Assessment of the Relationship between Clinical Manifestation and Pathogenic Potential of *Streptococcus pyogenes* Strains-Distribution of Genes and Genotypes of Toxins

Tomasz Bogiel 1,*, Alicja Domian 1, Zuzanna Dobrzyńska 1,2, Agnieszka Mikucka 1 and Eugenia Gospodarek-Komkowska 1

1. Introduction

*Streptococcus pyogenes* is classified as one of the most dangerous pathogens that causes infections only in humans. These Gram-positive bacteria mainly colonize the mucous membrane of the throat and skin, contributing to various forms of infections and their complications. These infections, which can be non-invasive or invasive, especially involve the respiratory tract, skin and soft tissue, and bloodstream infection [1]. Despite a progress in medicine, *S. pyogenes* can still be life-threatening. Therefore, knowledge of pathogenic potential of the species and supervision of diseases caused by these microorganisms is of high importance. This constitutes the basis for their control and prevention [2].

*S. pyogenes* strains determine their pathogenicity by numerous virulence factors synthesis. These include surface proteins (e.g., M protein, lipoteichoic acids, fibronectin-binding protein), enzymes (e.g., hyaluronidases, streptokinases), hemolysins (streptolysins S and O), or produced extracellular substances (e.g., cytolytic toxins, pyrogenic exotoxins) [2,3]. They play an important role in colonization, provide a line of defense against the host immune system, and influence the course of disease [2]. In this research we focused on pathogenic...
potential resulting from the following toxins synthesis: A, B, C, J, H, and K. These substances, encoded by speA, speB, speC, speJ, speH, and speK genes, belong to superantigens. They have the ability to stimulate the activity of T and B lymphocytes of the colonized organism [4,5]. The immune response is so strong that there is an excessive release of inflammatory mediators including INF-γ (interferon gamma) and TNF-α (tumor necrosis factor alpha). As a consequence, effusions, extensive organ damage, and the onset of shock can be observed [2,6]. In addition, superantigens can have mitogenic and pyrogenic effects, and also inhibit immunoglobulin synthesis. This distorts the proper functions of the host immune system, which promotes the entry of purulent streptococci into human cells [4,6]. Among the toxins mentioned above, SpeB serves as a potent cysteine protease involved profoundly in the virulence of S. pyogenes [2,3,5].

Improving our knowledge of the involvement of streptococcal toxins will allow us to determine the pathogenic potential of group A Streptococcus (GAS) strains, make a diagnosis easier and faster, or even prognostic the course of infection. This may enable the implementation of appropriate targeted therapy earlier. This, in turn, translates into the length and efficiency of healing process by shortening hospitalization time with minimizing side effects of treatment. In the long term, this may also contribute to reducing antibiotic resistance of S. pyogenes strains. Hence, the aim of this study was to assess, with the application of real-time PCR, the frequency of streptococcal toxins genes: speA, speB, speC, speJ, speH, and speK and to determine the genotypes distribution of S. pyogenes strains. An additional purpose of this study was to determine the virulence potential of individual S. pyogenes strains derived from different body sites to evaluate a relation between these genes and/or genotypes presence and the origin of S. pyogenes strains from specific types of clinical material.

2. Results

The conducted research enabled us to show the wide variety in the percentages of the examined genes and the distribution of genotypes, as shown in Table 1. The speB gene was observed with the highest frequency-145 (98.6%) strains, while the speK was the least frequent gene-12 (8.2%) strains.

Table 1. The percentages of genes presence and genotype distribution among the examined S. pyogenes strains (n = 147); (+)–gene presence confirmed, (−)–lack of a particular gene.

| Gene/Assigned Genotype Name | A  | B  | C  | D  | E  | F  | G  | H  | I  | J  | K  | L  | M  | N  | O  | P  | Q  | n  | %  |
|----------------------------|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|
| speB                      | +  | +  | +  | +  | +  | +  | +  | +  | +  | +  | −  | +  | +  | +  | +  | +  | +  | +  | 145 | 98.6 |
| speC                      | −  | +  | +  | −  | −  | +  | +  | −  | −  | −  | −  | −  | +  | −  | +  | −  | −  | +  | 69  | 46.9 |
| speJ                      | −  | −  | −  | +  | −  | +  | −  | −  | −  | −  | −  | −  | +  | +  | −  | −  | −  | −  | 42  | 28.6 |
| speH                      | −  | +  | −  | −  | −  | +  | −  | −  | −  | −  | −  | −  | +  | −  | −  | −  | −  | −  | 38  | 25.9 |
| speA                      | −  | −  | −  | −  | −  | −  | +  | −  | −  | −  | −  | −  | +  | −  | −  | −  | −  | +  | 19  | 12.9 |
| speK                      | −  | −  | −  | −  | −  | −  | −  | +  | +  | +  | +  | −  | −  | −  | +  | −  | −  | +  | 12  | 8.2  |
| n = 147                   | 32 | 25 | 23 | 14 | 13 | 13 | 9  | 3  | 3  | 2  | 2  | 2  | 2  | 1  | 1  | 1  | 1  | 1  | 12  | 8.2  |
| %                         | 21.8 | 17.0 | 15.6 | 9.5 | 8.8 | 8.8 | 6.1 | 2.0 | 2.0 | 1.4 | 1.4 | 1.4 | 0.7 | 0.7 | 0.7 | 0.7 |

The examples of amplification curves for each particular gene are presented in Supplementary Material–Figure S1, while the high resolution melting curves obtained to confirm amplification specificity can be found in Figure S2.

Based on the detected genes, 17 genotypes (A–Q) among the studied S. pyogenes strains were determined. The A genotype, which was devoid of each of the detected genes except the speB gene, was the most common-32 (21.8%) strains. Four (2.7%) strains presented distinctive genotypes (N–Q). The distribution of genes and genotypes among the
S. pyogenes strains was statistically compared with their theoretical distribution. In each case, the p-value was below 0.001, except the distribution of the speC gene (p = 0.458).

A statistically significant relationship was noted between the genes: speA and speC (p = 0.002), speA and speH (p = 0.013), speA and speI (p < 0.001), speC and speH (p < 0.001), speC and speI (p = 0.037), and speH and speI (p < 0.001).

The speA and speI genes were most common among the strains isolated from throat swabs—amongst 6 (31.6%) strains and 14 (33.3%) strains, respectively. In contrast, speB (60 strains, 41.4%), speC (31 strains, 44.9%), speH (20 strains, 52.6%) and speK (5 strains, 41.7%) genes dominated amongst the strains isolated from wound swabs. The percentages of GAS strains with the presence of particular genes, which were isolated from selected types of clinical material, are presented in Table 2.

**Table 2.** Occurrence of the virulence factor genes with respect to the origin of S. pyogenes strains from selected types of clinical material.

| Clinical Specimen Type | speA | speB | speC | speH | speI | speK |
|------------------------|------|------|------|------|------|------|
| Wound swab             | 3.0  | 60.0 | 41.4 | 20.0 | 12.6 | 5.1  |
| Throat swab            | 6.0  | 31.6 | 21.4 | 21.1 | 33.3 | 25.0 |
| Purulent material      | 2.0  | 10.5 | 12.4 | 23.7 | 4.8  | 0.0  |
| Ulcer swab             | 2.0  | 10.5 | 6.9  | 7.1  | 4.8  | 0.0  |
| Blood samples          | 1.0  | 5.3  | 6.2  | 4.3  | 2.6  | 1.0  |
| Nose swab              | 1.0  | 5.3  | 4.8  | 7.2  | 7.1  | 8.3  |

There was a statistically significant relation between the occurrence of: the speA gene and the origin of S. pyogenes strains from wound swabs (p = 0.033); the speI gene and S. pyogenes strains origin from throat swabs (p = 0.032); and the speH gene and the origin of S. pyogenes strains from purulent material (p = 0.012).

The A–D genotypes were the most prevalent among the strains isolated from wound swabs (43.8%, 52.0%, 47.8%, and 50.0%, respectively). In turn, genotypes E and F (both 38.5%) were dominant in strains isolated from throat swabs. The percentages of GAS strains with specified genotype (A–F), which were isolated from selected types of clinical material, are shown in Table 3.

**Table 3.** The distribution of genotypes A–F with respect to origin of S. pyogenes strains from selected types of clinical material.

| Clinical Specimen Type | A     | B     | C     | D     | E     | F     |
|------------------------|-------|-------|-------|-------|-------|-------|
| Wound swab             | 14    | 43.8  | 13    | 52.0  | 11    | 47.8  | 7     | 50.0  | 1     | 7.7   | 3     | 23.1  |
| Throat swab            | 3     | 9.4   | 3     | 12.0  | 4     | 17.4  | 4     | 28.6  | 5     | 38.5  | 5     | 38.5  |
| Purulent material      | 2     | 6.3   | 8     | 32.0  | 4     | 17.4  | 1     | 7.1   | 1     | 7.7   | 0     | 0.0   |
| Ulcer swab             | 7     | 21.9  | 0     | 0.0   | 0     | 0.0   | 1     | 7.1   | 2     | 15.4  | 0     | 0.0   |
| Blood samples          | 4     | 12.5  | 1     | 4.0   | 0     | 0.0   | 0     | 0.0   | 0     | 0.0   | 1     | 7.7   |
| Nose swab              | 1     | 3.1   | 0     | 0.0   | 2     | 8.7   | 0     | 0.0   | 1     | 7.7   | 2     | 15.4  |

The relationship between: the presence of A genotype, and the origin of GAS strains from ulcer swabs (p = 0.002), the presence of B genotype, and the origin of GAS strains
from purulent material \((p = 0.001)\), and the presence of E genotype, and the origin of GAS strains from wound swabs \((p = 0.024)\), has been demonstrated.

3. Discussion

*S. pyogenes* belongs to the group of bacterial pathogens contributing to the development of human infections, which significantly differ in location, course, prognosis, and mortality [7]. Therefore, it is important to conduct research on the pathogenic potential of *S. pyogenes*, consisting in the detection of the most important virulence factors or genes encoding them and the assessment of their occurrence depending on the location and form of infection.

The mentioned virulence factors genes of *S. pyogenes* are mostly pyrogenic toxins and proteases, among others: *speA*, *speB*, *speC*, *speJ*, *speH*, and *speK*, which encode streptococcal toxins designated as speA, speB, speC, speJ, speH, and speK, respectively. Streptococcal pyrogenic exotoxins are virulence factors produced exclusively by *S. pyogenes*. They are encoded by genes located on bacterial chromosomes or prophages [8,9]. Not all the genes of toxins are present in each GAS strain. Therefore, differences in the frequency of particular genes encoding these virulence factors can be observed and implicate the course of the disease and prognostic factors [10].

In the present research, the frequency of the *speA* gene was noted in 19 \((12.9\%)\) of the examined *S. pyogenes* isolates. There was also a statistically significant difference in the distribution of this gene between the studied GAS strains. In Yu and Ferretti [11] research, including 302 *S. pyogenes* isolates from patients from different countries, the percentage of strains with the presence of the *speA* gene was 14.9%. This result was similar to that obtained in our own research. The studies presented by Szczypa et al. [12] indicated the frequency of the *speA* gene in the examined strains at the level of 24.4%. The number of *S. pyogenes* isolates used in the above-mentioned research was, however, much lower (41 strains) than in the presented results of our own research. In the study by Li et al. [13], the gene determining the presence of pyrogenic toxin A was present in 34.3% of the tested isolates. This study included 271 GAS strains. The analysis of the results of our own and other authors, concerning the frequency of the *speA* gene in *S. pyogenes* strains, showed slight differences between the presence and the absence of this gene in terms of different forms of infections caused by the studied isolates.

The presence of the *speB* gene in the studied *S. pyogenes* strains was confirmed among 145 \((98.6\%)\) isolates in our own research. There was also a statistically significant difference in the distribution of the encoding speB gene between the studied GAS strains \((p < 0.001)\). Tyler et al. [14] noted the presence of the *speB* gene in 99.3% of *S. pyogenes* strains, causing invasive and non-invasive diseases. The result is relatively close to the results of our study. Interestingly, research presented by Wu et al. [15] showed the presence of the gene encoding speB only in 81.0% of GAS strains isolated from patients with scarlet fever and amongst 72.4% of strains isolated from patients with *acute pharyngitis* and *tonsillitis*. In the available literature, it is also possible to observe a higher percentage of GAS isolates carrying the *speB* gene in patients diagnosed with invasive forms of *S. pyogenes* infections [16]. Despite slight differences, the frequency of the *speB* gene was always high in the available literature.

In the presented studies, the frequency of the *speC* gene was found in almost half of the examined *S. pyogenes* strains \((46.9\%)\), and there was no statistically significant difference between the number of GAS strains with and without the gene encoding speC. In the research work of Yu and Ferretti [17], which included 315 isolates, similarly to our own studies, the gene encoding pyrogenic toxin C was present in 50.8% of *S. pyogenes* strains. In the study by Maripuu et al. [18], the presence of the *speC* gene was significantly higher than that recorded on the basis of our own research results. This gene was detected among 66.3% isolates causing bacteremia, Streptococcal Toxict Shock Syndrome (STSS), and *erysipelas* (the cited analysis included 92 GAS strains). This might indicate a greater share of strains with such a pathogenic potential in invasive infections [2]. The occurrence of *speC* was differently assessed by Li et al. [13]. The presence of that gene in the strains tested by them
reached 91.1%. The fact that as many as 219 isolates, out of 271 included in the study, were derived from patients with scarlet fever, could have an impact on such a high percentage of strains with speC gene.

In our research, the percentage of S. pyogenes strains with the spef gene was 28.6%. Statistically significant differences were also observed between the number of GAS strains with and without the gene encoding spef. A similar result (32.7%) was obtained by Friæs et al. [19], although the number of strains used by them for the analysis was almost three times higher. Among the genes of pyrogenic exotoxins investigated by them and encoded on the bacterial chromosome, i.e., spef, speG, and smeZ, the spef gene was the least frequent. In the study by Li et al. [20], the spef gene was present among isolates with a frequency of 22.2%. The researchers detected 11 genes encoding superantigens, of which spef belonged to the group of the three rarest genes encoding selected superantigens. On the other hand, Berman et al. [21] found the spef gene only in 11 (14.3%) strains, but the number of the tested isolates was over half lower than that in our own research. Taking into account 11 genes encoding superantigens they detected, spef was relatively rare, while six other genes, i.e., speG, smeZ, speI, speC, speA, and speH, were found more often.

Compared to the spef gene, the frequency of speH and speK genes in our own research differed to a greater extent from the results of other authors’ studies. In our studies, the speH gene was found in 38 (25.9%) examined strains, and the speK gene in 12 (8.2%). Statistically significant differences were observed between the number of GAS strains with and without the gene encoding speH and speK. The p-value was below 0.001 for both genes distribution. Friæs et al. [19] revealed the presence of the speH and speK genes in 82 (17.1%) and 118 (24.6%) GAS strains, respectively. Taking into account all the superantigen genes that they studied, which are encoded within prophages, speH and speK occurred relatively rarely. In Li et al. studies [20], the speH gene was noted in 155 (52.2%) strains of S. pyogenes, and the speK gene in only 2.0% of the isolates. It should be noted that speK was the least frequently detected gene, which was consistent with the results obtained in our own study. In the study by Berman et al. [21], the percentage of GAS strains with the speH gene was 36.4%. The researchers did not detect the speK gene in any of the S. pyogenes strains and the results obtained by them were similar to those developed by Li et al. [20].

Comparing the results of this study with the results of the above authors, it can be concluded that the reason for the differentiation of percentage of individual genes of toxin may be the very specificity of GAS strains from patients from different geographic regions, age of patients, or, particularly, symptoms of infection occurring in patients, but the above-mentioned aspects were not the subject of research of this work.

Of note, the speA, speC, speH, and speK genes are located within the prophages, which also had a key influence on the differences in their frequency [21,22]. According to the available literature, the frequency of pyrogenic toxins genes is also influenced by the presence of particular emm types. For example, the genes speA and spef are found much more often in strains of the emm1 type. The speC gene is the most common in GAS strains of the emm28 type, the speH gene in the GAS strains of the emm12 type, while the speK gene is associated with, among others, emm2 and emm3 types [18,23]. However, the emm types distribution of S. pyogenes was not the subject of this study.

Having data on the frequency of genes encoding virulence factors, it is possible to determine the patho-genetic profile (genotypes) of the studied GAS strains. In our own research, 17 different genotypes were observed. Their distribution differed significantly from the theoretical one, as indicated by the p-value, which was below 0.001. Namely, the number of S. pyogenes strain groups to which specific genotypes were assigned was quite diverse.

Compared to researches of other authors, the number of grouped genotypes was relatively small. This was due to the much greater number of virulence factor genes that have been detected in other studies. The more genes tested, the greater the diversity of the gene profiles. An important reason for obtaining a different number of genotypes was also the matter of the diversity of S. pyogenes strains used for the study. In our own
research, these strains were isolated from many types of clinical material, but throat and wound swabs were the most common. This means that the isolated GAS strains could show some similarity in the presence of the speA, speB, speC, speF, speH, and speK genes and, as a consequence, the convergence of genotypes. Moreover, most of the strains were isolated within one hospital, which probably resulted in a limited degree of possible geographic differentiation of the isolates.

Helal et al. [24] revealed large diversity of the genotypes they identified. They recorded 33 genotypes among only 38 S. pyogenes strains. They put together these genotypes on the basis of the genes speA, speC, speG, speH, speL, speM, smeZ, and ssa. Despite the small number of GAS strains studied by them, the number of genotypes was almost twice as high as in our own research. Abraham and Sistla [25] detected as many as 71 gene profiles, defined on the basis of the genes speA, speC, speG, speH, speL, speM, smεZ, and ssa, among 206 S. pyogenes strains. They isolated these strains from clinical material derived from patients with both non-invasive and invasive infections. It is noteworthy that the authors found the presence of 66 emm types of S. pyogenes, and showed a relationship between several of them and specific genotypes. The number of genotypes obtained during the research is undoubtedly influenced by the emm types among the S. pyogenes strains. Different emm types may have the same genotype and, vice versa, one emm type may be associated with different genotypes. Another factor contributing to the existence of different genotypes was horizontal gene transfer. The acquisition or loss of prophage by a bacterial cell favors its genetic variability. This translates into the differences in the frequency of particular gene profiles in the GAS population, and into intraspecific variation by generating new genotype variants [8,21,25,26].

Our own research shows that in S. pyogenes strains, the genotype speA (−) speB (+) speC (−) speF (−) speH (−) speK (−) (genotype A) was the most common (21.8%). This does not mean, however, that they only produced speB toxin in the infections pathomechanism. They could have other virulence factors genes that were not included in this study. When analyzing the presence of the speA, speB, speC, speF, speH, and speK genes, the percentage of genotype A in the research of other authors was negligible. Meisal et al. [27], who established gene profiles of 262 isolates causing invasive infections, detected genotype A in 15 (5.7%) GAS strains. Ibrahim et al. [28] and Maripuu et al. [18] found this genotype in only one strain (11.1% and 1.1%, respectively). This shows that the speB gene co-exists more often with the genes of pyrogenic exotoxins. Our research shows that it was most often the speC gene, which was confirmed by a fairly high percentage of genotype B (25 strains, 17.0%), C (23 strains, 15.6%), and F (13 strains, 8.8%). The simultaneous presence of these two genes was found in a total of 69 (46.9%) strains of S. pyogenes. However, there was no statistically significant relationship between the speC and speB genes (p = 0.531). A similar result was obtained by Commons et al. [23]. Almost half of the strains examined by them (53 isolates, 49.5%) were positive for the speB and speC genes, but the co-existence of the speB and speG genes was the most common (96 strains, 89.7%).

The genes investigated in the present study more often co-exist with genes of other virulence factors that have not been analyzed in our research. The speF gene is usually associated with the smeZ gene. Mostly, the speH and speL genes are also detected because they are encoded within the genetic material of the same bacteriophage ϕ370.2. On the other hand, the speK gene is usually associated with the speA gene [19,28]. In the case of our own research, only in two strains of S. pyogenes the simultaneous presence of the speA and speK genes was found. Furthermore, no statistically significant correlation between them was observed (p = 0.963), in contrast to the speA-speC, speA-speH, and speA-speF gene pairs for which the p-value reached 0.002, 0.013, and <0.001, respectively.

In our own research, the percentage of N-Q genotypes was the lowest. They were found once among the tested GAS strains. In the study by Berman et al. [21], the rarest gene profiles were completely different. The fact that the speA (−) speB (+) speC (−) speF (−) speH (+) speK (−) genotype (genotype G in our study), found by them in one strain, was...
detected by us in 9 (6.1%) GAS strains, is the evidence of the varied distribution of gene profiles among *S. pyogenes* strains.

*S. pyogenes* strains are isolated from many types of clinical material, such as throat and wound swabs or purulent material and blood [7,29]. By detecting genes encoding virulence factors it can be assessed whether their presence is related to the origin of GAS strains from a specific clinical material. In our study, the *speA* and *speJ* genes were most common (31.6% and 33.3%, respectively) among the strains isolated from throat swabs. Moreover, a statistically significant correlation was found between the presence of the *speJ* gene and the origin of GAS strains from these clinical materials (*p* = 0.032). The throat swab is usually taken in the case of *pharyngitis* or *tonsillitis*, which are classified as non-invasive infections [3]. Altun and Merici Yapıcı [30], who conducted studies among 200 patients with *pharyngitis* and *tonsillitis*, isolated only 15 GAS strains. The *speA* gene was present in three, and the *speJ* gene was present in only two of them. However, their research did not assess the relationship between the presence of the detected genes and the origin of *S. pyogenes* strains. Hamzah et al. [31] found the *speA* and *speJ* genes more frequently in GAS strains isolated from patients with non-invasive rather than invasive infections (65.0% vs. 35.0% for *speA* and 63.6% vs. 36.4% for *speJ*). According to them, this could be due to the overall greater number of GAS isolates that came from people with non-invasive infections. Detection of *speA* or *speJ* genes in GAS strains may also be associated with the potential development of invasive infections. Kittang et al. [32] demonstrated a relationship between these genes and the invasiveness of infections. In patients, from whom they isolated *S. pyogenes* strains, the following infections occurred: necrotizing fasciitis, STSS, *pneumonia*, meningitis, and *peritonitis*.

Although the *speB*, *speC*, *speH*, and *speK* genes, as well as A-D genotypes, dominated in strains isolated from wound swabs, no statistically significant correlation was found between the occurrence of these genes and gene profiles, and the origin of GAS strains from wound swabs. This could be due to the fact that the examined *S. pyogenes* strains, in general, were most often detected in this type of clinical material. The study by Strus et al. [9] shows that there is a relationship between the presence of the *speH* gene and the origin of the strains from this type of clinical material and wound infections in the patient. They explain that the reason for differences between the occurrence of superantigen genes and the type of clinical material and the form of infection, was the isolation of *S. pyogenes* strains from people from different countries and regions. On the other hand, in our own research, we found a correlation between the presence of the *speH* gene and the origin of the strains from purulent material (*p* = 0.012), and also the relation between the B genotype (in which the *speH* gene is found) and the origin of the strains from this type of material (*p* = 0.001). Purulent material culture is usually associated with an infection of the patient’s skin and subcutaneous tissue [33]. Therefore, it can be assumed that GAS strains with the *speH* gene contributed often to such infections.

There was also a relationship between A genotype and the origin of the strains from ulcer swabs. This may indicate that GAS strains with the *speB* gene only are a probable etiological factor of infections in patients with diabetes (diabetic foot infection) or patients with venous ulcers [34,35].

In conclusion, the results of this study show the diversity of *S. pyogenes* strains with regard to the presence of the *speA*, *speB*, *speC*, *speJ*, *speH*, and *speK* genes and their genotypes. Comparing our results with the available literature, the different values of the percentages of detected genes, as well as the number and percentages of gene profiles of the studied strains, were obtained, which confirms the validity of the above statement. The demonstrated relationships between the presence of some of the detected genes and genotypes, and the origin of *S. pyogenes* strains from specific types of clinical material, indicate that GAS strains isolated from them may be responsible for the occurrence of infection with a specific clinical manifestation. Different types of clinical material for microbiological investigation were used in the present study allowing GAS strains growth. It allowed for the assessment of their genetic diversity in relation to the origin of the isolates. Moreover, the assessment of
the frequency of occurrence of the detected genes and the determination of the *S. pyogenes* strains genotypes, allowed for the evaluation of their relative pathogenic potential. The detection of exotoxins genes, as well as other virulence factors, may also be useful in the development of vaccines. Based on these genes, the virulence factors of the specific GAS strains can be determined. The most common of them may be a potential antigen necessary to induce the immune response in humans, which requires further and more detailed studies [36].

4. Materials and Methods

4.1. Origin of the Strains and Their Selection Criteria

Initially, the study involved 168 clinical isolates of *S. pyogenes*, which were isolated between 2008 and 2020, mainly from wound swabs (38.7%), throat swabs (19.6%), and purulent material (11.9%) collected for routine diagnostic purposes. These strains came from the collection of the Microbiology Department of Ludwik Rydygier Collegium Medicum in Bydgoszcz Nicolaus Copernicus University in Toruń, Poland. The tested GAS strains were generally isolated from patients from surgical departments (32.1%) and outpatient clinics (15.5%). The detailed origin of the isolates included into the study is presented in the Supplementary Material: Tables S1 and S2.

Due to the fact that some GAS strains were derived from the same patients, 147 strains were eventually used to perform gene and genotype distribution calculations. Each of them was isolated from a different patient. An example of a gel showing PFGE patterns for the selected group of strains used to exclude duplicate isolates (data not shown) is shown in the Supplementary Material (Figure S3).

The reference *S. pyogenes* strains, derived from the German Collection of Microorganisms and Cell Cultures (DSMZ), served as positive control for a particular gene detection.

4.2. Bacterial Culture and Strain Identification

The tested and reference strains were plated on Columbia Agar with 5% sheep blood (Becton Dickinson, Franklin Lakes, NJ, USA). The culture was carried out at 37 °C for 18–20 h. The species affiliation of the tested and reference strains was confirmed by the (Matrix-Assisted Laser Desorption/Ionization Time-of-Flight Mass Spectrometry) MALDI-TOF MS method, using the MALDI Biotyper system (Bruker Daltonics GmbH & Co. KG, Bremen, Germany). This procedure was conducted according to the manufacturer’s protocol.

4.3. Bacterial Genomic DNA Isolation

Genomic DNA was isolated from the *S. pyogenes* strains in accordance with the instruction at the Centers for Disease Control and Prevention (CDC) website. In order to confirm the DNA isolation accuracy, the concentration of all DNA samples was evaluated spectrophotometrically (Photometer, Eppendorf, Germany). The DNA samples were stored at −20 °C before their further use in real-time PCR for the presence of the selected virulence factors genes evaluation.

4.4. Virulence Factor Genes Detection

The presence of the following genes: *speA*, *speB*, *speC*, *speH*, *speJ*, and *speK* was determined by real-time PCR method in the LightCycler 480 II Instrument (Roche Diagnostics, Basel, Switzerland). Positive and negative controls were used simultaneously and each time. The DNA isolated from *S. pyogenes*: DSMZ 20565 strain (for the *speA* and *speJ* genes), DSMZ 25932 strain (for the *speB*, *speC*, and *speH* genes) and DSMZ 11728 strain (for *speK* gene) served as PCR positive control. Reactions were performed with the application of molecular biology grade sterile water (EURx, Gdansk, Poland), F and R primers [9] (Genomed, DNA Sequencing and Oligonucleotide Synthesis Laboratory, Institute of Biochemistry and Biophysics of the Polish Academy of Sciences, Warsaw, Poland described in Table 4), and the 5x HOT FIREPol® EvaGreen® HRM Mix (no ROX) reaction mixture (Solis BioDyne, Estonia). The instruction attached to the reaction mixture was followed:
the reaction volume of one sample was 20 µL with 4 µL of HRM Mix added to reach the final concentration of 1x, both primers were used at the final concentration of 200 nM (5 µL of each at the initial concentration of 1 µM), the remaining volume was water-5 µL and DNA template (1 µL). The amplification program was initially optimized for a particular gene and finally consisted of: initial activation at 95 °C for 5 min, followed by 45 cycles (50 for speB gene) of amplification, each consisting of 10 s at 95 °C, 20 s (10 s for speJ and speK genes) at the annealing temperature (see Table 4), followed by 72 °C for 20 s (60 s for speH, speJ, and speK genes) (for the amplification results see Figure S1).

### Table 4. The real-time PCR primers specification and the annealing temperature applied in PCR amplification.

| PCR Primer Name | Gene Detected | Primer Sequence 5' → 3' | Tm (°C) | Annealing Temperature (°C) | Product Size (bp) |
|-----------------|---------------|-------------------------|---------|---------------------------|------------------|
| speA-F          | speA          | CTTAGAACCAAGAGAATGGC     | 49.7    | 52                        | 200              |
| speA-R          |               | ATAGGCTTGGATAACCATC      | 46.8    |                           |                  |
| speB-F          | speB          | TTCTAGGATACCTCACACC     | 49.7    | 55                        | 300              |
| speB-R          |               | ATTTGAGCAGTTGAGTAGC      | 49.7    |                           |                  |
| speC-F          | speC          | CATCATGAGGAAATACGC      | 49.7    | 55                        | 246              |
| speC-R          |               | TGTCAGAATTCGAGTCGC      | 49.7    |                           |                  |
| speH-F          | speH          | AAGCAATTCTTATAAATACCC    | 46.4    | 52                        | 630              |
| speH-R          |               | TTAGCTGATGGACACATCTACA  | 49.2    |                           |                  |
| speJ-F          | speJ          | GATAGTGAAAATATAGACGC    | 45.5    | 52                        | 639              |
| speJ-R          |               | GCTTCTATATTGTAGTCC      | 47.7    |                           |                  |
| speK-F          | speK          | GTGTGTCTTATGGACACGCC    | 54.4    | 56                        | 564              |
| speK-R          |               | GAAACATATATGCTCTAGAT    | 48.5    |                           |                  |

After the amplification reaction, high resolution melting curves protocol was applied (95 °C for 5 s, 65 °C for 60 s, and constant heating until reaching 97 °C, ramp rate 0.11 °C/s with 5 read-outs per each °C) and the real-time PCR products specificity was checked (for the melting curves results see Figure S2).

To verify sizes of the amplification products for the particular genes and prove the specificity of the applied real-time PCR methodology, conventional PCR and electrophoresis were applied. Their results, as the examples of the gel pictures showing electrophoretic resolution of targeted and specific amplification of bacterial DNA, were included into Supplementary Material (Figures S4 and S5). It visualizes, separately, the amplicons of the particular genes for the selected reference strains: amplification of speA gene (200 bp), speH gene (630 bp), speJ gene (639 bp) (Figure S4), and speB gene (300 bp), speC gene (246 bp), and speK gene (564 bp) (Figure S5).

### 4.5. Statistical Analysis

In order to compare the distribution of genes and genotypes among the tested S. pyogenes strains, with the theoretical distribution, the Chi-Square Goodness of Fit Test with α ≤ 0.05 was carried out. The GNU PSPP (data analysis software system) program was used for this purpose. Whereas, in order to assess the relationship between the presence of genes and genotypes, and the origin of S. pyogenes strains from specific types of clinical material, the Chi-Square Test of Independence with α ≤ 0.05 was applied. The calculations were performed in the STATISTICA 13.3 (data analysis software system) program.

### 5. Conclusions

Among the clinical S. pyogenes strains, there is considerable diversity in the occurrence of toxin genes. The fact that the speB gene and the speA (−) speB (+) speC (−) speJ (−) speH (−) speK (−) genotype dominated in the studied GAS strains, might be related with the preferential coding of this toxins on the bacterial chromosome of this species strains.
The predominance of the speA and speJ genes in strains isolated from throat swabs could potentially indicate a possible change in the clinical manifestation of the infection, from initially local to systemic. The relationship between the origin of S. pyogenes strains from purulent material and the more frequent occurrence of the speH gene among these strains indicates their molecular potential to cause skin and soft tissue infections.

Taking into account the pathogenic potential of a particular S. pyogenes strain and knowing the role of a particular toxin in the pathogenesis of S. pyogenes infections, we might be allowed to foresee the course of the infection. With the application of real-time PCR for particular toxins genes detection, as a standard microbiological procedure in the future, we may also be able to diagnose the infection easier, administer targeted antimicrobial therapy faster, and eventually decrease patients’ mortality rate.

**Supplementary Materials:** The following are available online at https://www.mdpi.com/article/10.3390/biomedicines10040799/s1: Figure S1. The examples of pictures showing results of real-time PCR (amplification curves) for detection of the following genes: speA, speB, speC, speH, speJ, speK. Figure S2. The examples of melting curves and melting peaks pictures showing specificity of the real-time PCR product for the confirmation of the following genes detection: speA, speB, speC, speH, speJ, and spek. Figure S3. An example of the gel picture showing electrophoretic resolution of bacterial DNA and the corresponding PFGE patterns for the selected group of strains. Figure S4. An example of the gel picture showing electrophoretic resolution of bacterial DNA amplification with conventional PCR, the study performed additionally to confirm the sizes of the amplification product for a particular genes and specificity of the applied real-time PCR methodology (speA, speH, and speJ genes). Figure S5. An example of the gel picture showing electrophoretic resolution of bacterial DNA amplification with conventional PCR, the study performed additionally to confirm the sizes of the amplification product for a particular genes and specificity of the applied real-time PCR methodology (speB, speC, and speK genes). Table S1. The detailed origin of the examined S. pyogenes strains (units). Table S2. The detailed origin of the examined S. pyogenes strains (clinical specimen).

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**Data Availability Statement:** The data presented in this study are not publicly available as a matter of confidentiality. However, these data are available upon request from the corresponding author.

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