# ARTICLE IN PRESS

YGENO-08670; No. of pages: 5; 4C:

Genomics xxx (2014) xxx-xxx



Contents lists available at ScienceDirect

# Genomics

journal homepage: www.elsevier.com/locate/ygeno



- Streptococcus pneumoniae Genome Database (SPGDB): A database for
- strain specific comparative analysis of Streptococcus pneumoniae genes
- and proteins
- Rayapadi Swetha <sup>a</sup>, Dinesh Kumar Kala Sekar <sup>b</sup>, Ekambaram Durga Devi <sup>b</sup>, Zaheer Zameer Ahmed <sup>b</sup>, Sudha Ramaiah <sup>a</sup>, Anand Anbarasu <sup>a,\*</sup>, Kanagaraj Sekar <sup>b</sup>
  - <sup>a</sup> Medical & Biological Computing Laboratory, School of Biosciences and Technology, VIT University, Vellore 632 014, India
  - b Laboratory for Structural Biology and Biocomputing, Supercomputer Education and Research Centre, Indian Institute of Science, Bangalore 560 012, India

#### ARTICLE INFO

#### Article history:

Received 19 May 2014 10

11 Accepted 22 September 2014

12 Available online xxxx

#### Keywords:

14

15 Genome

13

30 31

33

34

35

36

37

38

39 40

41

42

43 44

45

46 47

48

49

50

51 52

53

54

Annotation

17 Resistance 28

Pneumococci

Database

# 1. Introduction

Streptococcus pneumoniae or "pneumococcus" is the most common cause of pneumonia and related invasive diseases such as bacterial meningitis, sepsis, otitis media and sinusitis. The bacterium causes extreme morbidity and mortality worldwide, particularly in infants and in the elderly. It spreads through immediate contact with respiratory discharges from patients and healthy carriers [1-3]. In several developed countries, the burden of the pneumococcal disease is intensified with chronic diseases (sickle-cell disease, chronic renal failure, chronic liver disease and asplenia), HIV and Mycobacterial infection, as well as an aging population [4]. Pneumococci are also well-known for their inherent antibiotic resistance, principally affecting beta-lactam, macrolide and sulfonamide sensitivity, thereby, it results in treatment failures. In 2000, the World Health Organization (WHO) reported that 14.5 million cases of pneumococcal disease occurred, resulting in 826,000 deaths in toddler aged between 1 and 59 months [5]. Thus, the diseases caused by Pneumococci constitute a major global public health problem and it is being extensively studied at the genome level [6,7]. In recent decades, the increasing numbers of Pneumococci genomes being sequenced entails significant interest in comparing the genome of each strain with other strains. This provides insight into strain specific characteristics that may act as a significant role in virulence and antimicrobial resistance.

ABSTRACT

Streptococcus pneumoniae causes pneumonia, septicemia and meningitis. S. pneumoniae is responsible for significant mortality both in children and in the elderly. In recent years, the whole genome sequencing of various 20 S. pneumoniae strains have increased manifold and there is an urgent need to provide organism specific annotations to the scientific community. This prompted us to develop the Streptococcus pneumoniae Genome Database 22 (SPGDB) to integrate and analyze the completely sequenced and available S. pneumoniae genome sequences. Fur- 23 ther, links to several tools are provided to compare the pool of gene and protein sequences, and proteins structure 24 across different strains of S. pneumoniae. SPGDB aids in the analysis of phenotypic variations as well as to perform 25 extensive genomics and evolutionary studies with reference to S. pneumoniae. The database will be updated at 26 regular intervals and is freely accessible through the URL: http://pranag.physics.iisc.ernet.in/SPGDB/.

© 2014 Published by Elsevier Inc.

Hence, there is an urgent need to develop a database exclusively for 55 S. pneumoniae, which we have attempted in the form of Streptococcus 56 pneumoniae Genome Database (SPGDB). SPGDB provides links to tools 57 that facilitate the comparison between multiple genomes of *Pneumococci* 58 and to recognize the genome components of medical importance. The 59 database, SPGDB, provides a powerful, user-friendly interface to perform 60 various Boolean searches or sequence based searches. The database is 61 also interfaced with the genome map of pneumococcal strains and the 62 available three-dimensional structures of pneumococcal proteins in Pro- 63 tein Data Bank (PDB) [8]. This facilitates the users to explore functionally 64 active proteins involved in pneumococcal disease pathogenicity and 65 these structures can also be exploited further for structural analysis 66 based on the user requirements. Additionally, it provides detailed in- 67 formation on the characteristics, virulence factors, pathogenesis and 68 laboratory diagnosis of Pneumococci for researchers.

### 2. System design and implementation

The SPGDB database has been developed and hosted on Solaris server 71 and is powered by 2.66 GHz Xeon (R) processor with 4 GB FDIMM main 72 memory. The Solaris server was especially selected for its adaptability, 73 scalability and security. The entire data of SPGDB were stored and man- 74 aged in MySQL relational database. The search engine was written using 75 PERL/CGI and PERL/DBI modules. The front-end input data part was 76 coded in HTML, JavaScript and Ajax allows user-friendly web forms. 77 The complete genome of Pneumococcal strains available in NCBI genome 78

http://dx.doi.org/10.1016/j.ygeno.2014.09.012 0888-7543/© 2014 Published by Elsevier Inc.

Corresponding author at: VIT University, Tamil Nadu, India. Fax: +91 416 2243092. E-mail address: aanand@vit.ac.in (A. Anbarasu).

t1.2

t1.3

79

80

81

82

83

**Table 1**The number of occurrences of com-box in each strain of *Pneumococci* identified by 'DNA Motif search tool'

t1.5	Strain name	Number of occurrences
t1.6	Streptococcus pneumoniae 670-6B	12
t1.7	Streptococcus pneumoniae 70585	10
t1.8	Streptococcus pneumoniae A026	9
t1.9	Streptococcus pneumoniae AP200	9
t1.10	Streptococcus pneumoniae ATCC 700669	8
t1.11	Streptococcus pneumoniae CGSP14	13
t1.12	Streptococcus pneumoniae D39	7
t1.13	Streptococcus pneumoniae G54	8
t1.14	Streptococcus pneumoniae gamPNI0373	9
t1.15	Streptococcus pneumoniae Hungary19A-6	9
t1.16	Streptococcus pneumoniae INV104	4
t1.17	Streptococcus pneumoniae INV200	7
t1.18	Streptococcus pneumoniae JJA	7
t1.19	Streptococcus pneumoniae OXC141	6
t1.20	Streptococcus pneumoniae P1031	8
t1.21	Streptococcus pneumoniae R6	7
t1.22	Streptococcus pneumoniae SPN034156	4
t1.23	Streptococcus pneumoniae SPN034183	5
t1.24	Streptococcus pneumoniae SPN994038	5
t1.25	Streptococcus pneumoniae SPN994039	5
t1.26	Streptococcus pneumoniae SPNA45	5
t1.27	Streptococcus pneumoniae ST556	9
t1.28	Streptococcus pneumoniae Taiwan19F-14	8
t1.29	Streptococcus pneumoniae TCH8431/19A	9
t1.30	Streptococcus pneumoniae TIGR4	10

database [9] FTP site were obtained in Genome Feature Format (GFF3) and FASTA format and loaded into GBrowse. The database has been tested on multiple platforms (Windows, Linux and Solaris) with different web browsers. For better view of the database, the user can use recent versions of web-browsers. The database has been thoroughly validated

and produces the results quickly; however, it may vary depending on 84 the user network speed.

86

98

## 3. Complex, user-friendly search options

The SPGDB database provides a powerful and user-friendly search 87 engine. All annotations may be explored utilizing either a simple or 88 advanced Boolean-based search tools. In simple search, the user can 89 browse for different strains, genes and proteins of entire *Pneumococci* 90 by entering strain/gene/protein name in the text box, respectively. The 91 advance search has options to return list of proteins, localizing to a particular chamber. To serve downstream system level analysis, SPGDB 93 enables searching of proteins by COG category and pattern/profile. 94 Further, it facilitates the user to retrieve proteins based on status and 95 virulent genes, which was manually curated from various PUBMED 96 literatures.

#### 4. Facilitating sequence based DNA motif and BLAST searches

The DNA sequence motifs with biological function have become progressively necessary for the analysis of gene regulation [10] and they are 100 found non-randomly in the genome [11]. We have provided a search 101 tool in SPGDB that can be used to identify user-specified DNA motifs 102 like putative transcription factor binding sites and other interesting motifs within the pneumococcal strains. This search tool accepts an IUPAC 104 formatted stretch of DNA sequence with varying lengths and converts 105 the input sequence into a regular expression. Another tool, BLAST [12, 106 13] is also interfaced in SPGDB with which sequence similarity searches 107 can be performed for both protein and nucleotide sequences against a 108 specific or entire pneumococcal strains. The BLAST tool allows users to 109 set parameters like word size, gap open, extension penalty and substitution matrix. The results produced by both DNA motif and BLAST tool can 111

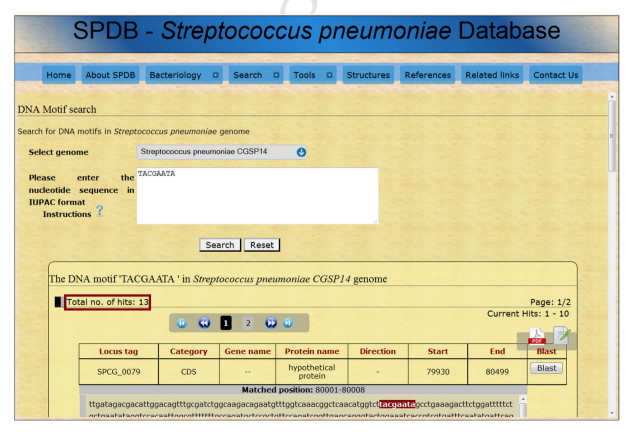
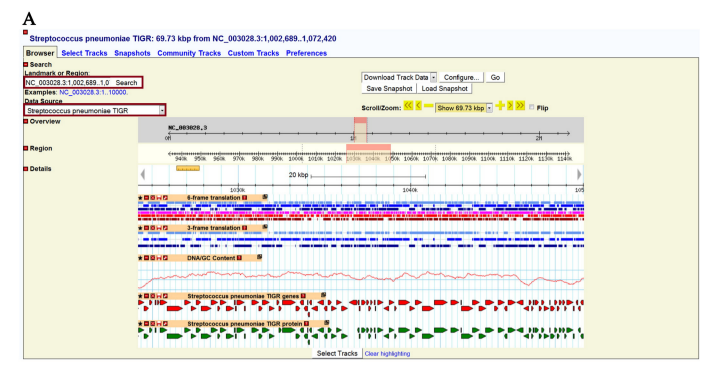


Fig. 1. The results of a com-box (TACGA{2}TA) searched in 'DNA motif search tool' against Streptococcus pneumoniae CGSP14.

125

R. Swetha et al. / Genomics xxx (2014) xxx-xxx





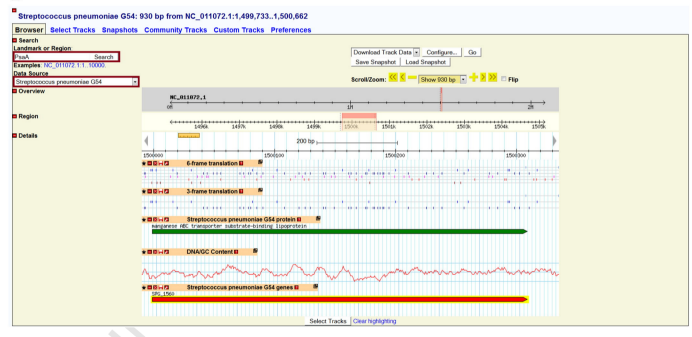


Fig. 2. A. The genes, proteins, GC content, 3-frame translation and 6-frame translation tracks of Streptococcus pneumoniae TIGR from 1,002,689 to 1,072,420 in GBrowse. B: The PsaA gene in Streptococcus pneumoniae G54 visualized in GBrowse.

be stored in the hard disk of a local computer as a text document or in a Portable Document Format (PDF) file.

### 4.1. Case study

112

113

114

115

116

117

118

119

 $120 \\ 121$ 

The com-box (also referred to as cin-box) is a sequence comprising of eight bases, TACGAATA. This sequence acts as a typical binding site for sigma factor, a bacterial transcription initiation factor [14]. The sequence of interest has been searched through the SPGDB DNA motif search tool against all pneumococcal strains and the results are shown in Table 1. The *S. pneumoniae* CGSP14 genome has the highest number of occurrences (13) of com-box compared to all complete genome

of pneumococcal strains (Fig. 1). This is just one example of how integration of this tool can led to new insights through the pneumococcal 123 genome analysis.

# **5.** Genome sequences utilizing GBrowse

In recent years, the genome content of *Pneumococci* has increased 126 drastically. It has to be accessible to researchers for easy interpretation 127 with the help of feasible and interactive viewer. To expedite this, a 128 platform-independent web based application, Generic Genome Browser 129 (GBrowse) has been incorporated in SPGDB. GBrowse was developed by 130 Stein et al. [15] of the Generic Model Organism System Database Project 131

(GMOD). The browser has the features like scroll, navigate and zoom in and out over the random regions of the genome. The user can fetch the region of genome or a landmark by specifying them in a search text box provided at the top left corner of the page. It displays five tracks (i) genes (ii) proteins (iii) GC content (iv) 3-frame translation and (v) 6-frame translation. The landmark on each track carries a link to the corresponding information on SPGDB database or NCBI [9]. Thus, the SPGDB GBrowse makes the end-user to easily view the genomic content of different strains of *Pneumococci*.

#### 5.1. Case study

Among the different pneumococcal strains, capsular serotype 4 clinical isolate of *S. pneumoniae* designated as TIGR4 (TIGR — The Institute of Genome Research) is highly virulent and invasive. The studies reported that the genome comprises of 2,160,837 base pairs with 39.7% of GC content [16]. The genome of this strain is visualized using GBrowse. Fig. 2A shows the various track of genome from the position 1,002,689 to 1,072,420 where the GC content is notably high.

The gene PsaA (Pneumococcal Surface Adhesin) was recognized as a potential adhesin and virulence factor in *Pneumococci*. It is a member of lipoprotein receptor-associated antigen I (LraI) family [17]. In GBrowse,

the PsaA gene (locus tag: SPG\_1560) of *S. pneumoniae G54* is searched 152 and it is found to be positioned from 123,363 to 125,492. It is observed 153 that the transcription of the gene occurs in 5′–3′ direction. In addition, 154 the page displays landmark of its corresponding protein, manganese 155 ABC transporter substrate-binding lipoprotein (Fig. 2B). When the 156 track is zoom out, the adjacent genes are found to be PsaC (locus tag: 157 SPG\_1559) and tpx (locus tag: SPG\_1561).

#### 6. Other utilities of SPGDB

As of August 20, 2014, 464 three dimensional structures of pneumococcal proteins are available in Protein Data Bank (PDB) [8]. Due to in161
creasing number of protein structures day by day, it is also necessary 162
to include these structures in SPGDB database. This can be useful to 163
the scientific community working on *Pneumococci* for the analysis of 164
functionally active proteins. These structures are visualized using the 165
interactive graphics JAVA based plug-in Jmol. Fig. 3 shows an example 166
of Jmol viewer displaying the three-dimensional structure of an enzyme, 167
phosphomevalonate kinase (PDB id: 1K47) [18].

The genome map, a pictorial representation of genomic sequence 169 data and its bioinformatics analysis, are downloaded for available pneu- 170 mococcal strains from the Genome Atlas Database [19]. The option to 171

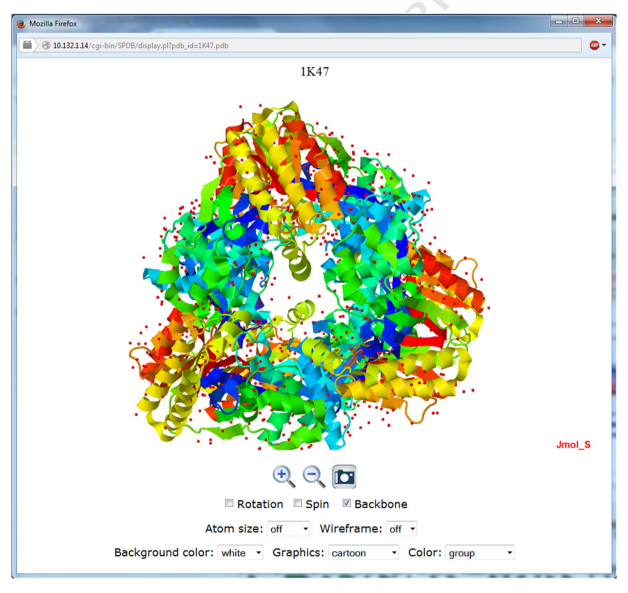


Fig. 3. The Jmol view of a three-dimensional crystal structure of phosphomevalonate kinase (PDB id: 1K47).

R. Swetha et al. / Genomics xxx (2014) xxx-xxx

view genome maps is incorporated in the database under 'Search' 172 menu. Additionally, the information on cultural characteristics, viru-173 lence factors, pathogenesis and laboratory diagnosis of Pneumococci 174 175 has been provided under 'Bacteriology' menu to render preliminary knowledge about the bacterium. The links for different resources related 176 to Pneumococci are provided in the 'Related links' menu. 177

#### 7. Conclusion

178

179

180

181

182

183

184

185

186

187

188

189

190

191

192

193

194

195

196

197 236

In summary, SPGDB database has been developed to deliver complete genomic and proteomic information of Pneumococci to the research community. By providing the dynamic search and browse tools, the database aims to act as not only an integrated resource for Pneumococci, but also as a versatile application platform for the genomic and comparative study of Pneumococci strains. The database will be updated periodically.

#### Acknowledgments

The authors gratefully acknowledge the facilities offered by the Supercomputer Education and Research Centre, Indian Institute of Science, Bangalore. The authors RS, SR and AA gratefully acknowledge the Indian Council of Medical Research (ICMR) for the research grant IRISD: 2014-0099 and also thank the management of VIT University for their support.

#### References

- [1] K.L. O'Brien, et al., Burden of disease caused by Streptococcus pneumoniae in children younger than 5 years: global estimates, Lancet 374 (2009) 893-902.
- T. Barichello, et al., Pathophysiology of acute meningitis caused by Streptococcus pneumoniae and adjunctive therapy approaches, Arq. Neuropsiquiatr. 70 (2012)

[3]	R. Pallares, et al., The epidemiology of antibiotic resistance in Streptococcus	198
	pneumoniae and the clinical relevance of resistance to cephalosporins, macrolides	199
	and quinolones, Int. J. Antimicrob. Agents 22 (2003) S15–S24.	200
F 43		~~~

[4] F. Blasi, et al., Understanding the burden of pneumococcal disease in adults. Clin. 201 Microbiol, Infect, 18 (2012) 7-14.

[5] The Wolrd Health Organization, Measuring Impact of Streptococcus pneumoniae 203 and Haemophilus influenzae Type b Conjugate Vaccination, WHO/IVB/12.0, 2012. 204 (www.who.int/vaccines-documents/) 205

R N Jones et al. Evolving trends in Strentococcus pneumonige resistance: implica- 206 tions for therapy of community-acquired bacterial pneumonia, Int. I. Antimicrob. 207 Agents 36 (2010) 197-204

E.S. Honsa, et al., The roles of transition metals in the physiology and pathogenesis of 209 Streptococcus pneumoniae, Front, Cell. Infect, Microbiol. 3 (2013) 92. 210

[8] H.M. Berman, et al., The Protein Data Bank, Acta Crystallogr. D Biol. Crystallogr. 58 211 (2002) 899-907 212

D.L. Wheeler, et al., Database resources of the National Center for Biotechnology 213 Information, Nucleic Acids Res. 35 (2007) D5-D12.

[10] P. D'haeseleer, What are DNA sequence motifs? Nat. Biotechnol. 24 (2006) 423-425. 215 [11] D. Halpern, et al., Identification of DNA motifs implicated in maintenance of bacterial 216 core genomes by predictive modeling, PLoS Genet. 3 (2007) 1614-1621.

S.F. Altschul, et al., Basic local alignment search tool, J. Mol. Biol. 215 (1990) 403-410. 218 M. Uthayakumar, et al., BSSB: BLAST Server for Structural Biologists, J. Appl. Crystallogr. 219

44 (2011) 651-654 [14] A. Dagkessamanskaia, et al., Interconnection of competence, stress and CiaR regulons 221

in Streptococcus pneumoniae: competence triggers stationary phase autolysis of ciaR 222 mutant cells, Mol. Microbiol. 51 (2004) 1071-1086. [15] L.D. Stein, et al., The generic genome browser: a building block for a model organism 224

system database, Genome Res. 12 (2002) 1599-1610. 225

[16] H. Tettelin, et al., Complete genome sequence of a virulent isolate of Streptococcus  $O_2$ pneumoniae, Science 293 (5529) 498-506. 227

J.W. Johnston, et al., Lipoprotein PsaA in virulence of Streptococcus pneumoniae: 228 surface accessibility and role in protection from superoxide, Infect. Immun. 72 (2004) 229 5858-5867 230

[18] M.J. Romanowski, et al., Crystal structure of the Streptococcus pneumoniae 231 phosphomevalonate kinase, a member of the GHMP kinase superfamily, Proteins 232 47 (2002) 568-571. 233

[19] P.F. Hallin, D.W. Ussery, Genome Atlas Database: a dynamic storage for bioinformatics 234 results and sequence data, Bioinformatics 20 (2004) 3682-3686. 235

202

208

214