



Genome-wide essential gene identification in *Streptococcus sanguinis*

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A clear perception of gene essentiality in bacterial pathogens is pivotal for identifying drug targets to combat emergence of new pathogens and antibiotic-resistant bacteria, for synthetic biology, and for understanding the origins of life. We have constructed a comprehensive set of deletion mutants and systematically identified a clearly defined set of essential genes for *Streptococcus sanguinis*. Our results were confirmed by growing *S. sanguinis* in minimal medium and by double-knockout of paralogous or isozyme genes. Careful examination revealed that these essential genes were associated with only three basic categories of biological functions: maintenance of the cell envelope, energy production, and processing of genetic information. Our finding was subsequently validated in two other pathogenic streptococcal species, *Streptococcus pneumoniae* and *Streptococcus mutans* and in two other gram-positive pathogens, *Bacillus subtilis* and *Staphylococcus aureus*. Our analysis has thus led to a simplified model that permits reliable prediction of gene essentiality.

The search for essential genes has long been a challenge. An essential gene is defined as one whose loss is lethal under a certain environmental condition. The identification of essential genes in bacteria promises to (i) identify critical genes and pathways for controlling pathogenic bacteria by identifying potential targets for antimicrobial drug development¹; (ii) reveal the minimal gene set for living organisms and to shed light on the origin of life^{2,3}; and (iii) reveal bacterial relationships during evolution^{4,5}. Several genome-wide mutant libraries of model microbes have been constructed^{3,6–22}. Although these libraries are invaluable for research in systems biology they show inconsistent essential gene results, even for closely related strains^{11,15,23}. This lack of consensus has prevented reliable prediction of essential genes or pathways in species that have not yet been examined.

To obtain a reliable account of essential genes, we turned to an opportunistic pathogen, *Streptococcus sanguinis* strain SK36, after completing its genome sequence²⁴. The streptococci encompass a large group of important human pathogens. Many *Streptococcus* species are responsible for infectious diseases, such as pneumonia, bacteremia, strep throat, rheumatic fever, scarlet fever, meningitis, infective endocarditis, and dental caries. *S. sanguinis* has long been recognized as a principal causative agent of infective endocarditis²⁵, and its virulence factors have been the subject of a number of investigations^{26–28}. It also initiates biofilm formation on tooth surfaces. The complete genome sequence provides an opportunity to greatly advance our understanding of this organism by enabling the construction of a comprehensive set of genome-wide mutants. *S. sanguinis* is in many ways an ideal candidate for such a study. The SK36 chromosome is 2.39 Mb and contains 2270 putative protein-coding genes, far fewer than that of most microbes used in previous genome-wide gene replacement mutagenesis studies; e.g., *Acinetobacter baylyi* with ~3310 genes²¹, *Bacillus subtilis* with ~4100 genes¹³, *E. coli* with ~4300 genes¹¹, and *S. cerevisiae* with ~6600 genes⁷. More importantly, *S. sanguinis* is highly competent. In our laboratory, up to 20% of *S. sanguinis* cells can be transformed by a simple method using ≤50 ng of DNA. Mutants of non-essential genes are therefore readily obtained, facilitating identification of essential genes. We report here identification of essential genes in *S. sanguinis* and a simplified picture of gene essentiality in streptococci and other bacteria that has emerged from our findings.

Results

Generation of *S. sanguinis* mutants. We set out to identify the essential genes of *S. sanguinis* by systematic gene replacement. Taking advantage of the completed *S. sanguinis* genome, we designed a PCR method (Figure S1A) to



precisely replace target genes with the *aphA-3* kanamycin resistance (Km^r) gene³⁰. The cassette used initially lacked a promoter to avoid potential dysregulation of downstream genes, which could complicate subsequent use of the mutant library in functional studies. To facilitate translation, we provided ribosome binding sites (RBS) for the *aphA-3* and adjacent downstream *S. sanguinis* genes. To ensure efficient homologous recombination, we created long (~1 kb) flanking sequences upstream and downstream of each targeted gene. Flanking sequences were limited to 1 kb based on two considerations: (i) the total length of the amplicon (~3 kb; 1 kb for the *aphA-3* gene plus 2 kb of flanking sequences) allowed ready PCR amplification; and (ii) the feasibility of sequencing flanking regions by Sanger's method from both ends (~1 kb). To test our approach, we randomly selected 10 and then 96 genes for mutagenesis by double cross-over homologous recombination. We successfully created 10 and 93 mutants, respectively, suggesting that the approach was efficient. We then designed and synthesized primers for replacement of 2266 of the 2270 ORFs in *S. sanguinis*, excluding only those four ORFs contained entirely within larger ORFs. In total, we synthesized over 10,000 primers in the construction of the 2266 gene replacement constructs (Table S1). These constructs were created by high-throughput PCR and purification in a 96-well format. Three PCR reactions were required for each replacement and one or more PCR reactions for mutant confirmation (Figure S1A and Figure 1). Over 9,000 PCR fragments were amplified for mutant creation and confirmation. To preclude false identification of genes as essential due to low transformation frequencies, we performed experiments to optimize the transformation efficiency of *S. sanguinis* SK36²⁶. Using our optimized method, up to 2×10^6 mutant colonies could be obtained from 10^7 bacterial cells for non-essential gene transformations.

In the initial round of replacements, we found that many of the genes for which mutants could not be obtained were annotated as

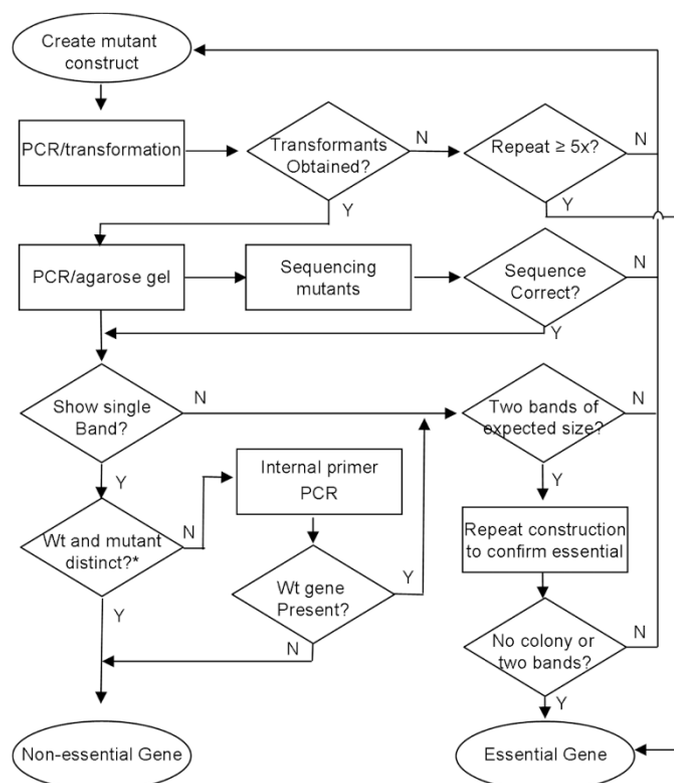


Figure 1 | Flowchart of mutant construction. Procedures for creation and identification of essential genes. Wt, wild type. *Wild-type and mutant band sizes expected to differ by ≥ 100 bp.

acquired via horizontal gene transfer or as encoding hypothetical proteins²⁴. We were curious as to the explanation for this finding. The expression of these genes was therefore examined by microarray analysis. Many had undetectable expression (Table S2). This led us to suspect that many of the unrecoverable mutants might have resulted from insufficient expression of the promoterless *aphA-3* gene. To account for this possibility, we created a second mutagenic construct that retained the native *aphA-3* promoter³⁰, and used it to re-mutagenize the 142 genes that had generated no recoverable mutants (Figures S1). We obtained over 60 additional mutants (Table S3), confirming our hypothesis. Lastly, the Km^r cassette-gene junctions of every mutant were sequenced for confirmation.

Identification of essential genes in *S. sanguinis*. Two types of essential genes were identified in *S. sanguinis*. The first type was genes whose attempted mutagenesis yielded no transformants. New amplicons for these genes were re-amplified and re-transformed in a second cycle. A non-essential gene, SSA_0169²⁸, was used as a positive transformation control to assess whether the failed replacements were due to essential target genes or due to low transformation efficiency of the competent cells. The genes that were not successfully mutagenized after five independent attempts were classified as essential (Figure 1). There were 60 essential genes of this type in *S. sanguinis* (Figure S1B).

For the second type of essential gene, mutant colonies produced double-bands in PCR amplifications using F1 and R3 flanking primers (Figure S1A). The size of one DNA band typically corresponded to the size expected for the replacement mutant while the other matched the wild-type gene. We interpreted this as indicating that these genes were also essential, such that selection for Km resistance resulted in duplication of the target gene²¹. To precisely identify double-band mutants, after sequence confirmation, we examined all PCR amplicons by 1% agarose gel electrophoresis for 4 hrs. Under this agarose gel electrophoresis condition, any amplicons with ≥ 100 bp difference were clearly identified. When bands resulting from amplification of the Km^r cassette and the wild-type gene were anticipated to differ by < 100 bp, an internal T1 primer was designed to determine whether a wild-type gene could be detected by PCR. Of 498 mutants examined with internal T1 primers, 57 produced an amplicon with the same size as in the wild type strain, indicating a “double-band” mutant. (Tables S1 and S3). We identified a total of 158 double-band essential genes (Figure S1B). Our final result was the identification of 218 essential genes including those that we could not mutagenize and those that gave rise to double-bands (Figure S1B, Table S3).

Essential genes associated with specific pathways. The distribution of essential genes in specific categories based on KEGG pathway maps³¹ was examined. The categories of translation, carbohydrate metabolism, nucleotide metabolism, and replication and repair had the greatest number of essential genes, with over 30 each (Figure S2). The categories with the highest percentage of essential genes were translation (76%), transcription (67%), protein folding, sorting and degradation (57%), and glycan biosynthesis (50%).

We next assigned genes to biochemical pathways based on their annotations in KEGG (see Figure 2 for an overview and Table S4 for a detailed analysis). The essential genes were clustered in many pathways: (i) glycolysis, (ii) pentose phosphate pathway (PPP), (iii) peptidoglycan biosynthesis, (iv) terpenoid backbone biosynthesis, (v) glycerophospholipid metabolism, (vi) glycerolipid metabolism, (vii) fatty acid biosynthesis, (viii) nucleotide biosynthesis, (ix) metabolism of cofactors and vitamins including folate biosynthesis, (x) energy metabolism (production of ATP, NADH and NADPH), (xi) DNA replication, (xii) transcription, (xiii) protein biosynthesis, (xiv) GTP-binding proteins and (xv) cell division.

These results were in keeping with expectation, given our current understanding of bacterial metabolism. Among the metabolic

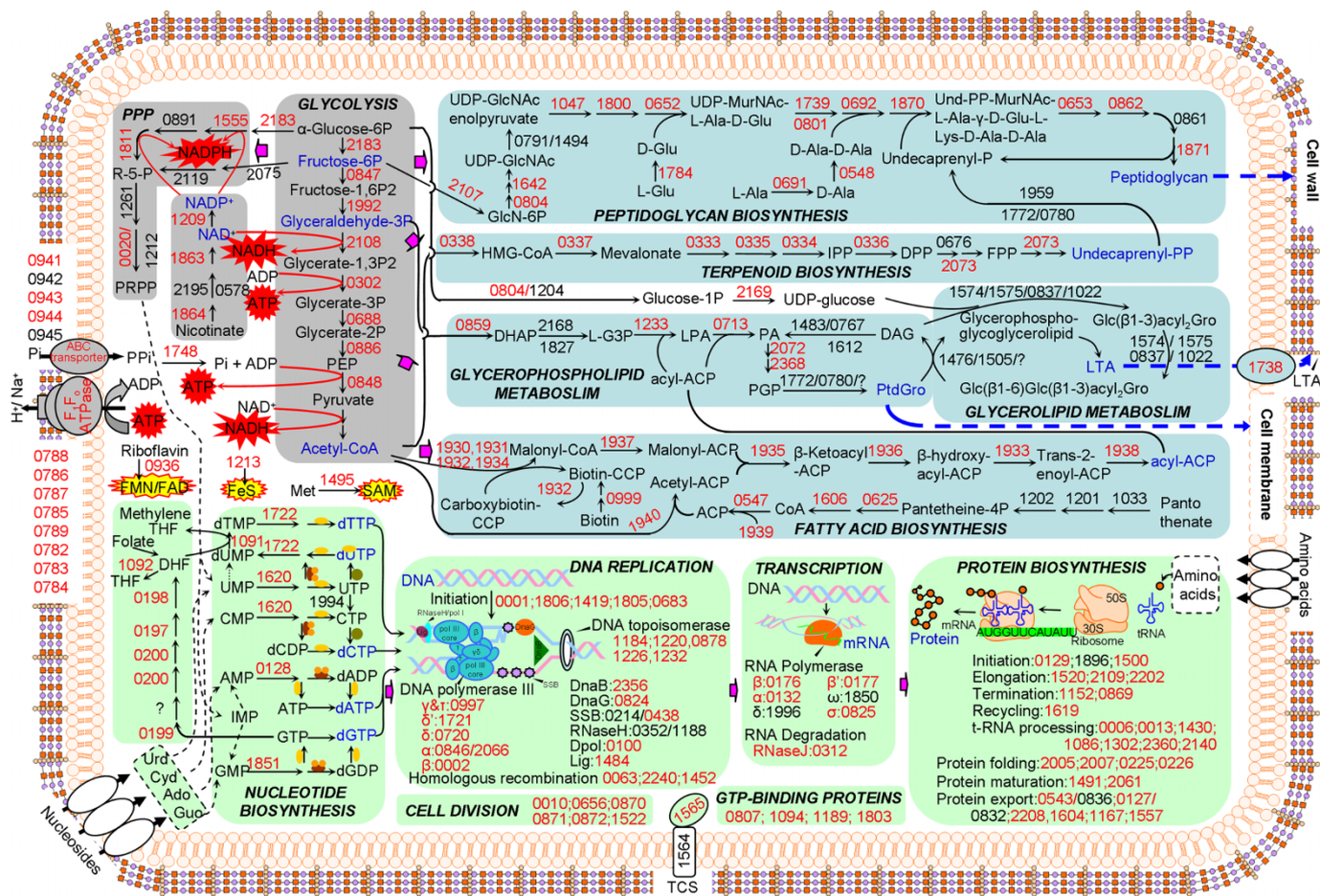


Figure 2 | Concise pathways of *S. sanguinis* SK36 essential genes. The three functions associated with essential pathways are indicated by color: pale blue, cell envelope; gray, energy production; light green, processing of genetic information. Numbers are SSA#; red number, essential gene; black number, nonessential gene; solid arrow, an enzymatic reaction; dashed arrow, multistep pathway; blue dashed arrow, products involved in cell wall and membrane formation; block arrow, product from one pathway serving as input to another pathway; oval with bold arrow, transporter; slash, separating paralogs. Ado, adenosine; Cyt, cytidine; DAG, 1,2-diacylglycerol; DHAP, dihydroxyacetone phosphate; DHF, dihydrofolate; L-G3P, sn-glycerol 3-phosphate; Guo, guanosine; LPA, lysophosphatidic acid; LTA, lipoteichoic acid; PA, phosphatidic acid; PGP, phosphatidylglycerophosphate; PPP, pentose phosphate pathway; TCS, two-component system; THF, tetrahydrofolate; Urd, uridine. In nucleotide biosynthesis, symbols are used to represent essential genes that participate in multistep reactions: SSA_1865 (green circle), SSA_0848 (yellow oval), SSA_2263, SSA_0771 (orange circles) and SSA_0768 and SSA_0770 (brown oval). Essential genes associated with ribosome and aminoacyl-tRNA biosynthesis are not indicated here, nor are SSA_0575, SSA_0800 and SSA_1903, which could not be categorized into these pathways.

pathways containing essential genes, glycolysis and pyruvate oxidation from α -D-glucose 6-phosphate to acetyl-CoA play a pivotal role because they generate energy and provide input molecules for other essential metabolic pathways. Terpenoid backbone biosynthesis feeds into the pathway for synthesis of peptidoglycan, which is the main component of the cell wall. Glycerophospholipid metabolism in conjunction with glycerolipid produces lipoteichoic acid (LTA), a component of the cell wall in gram-positive bacteria. Phospholipids biosynthesized from glycerophospholipid metabolism and fatty acid biosynthesis compose the cell membrane. The nucleotides synthesized from PRPP derived from the pentose phosphate pathway (PPP) or from nucleosides imported from the growth medium supply materials for DNA replication and transcription. Several vitamins and cofactors including folate, biotin, pantothenate, nicotinate, riboflavin, SAM and FeS clusters are involved in the pathways above. ATP and NADH/NADPH produced from glycolysis, pyruvate oxidation and PPP provide energy.

We then analyzed essential genes by network functions using systems biology. All essential genes and their related functions were linked together in pathways and studied as a whole. This picture was simplified dramatically when it became apparent that all of these

pathways could be linked to three basic biological functions: maintenance of the cell envelope, energy production and processing of genetic information. Remarkably, we found only three essential genes (none of which have been assigned exact functions) that could not be linked to these three functions (SSA_0575, SSA_0800, and SSA_1903; Table S4). SSA_0575 is annotated as a haloacid dehalogenase-like hydrolase, SSA_0800 as a glutamine amidotransferase and SSA_1903 as a conserved hypothetical protein.

Identification of additional essential genes in minimal medium.

From our analysis, we found that the three functions of maintenance of the cell envelope, energy production and processing of genetic information, and no others, are essential to *S. sanguinis*. Based on the genome sequence, we predicted that *S. sanguinis* possesses *de novo* synthesis pathways for all amino acids and nucleotides from glycolysis and PPP (Figure 3A). We identified 96 genes responsible for biosyntheses of 19 amino acids starting from glycolysis, leaving only the undetermined gene responsible for L-asparagine biosynthesis from L-aspartate (Figure 3A). Two pathways each were present for biosyntheses of L-glutamate, L-serine and L-glycine and L-threonine. Yet, in our initial screen, we found few essential genes related to

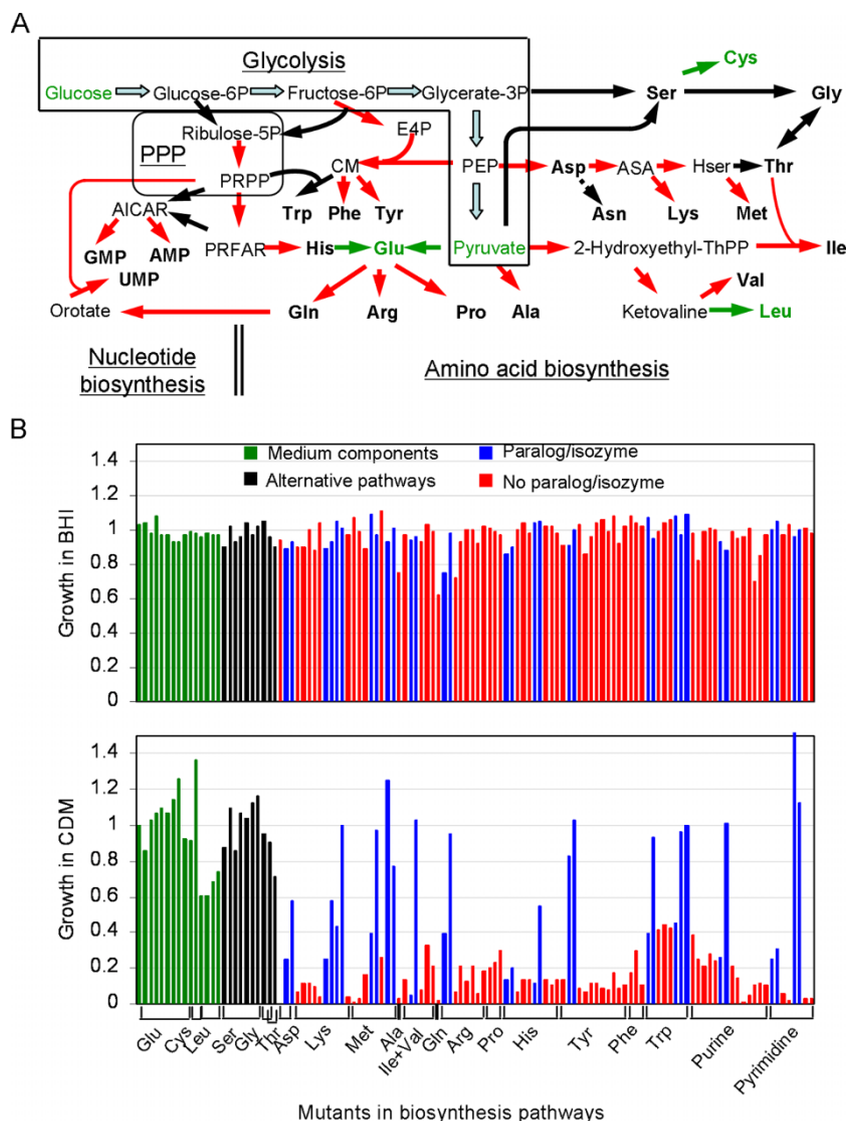


Figure 3 | Growth of *S. sanguinis* mutants related to amino acid and nucleotide biosyntheses in BHI and CDM media. (A) amino acid and nucleotide biosynthesis pathways in which mutants grew (black arrow) or did not grow (red arrow) in CDM medium. Components of the CDM medium are indicated in green. PPP, pentose phosphate pathway, dashed arrow, undetermined gene responsible for asparagine biosynthesis. Note the two pathways for serine, glycine, threonine, glutamate and AICAR biosyntheses. ASA, L-aspartate 4-semialdehyde; CM, chorismate; E4P, D-Erythrose 4-phosphate; Hser, homoserine or o-phospho-L-homoserine; PRFAR, phosphoribuloyl-formimino-AICAR-phosphate. (B) growth of mutants in BHI and CDM media. Relative growth was represented by mutant $OD_{450} / SK36 OD_{450}$ after 2d anaerobic incubation.

amino acid synthesis or nucleotide synthesis from PRPP to UMP/GMP/AMP, suggesting amino acids and nucleotide precursors were provided by the rich brain heart infusion (BHI) medium. To examine this possibility, we compared the growth of strains with mutations in genes responsible for amino acid and nucleotide *de novo* synthesis in a chemically defined medium (CDM) with that in BHI, as described previously²⁷ (Figure 3B; Table S5). CDM contains only 3 amino acids (L-glutamate, L-cysteine and L-leucine) and lacks nucleosides and nucleobases. All of these mutants grew to levels similar to SK36 in BHI medium (Figure 3B). However, the growth of the mutants was significantly lower than that of SK36 in CDM medium, with the following exceptions: (i) the mutants whose deleted genes were involved in the synthesis of the amino acids that were supplied in CDM medium (Glu, Cys and Leu); (ii) the mutants possessing alternative pathways (Ser, Gly and Thr); and (iii) the mutants whose replaced genes had paralogs or isozymes. Although dramatically reduced, some of the mutants still grew to low levels, perhaps benefitting from carry-over of nutrients from the inocula. Similar results were obtained with 23 mutants associated with nucleotide

biosynthesis from PRPP and/or L-glutamine (Figure 3B). The combined results suggest that *S. sanguinis* does indeed obtain required amino acids and nucleotide precursors from the BHI medium. Provision of nutrients by BHI appears to explain another result that initially appeared discrepant with our model. We predicted that SSA_1201, SSA_1202 and SSA_1033 would be essential because they were required for biosynthesis of CoA from pantothenate, but mutants of these three genes were obtained (Figure 2). We found, however, that these mutants exhibited minimal or undetectable growth in CDM, suggesting pantetheine-4P is provided by BHI (Table S5).

Examination of paralogs and isozymes with double-knockout mutants. In our analysis, we also found that some genes predicted to be critical in one of these three functions were not identified as essential. We considered that their functions may be performed by genes encoding paralogs or isozymes with similar functions. To examine this possibility, we identified gene paralogs based on protein sequence homology (Table S3). We also examined the genome

Table 1 | Double-gene knockouts in *S. sanguinis*.

Double mutant	Donor	Product Name	Acceptor	CFU	Gel	Essential
Ssx_0791::1494	SSA_0791	UDP-N-acetylglucosamine 1-carboxyvinyltransferase	Ssx_1494	0		yes
Ssx_0578::2195	SSA_0578	nicotinic acid mononucleotide adenyltransferase	Ssx_2195	0		yes
Ssx_0352::1188	SSA_0352	ribonuclease HIII	Ssx_1188	0		yes
Ssx_2168::1827	SSA_2168	NAD(P)H-dependent glycerol-3-phosphate dehydrogenase	Ssx_1827	>5000	1-band	No
Ssx_0169::1494	SSA_0169	hypothetical protein	Ssx_1494	>5000	1-band	No
Ssx_0169::2195	SSA_0169	Donor Product Name	Ssx_2195	>5000	1-band	No
Ssx_0169::1188	SSA_0169	hypothetical protein	Ssx_1188	>5000	1-band	No
Ssx_0169::1827	SSA_0169	hypothetical protein	Ssx_1827	>5000	1-band	No

All mutants with double crossover replacement are named Ssx. The number indicates the replaced gene.

annotation and literature for potential isozymes. When we re-examined the linked essential genes in the network, we found paralogs or isozymes in every case in which a series of linked essential genes was interrupted by a “non-essential” gene in the same pathway (Figure 2; for example, paralogs SSA_0791/SSA_1494 involved in peptidoglycan biosynthesis, isozymes SSA_0578/SSA_2195 involved in NAD⁺/NADP⁺ biosynthesis). This suggested that these non-essential genes might substitute for one another in performing essential functions. To test this hypothesis, we selected four pairs of paralogous or isozyme genes for double-gene knockouts to examine essentiality: SSA_0791/SSA_1494, SSA_1827/SSA_2168, SSA_0578/SSA_2195, and SSA_0352/SSA_1188. As a control, double mutants were created by combining the replacement of each gene of interest with a replacement of the SSA_0169 gene, which is a hypothetical gene with no known function²⁸. As we anticipated, double mutants could not be constructed for the gene pairs SSA_0791/SSA_1494, SSA_0578/SSA_2195, or SSA_0352/SSA_1188, whereas SSA_0169 double mutants were readily obtained (Table 1). Unexpectedly, the double mutant SSA_1827/SSA_2168 was viable. This may result from the existence of an alternative pathway (composed of SSA_0049/SSA_0050, SSA_0287 and SSA_1826) in glycerolipid metabolism.

Identification of essential genes in other streptococcal species. To validate the hypothesis that essential genes are associated with these three functions, we examined representative genes in two more pathogenic streptococcal species. We first selected 8 genes in *S. pneumoniae* strain TIGR4 that were not previously identified as essential in either of two *S. pneumoniae* genome-wide essential gene screens^{10,17}. These 8 genes are predicted to be involved in maintenance of the cell envelope or in energy production (SP_0382, SP_0383, SP_0384 and SP_0262, involved in terpenoid backbone biosynthesis; SP_1511, SP_1513 and SP_1514 encoding F₁F_o-ATPase subunits; and SP_0261 involved in glycerophospholipid metabolism). We also selected one gene (SP_0489) as a control that encodes a protein unrelated to these three essential functions whose ortholog was non-essential in *S. sanguinis*. The *S. pneumoniae* genes were

mutagenized and then characterized in the same manner as for *S. sanguinis*. As shown in Table 2, none of the eight genes could be mutagenized, although the control produced numerous transformants. The process was repeated for another pathogen, *S. mutans* strain UA159. The orthologs of the above 8 genes in *S. mutans* were selected and examined for gene essentiality, yielding identical results (Table 2). Again, all tested genes were identified as essential in *S. mutans*. These results suggested the accuracy of our essential gene predictions for other streptococci.

We used comparative genomics³² to examine the conservation among streptococcal species of the genes we identified as essential. We downloaded 49 complete streptococcal genomes publicly available from the NCBI database and compared them with the *S. sanguinis* essential genes. The vast majority of *S. sanguinis* essential genes (202 of 218) had orthologs in all of the other 48 streptococcal genomes (Table S6A). Thus, in agreement with expectation, most of the genes we identified as essential were highly conserved within streptococci.

Comparison with essential genes in other gram-positive bacteria.

We then compared essential genes of *S. sanguinis* with those in other species that have been examined experimentally. All essential genes of the 13 bacterial species presently included in the Database of Essential Gene (DEG)³³ were collected and searched against the *S. sanguinis* protein database by BLASTP to find homologs (Table S7). The sequence comparison suggested that there were many differences. In the other species, about half (33 to 66%) of the essential genes with homologs in *S. sanguinis* matched *S. sanguinis* essential genes, with the rest matching non-essential *S. sanguinis* genes.

To find the reason for this inconsistency, we analyzed more carefully the other two gram-positive species in the database, *Staphylococcus aureus* NCTC8325 and *B. subtilis* 168 (Figure 4 and Table S8), that were subjected to genome-wide screens^{13,23} in similar nutrient-rich media (BHI and LB). Some essential genes in *S. sanguinis* had homologs identified as non-essential in *S. aureus* and

Table 2 | Confirmation of essential genes in *S. pneumoniae* and *S. mutans*.

SP_ID*	Product Name	CFU [†]	Gel [‡]	Essential	SMU_ID*	CFU	Gel [‡]	Essential
SP_0382	diphosphomevalonate decarboxylase	0		yes	SMU.937	0		yes
SP_0383	phosphomevalonate kinase	0		yes	SMU.938	0		yes
SP_0384	isopentenyl pyrophosphate isomerase	0		yes	SMU.939	0		yes
SP_1514	FOF1 ATP synthase subunit C	2	2-band	yes	SMU.1534	1	2-band	yes
SP_1513	FOF1 ATP synthase subunit A	2	2-band	yes	SMU.1533	0		yes
SP_1511	FOF1 ATP synthase subunit delta	6	2-band	yes	SMU.1531	1	2-band	yes
SP_0262	phosphatidate cytidyltransferase	0		yes	SMU.1785	0		yes
SP_0261	undecaprenyl pyrophosphate synthase	0		yes	SMU.1786	1	2-band	yes
SP_0489	PAP2 family protein	242	1-band	no	SMU.1702c	874	1-band	no

*Gene identifiers. IDs in the same row indicate orthologous genes

[†]Number of transformants obtained from attempt to mutagenize the gene indicated

[‡]Appearance of PCR products resulting from amplification of the target gene locus from selected transformants

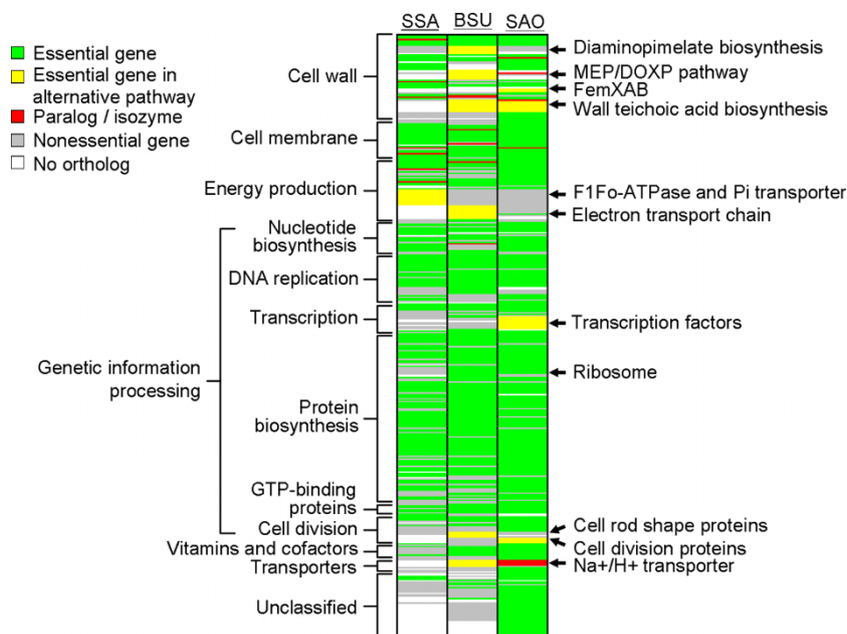


Figure 4 | Essential gene comparisons among *S. sanguinis*, *S. aureus* and *B. subtilis*. Essential gene comparisons were based on sequence homologs, and genes were grouped according to pathways or functions. SSA, *S. sanguinis* SK36 genes; BSU, *B. subtilis* 168 genes; SAO, *S. aureus* NCTC8325 genes. Colors represent gene categorizations, as indicated in the figure.

B. subtilis, and vice versa although many essential genes in *S. sanguinis* were shared with these two species. However, we found that most of the differences could be explained by gene paralogs, isozymes, or alternative pathways. As shown in Figure 4, *B. subtilis* synthesizes peptidoglycan from UDP-N-acetylmuramoyl-L-alanyl-D-glutamate via an alternative pathway using meso-2,6-diaminopimelate rather than L-lysine that results in the genes responsible for producing meso-2, 6-diaminopimelate from L-aspartate 4-semialdehyde via lysine biosynthesis pathway being essential. *S. aureus* cross-links peptidoglycan via the essential *femXAB* operon rather than the streptococcal *murMN*. The essential mevalonate pathway in the terpenoid backbone pathway in *S. sanguinis* and *S. aureus* is replaced by an essential MEP/DOXP pathway in *B. subtilis*. In *S. sanguinis*, LTA and wall teichoic acid (WTA, a component of cell wall in gram-positive bacteria) are likely synthesized by the same set of enzymes because both have identical components in *S. sanguinis*³⁴ and repeating units with identical structures in *S. pneumoniae*³⁵, whereas *S. aureus* and *B. subtilis* use a set of enzymes different from LTA synthesis to produce wall teichoic acid, and the genes encoding them are essential. *S. sanguinis* lacks a respiratory chain and undergoes fermentation to produce acids such as lactate and acetate, and the F_1F_0 -ATPase uses ATP hydrolysis to pump intracellular protons out. This is indispensable for creating a protonmotive force for a variety of transport processes and to maintain pH homeostasis, which likely results in the genes encoding F_1F_0 -ATPase being essential in *S. sanguinis*. In contrast, *S. aureus* and *B. subtilis* use the electron transport chain to generate protonmotive force, and the genes associated with its biosynthesis (i.e. menaquinone) are essential. *B. subtilis* and *S. aureus* likely maintain cellular pH homeostasis via a Na^+/H^+ transporter, the genes for which are essential in *B. subtilis* and paralogous in *S. aureus*. *S. sanguinis* lacks this transporter. In genetic information processing, the set of essential genes in *S. sanguinis* is smaller than those in other two species, but their essential pathways are still identical.

Discussion

Accurate prediction of essential genes is important for identification of drug targets to combat the emergence of pathogens and antibiotic-resistant bacteria²⁰, especially for serious infectious agents for which

there is no research model system available. In addition, the rational design of bacterial cells through synthetic biology, as is currently possible, requires an understanding of the minimal gene set for further advances^{3,36}. However, the lack of consensus among experimental essential gene lists has prevented prediction of essential genes in other species. In our essential gene comparisons, we found significant differences of essential genes identified for *S. sanguinis* and those identified in previous studies with other species. We suggest several possible explanations for these differences. (i) We would expect some different genes due to genetic inheritance among species. For example, the essential genes for cell wall composition in gram-positive bacteria are different from those in gram-negative bacteria. (ii) Identified essential genes using single-gene knockouts will vary when the bacteria being compared differ in protein paralogs, isozymes or alternative pathways for essential functions. These cases can be tested by creating double-gene knockouts, as we have demonstrated (Table 1). (iii) Differences in screening conditions may have caused differences in essential gene identification, although the most favorable environmental conditions for each bacterium were used in most screens. As we found for *S. sanguinis* above, the use of a minimal medium obviously results in the identification of additional essential genes. (iv) False-negative results may have been obtained in some cases due to partial gene inactivation, which is known to occur frequently in random insertion mutagenesis³⁷. (v) Some genes may have been falsely identified as essential because of genetic system limitations. Many bacterial species have low transformation frequencies. In these cases, even a moderate reduction in plating efficiency may result in lack of mutant recovery, resulting in categorization of the gene as “essential” in a large-scale, genome-wide mutagenesis. (vi) Finally, insufficient expression of the selection marker, as occurred with our promoterless Km^r gene, may result in false-positive identification of essential genes.

Many *in silico* methods have also been established to predict essential genes. These methods include ortholog identification²³, genomic intrinsic feature analysis³⁸, gene evolution rate³⁹, phylogenetic conservation⁴⁰, network analysis⁴¹ and machine learning based integrative approaches^{38,42}. Of these, essential gene prediction via phyletic conservation is the most commonly used. Our results indicate that the vast majority of *S. sanguinis* essential genes are indeed conserved



in closely related species (Table S6A). We also found that essential genes had greater sequence conservation among the streptococci than non-essential genes. The average identity for the essential gene protein sequences was 79.71% while for the non-essential gene protein sequences, it was 58.96% (Table S6B). It should be noted, however, that phyletic conservation alone was a poor predictor of gene essentiality. We found 787 non-essential *S. sanguinis* genes with orthologs in all 48 streptococcal genomes. Inclusion of bacteria from other genera in our analysis would have reduced the number of conserved genes identified, likely increasing the specificity of the phyletic method for essential gene prediction. However, the sensitivity of this method then decreases, as we found a number of essential *S. sanguinis* genes with orthologs that can be detected in different genera by identity of annotated function, but not by sequence similarity (data not shown).

Using our findings, we believe that most essential genes in other bacteria can now be predicted based on their genome annotations. We have established a model of essential pathways (Figure 5), in which the essential genes are linked by crucial chemical compounds. Although essential genes may differ among species due to different final products or alternative pathways, we propose they will contribute to the three basic functions of maintenance of the cell envelope, energy production, and processing of genetic information. We used this model for predicting essentiality of paralogs and isozymes by double-gene knockouts in *S. sanguinis* and for predicting essential genes in *S. mutans*, *S. pneumoniae*, *B. subtilis* and *S. aureus*. Our predictions were largely confirmed.

This model can also be used to explain many of the apparent inconsistencies observed previously among different organisms that were not predictable based on gene sequence conservation alone. Bacteria can be categorized into gram-negatives, gram-positives

and Mycobacteria due to the differences in their cell envelope compositions. The greatest difference is that gram-negative bacteria contain an outer membrane with lipopolysaccharides while LTA and WTA are found in gram-positive bacteria and mycolyl-arabinogalactan-peptidoglycan complex in Mycobacteria. This leads to different sets of essential genes responsible for the respective envelope components. In the gram-negative bacteria *E. coli*¹⁸ and *Pseudomonas aeruginosa*¹⁹, the genes responsible for lipopolysaccharide biosynthesis have been demonstrated essential. Many genes responsible for biosynthesis of mycolyl-arabinogalactan-peptidoglycan complex were identified as being essential in *Mycobacterium tuberculosis*¹⁴. In *B. subtilis* and *S. aureus*^{13,23}, the genes responsible for the biosynthesis of WTA and LTA were essential. Our results indicated most *S. sanguinis* orthologs of genes responsible for LTA biosynthesis in *S. aureus* were nonessential. Although this can be explained by the existence of paralogs (Figure 2), we cannot exclude the possibility of an alternative biosynthetic pathway for *S. sanguinis* LTA synthesis, as glycerophosphate residues in the LTA of *S. sanguinis* DSM 20567 may be substituted with D-alanine ester, α -D-glucosyl and α -isomalto-oligosaccharide residues⁴³.

Most bacteria possess a respiratory chain that is important for energy production and maintenance of redox balance. It has been demonstrated that the genes involved in synthesis of electron transport chain components, such as CoQ, menaquinone or heme, are essential in *E. coli*¹⁸, *P. aeruginosa*¹⁹, *B. subtilis* and *S. aureus*^{13,23}. Although an important function of electron transport is generation of ATP via the F_1F_0 -ATPase, there was only one instance of an F_1F_0 -ATPase gene being identified as essential in previous studies performed with the latter two species^{9,13,23}. One possibility is that these genes were missed. Most replacements of F_1F_0 -ATPase genes in *S. sanguinis*, *S. pneumoniae*, and *S. mutans* generated “double-band”

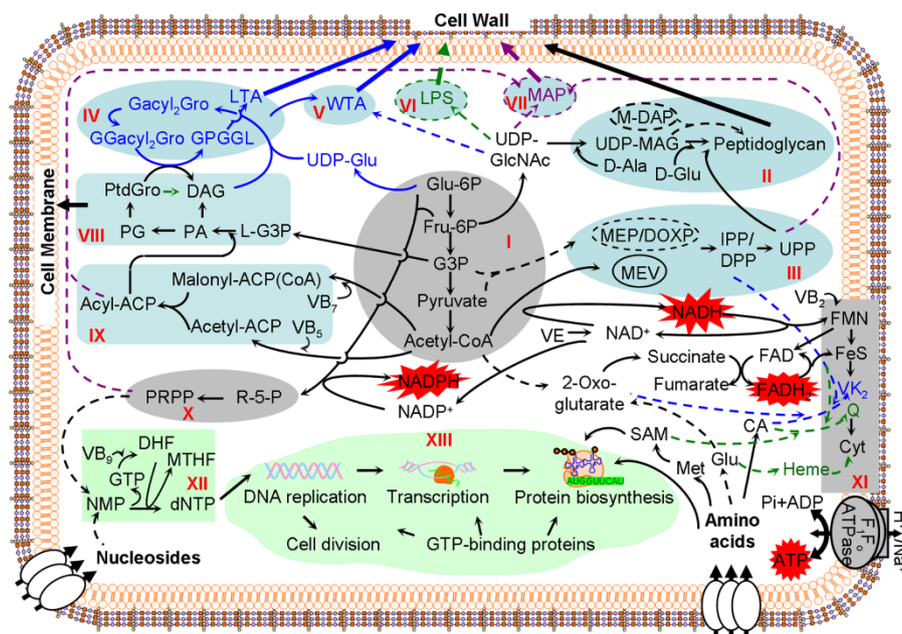


Figure 5 | Deduced essential pathways based on *S. sanguinis* essential genes. Colors indicate the three functions associated with essential pathways, as in Figure 2. I, glycolysis and pyruvate oxidation; II, peptidoglycan biosynthesis; III, terpenoid backbone biosynthesis; IV, glycerolipid metabolism (LTA biosynthesis); V, WTA biosynthesis; VI, lipopolysaccharide biosynthesis; VII, MAP biosynthesis; VIII, phosphoglycerolipid metabolism; IX, fatty acid biosynthesis; X, electron transport chain; XI, pentose phosphate pathway; XII, nucleotide biosynthesis; XIII, genetic information processing including DNA replication, transcription, protein biosynthesis, GTP-binding proteins and cell division. Black arrow, all bacteria; black dashed arrow or oval, alternative pathways; blue arrow, gram-positive bacteria only; green arrow, gram-negative bacteria only; violet arrow, mycobacteria only; bold arrow, synthesis of cell envelope. CA, chorismate; Cyt, cytochrome; DAG, 1,2-diacylglycerol; DG-DAG, diglucosyl-diacylglycerol; G3P, D-glyceraldehyde 3-phosphate; GPGGL, glycerophosphoglycerolipid; L-G3P, glycerol-3-phosphate; LPS, lipopolysaccharide; Malonyl-ACP(CoA), malonyl-ACP or malonyl-CoA; M-DAP, meso-2,6-diaminopimelate; MEV, mevalonate; MG-DAG, monoglucosyl-diacylglycerol; PA, phosphatidate; PG, phosphatidylglycerophosphate; R-5-P, D-Ribulose 5-phosphate; PPP, pentose phosphate pathway; PtdGro, phosphatidylglycerol; UDP-MAG, UDP-N-acetylmuramoyl-L-alanyl-D-glutamate; UPP, undecaprenyl diphosphate.



mutants (Table 2; Table S3). This would lead to identifying F_1F_0 -ATPase genes as nonessential in studies in which the presence of double-bands was not investigated. Nevertheless, the ability of these species to grow anaerobically indicates that ATP generation by F_1F_0 -ATPase is not essential, suggesting that electron transport is required for other purposes, such as secondary active transport. In *S. sanguinis*, all eight subunits of the F_1F_0 -ATPase were identified as essential. Since streptococci lack a respiratory chain, the essential function of the F_1F_0 -ATPase in these species is also not generation of ATP and is, instead, likely to be export of protons using energy from ATP hydrolysis. A previous analysis of essential genes in *S. pneumoniae* identified five of the eight F_1F_0 -ATPase subunit genes as essential^{10,17}. Here, we demonstrated the other three genes encoding F_1F_0 -ATPase components were also essential in *S. pneumoniae*, as were their orthologs in *S. mutans* (Table 2).

In conclusion, by choosing an ideal test organism and by employing exhaustive measures to avoid false-positive and false-negative identifications, we have reliably identified the essential genes of *S. sanguinis*. The validity of our findings is suggested by the virtually perfect association between our list of essential genes and the list of genes expected to be required for the three functions of cell envelope maintenance, energy production, and processing of genetic information. Although *S. sanguinis* is important in its own right, the relative ease with which these results can be used to identify essential genes in other prominent streptococcal pathogens including *S. pyogenes* and *S. pneumoniae* lends increased importance to this work. Moreover, our study suggests that with minimal additional effort, our results can be used to predict essential genes for most bacteria for which an annotated genome sequence is available.

It should be noted that although we examined all protein coding regions for their essentiality, it is well known that some non-coding regions are also essential if they contain DNA sequences for important biological functions. Such sequences include the chromosomal origin of replication, promoters, tRNAs, rRNAs and perhaps small RNAs.

Finally, although the focus of the current study was the identification of essential genes, the study has also produced an ordered, comprehensive library of non-essential gene mutants of *S. sanguinis* SK36. The design of the mutagenic constructs to (i) ensure near-complete deletion of each gene; (ii) retain expression signals for adjacent genes; and (iii) introduce a promoter only in cases where it was required, in combination with the care with which each mutant was characterized ensures that the library will be of great value. We are currently employing the mutants for a number of investigations of gene function involving conventional and systems biology approaches.

Methods

Bacterial strains and growth. *S. sanguinis* strain SK36 was grown at 37°C in BHI broth (BD) as described previously²⁷. *S. pneumoniae* TIGR4 and *S. mutans* UA159 strains purchased from ATCC were grown in Todd Hewitt (TH; BD) broth plus 0.5% yeast extract and in BHI broth under microaerobic conditions (6% O₂, 7.2% CO₂, 7.2% H₂ and 79.6% N₂).

Primer design and PCR. We developed a recombinant PCR method for in vitro creation of linear constructs for the replacement of every protein-coding gene in the *S. sanguinis* SK36 genome (Figure S1A). Based on the complete *S. sanguinis* SK36 genome sequence²⁴, three sets of primers (F1/R1, F2/R2 and F3/R3) were designed to amplify the *S. sanguinis* sequence upstream from each targeted gene, the *aphA-3* gene, encoding Km^{r45} and the *S. sanguinis* sequence downstream from each targeted gene, respectively.

For most of the mutagenized genes, the R1 and F3 primers were designed to delete the coding region from 6 bp after the start codon to 30 bp before the stop codon. Stop codons were inserted in all three frames to prevent fusion of the N-terminus of the targeted open reading frame with the Km^r protein. The last 30 bp were retained to preserve potential ribosomal binding sites used by adjacent downstream genes. The upstream retained region was extended from 6 bp to 100 bp when two neighboring genes were located head-to-head in opposite orientation to prevent deletion of potential promoters for flanking genes. Primers R1 and F3 contained 25-nt sequences that are complementary with the antibiotic selection cassette at their 5' ends. The P1,

P2, and various T1 primers were designed for sequencing to confirm mutants. The sequence of every primer is documented in Table S1.

Three PCR amplicons were created using F1/R1, F2/R2 and F3/R3. All PCR reactions were performed at 94°C for 1 min, and 30 cycles of 94°C for 30 sec, 54°C for 30 sec and 68°C for 1.5 min. After DNA purification by PureLink 96 PCR purification kits (Invitrogen), the three PCR amplicons were combined in equal amounts in one tube and amplified again using the F1 and R3 primers to obtain the final linear recombinant PCR amplicon. Conditions were 94°C for 2 min, 30 cycles of 94°C for 30 sec, 55°C for 30 sec and 68°C for 3.5 min, and finally 68°C for 4 min. High-fidelity Platinum[®] Taq DNA polymerase (Invitrogen) was used in all reactions.

Promoterless and promoter-containing cassette construction. We initially created a promoterless Km^r cassette⁴⁶ to eliminate possible polar transcriptional effects on neighboring genes. To address instances of poor expression of the *aphA-3* gene, we created a second construct in which the native promoter of *aphA-3* gene³⁰ was included in the Km^r cassette. Apart from the design of new R1_Promoter and F2 primers to include the promoter, all other construction steps were the same as for the promoterless constructs.

Gene replacement and mutant storage. The above linear PCR amplicons (~50 ng) were directly transformed into *S. sanguinis* as described previously²⁶. Allelic exchange mutants generated by double cross-over homologous recombination were selected by two-day microaerobic incubation on BHI agar plates containing 500 µg/ml kanamycin. For each replacement mutant, one colony was randomly picked and cultured in BHI with Km. To determine whether the mutant contained the expected gene replacement, colony PCR was performed using F1 and R3 primers. The PCR amplicon was examined by 20-cm long agarose gel electrophoresis. The amplicon was further confirmed by Applied Biosystems Big Dye terminator DNA sequencing. The sequencing confirmation was performed using the P1 primer, which binds to the Km^r cassette (Figure S1A). The mutants which gave rise to a DNA amplicon of the expected size by long gel electrophoresis and with the expected junction sequence determined from the P1 primer or the T1 primer were collected as the final confirmed mutants. These were retained and cryopreserved in 30% glycerol at -80 °C.

The methods for construction of gene replacement amplicons with the promoter-containing Km^r cassette for *S. mutans* UA159 and *S. pneumoniae* TIGR4 were the same as for *S. sanguinis*. Transformation of *S. mutans* UA159 was conducted similarly to that of *S. sanguinis* except *S. mutans* CSP was used. Transformation of *S. pneumoniae* TIGR4 was performed as described by Bricker and Camilli⁴⁷.

Microarray analysis. *S. sanguinis* cultures at late log phase were used for microarray analysis. RNA from each of three independent samples was isolated by RNeasy mini kit (Qiagen, Valencia, CA). Spotted microarray slides were obtained from the Pathogen Functional Genomics Resource Center at JCVI. The microarray was performed according to the manufacturer's protocol. Each sample was divided into two parts and labeled separately by Cy5 and Cy3 dyes for microarray. The microarray data were analyzed using the programs Spotfinder and Midas to obtain the expression ratio of each gene labeled with each dye. All ratios were within the range of 0.6 to 1.5, indicating consistency in the labeling and analysis. Additionally, the ratio of dye intensity to background in a microarray slide was obtained after Spotfinder analysis. Absolute expression of each gene was represented by the average ratio of dye intensity to background for each slide. The microarray data have been deposited in the NCBI Gene Expression Omnibus (GEO) with record GSE25340.

Growth comparison of mutants in CDM and BHI media. Selected mutants were cultured overnight at 37°C in 96-well blocks with 1 ml BHI broth under anaerobic conditions (10% CO₂, 10% H₂ and 80% N₂ with a palladium catalyst). The overnight cultures were inoculated into 1 ml either CDM²⁷ or BHI medium in 96-well plates by dipping with multichannel tips. The inocula were incubated for 2 d under the same conditions as above. Cultures were then mixed by pipetting several times with a multichannel pipette, and 200 µl cultures were transferred into 96-well plates for measuring OD₄₅₀. The relative growth of each mutant was calculated as mutant OD₄₅₀/SK36 OD₄₅₀.

Gene function analysis. The pathway distributions of essential genes were analyzed via KEGG. KEGG annotations were downloaded to a local computer from /pub/kegg/genes/organisms/ssa/. The assigned KO numbers and path numbers of *S. sanguinis* genes were extracted. Multiple KO numbers or path numbers from KEGG were often assigned for a single gene if it was involved in different pathways. To view the essential pathways, these genes were assigned to the pathway with the greatest percentage of essential genes. To study essential genes as a whole, as many essential genes as possible were linked together via pathways. Based on a product in one pathway being the substrate of another pathway, the essential pathways were linked together and then integrated into their possible functions.

Comparative genomics. Comparative genomic analyses were performed to identify conserved proteins in *S. sanguinis* as previously described²⁴. We downloaded 48 other completed streptococcal genomes and their annotations from public databases. *S. sanguinis* proteins were compared to other streptococcal protein databases by BLASTP. Significant matches ($E < 1e-5$) were analyzed to find homologs in other streptococcal genomes. The *S. sanguinis* database was also BLASTP searched against itself to identify paralogs. Protein sequence conservation was calculated by percent amino acid identity of orthologs. Essential genes in *Staphylococcus aureus* NCTC



8325, *Bacillus subtilis* 168 and *S. pneumoniae* were obtained from the Database of Essential Genes (DEG)³³. The *S. sanguinis* orthologs were identified using BLASTP.

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Author contributions

PX conceived the main idea and the conceptual design of the experiments. PX, XG, LC, XW, YD, JX, JP, VS, MT and KE constructed mutants; PX designed primers and performed bioinformatics analysis; XG established the high throughput transformation system and performed microarray and gene functional analysis; XW performed double-knockout and mutant growth comparison; PX, XG, LC, JX, TK and DB analyzed the data. PX, XG, TK, DB and GB wrote the manuscript.

Additional information

Supplementary information accompanies this paper at <http://www.nature.com/scientificreports>

Competing financial interests: The authors declare no competing financial interests.

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