## The Gamma-Hemolysin Locus of *Staphylococcus aureus* Comprises Three Linked Genes, Two of Which Are Identical to the Genes for the F and S Components of Leukocidin

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The Staphylococcus aureus gamma-hemolysin comprises two polypeptides, whereas the gamma-hemolysin locus (hlg) contains three open reading frames. The hlgA and hlgB genes encode the  $\gamma 1$  and  $\gamma 2$  components, respectively. The HlgB protein ( $\gamma 2$ ) has 27% residue identity with S. aureus alpha-toxin. Surprisingly, hlgB and hlgC are 98.5 and 99.1% identical to the lukF and lukS genes, respectively, encoding the F and S components of the Panton-Valentine leukocidin.

Staphylococcus aureus can express five cytolytic toxins, the alpha-, beta-, delta-, and gamma-toxins and a toxin that acts on leukocytes, the Panton-Valentine leukocidin (9, 14). The leukocidin is composed of two proteins, the F and S components (10). There is firm evidence that gamma-toxin is also composed of two proteins (1, 5, 6, 13). We previously reported the cloning and analysis of the S. aureus gammatoxin locus (2). Nonhemolytic transposon insertion mutants fell into two groups on the basis of in vitro complementation tests. The corresponding genes hlgA and hlgB expressed proteins with molecular weights of 32,000 and 36,000, respectively, in minicells.

(A preliminary report of part of this work has been presented previously [4].)

In this report we present the DNA sequence of the hlg locus. Deletions in hlg plasmids were generated with BAL 31. Sequencing reactions on plasmid templates employed T7 polymerase. Sequence analysis and data base searches were performed with the University of Wisconsin Genetics Computer Group package (3) on a VAX computer. For amplification by the polymerase chain reaction, standard reaction conditions (7) for 30 cycles and AmpliTaq polymerase were used. The assays for gamma-hemolysin, polyacrylamide gel electrophoresis, and immunoblotting were as described previously (2). Antiserum to the  $\gamma 1$  component of gamma-hemolysin (1) was donated by M. Clyne, and antileukocidin serum was obtained from C. Adlam.

The 3,797-bp sequence of the *hlg* locus was determined from pJC01 and pJC08 (2). Surprisingly, three open reading frames, designated *hlgA*, *hlgB*, and *hlgC*, were identified. Each is preceded by a potential ribosome binding site and encodes a protein with a putative signal sequence. The mature forms of the HlgA, HlgB, and HlgC proteins have molecular weights of 31,925, 34,123, and 32,551, respectively, and pI values of 9.43, 9.08, and 9.01. The position of the *hlgB* gene corresponds to the region that encodes the HlgB protein (2). The putative ribosome binding site lies

within an open reading frame, designated *hlgC*, upstream from *hlgB*. The *hlgC* gene has a potential ribosome binding site and also a promoter (Fig. 1). It is likely that these two genes are cotranscribed. The *hlgA* gene spans a region defined by several transposon insertions. A putative promoter and a ribosome binding site are located 5' to the coding sequence, and a transcriptional termination sequence occurs 3' to the coding sequence. Thus, *hlgA* is likely to be monocistronic.

Beginning at residue 30 of the translation product of hlgA, a sequence of amino acids that corresponds exactly to the amino-terminal sequence of the y1 component of gammatoxin (1) was identified. This demonstrates unambiguously that  $\gamma 1$  is specified by hlgA. The HlgA ( $\gamma 1$ ) and HlgC ( $\gamma 2$ ) proteins have 70% residue identity, and each has approximately 30% identity with HlgB (Fig. 2). Additionally, there is 27% residue identity between HlgB and S. aureus alphatoxin, and a lower level of similarity was noted between alpha-toxin and either HlgA or HlgC (Fig. 2). Surprisingly, the hlgB and hlgC genes were found to be 98.5 and 99.1% identical to the recently published sequences of the leukocidin F and S components, respectively (Fig. 1). The luk genes were cloned by using oligonucleotide probes derived from amino-terminal sequences of the purified F and S components (11, 12). Furthermore, antileukocidin serum inhibited the gamma-hemolytic activity expressed by strains PG23 (a toxin shock syndrome isolate [1]) and Smith 5R in agarose diffusion tests (data not shown). The reported pI values of 9.08 and 9.39 for the leukocidin F and S components, respectively, are similar to the values predicted from the hlgB and hlgC sequences presented here.

The three hlg open reading frames were cloned separately into plasmid vectors. The hlgB gene was cloned on a Sau3A fragment into pK18, forming plasmid pHLGB. The hlgA and hlgC genes were cloned after specific amplification by polymerase chain reaction primers that incorporated BamHI recognition sites, and the recombinant plasmids were verified by sequence determination. The amplified products were cloned in pK18, forming pHLGA and pHLGC, respectively. Lysates of cells harboring pHLGA contained a  $32,000-M_r$  protein that reacted with the anti- $\gamma 1$  serum in Western immunoblots (data not shown). No immunoreactive

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TGTCTCTTCAATACATGTTGATAGTAATTAACTTTTAACGAACAGTTAA<u>TTCGAA</u>AACGCTTACAAATGGAT<u>TATTAT</u>ATATATAGAACTTAAAATTAAATAGAAAGAAAGTGATTTCTAT -35 -10 121 GATTAAAAATAAATATTAACAGCAACTTTAGCAGTTGGTTTAATAGCCCCTTTAGCCAATCCATTTATAGAAATTTCTAAAGCAGAAAATAAGATAAGAAGATATCGGCCAAGGTGCAGA 241 361 481 TAATTATCTTCCTAAAAAATTGATTCAGCAGATGTTAGTCAGAAATTAGGCTATAATATCGGCGGAAACTTCCAATCAGCGCCATCAATCGGAGGCAGTGGCTCATCAACTACTC 601 721 ATATGATCAATACTTATTTGCACAAGACCCAACTGGTCCAGCAGCACGAGACTATTTCGTCCCAGATAATCAACTACCTCCTTTAATTCAAAGTGGCTTTAATCCATCATTTATTACAAC 841 ATTGTCACACGAAAGAGGTAAAAGGTGATAAAAGCGAGTTTGAAATCACTTACGGCAGAAACATGGATGCTACATATGCTTACGTGACAAGACATCGTTTAGCCGTTGATAGAAAACATGA 961 TGCTTTTAAAAACCGAAACGTTACAGTTAAATATGAAGTGAACTGGAAAACACATGAAGTAAAAAATTAAAAGCATCACACCTAAGTAAACAGTTACAATCATCTTAAAAAAACCTGGGACA STOP hlgA terminator 1081  $\tt CTTCATACT\underline{TGTCTCAGGATTTTTTAACAATTGAATCAGCCTCATAACATTAAATTATTTTTTTCGTACATTAAATTAATAATAATAACAACTGATTTTTTTATAAGAATAAAGTATCGAACCA$ 1321 TGTATTTTACAAACATTAATTAAAAAAAGAAAGCAAGACATTCGTGCAATCCGTTACCTTAAATTGTTTACCAACAATACCAAGGTTTTATTATTATTATTCTCACAAAATTAG 1441 AGTTGATAA<u>TGGTCA</u>ATTGTTAATGAACAGTTAAT<u>TATAAT</u>AACGCCCAA -10 1561 RBS hlgC→ 1681 TTATTAGAAAATGCTAAAGCTGCTAACGATACTGAAGACATCGGTAAAGGAAGCGATATAGAAATTATCAAAAGGACAGAAGATAAAAACAAGTAATAAATGGGGCGTGACTCAAAATATT 1801 2041 AATATCGGTGGTAATTTCCAATCAGCCCCATCACTCGGTGGTAATGGATCATTTAACTATTCTAAATCGGATTAGCTATACACAAAATTATGTAAGTGAAGTAGAACAAAACTAA 2161 2281 2401 2521 AATTGGAAGACTCATGAAATCAAGGTGAAAGGA Gaattgatatgaaa<mark>at</mark>gaataaattagtcaaa RBS? STOP hlgC hlgB 2641 TGAAGGTAAAATAACACCAGTCAGCGTAAAAAAAGTCGATGACAAAGTTACTTTATACAA 2881 TAAATACAATGTATCTATAAGCTCACAATCTAATGATTCAGTAAACGTCGTTGATTATGCACCAAAAAATCAAAATGAAGAGTTTCAAAATACTTTAGGCTATACATTTGGTGG 3001 TGACATTAGTATCTCTAATGGTTTATCTGGTGGACTTAATGGAAATACAGC 3121 AAATGTTGGCTGGGGAGTTGAAGCACATAAAATTATGAATAATGGTTGGGGACCTTATGGAAGAGATAGCTTCCACCCAACATATGGTAATGAACTCTTCTTAGCTGGCAGACAAAGCAG 3241 CAGA TTCTACTGGGCAGGCGCAAATTATAAAAACTTTA 3481 STOP hlaB terminator GTAATACTGCCATTCTTTTCTCAATGTGAGATATAAAGGAATAGCTACAATTAAAGTGAATATTĀGCCTGGAATCGCGTTTAACAACACTCCCACACAGGTAAATTAAAATAATAGT

FIG. 1. Nucleotide sequence of the gamma-hemolysin locus of S. aureus Smith 5R. Possible promoter and terminator sites and ribosome binding sites (RBS) are indicated. The translation initiation codon and termination codon for each open reading frame are in boldface type. Divergences between hlgC and lukS and between hlgB and lukF are indicated by the residues present in these positions in the luk genes shown below the hlg sequence. Residues of hlg not present in the luk sequence are represented by hyphens.

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Hla 1 HlgA 1 HlgB 1 HlgC 1	ADSDINIKTGTTDIGSNTTVKTGDLVTYDKENGMHKKVFYSFIDDKNAIKKLLVIRTKGTIAGQYRVYSEEGANKSGLAWPSAFKVQLQLPDNEV ENKIEDIGQGAEIIKRTQDITSKRLAITQNIQFDFVKDKKYNKDALVVKMQGFISSRTTYSDLKKYPY-IKRMIWPFQYNISLKT-KDSN AEGKITPVSVKKVDDKVTLYKTTATADSDKFKISQLITFNFIKDKSYDKDTLVLKATGNINSGFVKPNPNDYDFSKLYMGAKYNVSISQSNDS ANDTEDIGKGSDIEIIKRTEDKTSNKWGVTQNIQFDFVKDKYNKDALLLKMQGFISSRTTYYNYKKTNH-VKAMRWPFQYNIGLKT-NDKY . * * * * * * * * * * * * * * *
Hla 96 HlgA 89 HlgB 95 HlgC 91	AEISDYYPRNSIDTKEYMSTLTYGFNGNUTGDDTGKIGGLIGANUSIGHTLKYVQPDFKTILESPTD-KKUGWKUIFNNMUNQNWGPYDRDSWNPUYGNQ UDLINYLPKNKIDSADVSQKLGYNIGGNFQSAPSIGGSGSFNYSKTISYNQKNYVTEVESQNS-KGVKWGVKANSFVTPNGQVSAYDQY VNVUDYAPKNQNEEFQVQNTLGYTFGGDISISN-GLSGGLNG-NTAFSETINYKQESYRTTLSRNTNYKNUGWGYEAHKIMNNGWGPYGRDSFHPTYGNE VSLINYLPKNKIESTNYSETLGYNIGGNFQSAPSLGGNGSFNYSKSISYTQQNYVSEVEQQNS-KSVLWGVKANSFATESGQKSAFDSD  * * * * * * * * * * * * * * * * * * *
Hla 191 HlgA 178 HlgB 194 HlgC 180	LFMKTRNGSMKAADNFLDPNKASSLLSSGFSPDFATVITMDRKASKQQTNIDVIYERVRDD-YQLHWTSTNWKGTNTKD-KWTDRSSERYKIDWEK LFAQ-DPTGPAARDYFVPDNQLPPLIQSGFNPSFITTLS-HERGKGDKSEFEITYGRNMDATYAYVTRHRLAVDRKHDAFKNRNVTVKYEVNWKT LFLAGRQSSAYAGQNFIAQHQMPLLSRSNFNPEFLSVLS-HRQDGAKKSKITVTYQREMDL-YQIRWNGFYWAGANYKN-FKTRTFKSTYEIDWEN LFVGYKPHSKDPRDYFVPDSELPPLVQSGFNPSFIATVS-HEKGSSDTSEFEITYGRNMDVTHAIKRSTHYGNSYLDGHRVHNAFVNRNYTVKYEVNWKT **
Hla 286 HlgA 272 HlgB 288 HlgC 280	EEMTN HEVKIKSITPK HKVKLLDTKETENNK HEIKVKGQN

FIG. 2. Comparison of the amino acid sequences of the processed forms of Hla (S. aureus alpha-toxin), HlgA, HlgB, and HlgC. The multiple alignment was produced by the Clustal V program (8) with the default parameters. Identical residues are indicated by asterisks, and conserved substitutions are indicated by dots. Spaces have been introduced to maximize alignment.

product was detected in extracts of cells harboring pHLGB or pHLGC with antisera against two purified components of gamma-hemolysin (1), possibly because of the low level of expression. Whole-cell lysates of Escherichia coli containing these plasmids were tested for hemolytic activity in diffusion tests in rabbit erythrocyte agarose plates. None was hemolytic alone, but the pHLGB extract acted synergistically with pHLGA or pHLGC samples to lyse the erythrocytes (Fig. 3). Hemolysis produced between pHLGA and pHLGC was slower to develop and resulted in a zone of lysis that was more opaque than that formed between pHLGB and pHLGA lysates. Dextran and dextran sulfate inhibited both synergistic reactions. This inhibition is a characteristic of gamma-hemolysis (9). In addition, hemolysis did not occur in agar incorporating the erythrocytes, presumably because of inhibition by the sulfonated polymers present.

All available evidence suggests that gamma-hemolysin and leukocidin are encoded by the same locus. The relative proportions of the polypeptide components in culture supernatants could be crucial in determining the type of cytolytic activity expressed by a particular strain. It is not yet known whether the strain from which the leukocidin genes were cloned has a functional *hlgA* gene and expresses gamma-hemolysin as well as leukocidin. Similarly, it is not known

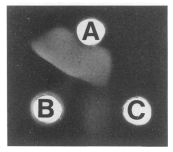


FIG. 3. Synergistic hemolysis between components of gamma-hemolysin. Extracts of *E. coli* cells harboring pHLGA (well A), pHLGB (well B), or pHLGC (well C) were applied to wells in agarose plates containing rabbit blood.

whether gamma-toxin-producing strains also express leukocidin activity.

The nucleotide sequence reported herein has been submitted to GenBank under accession number L01055.

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