be deduced, although relationships between more distantly related isolates are likely to be obscured by a history of recombination (16). The population genetic structure of GAS, based on neutral housekeeping loci, will provide a framework upon which to measure the distribution of adaptive loci. This, in turn, should provide new insights into the molecular basis for biological diversity among GAS, as well as the role of cell surface antigens in structuring the population (19, 20).

ACKNOWLEDGMENTS

We thank Yury Nunez, Eric Peterson, and Michelle Benitez for expert technical assistance, Susan Hollingshead (UAB) for supplying the MGAS strains, and Jim Hadler and Nancy Barrett (CT DOH) for providing the invasive isolates collected in Connecticut during 1998 (CT98 series) and the *emm*-typing data. We also acknowledge the Streptococcal Genome Sequencing Project funded by USPHS/NIH grant AI-38406 and the work performed by B. A. Roe, S. P. Linn, L. Song, X. Yuan, S. Clifton, R. E. McLaughlin, M. McShan, and J. Ferretti.

This work was supported by grants from the Wellcome Trust (to B.G.S.), the National Institutes of Health (AI-28944 to D.E.B. and GM-60793 to D.E.B. and B.G.S.), the American Heart Association (grant-in-aid to D.E.B.), and a Brown-Coxe Postdoctoral Fellowship (to A.K.). M.C.E. is a Royal Society University Research Fellow. D.E.B. is an Established Investigator of the American Heart Association.

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