Abstract

Diseases caused by *Streptococcus pyogenes* (Group A streptococcus, GAS) range from superficial infections such as pharyngitis and impetigo to potentially fatal rheumatic heart disease and invasive disease. Studies spanning emm-typing surveillance to population genomics are providing new insights into the epidemiology, pathogenesis, and biology of this organism. Such studies have demonstrated the differences that exist in the epidemiology of streptococcal disease between developing and developed nations. In developing nations, where streptococcal disease is endemic, the diversity of GAS emm-types circulating is much greater than that found in developed nations. An association between emm-type and disease, as observed in developed countries is also lacking. Intriguingly, comparative genetic studies suggest that emm-type is not always a good predictor of the evolutionary relatedness of geographically distant isolates. A view of GAS as a highly dynamic organism, in possession of a core set of virulence genes that contribute to host niche specialization and common pathogenic processes, augmented...
by accessory genes that change the relative virulence of specific lineages is emerging. Our inability to definitively identify genetic factors that contribute to specific disease outcome underscores the complex nature of streptococcal diseases.

Contents

1 Introduction .................................................................................................................. 30
2 emm-Gene-Based Molecular Epidemiology .............................................................. 31
3 Non-emm-Gene Virulence Factors and GAS Epidemiology ..................................... 34
4 Multilocus Sequence Typing ...................................................................................... 38
5 Genomic Studies of Group A Streptococcus ............................................................... 39
6 The Role of Other β-Hemolytic Bacteria in GAS-Associated Diseases .................. 40
7 Conclusion ................................................................................................................... 41
References ...................................................................................................................... 42

1 Introduction

Diseases associated with infection by Streptococcus pyogenes (group A streptococcus, GAS) have been so prevalent in history that many have been given easily recognizable common names (e.g., strep throat, scarlet fever, school sores, childbed fever). While the incidence of most diseases has declined markedly in countries with high per capita income, countries and regions of low income continue to suffer a high burden of these and other GAS associated diseases. Infection by GAS has been estimated to result in half a million deaths each year (Carapetis et al. 2005), and thereby position GAS as one of the top ten bacterial killers of humans. The majority of these deaths follow the development of rheumatic heart disease (RHD) and occur in developing nations. In more affluent countries, with better access to healthcare and antibiotic treatment, the prevalence of RHD is much lower, the majority of deaths attributed to the clinical manifestations associated with streptococcal invasive disease.

The burden of GAS-related diseases, differences in the geographic occurrences of these diseases, environment and the preferred body sites colonized by the organism, and accompanying changes in disease outcomes have all impelled continuing molecular and epidemiological studies of this organism. The aims of these studies have been to develop a greater understanding of molecular differences that underpin relative differences in virulence of discrete GAS lineages, identify plausible microbiological based rationale for differences in the epidemiology of GAS in different geographic locations, and predict the effectiveness of putative vaccine candidates (Smeesters et al. 2009; Steer et al. 2009d). Almost all GAS epidemiological studies use the M-protein or its genetic counterpart, the emm-gene as the basis for discriminating between GAS lineages. Here, we
describe how these studies, the analysis of other genetic markers and genomics has changed our view of the epidemiology and biology of this organism.

2 *emm*-Gene-Based Molecular Epidemiology

Rebecca Lancefield first reported serotypic diversity in GAS more than 80 years ago. Unlike other bacterial species in which diversity was based on variation in capsular antigens, the serotypic variation that Lancefield observed was based on variation in the N-terminal region of the surface exposed M-protein (Cunningham 2000; Fischetti 1989). Subsequent work in the 1950s showed that the presence of type specific antibodies in animal and human serum was responsible for immunity against the homologous emm-type but did not protect against heterologous emm-types (Lancefield 1962). Thus, M-protein-based serotype diversity was given a functional credence, and became the basis of the GAS typing scheme (Lancefield 1962).

Serotype-based M-typing has given way to nucleotide-based procedures (emm-typing) that targets the nucleotide sequence corresponding to the hyper-variable amino terminal region of the M-protein. More than 200 different emm-sequence types have now been reported (Beall et al. 1996; Facklam et al. 2002). Although undergoing several minor revision since it’s introduction, the basic premise of emm-typing remains the same (http://www.cdc.gov/ncidod/biotech/strep/strepblast.htm). In both its serological and nucleotide-based incarnations, emm-typing has been extensively used to examine both geographic strain distribution and disease association. With the large number of emm-type-based surveillance studies carried out in developing countries over the past two decades it has become clear that the epidemiology of GAS differs between developing and developed regions. Excellent systematic reviews summarizing the global molecular epidemiology of GAS have been published separately by Smeesters and Steer in 2009 (Smeesters et al. 2009; Steer et al. 2009d). In low-income regions where streptococcal infections and disease are endemic, the diversity of circulating emm-types is high. This has been demonstrated globally in distinct area such as India, Fiji, Ethiopia, and Brazil (Abdissa et al. 2006; Dey et al. 2005; Smeesters et al. 2006, 2008, 2010a; Steer et al. 2009e; Tewodros and Kronvall 2005). These studies also demonstrate that no one emm-type is dominant in these regions. Steers review reported the greatest diversity of emm-types to be found in Pacific Regions and Africa. Figure 1, depicting the emm-type diversity as determined using Simpson Reciprocal Index, clearly demonstrates the differences in diversity between countries. The pool of emm-types recovered in separate studies in low-income regions also differs (Smeesters et al. 2009; Steer et al. 2009d). For example, only one third of the emm-types recovered in two Ethiopian studies (Abdissa et al. 2006; Tewodros and Kronvall 2005) were described in a separate Fijian study (Steer et al. 2009e). Taken together the data appear to fit a model where the number of circulating emm-types is high, and subject to temporal flux. Few longitudinal studies, examining the rate of emm-type replacement have been
carried out in endemic regions (Steer et al. 2009b). There is also lack of surveillance and prospective surveys in some areas of the world where the burden of streptococcal disease is high (Carapetis et al. 2005). This is particularly true for Africa, South America, and some locations in Asia (Smeesters et al. 2009; Steer et al. 2009d).

In developed countries, fewer emm-types appear to circulate, with even fewer dominant. Steer et al. (Steer et al. 2009d) reported that 25 emm-types accounted for more than 90% of all isolates recovered in developed nations, with 146 different emm-types accounting for the remaining 10%. Recovery of these predominant emm-types in multiple studies also suggests that they are consistently present in the wider population (McNeil et al. 2005; Shulman et al. 2009; Smeesters et al. 2006). Longitudinal studies carried out in smaller geographic regions show that different emm-types are recovered at different time points (Shulman et al. 2009), demonstrating that local temporal flux in circulating emm-types occurs.

Molecular techniques, such as vir-typing (Gardiner et al. 1995) and emm pattern-typing (Hollingshead et al. 1993) represent alternate methods to categorize GAS based on variation in the emm-gene and surrounding DNA. Emm pattern typing categorizes emm-types into three distinct patterns based on chromosomal architecture (pattern types A-C, D, and E). The patterns are determined by the presence and arrangement of emm and emm-like genes, as determined by specific PCR reactions (Hollingshead et al. 1993). Emm-type correlates well with specific emm pattern type (McGregor et al. 2004b). In general emm pattern-type correlates with host colonization site. Pattern A-C strains are usually associated with throat colonization, pattern D strains mainly recovered from superficial skin infection, while pattern E represents a “generalist” group associated with both tissue sites (Bessen and Lizano 2010). Similar to emm-typing, exceptions to the emm pattern type/tissue tropism have been reported, especially in low-income populations.
(Bessen and Lizano 2010). Only about 20 % of the 223 known emm-types belong to pattern A-C type. These emm-types predominantly circulate in high-income countries. The remaining 80 % of emm-types are equally distributed between the pattern D and E groups, and are mostly found in developing countries. Of note, emm-types belonging to emm pattern A-C, which contain the so-called ‘rheumatogenic’ and ‘invasive’ M-types have been most extensively studied in relation to disease outcome (Erdem et al. 2005; Fischetti 1989; Smeesters et al. 2010b).

Certain emm-types, belonging to pattern A-C, are predominantly recovered from the throat rather than skin in developing countries (Shulman et al. 2004). As rheumatic fever (RF) is considered to follow an episode of pharyngitis, these types have also come to be known as ‘rheumatogenic’ emm-types (Bisno et al. 2003; Martin and Barbadora 2006; Stolleraman 2001). Other emm-types are more commonly associated with skin infection and glomerulonephritis, and have come to be known as ‘nephritogenic’ emm-types. A third group of emm-types (e.g., emm1, emm12, emm3, and emm18) associated with the outbreak of severe invasive disease in the USA and Europe over the past two decades are known as ‘invasive’ emm-types (O’Loughlin et al. 2007; Vlaminckx et al. 2007). Interestingly recent epidemiological studies carried out in geographic regions where RHD and streptococcal infections are endemic have failed to recover significant numbers of these rheumatogenic emm-types (Bessen et al. 2000; Steer et al. 2009c). In fact, epidemiological studies in tropical regions, where the incidence of RF, but the incidence of pharyngitis is low is challenging the dogma that there is an association between GAS, throat infection, and RF (Bessen et al. 2000; Parks et al. 2012). Unique emm-types have also been implicated in RF in Hawaii (Erdem et al. 2007). Similarly, ‘invasive’ emm-types while present are not common in low income countries (Steer et al. 2009a).

Emm-typing has served the scientific community well for several decades, and is still relevant to tracking disease outbreaks. However, the differences in emm-type diversity and disease associations have created challenges for the streptococcal community when attempting to extrapolate data and interpretation of results from developed nations to developing nations. It should be acknowledged that comparisons of emm-type prevalence in different locations can be complicated by the different methods used to acquire isolates. Some studies represent single timepoint surveillance, whereas others analyses isolates collected over time (Shulman et al. 2009). Isolates recovered during epidemic outbreaks are also likely to differ from those recovered during non-disease related surveillance in the same location. Nevertheless, the data from different studies conducted in endemic and non-endemic regions collectively suggest differences in diversity observed between these groups are real.

Emm-type is also assumed to be marker for evolutionary relatedness, and by extension, similarity in pathogenic potential of isolates. While this is undoubtedly true for isoaltes collected during clonal outbreaks, it is less so for geographically or temporally unrelated strains. As described in more detail below, evidence for lateral gene transfer (LGT) and recombination of DNA, including the emm-gene is strong. The section of the emm-gene used for emm-typing encodes the region of...
the corresponding protein that may be targeted by the host immune system, and is therefore likely to be under strong immune selection pressure. Replacement of this region, through recombination involving part or all of the emm-gene is an efficient method for escaping the host immune response (Panchaud et al. 2009; Whatmore and Kehoe 1994). In countries where streptococcal disease is endemic, and emm-type diversity is high, the conditions for LGT of the emm-gene is much greater than non-endemic countries.

3 Non-emm-Gene Virulence Factors and GAS Epidemiology

GAS pathogenesis depends on the coordinated regulation of a complex repertoire of virulence factors. While host factors undoubtedly contribute to variation in susceptibility to GAS, the frequent association of specific emm-types with disease in developed countries lends itself strongly to the argument that specific bacterial characteristics are associated with defined epidemiologies. The most comprehensive body of work examining association between non-emm genetic factors and GAS disease has occurred in the area of serious invasive disease. However, despite many years of research, the identification of a defined subset of invasive disease related virulence factors remains elusive.

There is increasing evidence that in populations where GAS is endemic, despite the diversity of emm-types, there is conservation of certain virulence factors linked to tissue tropism. While emm pattern typing has been useful for classification of GAS into these three patterns, several elegant epidemiological studies have led to the identification of a specific subset of genes which may confer GAS with a tissue specific phenotype (Fig. 2) (Bessen et al. 2005; Kalia and Bessen 2004; Kratovac et al. 2007). Linkages to tissue-specificity are particularly strong for genes encoding the colonization factors serum opacity factor (sof), collagen-binding protein (cpa), and fibronectin binding proteins (prtF1) and (fbaA), which have a defined pattern of presence or absence in strains associated with distinct tissue sites (Kratovac et al. 2007). There is also significant historical evidence to implicate these colonization factors in tissue tropism, and most recently, comparative genomic hybridization analysis of 96 GAS isolates revealed that a major defining factor of tissue tropism in skin versus throat isolates is the presence or absence of genes encoding fibronectin binding proteins (Bessen et al. 2011).

Historically, SOF has provided the basis for an alternative GAS typing scheme. Streptococci of differing emm-types produce distinct variants of SOF, and type specific antibodies can be used to inhibit serum opacification by SOF variants (Maxted et al. 1973). There is a strong correlation between emm-type and SOF phenotype, and SOF antisera has been used as a method of GAS typing in the past. A comprehensive study of GAS isolates in 2000 showed that SOF is a useful predictor of emm-type, and that discordance between sof and emm is rare. However, GAS strains of the same emm-type do not always contain the same sof allele, and strains with diverse emm-types can have highly similar SOF genes (Beall et al.
No correlation has been found between various sof alleles and different disease states, or geographical locations. A comparison of published studies from around the world shows that most reported sof/emm combinations are distributed globally (Dhanda et al. 2011; Goodfellow et al. 2000; Johnson et al. 2006; Sakota et al. 2006). This lack of geographical segregation may, in part, be explained by the linkage between sof and the “generalist” tissue phenotype, that is, isolates which are equally associated with skin and pharyngeal infection, which represent approximately 50% of GAS isolates globally (Bessen and Lizano 2010). Typically, pharyngeal specialists are opacity factor negative, and harbor a prtF1 allele (Courtney and Pownall 2010; Cunningham 2000; Kratovac et al. 2007). In contrast, skin specialists are more likely to be endowed with fbaA. In a study of GAS isolates from the Northern territory of Australia, where skin infection predominates, 92% of isolates were found to be fbaA positive (Ramachandran et al. 2004). Similarly, cpa is more frequently associated with skin trophic isolates than strains which would be regarded as throat specialists (Kratovac et al. 2007). The differential distribution of these genes between skin versus throat versus generalist strains largely parallels the architecture of the overall mga regulon.

Additionally, a number of genes appear to have evolved into discrete phylogenetic lineages associated with tissue site preference. The regulatory gene mga is present in the GAS genome in one of two allelic forms, mga1, which is predominately associated with throat specialist strains, and mga2, which appears to be found in all emm pattern D and E strains. These allelic forms are mutually exclusive (Bessen et al. 2005). Similarly, rofA/nra encodes as transcriptional regulator which regulates expression of pillus structural genes (Kreikemeyer et al. 2003). rofA is associated with emm pattern A-C and E strains (throat specialists and generalists), skin specialists typically harbor nra (Bessen et al. 2005).

The plasminogen activator, streptokinase, has been found in all GAS strains screened to date. Streptokinase is comprised of three distinct domains, α, β, and γ. There is significant variability within the β domain of ska, and 3 distinct ska alleles.
have been described—type 1, type 2a, and 2b (Kalia and Bessen 2004). There is 
strong genetic linkage between the gene encoding the plasminogen binding 
M protein (PAM) and type 2b ska alleles, which appear to be almost exclusively 
coinherited (Kalia and Bessen 2004; McArthur et al. 2008). Isolates harboring this 
allele are typically regarded as “skin trophic”. In contrast, cluster 2a alleles are 
almost exclusively associated with isolates that would be regarded as throat-
trophic based on emm pattern A-C, while type 1 ska alleles are found in both emm 
pattern A-C and E backgrounds (Kalia and Bessen 2004).

These preferences for tissue site, and distribution of virulence factors, are not 
absolute. In geographical regions where GAS infection is endemic, the demarca-
tion line between emm-type and tissue site preference is less well defined. Pop-
ulation surveillance of GAS isolates in Nepal found that 19 % of isolates 
associated with skin infection display a genetic architecture usually associated 
with pharyngeal specialists (Sakota et al. 2006), while in an Ethiopian study, 28 % 
of isolates associated with tonsillitis harbored an emm pattern D, or skin specialist 
genomic arrangement (Tewodros and Kronvall 2005). Similarly, GAS with an 
emm pattern D, or skin trophic chromosomal arrangement harboring ska type 1 
alleles have been reported (Kalia and Bessen 2004). This underscores the high 
degree of genetic diversity among GAS isolates. In areas where GAS is endemic, 
recombination between skin and throat specialists appears to be common (Bessen 
et al. 2011; Kalia et al. 2002). Given these findings, it will remain important to 
monitor the relationships between virulence factor distribution and tissue tropism 
in epidemiological studies.

Significant intra-emm-type genetic variation, particularly with respect to reg-
ulatory gene profile, phage content, and toxin profile have been observed in the 
dominant emm-types associated with invasive disease in developed countries. 
Perhaps the most striking example of this is the reported difference in transcription 
profile between invasive and colonizing GAS strains of the same emm-type in 
outbreaks of GAS infection (Johnson et al. 1992; Marcon et al. 1988; Sumby et al. 
2006). Expression microarray analysis of M1 isolates from the United States, 
Canada, and Finland revealed the existence of distinct non-invasive and invasive 
transcriptome profiles within a group of clinical M1 strains (Sumby et al. 2006). 
These transcriptome profiles have since been linked to mutations in the control of 
virulence regulatory sensor kinase (covRS; alternatively designated csrRS), which 
is responsible for the regulation of approximately 10 % of the GAS genome 
(Graham et al. 2002; Sumby et al. 2006). Acquisition of mutations in covRS which 
inactivate the ability of this system to negatively regulate the GAS genome during 
infection results in upregulation of capsule, loss of SpeB expression and increased 
disease severity in animal models of infection (Walker et al. 2007). Acquisition of 
these mutations appears to be dependent on the expression of specific virulence 
factor genes, including emm, hasA, and the phage encoded sda1 (Cole et al. 2011). 
The presence of mutations in covRS associated with clinical invasive GAS isolates 
have been extensively reported in studies of isolates from developed nations, and 
these mutations are not restricted to emm1 GAS. A recent study of isolates from 
Canada that compared the genome sequences of GAS serotype M3 isolates from
human pharyngitis cases and from human invasive disease reported that covS mutations occur with a higher frequency in invasive-disease isolates than in pharyngeal isolates (Shea et al. 2011). Several studies report covS mutations in GAS isolates recovered from patients with severe STSS (Ato et al. 2008; Ikebe et al. 2010). Similarly, a Hawaiian emm81.0 GAS isolate from the throat that had spread to the bloodstream was shown to have acquired a mutation in covS (Garcia et al. 2010).

While these mutations can be linked to “hypervirulence”, the association is not absolute. In a retrospective study of GAS clinical isolates from Chile, 77/110 isolates were found to have SNPs in the covRS regulon, irrespective of their site of isolation. However, only two of these isolates were found to express the high levels of capsule associated with hypervirulence as a result of mutations in covS (Wozniak et al. 2012). In contrast, screening of isolates from the Northern Territory of Australia, where skin infection predominates, found a much lower frequency of covRS mutation. Of the invasive (n = 12) and superficial (n = 13) Northern Territory isolates screened, only one was found to contain a covS mutation (Maamary et al. 2010). Whether this lower frequency of mutation is common in regions such as the Northern Territory, where emm-types are diverse, and skin infection is endemic, is unclear due to a lack of data on the covS status of clinical isolates from these and other similar regions.

Studies of other virulence markers have been used to track the origins of outbreak strains, and may prove useful in characterizing newly emergent strains of GAS. Exotoxin profile has been used as a means of differentiating GAS isolates in a number of studies. To date, 11 distinct superantigens have been identified. Three of these are chromosomally encoded (SpeG, SpeJ, and SMEZ), while the remaining eight (SpeA, SpeC, SpeH, SpeI, SpeJ, SpeK, SpeL, SpeM, and SSA) are located on temperate phage. Exchange of mobile genetic elements (MGE) such as phage contributes significantly to the genetic diversity seen within the GAS species, and is thought to be involved in the emergence of highly successful virulent clones (Lintges et al. 2010; Maamary et al. 2012; Maripuu et al. 2008). Superantigen profiling may therefore prove useful as a marker for the presence and transfer of prophages that encode additional genes playing a role in GAS pathogenesis or tissue tropism.

A high level of allelic variation within certain superantigen genes has been reported (Bianco et al. 2006; Talkington et al. 1993), and this, combined with the fact that studies of superantigen distribution from different groups often use different primer sets, confounds the comparison of superantigen profiling by different groups. However, a 2012 study of the distribution of superantigens amongst 480 diverse GAS isolates from Portugal has reported the use, for the first time, of a defined primer set enabling amplification of all reported superantigen alleles to date (Friaes et al. 2012). Using this technique, 11 different superantigen profiles were identified. Interestingly, this study found that none of the individual superantigen genes could be linked with a specific emm-type, but rather, superantigen profile shows a strong association with emm-type. Superantigen profile could not be accurately predicated using alternative typing methods (other than PFGE
subtyping), leading to the conclusion that superantigen profile may be a useful predictor of emm-type, but that superantigen profile cannot be inferred from emm-type (Friaz et al. 2012). This assertion is supported by previous findings from numerous studies undertaken in various countries suggesting a link between emm-type and superantigen profile (Commons et al. 2008; Le Hello et al. 2010; Schmitz et al. 2003; Vlaminckx et al. 2003).

The reported circulation of superantigens within the streptococcal population varies temporally. Screening of GAS isolates associated with a reemergence of invasive disease in Denmark between 1999 and 2002 found that the frequency of emm1 isolates harboring speA decreased from 94% in 1999 to 71% in 2002, while the emm1-specific prevalence of speC increased from 25 to 53% over the same period (Ekelund et al. 2005). Similarly, a comprehensive study of GAS isolates from Melbourne, Australia indicates that strains before the mid-1980s do not typically harbor speK, which is found in the genome of a high proportion of contemporary emm3 isolates (Commons et al. 2008). This highlights the potential usefulness of superantigen profiling to monitor the emergence of highly successful GAS clones during epidemics. There is evidence to suggest that superantigen prevalence in the GAS population differs geographically (Commons et al. 2008; Proft et al. 2003). However, cross-study comparison is, as stated previously, confounded by the use of different primer sets between studies.

4 Multilocus Sequence Typing

As virulence factors may be under strong selection pressure, and as is the case with superantigens, be present on mobile genetic elements (MGEs), they are not an ideal target for determination of evolutionary relationships between GAS strains. Multilocus Sequence Typing (MLST) is nucleotide-based method for investigating relationships among bacteria of the same species that targets seven selection neutral house-keeping genes. The genes used in the GAS MLST scheme (http://spyogenes.mlst.net/) are glucose kinase (gki), glutamine transport ATP-binding protein (gtr), glutamate racemase (murI), DNA mismatch repair (mutS), transketolase (recP), xanthine phosphoribosyltransferase (xpt), and acetoacetyl-CoA thiolase (yql). The combination of seven allelic variants is used to denote a specific multilocus sequence type (ST). More than 600 STs are listed on the S. pyogenes MLST website, indicating that ST is more discriminatory than emm-type.

A strong linkage between emm-type and ST has been reported for isolates from endemic countries (Enright et al. 2001). However a subsequent study by McGregor et al. (McGregor et al. 2004a) examining STs of isolates recovered from the Indigenous population of a remote Australian island (representing a low income region) reported a weaker relationship between emm-type and ST combinations present in isolates from the island and emm-type/ST combinations from other populations. The data can be interpreted as providing strong circumstantial evidence for LGT of the emm-gene in regions where GAS diversity is high. Further
complicating interpretation of MLST data was the observation of LGT involving housekeeping present on the ‘core’ genome (McGregor et al. 2004b). The proportion of emm-types associated with distant genetic backgrounds, as determined by MLST, were found to be much high for skin specialists (pattern D strains) and generalists (pattern E strains) than throat specialists (pattern A-C), suggesting that recombination events in latter occur at a lower frequency (Bessen et al. 2008).

5 Genomic Studies of Group A Streptococcus

It is becoming clear that emm-typing provides limited value when attempting to identify genetic factors linked to niche specialization of specific disease outcomes. Other approaches, examining the presence or absence of multiple genetic factors has been one pathway taken in attempt to provide a more thorough analysis of streptococcal diseases (Bessen et al. 2011; McMillan et al. 2006). Genomic studies are also proving to be pivotal in understanding the level of diversity that exists in the streptococcal population. The first M1 GAS genome was published in 2001 (Ferretti et al. 2001), with genome from other emm-types following soon after (Beres et al. 2002; Green et al. 2005; Smoot et al. 2002). Comparative analyses of these genomes clearly demonstrated the importance of MGEs, LGT, and recombination in generating diversity between emm-types. The majority of GAS genomes possess multiple bacteriophage, which in turn often carry virulence genes predicted to change the virulence of a lineage. Integrative conjugative elements (ICE) and remnant MGEs are also common (Green et al. 2005; Maamary et al. 2012; Smoot et al. 2002; Sumby et al. 2005).

However, it has been the intra-emm-type genomic studies utilizing multiple isolates with defined clinical histories that are providing the greatest insight into how strains with altered capacity to cause disease may evolve (Ben Zakour et al. 2012; Fittipaldi et al. 2012; Maamary et al. 2012; Sumby et al. 2005; Tse et al. 2012). These studies suggest that the acquisition of new MGEs, resulting in the elaboration of novel virulence gene repertoire is an important facet underlying the changes in virulence (Maamary et al. 2012; Sumby et al. 2005). Genomic analysis of the recent invasive disease outbreak associated with GAS emm59 provides an illustrative example of the value of population-based genomics. Prior to 2005, emm59 was not considered to be an emm-type associated with major outbreaks of invasive disease. However, in the second half of the decade, more than 500 invasive disease cases associated with this emm-type were reported in Canada (Tyrrell et al. 2010). Comparison of the core genome of outbreak isolates with historical and temporally distinct clones found the outbreak isolates to form a genetically distinct group (Fittipaldi et al. 2012). However, differences in the core genome between outbreak and non-outbreak isolates were remarkably small. While differences in MGE content between the outbreak strains and non-outbreak strain were apparent, as were difference in biological properties, the identity of putative genes within the MGEs that contribute to the relative differences in virulence could not be identified.
The first genomic analysis of GAS emm12 isolates involved in a scarlet fever outbreak were also recently reported (Tse et al. 2012). Unlike the emm59 outbreak, this outbreak appeared to be multiclonal. A novel ICE and bacteriophage were found to be distributed throughout the different clonal lineages involved in the outbreak. The latter contains genes encoding superantigens, and were hypothesized to be one of the reason for the observed increase in virulence.

6 The Role of Other β-Hemolytic Bacteria in 'GAS-Associated Diseases

*Streptococcus dysgalactiae* subsp. *equisimilis* (SDSE) is generally considered as a commensal organism with occasional potential to cause diseases in humans. Over the past few decades there have been several reports of SDSE causing diseases that are normally attributed to GAS. The list of diseases includes tonsillitis, skin infections, post streptococcus arthritis, pleuropneumonia, meningitis, endocarditis, puerperal septicaemia, necrotizing fasciitis, toxic shock syndrome (Jensen and Kilian 2012).

However, a possible link between SDSE and RF/RHD is still deemed as circumstantial, and controversial, with a general recalcitrance among the medical community against the view that β-hemolytic streptococci other than GAS (for example SDSE) could also be associated with RF/RHD. A major problem in definitively demonstrating a link between SDSE and RF in clinical settings is the long lag between infection and RF, and possible requirement for multiple episodes of infection, both of which call for well-controlled follow up studies. Such studies are near impossible as SDSE and GAS do cocolonize at the same tissue sites. Animal models of RHD are also in their infancy (Lymbury et al. 2003).

Despite the resistance to the accommodation of alternative views, several observations have strongly supported possible involvement of SDSE in the pathogenesis of RF/RHD. We and others (Bramhachari et al. 2010; McDonald et al. 2007) have observed that the GAS throat isolation rate in some communities is not commensurate with the RF/RHD burden. However, in these same population SDSE isolation rates from the throats are high. Indeed SDSE, but not GAS, has been recovered from an Indigenous Australian child after recurrent severe pharyngitis which was followed by RF in this patient (Davies et al. 2005) suggesting a role for SDSE in the pathogenesis of RF. All SDSE strains also express the M protein which may elicit heart-tissue cross-reactive autoantibodies (Haidan et al. 2000). In this study purified F(ab’)2 fragments from the Indigenous Australians reacted with surface protein extracts of SDSE, but not with those of GAS skin isolates (Haidan et al. 2000). RF/RHD associated GAS M types are also reported to possess collagen binding motifs in their M proteins suggesting a possible role for this interaction in the pathogenesis of RF/RHD (Dinkla et al. 2003) Interestingly, similar motifs were also seen in some SDSE M proteins (Dinkla et al. 2007). Collectively, these above
observations strongly suggest SDSE possessed many of the same characteristics as GAS that is linked to the pathogenesis of RF/RHD.

Phylogenically SDSE and GAS are closely related to each other (Lefebure et al. 2012). While there are conspicuous absences of some virulence genes in SDSE, up to half of GAS genes encoding virulence factors or surface associated proteins could be present in SDSE (Davies et al. 2007). Intra and Interspecies LGTs in GAS and SDSE have been long recognized (Sriprakash and Hartas 1996; Towers et al. 2004) with phages and ICEs contributing significantly to the population structure of these streptococci. Clonal relationships revealed that recombination far outweighs mutation in generating clonal variants. While both replacement and additive changes occur through LGT, characteristics that influence host-pathogen interactions are likely to be due to additive LGT (Choi et al. 2012).

Interestingly, cross-species LGTs between GAS and SDSE seem to be predominantly unidirectional; from the former to the latter (Choi et al. 2012). This seems to be true even for selectove-neutral housekeeping genes (McMillan et al. 2010). However, although the same emmSTs were found in different clonal complexes of GAS and SDSE suggesting occurrence of LGT of the emm-gene within the species, cross-species replacement of this gene was rarely observed. Likewise, SDSE with group A carbohydrate was found rarely. All SDSE isolates are speB-negative (Bramhachari et al. 2010) and no instance of additive LGT for this locus was ever reported to our knowledge. These observations, and mechanisms such as general resistance to phage mediated changes in SDSE due to clustered regularly interspaced short palindromic repeats (Shimomura et al. 2011) suggest that despite extensive ongoing LGTs, species demarcation between SDSE and GAS is unlikely to blur. However, with extensive additive LGTs into SDSE and sharing of many virulence characteristics the epidemiology of GAS diseases may be blurred in regions of high endemicity.

7 Conclusion

In a relatively short time, studies of the molecular epidemiology of GAS have progressed from the study of single genes, to population-based genomic comparisons. These more comprehensive studies have led to a view of GAS as highly dynamic organism. In endemic regions, where the number of circulating emm-types is high, no one lineage is associated with disease. SDSE is also abundant in these regions. In non-endemic countries, a small pool of emm-types that fluctuates at the local level is present. Ongoing LGT and recombination occasionally give rise to new lineages whose novel genetic repertoire may increase their virulence and disease causing potential. Mutation of core genes is a second mechanism that can also change the virulence properties of streptococci. Our inability to identify or validate the role of genes is specific disease outcomes, highlights the complex nature of streptococcal virulence, and suggests that different genetic repertoires may be responsible for the same disease in different lineages. Changes in
accessory virulence gene repertoire occur against a background of core virulence genes that are associated with niche specialization and core pathogenic processes. For a complete understanding of streptococcal functional genomics of large global collections of streptococci will be essential.

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