Function of the central domain of streptokinase in substrate plasminogen docking and processing revealed by site-directed mutagenesis

ANITA CHAUDHARY, 1, 3 S. VASUDHA, 1 K. RAJAGOPAL, 1 SNEHA SUDHA KOMATH, NANDITA GARG, 2 M. YADAV, SHEKHAR C. MANDE, AND GIRISH SAHNI

Institute of Microbial Technology, Sector 39-A, Chandigarh-160036, India

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Abstract
The possible role of the central β-domain (residues 151–287) of streptokinase (SK) was probed by site-specifically altering two charged residues at a time to alanines in a region (residues 230–290) previously identified by Peptide Walking to play a key role in plasminogen (PG) activation. These mutants were then screened for altered ability to activate equimolar “partner” human PG, or altered interaction with substrate PG resulting in an overall compromised capability for substrate PG processing. Of the eight initial alanine-linker mutants of SK, one mutant, viz. SKKK256,257AA (SK-D1), showed a roughly 20-fold reduction in PG activator activity in comparison to wild-type SK expressed in Escherichia coli (nSK). Five other mutants were as active as nSK, with two [SKRE248,249AA and SKKK251,252AA, referred to as SK(C) and SK(H), respectively] showing specific activities approximately one-half and two-thirds, respectively, that of nSK. Unlike SK(C) and SK(H), however, SK(D1) showed an extended initial delay in the kinetics of PG activation. These features were drastically accentuated when the charges on the two Lys residues at positions 256 and 257 of nSK were reversed, to obtain SKKK256,257EE [SK(D2)]. This mutant showed a PG activator activity approximately 10-fold less than that of SK(D1). Remarkably, inclusion of small amounts of human plasmin (PN) in the PG activation reactions of SK(D2) resulted in a dramatic, PN dose-dependent rejuvenation of its PG activation capability, indicating that it required pre-existing PN to form a functional activator since it could not effect active site exposure in partner PG on its own, a conclusion further confirmed by its inability to show a “burst” of p-nitrophenol release in the presence of equimolar human PG and p-nitrophenyl guanidino benzoate. The steady-state kinetic parameters for HPG activation of its 1:1 complex with human PN revealed that although it could form a highly functional activator once “supplied” with a mature active site, the $K_m$ for PG was increased nearly eightfold in comparison to that of nSK-PN. SK mutants carrying simultaneous two- and three-site charge-cluster alterations, viz., SKKK248,249AA and SKKK251,252AA [SK(CH)], SKKK251,252AA [SK(FH)], and SKKK248,249AA,FKKK272,273AA [SK(CFH)], showed additive/synergistic influence of multiple charge-cluster mutations on HPG activation when compared to the respective “single-site” mutants, with the “triple-site” mutant [SK(CFH)] showing absolutely no detectable HPG activation ability. Nevertheless, like the other constructs, the double- and triple-charge cluster mutants retained a native like affinity for complexation with partner PG. Their overall structure also, as judged by far-ultraviolet circular dichroism, was closely similar to that of nSK. These results provide the first experimental evidence for a direct assistance by the SK β-domain in the docking and processing of substrate PG by the activator complex, a facet not readily evident probably because of the flexibility of this domain in the recent X-ray crystal structure of the SK-plasmin light chain complex.

Keywords: plasminogen; plasminogen activation mechanism; site-specific mutagenesis; streptokinase; substrate plasminogen

Reprint requests to: Girish Sahni, Institute of Microbial Technology, Sector 39-A, Chandigarh, 160036 India; e-mail: sahni@imtech.ernet.in.

1These authors have contributed equally in this work.
2Present address: Torrent Pharmaceuticals Ltd., Gandhinagar, Ahmedabad, India.
3Visiting Fellow of the Department of Biotechnology, Government of India on deputation from the Department of Environmental Sciences, Indian Agricultural Research Institute, Pusa, New Delhi, India.

Abbreviations: SK, Streptococcus equisimilis streptokinase; PG, plasminogen; PN, plasmin; HPG, human plasminogen; HPN, human plasminogen; BSA, bovine serum albumin; CD, circular dichroism; SD, standard deviation; TPA, tissue plasminogen activator; NPG, p-nitrophenyl p-guanidinobenzoate; IPTG, isopropyl-1-thio-β-D-galactopyranoside; ORF, open reading frame; PBS, phosphate buffered saline; PCR, polymerase chain reaction; UK, urokinase.

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Plasminogen activators such as streptokinase, tissue plasminogen activator, and urokinase are widely used as thrombolytic agents for the treatment of diverse circulatory disorders, including myocardial infarction (ISIS-3, 1992). However, unlike UK and TPA, which are proteases, SK is enzymatically inert. It acts by first forming a noncovalent, high-affinity complex with HPG. This complex is then believed to undergo an intramolecular conformational change that exposes the active site in the zymogen. The active site within SK-HPG now becomes capable of acylation and can be titrated by hydrolysis of p-nitrophenyl p-guanidinobenzoate (p-NPBG) to p-nitrophenol as a sharp “burst” (McCintock & Bell, 1971), or catalyze the processing of small-MW amidolytic substrates. This enzymatically active, “virgin” complex also rapidly develops a capability to catalyze the specific transformation of substrate molecules of HPG to HPN by cleavage at the Arg561–Val562 peptide bond (reviewed by Castellino, 1981). An understanding of the structural processes involved in, first, the exposure of the active site in the virgin SK-HPG complex, followed by its transformation to a highly specific protease (either “activated” SK-HPG, or SK-HPN) that, unlike free plasmin, displays a very high substrate preference for HPG (Markus & Werkheiser, 1964), is crucial to the design of improved SK-based thrombolytic agents (Marder, 1993).

SK has been shown to be composed of three distinct domains, designated α, β, and γ separated by two coiled coils, and small regions at the N- and C-termini of the protein with disordered, flexible structures (Conejero-Lara et al., 1996; Parrado et al., 1996; Wang et al., 1998). While the regions at the two ends are known to be dispensable for the biological activity of SK, selective deletions of α or γ domains, or parts thereof, lead to drastic reduction in HPG activation ability, signifying that all three domains are vital for the functioning of the molecule (Malke et al., 1987; Fay & Bokka, 1998). However, the exact manner in which each domain contributes toward the high rates of PG activation characteristic of full-length SK remains to be clarified. Although it has been demonstrated that the three domains retain significant amounts of their original native-like structure when individually isolated, none displays significant HPG activator activity on its own (Parrado et al., 1996). In contrast to the absence of any appreciable HPG activator activity in isolated domains, the characteristic high-affinity binding of SK with HPG is relatively well preserved in the β-domain, and to a lesser degree, in fragments derived from the α-domain (Nihalani & Sahni, 1995; Reed et al., 1995; Rodriguez et al., 1995; Conejero-Lara et al., 1996, 1998; Nihalani et al., 1998). More recently, using physico-chemical approaches, it has been elegantly demonstrated that the β-domain contributes in a major way toward the generation of the high-affinity interactions between SK and HPG that lead to the formation of the equimolar activator complex between the two proteins (Conejero-Lara et al., 1998). A relatively small locus in the β-domain (residues in and around 230–290, termed the “core” region) has earlier been implicated in the interaction of the activator complex with substrate HPG through a Peptide Walking approach (Nihalani et al., 1997). Therefore, the exact role of this domain, particularly in terms of the loci/epitopes that may be involved in interacting with HPG to translate the primary event in SK-HPG interaction, namely the avid binding between these two proteins to the catalytic, high efficiency conversion of substrate molecules of HPG to HPN by the activator complex needs to be explored further.

Despite the compelling biochemical evidence implicating the β-domain in both activator complex formation and in the interaction of this complex with substrate HPG, the recent X-ray diffracton structure of the SK-plasmin light chain complex does not reveal an overt structural involvement of this domain in activator complex formation (Wang et al., 1998). However, what is fairly discernible is that SK and PG together (i.e., the activator complex) provide a perfect docking site for substrate PG to be positioned optimally for processing by the active site. Computer modeling studies with substrate microplasminogen artificially docked onto the SK-plasmin light chain complex, indeed, suggest several potential contacts between the substrate and the α- and β-domains of the SK of the activator complex. This observation thus reinforces a proposed model of SK action (Nihalani et al., 1998) similar to the “protein co-factor assisted” mechanism recently forwarded to explain the switch of the relatively nonspecific, trypsin-like substrate specificity of the plasmin active site into a highly specific substrate preference upon association with SK (Esposin & Mather, 1998). In the present paper, using site-directed mutagenesis as a tool, we present experimental evidence for an important role of the β-domain in substrate docking and processing by the SK-plasmin(ojen) activator complex, and probably in zymogen activation as well.

**Results**

Construction, purification, and functional characterization of alanine-linker charge-cluster mutants of the “core” region of the beta-domain of SK

The sequence of residues 230–290 of SK from *Streptococcus equisimilis* (termed the “core” region) that has been previously implicated in the interaction of SK with HPG by Peptide Walking studies (Nihalani et al., 1997) is depicted in Figure 1. This region has earlier been implicated in the interaction of SK with HPG by Peptide Walking studies (Nihalani et al., 1997) is depicted in Figure 1. This region

![Fig. 1. Primary structure of the target sequence for mutagenesis in the beta-domain of SK. The amino acid sequence of the residues 230–290 of the beta-domain of SK from S. equisimilis H46A is shown. The charged cluster residues that were mutated to alanines in this study are indicated in italics. The sites whose charges were reversed by mutation are underlined.](image-url)
of primary structure of SK shows several positively and negatively charged clustered residues. To delineate the possible role of these side chains, site-specific mutations of eight such clustered charges (two residues per site) to alanine residues were carried out using a PCR-based “megaprimmer” approach (Sarkar & Sommer, 1990; Smith & Klugman, 1997). The mutated cassettes were then docked back into the native SK gene in the expression plasmid, and the genes expressed intracellularly in Escherichia coli and purified to homogeneity (see Table 1 for sequence of primers used for mutagenesis, and Materials and methods for details). The native-like recombinant SK (termed nSK) expressed in the heterologous host, after purification, was found to exhibit several of the known characteristic features of native SK from S. equisimilis, e.g., a MW of 47 kD, native N-terminal sequence except for the presence of a Met residue at the N-terminus (as expected from the gene sequence), high-affinity binding with HPG, and a specific activity for HPG activation of around 1.0 × 10^3 IU per mg protein. In case of all the mutants constructed (Table 2), these criteria were essentially found to be native like, except for reduction, to varying extents, in the specific activities for HPG activation of selected mutants. For simplification, the various SK mutants described in this study have been abbreviated with different suffixes, e.g., SK(C), SK(D1), etc. (see Table 2). Of all the eight initial alanine-linker mutants, only one (SKKK256.257AA), referred to as SK(D1), showed a major diminution in PG activation ability, to a level approximately 5% that of nSK. Two other mutants, viz., SKKK244.246AA and SKKK281.282AA, abbreviated as SK(C) and SK(H), respectively, showed relatively marginal decreases in specific activity for HPG activation, while the remaining were found to be totally unaltered. Remarkably, all the mutants, including those with lowered activity, retained near-native HPG binding affinities as measured by a solid phase radio-assay employing 125I-labeled HPG.

The mutant SK(D1) displayed kinetics of HPG activation that were distinctly different on a qualitative basis when compared to the other mutants, viz., SK(C) and SK(H), that had detectable decrease in activity as well. In the case of SK(D1), the kinetics of PG activation (Fig. 2A) was noticeably slower, especially in the initial phases. The progress curves generated with SK(D1) demonstrated a perceptible lag even when relatively high concentrations of the mutant were used in the assay, while nSK demonstrated a clear-cut absence of such a lag even when assayed using low amounts (Fig. 2A). Mutants SK(C) and SK(H) also showed the absence of any lag. The overall specific activity of SK(D1) for HPG activation (computed from plots of change in absorbance at 405 nm vs. t^6; see Materials and methods) was determined to be

<table>
<thead>
<tr>
<th>Charge-cluster mutation</th>
<th>Oligonucleotide sequence</th>
<th>R.E. site</th>
</tr>
</thead>
<tbody>
<tr>
<td>DE 238,240 AA</td>
<td>5′-ATTTTACcATGGGGcAAAGcGTACTTACC-3′</td>
<td>Nco I</td>
</tr>
<tr>
<td>RK 244,246 AA</td>
<td>5′-AAGGGTTTACgTACgtGTAeAAACGGAAC-3′</td>
<td>Bsi W1</td>
</tr>
<tr>
<td>RE 248,249 AA</td>
<td>5′-CGTGTTAAAAAAATGtGcCAAGTCTTATGAT-3′</td>
<td>Fsp I</td>
</tr>
<tr>
<td>KK 256,257 AA</td>
<td>5′-GCTTATAGAAATATGCcAAGcATCGTGGTCTGAAATGAA-3′</td>
<td>Vsp I</td>
</tr>
<tr>
<td>KK 256,257 EE</td>
<td>5′-AGGATCAATgAAATIGTCTC6iAAATGAAGAAATA-3′</td>
<td>Bsm AI</td>
</tr>
<tr>
<td>EE 262,263 KK</td>
<td>5′-AAAAATCCTGGTCAATGcGcAAATAAACACT-3′</td>
<td>Alw 26I</td>
</tr>
<tr>
<td>EE 262,263 AA</td>
<td>5′-ATAAAAATCTCGGgC6TcAGAcGcAAATAAACACT-3′</td>
<td>Bsp 126I</td>
</tr>
<tr>
<td>EK 272,273 AA</td>
<td>5′-ACCCTGATCCTGAcegG6TcATCGTCCCT-3′</td>
<td>Pst I</td>
</tr>
<tr>
<td>KK 278,279 AA</td>
<td>5′-AGAAATATTACGTCCTGcATcGc6AcGcAAACTG-3′</td>
<td>Bsa AI</td>
</tr>
<tr>
<td>EE 281,282 AA</td>
<td>5′-AAAAAGGAtgGcGGGCGTGATGCcC-3′</td>
<td>Nar I</td>
</tr>
<tr>
<td>Upstream primer</td>
<td>5′-ATTATGAACGTGACTCCTCAATCGTC-3′</td>
<td>Bse RI</td>
</tr>
<tr>
<td>Downstream primer</td>
<td>5′-ATAGGCTAAATGAATgTGACATCCTCCTCC-3′</td>
<td>Bsm I</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Type of SK</th>
<th>Short name</th>
<th>Specific activity (× 10^3 IU/mg)</th>
<th>Dissociation constant [Kd] of SK-HPG complex (nM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild-type SK</td>
<td>nSK</td>
<td>1.00 ± 0.15</td>
<td>1.07 ± 0.47</td>
</tr>
<tr>
<td>SKkk238,240AA</td>
<td>SK(A)</td>
<td>1.00 ± 0.16</td>
<td>1.03 ± 0.17</td>
</tr>
<tr>
<td>SKkk244,246AA</td>
<td>SK(B)</td>
<td>1.00 ± 0.10</td>
<td>1.18 ± 0.17</td>
</tr>
<tr>
<td>SKkk248,249AA</td>
<td>SK(C)</td>
<td>0.45 ± 0.09</td>
<td>1.48 ± 0.54</td>
</tr>
<tr>
<td>SKkk256,257AA</td>
<td>SK(D1)</td>
<td>0.04 ± 0.01</td>
<td>1.83 ± 0.04</td>
</tr>
<tr>
<td>SKkk262,263AA</td>
<td>SK(E)</td>
<td>1.00 ± 0.12</td>
<td>1.46 ± 0.14</td>
</tr>
<tr>
<td>SKkk272,273AA</td>
<td>SK(F)</td>
<td>0.90 ± 0.05</td>
<td>1.15 ± 0.19</td>
</tr>
<tr>
<td>SKkk278,279AA</td>
<td>SK(G)</td>
<td>1.00 ± 0.05</td>
<td>1.01 ± 0.12</td>
</tr>
<tr>
<td>SKkk281,282AA</td>
<td>SK(H)</td>
<td>0.70 ± 0.12</td>
<td>1.29 ± 0.05</td>
</tr>
</tbody>
</table>

*Specific activity for the HPG activation was measured as detailed in Materials and methods. Values shown are mean of at least three independent experiments. Dissociation constants were measured using solid phase radio-assay employing 125I-labeled HPG.
barely one-twentieth as compared to nSK. The cause of this “latent” activation became apparent when the time course of amidolytic activation by equimolar quantities of SK(D1) and HPG was examined (Fig. 2B), which clearly showed significantly slow activation kinetics in case of SK(D1), with appearance of full amidolytic ability requiring ∼25–30 min. On the other hand, nSK-HPG equimolar mixtures generated amidolytic activity rapidly, with saturation being attained within 7–8 min after initiation of the reaction. The much slower amidolytic activation suggested that the delayed substrate HPG activation kinetics observed with this mutant likely arose from a slower exposure/maturation of the plasminogen active site in the SK(D1)-HPG activator complex.

Charge-reversal mutations at positions 256 and 257 in SK drastically compromise the HPG activation as well as amidolytic activation properties of the molecule

To further explore the role of the positively charged lysine residues of native SK at positions 256 and 257, these were altered to negatively charged glutamic acid residues by site-directed mutagenesis. The resultant mutant [SKKK256.257AA, abbreviated SK(D2)] was purified, and its HPG activation as well as amidolytic activation characteristics with equimolar partner HPG were examined. The construct showed a highly diminished overall rate of HPG activation (Fig. 3A), which was determined to be ∼500 L.U./mg, which is roughly 200-fold lower than that of nSK, and nearly 10 times lower than that of SK(D1). In addition, the progress curves of HPG activation generated with SK(D2) showed markedly prominent lag periods (Fig. 3A) when compared to that of SK(D1) (Fig. 2A). Also, unlike SK(D1), SK(D2) showed much slower amidolytic activation with 1:1 molar HPG (less than 10% amidolytic activity as compared to nSK after 20 min; see Fig. 3B), suggesting that the substitution of negatively charged side chains at positions 256 and 257 of SK had adversely affected the proteins’ ability to rapidly form a stoichiometric complex with HPG, or grossly undermined the ability of the complex, once formed, to initiate active site exposure in “partner” HPG. However, the apparent affinity of SK(D2) for HPG, when determined by the solid phase radioassay, was found to be virtually unaltered (approximately 1.5-fold decrease compared to nSK; K_f = 1.8 nM), indicating that its observed inability to activate equimolar zymogen did not emanate from a correspondingly lowered affinity between the complexing partners. The survival of the capability in SK(D2) to bind HPG was confirmed by examining the binding of this mutant with HPG-agarose, which clearly demonstrated that this mutant could indeed bind with this affinity matrix in a manner that was indistinguishable from that of nSK (see Fig. 4B). These results strongly indicated that the “biochemical lesion” in SK(D2) was not an inability to stoichiometrically complex with HPG, but at the post-complexation step of active site exposure in partner HPG. To explore this further, we tested the ability of this mutant to give the characteristic “burst” of p-nitrophenol release with equimolar HPG after reaction with the active site acylating agent, NPGB (Chase & Shaw, 1969). It is known that the SK-HPG virgin complex can undergo acylation at the cryptic active site with this “pseudo-substrate” even before the conversion of the HPG to HPN, while HPG or SK, individually, are unreactive with the reagent (McClintock & Bell, 1971; Reddy & Markus, 1972; Wohl, 1984). The results (see Fig. 3B, inset) showed clearly that SK(D2) failed to show any significant release of p-nitrophenol, while a characteristic burst was observed in case of nSK [as well as with SK(D1)], indicating that SK(D2) could not engender a NPGB-reactive active center in equimolar HPG.

The presence of a distinct delay in the activation of partner HPG2, on the one hand, and the presence of a highly diminished HPG activator activity in SK(D2) prompted us to explore whether this phenotypic deficiency could be overcome by providing a “preformed” active site in the form of HPN. For this purpose, we added
trace quantities of HPN into the PG activation reactions of SK(D2) and then measured the rates of HPG activation using the spectrophotometric method (Fig. 3C). The results demonstrated a remarkable rejuvenation of PG activator activity in the reactions as a function of increasing HPN. By contrast, when the activation reactions were carried out using HPG preparations whose intrinsic HPN content was decreased by passage through soybean trypsin inhibitor-agarose (see Materials and methods), the activator activity was seen to be perceptibly decreased. These results clearly indicated that a rate-limiting factor in the generation of activator activity by SK(D2) was the conversion of the initial SK(D2)-HPG complex to an enzymatically competent state possessing amidolytic and PG activation functions. This deduction was fully confirmed when SK(D2) was pre-incubated briefly with equimolar HPN, and the SK(D2)-HPN complex assayed as before for activator activity in the presence of excess HPG (1 µM). In this case
(Fig. 3C), a remarkably high activity was seen to be engendered. However, we noticed that the specific activity of SK\(\text{D2}\) complexed with HPN, although considerably resuscitated, was still perceptibly less than that of nSK-HPN, indicating that this mutant, despite being supplied with a fully formed active site, was still catalytically less active than nSK under the experimental conditions. Nevertheless, the foregoing results on the rejuvenation of SK\(\text{D2}\) with HPN clearly showed that a principal “block” in the action of SK\(\text{D2}\) was at the level of the formation of a functional activator after complexation with equimolar HPG.

The steady-state kinetic parameters for the catalytic processing of HPG and amidolytic peptide substrate by the equimolar complexes of SK\(\text{D1}\) and SK\(\text{D2}\) with HPN are presented in Tables 3 and 4, respectively. It is evident from these data that both mutants possess near-native ability for HPG processing under substrate-saturating conditions after being “partnered” with HPN. However, the \(K_m\) values for substrate HPG in case of SK\(\text{D1}\) and SK\(\text{D2}\) are approximately four- and eightfold higher, respectively, when compared to nSK. It is worth mentioning here that the mutation of two tandem lysine residues\(\text{KK 272,273}\) situated close-by in the beta domain, to alanine residues, had not resulted in any perceptible change in the biological activity of the resultant mutants\(\text{see Table 2}\).

Table 3. Steady-state kinetic parameters for HPG activation by equimolar complexes of mutant SK and HPN\(^a\)

<table>
<thead>
<tr>
<th>Activator protein</th>
<th>(K_m) ((\mu)M)</th>
<th>(k_{cat}) (min(^{-1}))</th>
<th>(k_{cat}/K_m) (min(^{-1})/(\mu)M)</th>
</tr>
</thead>
<tbody>
<tr>
<td>nSK-HPN</td>
<td>0.45 ± 0.05</td>
<td>10.75 ± 0.32</td>
<td>23.89</td>
</tr>
<tr>
<td>SK(\text{D1})-HPN</td>
<td>2.10 ± 0.20</td>
<td>08.06 ± 0.64</td>
<td>3.84</td>
</tr>
<tr>
<td>SK(\text{D2})-HPN</td>
<td>4.00 ± 0.10</td>
<td>09.68 ± 0.51</td>
<td>2.42</td>
</tr>
</tbody>
</table>

\(^a\)The kinetic parameters for HPG activation were determined at 22 °C as detailed in Materials and methods. The data represent the mean of three independent determinations.

Table 4. Steady-state kinetic parameters for amidase activity of equimolar complexes of mutant SK with HPN\(^a\)

<table>
<thead>
<tr>
<th>Protein</th>
<th>(K_m) (mM)</th>
<th>(k_{cat}) (min(^{-1}))</th>
<th>(k_{cat}/K_m) (min(^{-1})/mM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HPN</td>
<td>0.18 ± 0.04</td>
<td>300 ± 20</td>
<td>1,666.67</td>
</tr>
<tr>
<td>nSK-HPN</td>
<td>0.55 ± 0.02</td>
<td>378 ± 10</td>
<td>687.27</td>
</tr>
<tr>
<td>SK(\text{D1})-HPN</td>
<td>0.91 ± 0.02</td>
<td>359 ± 16</td>
<td>394.51</td>
</tr>
<tr>
<td>SK(\text{D2})-HPN</td>
<td>0.80 ± 0.10</td>
<td>351 ± 14</td>
<td>438.75</td>
</tr>
</tbody>
</table>

\(^a\)For determination of the amidolytic parameters, SK-HPN were pre-complexed in equimolar ratio and an aliquot of this mixture was assayed for amidolysis at varying concentrations of Chromozyme PL (see Material and methods). The data represent the mean of three independent determinations.
residues at positions 256 and 257 are specifically involved in the interaction of the SK-plasmin(ogen) activator complex with substrate PG. To further examine this aspect i.e., whether the observed effect on substrate-activator complex interactions was position-specific, rather than that arising from reversing the charges of two tandem residues in this segment of the primary structure of SK leading to a general “charge-destabilization” of this segment, we reversed the charges on two other such residues (EE 262,263) lying in the immediate vicinity of residues 256 and 257. The resultant mutant (SK_EE262,263KK) was then examined for its functional properties after purification. The results (data not shown) revealed no perceptible alteration in the specific activity of its HPG activator activity nor its ability to activate zymogen, suggesting strongly that the two lysine side chains at positions 256 and 257 were specifically involved in promoting the interaction of the activator complex with substrate PG.

In contrast to an increased $K_m$ for substrate PG, the $K_m$ values of the HPN complexes of SK(D1) and SK(D2) for the low-MW amido-lytic substrate (Chromozyme PL) were close to that of nSK, and the turnover rates ($k_{cat}$ values) were also similar, suggesting that the basic characteristics of the HPN active site in terms of their ability to catalyze the scission of specific peptides, upon complexation with SK(D1) and SK(D2), were unchanged. It should be mentioned here that one prominent and easily detectable consequence of the binding of SK with HPN is a roughly fivefold elevation of its $K_m$ for amido-lytic substrates without a concomitant alteration of the $k_{cat}$ values (Robbins et al., 1981), which has been interpreted to reflect a decreased accessibility of the substrate for the active center in the plasmin moiety of the activator complex, probably due to steric hindrance arising from SK binding in the vicinity of the active site (Wohl, 1984). The isolated SK $\beta$-domain (fragment 143–293) also brings about a similar alteration in $K_m$ values of HPN for the amido-lytic substrate, as reported earlier (Nihalani et al., 1997). Table 4 shows that both mutants, like nSK, brought about the characteristic increase in the $K_m$ for amido-lytic substrate over that of free HPN. These results indicated that despite the alteration at residues 256 and 257 the binding of the mutants with HPN resulted in a native-like inhibition of the accessibility of the amido-lytic substrate for the active site.

The simultaneous introduction of multiple charge-cluster to alanine mutation in the “core” region of the $\beta$-domain destroys substrate processing and zymogen activation properties without significantly affecting high-affinity interactions of SK with partner PG

The single charge-cluster sites whose mutations had led to a detectable reduction in terms of specific activities for HPG activa-

<table>
<thead>
<tr>
<th>Activator protein</th>
<th>$K_m$ ($\mu$M)</th>
<th>$k_{cat}$ (min$^{-1}$)</th>
<th>$k_{cat}/K_m$ (min$^{-1}$/$\mu$M)</th>
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<tbody>
<tr>
<td>nSK-HPN</td>
<td>0.45 ± 0.05</td>
<td>10.75 ± 0.32</td>
<td>23.89</td>
</tr>
<tr>
<td>SK(CH)-HYP</td>
<td>0.50 ± 0.07</td>
<td>2.36 ± 0.41</td>
<td>4.73</td>
</tr>
<tr>
<td>SK(FH)-HYP</td>
<td>0.33 ± 0.08</td>
<td>2.45 ± 0.25</td>
<td>7.42</td>
</tr>
<tr>
<td>SK(CFH)-HYP</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

The kinetic parameters for HPG activation were determined as described in Materials and methods. The data represent the mean of three independent determinations.

No detectable activity, over controls, could be measured.
could bind with HPG with a high affinity (Table 7; Fig. 4A), with the dissociation constants of their 1:1 complexes with HPG continuing to be in the nanomolar range despite the double/triple site mutations. In case of the triple-site mutant [SK(CFH)], a relatively modest (approximately threefold) reduction in affinity for HPG was evident. However, this diminution in binding affinity of this mutant does not appear to be commensurate with the drastic reduction in its biological activity. It may be mentioned that the latter was measured in the presence of HPG at concentrations several orders of magnitude higher than the nanomolar range of the dissociation constant between SK and HPG. The survival of a near-native avidity between SK(CFH) and HPG was also confirmed when the binding characteristics of this mutant onto HPG-agarose was studied, which showed that SK(CFH) could indeed bind tightly with the affinity matrix in a manner indistinguishable from that of nSK (Fig. 4B). Similar results were obtained for the double charge-cluster mutants (data not shown). These observations indicated that the mutations in the core region of the β-domain had led specifically to a reduction in the ability of these proteins to process substrate molecules [and in case of SK(CFH) and SK(D2), zymogen activation of partner HPG as well] without significantly altering their ability to bind tightly with HPG to form the initial 1:1 complex.

The double- and triple-site mutants were then assessed for their amidolytic parameters to explore whether the reduced catalytic efficiency of their HPN complexes for HPG substrate was reflected in a similar change with respect to the small-MW amidolytic substrate as well (Table 6), presumably due to alterations brought about in the active site of HPN. It was observed that, if anything, the $k_{cat}$ for amidase activity had marginally increased as a result of these mutations, indicating that the primary specificity of the HPN active site was basically unaltered upon complexation with the mutants. Remarkably, however, these mutants lacked an ability to effect an increase in the $K_m$ of HPN for the amidolytic substrate, a property that is characteristic of the binding of native SK with free HPN (Robbins et al., 1981; Wohl, 1984). Thus, it appears that these mutants, although exhibiting near-native affinities for HPN (Table 7), failed to engender the structural alteration(s) in partner HPN that result in a reduced access of the small-MW peptide substrate to the active site due to steric hindrance. Interestingly, in terms of this effect, the single-site charge cluster mutant, SK(C), was similar to the double- and triple-site charge cluster mutants, whereas SK(F) and SK(H), like nSK, increased the $K_m$ of HPN for amidolytic substrate.

With a view to explore whether the altered functionalities of the mutants were reflected in their folding characteristics, in particular their secondary structural features, their far-UV CD spectra were recorded. The spectrum for nSK was closely similar to the one reported previously for this protein (Radek & Castellino, 1989). The CD spectrum of SK(CFH) revealed (Fig. 5) a remarkable similarity to that of the native protein, showing that the overall folding of this mutant despite the triple-charge-cluster mutations, was essentially preserved. The spectra of SK(D2), SK(CH), and SK(FH) were nearly identical to that of nSK (data not shown).

### Table 6. Steady-state kinetic parameters for amidase activity of equimolar complexes of mutant SK with HPN

<table>
<thead>
<tr>
<th>Protein</th>
<th>$K_m$ (mM)</th>
<th>$k_{cat}$ (min⁻¹)</th>
<th>$k_{cat}/K_m$ (min⁻¹/mM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HPN</td>
<td>0.18 ± 0.04</td>
<td>300 ± 20</td>
<td>1,666.67</td>
</tr>
<tr>
<td>nSK-HPN</td>
<td>0.55 ± 0.02</td>
<td>374 ± 10</td>
<td>680.00</td>
</tr>
<tr>
<td>SK[C]-HPN</td>
<td>0.16 ± 0.05</td>
<td>467 ± 32</td>
<td>2,918.75</td>
</tr>
<tr>
<td>SK[F]-HPN</td>
<td>0.60 ± 0.06</td>
<td>462 ± 29</td>
<td>770.00</td>
</tr>
<tr>
<td>SK[H]-HPN</td>
<td>0.54 ± 0.10</td>
<td>385 ± 25</td>
<td>712.96</td>
</tr>
<tr>
<td>SK[CH]-HPN</td>
<td>0.19 ± 0.03</td>
<td>561 ± 36</td>
<td>2,952.63</td>
</tr>
<tr>
<td>SK[FH]-HPN</td>
<td>0.25 ± 0.07</td>
<td>274 ± 13</td>
<td>1,096.00</td>
</tr>
</tbody>
</table>

### Table 7. Dissociation constants for HPG-binding by double- and triple-charge-cluster SK mutants

<table>
<thead>
<tr>
<th>Type of SK</th>
<th>Short name</th>
<th>Dissociation constant [$K_d$] of SK-HPG complex (nM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild-type SK</td>
<td>nSK</td>
<td>1.07 ± 0.47</td>
</tr>
<tr>
<td>SK[E248,249AA, EK248,252AA]</td>
<td>SK(CH)</td>
<td>1.73 ± 1.10</td>
</tr>
<tr>
<td>SK[E272,273AA, EK281,282AA]</td>
<td>SK(FH)</td>
<td>1.29 ± 0.15</td>
</tr>
<tr>
<td>SK[E248,249AA, EK272,273AA, EK281,282AA]</td>
<td>SK(CFH)</td>
<td>3.10 ± 0.94</td>
</tr>
</tbody>
</table>

*The amidolytic parameters were determined as outlined in Materials and methods. The data represent the mean of three independent determinations.

---

The figure shows the far-UV CD spectra in the wavelength range of 197–250 nm as outlined under Materials and methods. The curves shown are: nSK (closed squares) and SK(CFH) (open circles). The CD spectra of SK(FH) and SK(CH) were found to be closely similar to that of SK(CFH).
These results thus suggest the absence of grossly misfolded structures in the mutated SK derivatives. It should be mentioned, however, that the native-like CD spectra do not necessarily rule out the presence of conformational alterations in the vicinity of the mutations since the spectral contributions arising from these changes could easily have been “averaged out” by the spectral contributions emanating from the remainder of the molecule.

Discussion
Perhaps the most intriguing question associated with the mode of action of PG activators in general, and the SK-plasminogen activator complex in particular, is the structural basis of their inordinate high substrate specificity. In the case of the latter, the identification of the mechanism(s) underlying the conversion of the relatively nonspecific, trypsin-like substrate specificity of plasmin to that of a highly PG-specific protease upon complexing with SK still constitutes a challenge despite the recent availability of the three-dimensional structure of the SK-human plasmin light chain complex (Wang et al., 1998). Conceptually, SK could produce a change in the specificity of the plasminogen active site by changing its conformation/geometry per se, or by providing altered secondary substrate binding subsites (S2, S3, etc.) into the active center, or by improving substrate presentation, through a more optimal docking, of PG at the active site. It is also possible that more than one of the above possibilities is operative together in the functioning of SK. At present there is no direct evidence for the first two possibilities. However, the structure of the SK-plasmin light chain complex strongly supports the third possibility, which we had also advanced earlier on the basis of biochemical studies (Nihalani et al., 1998), that SK “switches” the substrate specificity of plasmin because the SK-plasminogen complex provides two substrate-specific sites for the optimal binding of substrate PG onto the complex. Indeed, modeling studies using the crystal structure data on the SK-HPN light chain complex (Fig. 6) indicate several potential contacts in the α- and β-domains of SK between docked substrate and the complex and that the SK-HPG interface provides a “near perfect” concavity for substrate positioning wherein the scissile peptide bond in the latter becomes ideally amenable to cleavage by the active center of the activator complex (Esmon & Mather, 1998; Wang et al., 1998). In the present study, a direct experimental evidence of the involvement of the β-domain in such a “protein cofactor mediated” catalytic assistance and substrate docking mechanism for SK action is being reported.

Our mutagenesis studies show that the replacement of the two tandem Lys residues at positions 256 and 257 in the core region with either Ala or Glu residues drastically affects the HPG activation properties of the molecule. The involvement of these residues in SK functioning has been shown in an earlier study as well (Lin et al., 1996). In that study, the Lys residues were altered to Ala residues, and a decrease in HPG activation kinetics by the mutant was noted. However, in the present investigation, by mutating the lysyl to glutamyl side chains, a structurally much more nonconservative alteration, greater insight into the structure-function interrelationship of residues 256 and 257 could be obtained. The presence of two consecutive negative charges at this locus turned out to be highly detrimental to the functioning of the molecule, which now showed a very low overall HPG activation capability even though its apparent affinity for HPG to form a 1:1 complex was largely unaffected. The slow activation kinetics of HPG by this mutant was attributable to a highly compromised ability for effecting zymogen activation in partner HPG. In the case of the relatively less drastic mutation KK256,257AA [SK(D1)], the property of zymogen activation was slowed to a great extent compared to the native protein. The underlying “biochemical lesion” that was caused by mutation at Lys256 and Lys257 in SK, therefore, seemed to be related to the first step(s) after complexation with HPG that lead to the exposure of the active site in the latter. Therefore, when the charges of the two residues were reversed [as in SK(D2)], there was a highly enhanced delay in exposure of the active site in partner HPG.

In the case of native SK, the nascent 1:1 SK-HPG complex is believed to undergo a rapid conformational change to an “activated” form that displays both amidolytic and esterolytic capability even before any proteolytic cleavage has taken place through a process that is presently unclear (Esmon & Mather, 1998). It also remains unclear whether it is this “virgin” complex, or the SK-HPN complex (to which it rapidly converts), that possesses the earliest PG activator ability (Davidson et al., 1990; Shi et al., 1993). Whatever the exact mechanism of this zymogen activation in the SK-HPG complex, our results show that the complex between SK(D2) and HPG is drastically compromised with respect to this transformation, as indicated by an inability to participate in the “NPGB burst” reaction and an enhanced delay in amidolytic activation of partner HPG. However, the 1:1 complex of SK(D1) with HPG was able to react with NPGB and also generate amidolytic activity albeit slowly (Figs. 2, 3). Thus, the rate-limiting factor in the HPG activation process by these two mutants seems to be a delay, to varying extents, in the generation of an amidolytically competent SK-plasminogen complex. Indeed, in perfect accord with this proposed scenario, the apparent HPG activation capability of both mutants increased dramatically once a “fully mature” partner, in the form of HPG, was provided. The kinetic analyses of the now “rejuvenated” activator complexes (formed with HPN instead of HPG) revealed, however, that their interaction with substrate HPG was significantly reduced, as indicated by a nearly fourfold increased $K_m$ for HPG in the case of SK(D1), and a eightfold increased $K_m$ in the case of SK(D2).

Our results on the involvement of Lys256 and Lys257 of SK in substrate recognition represent the first experimental demonstra-
tion of the direct involvement of the β-domain in providing assistance in the sequestering of substrate PG for processing by the activator complex. These findings provide, at least in part, a rationale for the observation that consequent to SK binding, the $K_m$ of the HPN active site that is measurable in the millimolar range for unconstrained, small-MW substrates is found to be several orders of magnitude lower when tested against HPG as substrate (Robbins et al., 1981). The complementary loci in substrate PG with which the side chains of residues 256 and 257 make contact in this process of increasing the efficacy of enzyme-substrate interactions, however, are presently unclear. The kringle domains of HPG, due to the presence of lysine binding property in these structures, immediately offer themselves as probable candidates, especially since these have been implicated clearly both in 1:1 interaction between SK and HPG as well as in the interaction of the activator complex with substrate HPG (Shi & Wu, 1988; Conejero-Lara et al., 1998; Wang et al., 1998; Young et al., 1998).

However, precise structural information on this important aspect is sorely lacking at present, particularly as the recently published X-ray diffraction data employed crystals of complexes between SK and microplasmin, which is devoid of the kringle domains (Wang et al., 1998). Our results, nevertheless, point unmistakably to a direct role of this region, in particular of the two lysine residues at position 256 and 257, probably in the docking of substrate PG during its conversion to product (plasmin) as a result of the highly specific proteolytic cleavage at the scissile peptide bond by the SK-plasmin(o)gen activator complex. The observation that alterations in these residues lead to an apparent decreased substrate affinity is entirely in keeping with this inference, even though the identity of the complementary site(s) in substrate PG with which these residues interact is presently unclear. In addition, these residues seem to be important in thezymogen activation process as well. The molecular details of the process whereby SK binding to PG leads to active site exposure even before any proteolytic scission has taken place has not been elucidated. Wang et al. (1998) hypothesize that Lys698 of HPG, which is located close to the activation pocket, can replace the α-amino group of Val562 (hence, forming the critical salt bridge to Asp740 during the zymogen activation process) in the SK-HPG virgin complex. A mechanism such as this, or alternate ones (Esmon & Mather, 1998; Khan & James, 1998), presume an underlying conformational change in the activation pocket of HPG subsequent to SK binding that activates the hitherto latent active site without scissile peptide bond cleavage. It appears that the placement of two negative charges at 256 and 257 of SK strongly hinders this conformational change. At the same time, the fact that the substitution of alanines instead of glutamic acid residues at these positions had a much milder effect on thezymogen activation ability of the molecule suggests that the two Lys residues in nSK are not involved inzymogen activation per se, since had it been otherwise, then irrespective of the type of mutation, the effect would have been a drastic decrease inzymogen activation ability. Wang et al. (1998) suggest that the binding of the γ-domain of SK to the autolysis loop region of HPG may cause the conformational change necessary to form a salt linkage between the Asp740 and Lys698 of HPG. An inspection of the recently elucidated three-dimensional structure of the SK-HPN light chain complex (Fig. 6) shows that residues 256 and 257 occur in a highly flexible region of the β-domain. This region is encompassed by a long loop (residues 248–259), which was not observable clearly in the experimental electron density map due to positional disorder. Moreover, the β-domain itself exhibits high mobility as evidenced by its large thermal factors probably due to the absence of the kringles (Wang et al., 1998). In the SK-HPN light chain crystal structure, the tip of the disordered loop encompassing residues 256–257 in the β-domain is more than 50 Å away from the Arg561 of the partner microplasmin (see Fig. 6). Thus, unless one invokes the possibility that the β-domain’s juxtaposition with the rest of the activator complex is actually different in the full-fledged ternary complex, viz., HPG.SK.HPG, occupying either a position more proximal to the active site area or indirectly influencing the conformational changes leading to active site exposure, it is difficult to make more definitive deductions at this juncture regarding the structural basis of Lys256,257 in activating partner HPG.

Unlike the crystal structure of the SK-plasmin light chain complex, the X-ray diffraction data on the closely similar PG activator protein, staphylokinase (SAK) complexed simultaneously with substrate and partner HPG (both in the form of the HPN light chain), appears to provide a better glimpse of the positioning of the substrate with respect to the plasmin active site. Interestingly, SAK displays a marked structural similarity to the α-domain of SK (Wang et al., 1998). Besides this structural similarity, both SK and SAK have common functional similarities as well, in that both are intrinsically inert bacterial proteins that activate PG “indirectly” by first forming proteolytically functional activator complexes with partner plasmin(o)gen. However, unlike SK, SAK requires a preformed active site to generate a functional activator, whereas SK can expose the active site in partnerzymogen directly (Collen et al., 1993). The lack of a zymogen activation ability in SAK seems to be commensurate with its smaller size (approximately one-third of SK) since it lacks the counterparts to the β- and γ-domains that SK possesses. Besides the additional attribute of autocatalytic zymogen activation, the presence of all three domains in SK confers a catalytic advantage onto this activator protein over SAK, which is exemplified by the fact that the SK-HPN complex possesses a substrate HPN turnover rate that is significantly higher than that of SAK-HPN (Robbins et al., 1981; Radek et al., 1993). The present study suggests that at least a major part of this additional “catalytic power” emanates from the crucial assistance in substrate binding and/or docking that the β-domain provides, and which probably plays a crucial role in enhancing overall substrate processivity as well. The crystal structure of the SAK-plasmin light chain complexed with substrate microplasmin (Parry et al., 1998) strongly suggests substrate docking, on the one hand, and subtle alterations of the secondary S3/S4 subites of the active center of plasmin by the bound SAK, on the other, as being responsible for the switch in the substrate specificity of the bound HPN. If one assumes that the α-domain of SK retains a high level of functional analogy with SAK, it should not seem surprising that the intrinsic catalytic power of SAK is much less than that of SK because it has to generate this activity entirely on its own while in the case of SK, both the α- and β-domains (and probably the γ-domain as well) interact together to engender much higher rates of HPG activation. Our results demonstrate that at least one of the major factors behind the contribution of the β-domain toward the higher catalytic efficiency observed with SK-HPN arises directly from the locus encompassing the residues 256 and 257, since alterations of these residues led to a perceptible increase in $K_m$ for substrate PG.

On the other hand, mutants altered simultaneously at several residues around this site [as in case of SK(CH) and SK(FH1)] resulted not so much in an increased $K_m$ but in an overall reduced
substrate processivity \((k_{cat})\) by their activator complexes. In case of the triple charge-cluster mutant, SK(CFH), which did not display any significant ability to activate partner HPG, virtually no PG activator activity could be detected even when a preformed active site, in the form of HPN, was supplied to form the 1:1 partner. These mutants, including SK(CFH), appeared to be native like by CD spectral analysis and showed dissociation constants for HPG that, like that of nSK, were in the nanomolar range of concentration. At first sight, the most straightforward explanation for this phenomenon is that the charges in this segment interact directly with residues involved in catalysis. However, this possibility appears somewhat remote since the \(k_{cat}\) values for amidolysis of 1:1 complexes of the mutants with HPN were not decreased, indicating an essentially unaltered active site topology. Thus, as previously discussed, a rational explanation of the lowered \(k_{cat}\) values obtained with mutants SK(CH) and SK(FH) in the absence of any perceptible changes in \(K_m\) values cannot be forwarded at this juncture and will probably have to await the availability of future crystallographic data (such as that gleaned from HPG.SK.HPG ternary complexes) offering much greater structural detail than the one presently available. The \(\beta\)-domain in this structure has demonstrably increased average thermal factor probably due to its enhanced mobility arising from an absence of stabilizing interactions that are likely kringle-mediated. Nevertheless, despite the absence of a directly verifiable structural explanation, it is tempting to speculate on possible alternate mechanisms that may help explain the observed results. Thus, one can envision a scenario of a protein-cofactor mediated catalysis wherein both substrate docking and product release are assisted through selective interaction of the \(\beta\)-domain of the activator complex with discrete structural elements, such as the kringles, in the substrate PG. The cardinal role played by the kringle domains of PG in substrate recognition by the activator complex is evident from the observed decrease in microplasminogen activation, which is devoid of kringles, by SK (Shi & Wu, 1988; Shi et al., 1990), and also by the significant fall in the formation of ternary complex (HPG.SK.HPG) in the presence of \(\epsilon\)-amino-n-caproic acid (Young et al., 1998), an agent that binds to kringles and alters their conformational states. It is well recognized that the scission of the Arg561–Val562 peptide bond of HPG to HPN generates a large conformational change in the latter, particularly in terms of the three-dimensional arrangement of the kringle domains (Mangel et al., 1990; Marshall et al., 1994; Bock et al., 1996). Thus, in this model, kringle-mediated recognition of the substrate by the core region of the \(\beta\)-domain of the SK in the activator complex would enhance enzyme-substrate collision and docking, and following conversion of substrate HPG to HPN, the facilitated release of the product (plasmin) as well. The latter event would accrue directly as a result of reduced affinity of the activator for HPN consequent to conformational changes not only at the active site but also in and around the interacting kringle(s) as well. Thus, SK might serve to enhance the affinity of substrate-activator complex interactions by “exploiting” elements of the substrate molecule that are destined to undergo the most prominent conformational changes upon conversion to product (e.g., the scissile peptide bond region itself and the kringle domain conformations) so that the product can be released readily from the activator complex upon completion of catalysis. Kringle 5 is closest to the scissile peptide bond region and therefore a likely candidate mediating this interaction. The fact that the two- and three-site charge-cluster mutants have impaired ability, unlike nSK, to effect the characteristic increase in the \(K_m\) for the amidolytic substrate in their 1:1 complexes with HPN signifies an altered mode of binding of their \(\beta\)-domain in the vicinity of the active site, which might be responsible not only for an improper positioning of substrate PG, but also in failing to recognize, to varying extents, the kringle-mediated structural changes that help in product release. In case of SK(CFH), the impairment was not only at the level of HPG processing but in terms of zymogen activation as well, indicating that the triple-site charge-cluster mutations had resulted in a relatively greater structural destabilization in this segment of the \(\beta\)-domain. Indeed, the nearly two- to threefold decreased affinity of complex formation observed with these mutants may signify a subtly altered mode of interaction with partner HPG due to local structural disturbances, which might affect the catalytic properties of the complex without a concomitantly drastic decrease in the ability to form the stoichiometric activator complex. Thus, on the whole, the present experimental observations suggest that the \(\beta\)-domain of SK is involved in both the docking of substrate PG and its release as product.

It is intriguing that despite the six simultaneous point-mutations in SK(CFH), its 1:1, high-affinity interaction with HPG was unchanged, and its overall structure, as evidenced by CD, retained a native-like folding. At first sight, this might appear to be at variance with our earlier conclusions, based on a Peptide Walking approach (Nihalani et al., 1997), that this locus, besides participating in substrate recognition, is intimately involved in 1:1 interaction between SK and HPG. In that study, partially overlapping peptides derived from the sequence 230–290 of SK were found to inhibit strongly the formation of a functional activator complex when added prior to complexation between equimolar SK and HPG. Conversely, peptides derived from the other regions of the \(\beta\)-domain primary structure failed to show this effect. Although the present study substantially validates our earlier conclusion that a substrate-specific binding locus is present in the region in and around residues 250–270 of SK, an overt effect of the mutations in the 230–290 region on 1:1 interactions between SK and HPG was not detected. However, it is conceivable that intermolecular contacts other than those involving charged side chains are responsible for this high-affinity interaction. Indeed, a few distinct patches of clustered apolar residues are present in the core region of the \(\beta\)-domain of SK, that are potentially capable of providing intermolecular hydrophobic contacts with similar residues in HPG. We have recently observed that alanine linker mutagenesis of these patches results in a drastic reduction in the activity of the mutants (A. Chaudhary & G. Sahni, unpubl. results). The detailed functional effects of alterations, such as these or of other potential contacts between SK and HPG as revealed by the recently published X-ray diffraction data, are currently being investigated in detail. It is hoped that the availability of SK mutants specifically altered in terms of their 1:1 interaction with HPG, or with respect to substrate and/or product affinity, or turnover rates, followed by their high-resolution structural evaluation will pave the way for a clearer understanding of the molecular mechanism of SK action.

Materials and methods

Proteins and reagents

Glu-HPG was procured from Boehringer-Mannheim (Mannheim, Germany). Where albumin-free HPG was specifically needed, Glu-HPG was purified from human plasma by affinity chromatography on lysine-agarose in the presence of proteases inhibitors as de-
scribed (Deutsch & Mertz, 1970). Both preparations contained less than 2% of lys-HPG and less than 0.01% free HPN. Where needed, HPG was converted to HPN by the action of agarose-immobilized UK (300 Plough units/mg Glu-HPG) in 50 mM Tris-CI buffer, pH 8.0, containing 10 mg/mL HPG, 25 mM L-lysine and 25% glycerol for 12 h at 22°C. UK was covalently immobilized onto cross-linked agarose (Sepharose 6B-CL; Pharmacia Ltd., Uppsala, Sweden) by reductive amination (Stults et al., 1989). Soybean trypsin inhibitor agarose (~3 mg STI/mL agarose), used to prepare HPN-deficient HPG, and HPG-agarose (~2 mg HPG/mL agarose) were also prepared by covalent immobilization of the respective proteins using reductive amination chemistry. The plasmin content in HPG preparations, as well as the number of active sites generated in HPN, were determined by active-site titration with the acylating agent, NPGB (McClintock & Bell, 1971). Protein concentrations were determined by the method of Bradford (1976). S. equisimilis SK (W.H.O. standard; 700 I.U. per vial), used for the determination of specific activities of SK or its mutant forms, was procured from National Institute of Biological Standards and Control, South Mihu, Hertfordshire, U.K. Chromogenic plasmid substrate, tosyl-Gly-Pro-Lys-anilide (Chromozyme PL) was obtained from Boehringer-Mannheim. The T7 RNA polymerase promoter-based expression vector, pET23(d) used for the intracellular expression of SK in E. coli, was a product of Novagen Inc. (Madison, Wisconsin). STI was procured from Sigma Chemical Co. (St. Louis, Missouri). Thermostable DNA polymerase (pfu) with proof-reading activity, plasmid Bluescript II KS, and E. coli XL-Blue were procured from Stratagen Inc. (La Jolla, California). Restriction endonucleases and other enzymes used for rDNA experiments were procured from New England BioLabs (Beverly, Massachusetts) or Promega Inc. (Madison, Wisconsin). Oligonucleotide primers were either synthesized in-house on an Applied Biosystems (Foster City, California) DNA synthesizer model 492, or custom-synthesized by Ransom Hill Biosciences Inc. (Ramona, California). N-terminal protein sequencing was done on a Perkin Elmer/Aplied Biosystems sequencer, model 476 A. PCR-generated DNAs from agarose gels, and plasmid DNAs were routinely purified using kits from Qiagen Inc. DNAs from agarose gels, and plasmid DNAs were routinely purified using kits from Qiagen Inc. DNA sequencing to confirm the incorporation of the desired mu-

cluster sites (see Fig. 1) mutated either to Ala residues (in most cases) or, in a few instances, to a complementary side-chain to effect charge-reversal (e.g., Lys to Glu, and vice-versa), with each mutant type representing the incorporation of two, four, or six simultaneous point mutations, respectively. In Table 1 are also shown the diagnostic R.E. sites introduced by translationally silent mutagenesis (Raghava & Sahni, 1994) in each primer to aid the screening of the clones. The sequence of the two flanking primers (termed the “upstream” and “downstream” primers) in the amplification reaction are also shown. Each of these carried an internal R.E. site unique to the SK ORF [as also to the plasmid pET23(d)-SK], viz., Bse I and Bsm I to facilitate the re-ligation of the mutant PCR blocks back into the SK ORF in the expression vector. Briefly, the first PCRs (to generate the different mutant megaprimer) were carried out in a volume of 100 μL and contained (final reaction conditions are given) 200 μM dNTPs, 100 ng of template DNA, pET23(d)-SK, the T7 polymerase based expression vector in which the full-length SK gene had been cloned; see Nihalani et al., 1998 and references cited therein), 20 pmol each of the mutagenic and downstream primers, 2.5 U of standard cloned pfu thermostable DNA polymerase, and 10 μL of 10X pfu reaction buffer. The PCR cycling conditions were in accordance with the denaturation temperatures of different primers, calculated using the computer program, Oligo (version 4.0). In general, all reactions were started with a “hot start” (94°C, 5 min), followed by a denaturation (94°C for 45 s), annealing (50°C for 1 min) and extension (72°C, 1 min) phases. After a total of 30 cycles, a 10 min period at 72°C was given for a final extension of partially finished daughter DNA molecules, and the reactions were terminated, and then processed through Qiagen PCR purification columns. The relevant DNA cassettes were then purified by agarose gel electrophoresis, by excising the required DNA band and further purifying by Qiagen gel extraction kit. The amplified DNAs was then used as megaprimer for a second PCR (referred to as PCR-II) either with wild-type SK gene as the template, or with full-length extended PCR products carrying different mutations as the template to generate double- and triple-charge cluster mutant DNAs (see below). The full-length, extended DNA cassettes were then cloned back into pET23(d)-SK expression vector at the BseRI and Bsm I sites. The PCR-II pre-mixes (100 μL) contained: 60 ng template [pET23(d)-SK], 200 μM dNTPs, 10 μL of 10X pfu buffer, and varying concentrations of megaprimer (200–600 ng). The mixes were held for 5 min at 95°C (“hot start”) and the reaction initiated with addition of 5.0 U of pfu DNA polymerase per reaction. This was followed by seven cycles of denaturation (94°C, 45 s) and extension (72°C for 3 min) to ensure build-up of mutated strand DNA and to effect megaprimer extension. At the end of this phase of the PCR, 20 pmol of upstream primer was added, and thermal cycling continued as follows, for 10 cycles: 94°C for 45 s, 51°C for 1 min, and 72°C for 1 min. After the last cycle, 20 pmol of downstream primer was added, and cycling continued as before for 15 cycles, followed by a final extension at 72°C for 10 min. The extended PCR products (504 bp) were then purified on agarose gels and the DNA bands isolated after excision of gel blocks. These were digested with BseRI and BsmI restriction enzymes and ligated with similarly digested pET23(d)-SK plasmid DNA, and transformed into E. coli XL-Blue electrocompetent cells. The desired clones were selected by screening mini-prep plasmid DNAs for the diagnostic R.E. sites incorporated through the PCR primers (Table 1), and the positive clones were subjected to automated DNA sequencing to confirm the incorporation of the desired mu-
tation, as also to rule out the presence of any unwanted mutations due to the PCR amplification. For the construction of double and triple charge-cluster mutants, the same overall strategy was followed except that complementary megaprimers and templates were used for the extension step in the PCR-II step e.g., for the construction of the double charge-cluster mutant SK\textsubscript{RE248.248AA,EK281.282AA}, the megaprimer carrying the mutations RE248.248AA was used with the full-length extended PCR product carrying the mutations EK281.282AA. Similarly, for the construction of the double charge-cluster mutant SK\textsubscript{EK272.273AA,EK281.282AA}, the extended PCR product containing the mutations EK281.282AA was employed as the template, with the megaprimer carrying the mutations EK272.273AA used to prime the reaction at the PCR-II step. In case of the triple charge-cluster mutants containing the mutations EK281.282AA was employed as the template for the extension step in the PCR-II step e.g., for the construction strategy was as follows. First, a megaprimer containing the double charge-cluster mutations EK272.273AA and EK281.282AA was constructed using as template the purified PCR product carrying the single cluster mutant EK281.282AA, and carrying out another PCR using the upstream primer bearing the EK272.273AA mutation along with the downstream BsmI-site containing primer. This double charge-cluster megaprimer was then isolated, and used along with the full-length extended PCR product with the mutation RE248.248AA (used as template in PCR-II), to generate PCR amplified DNA block containing three simultaneous charge-cluster mutations. All the purified mutant PCR DNAs were then digested with Bse RI and Bsm I restriction enzymes, and cloned into similarly digested plasmid vector pET23(d)-SK to obtain the desired mutant SK clones. All mutations were confirmed by DNA sequencing of the complete SK gene in the expression vector using the T7 promoter and terminator sequencer primers. This also established the absence of any unwanted mutation introduced in the SK gene during mutant construction particularly as a result of the PCR amplification.

Expression and purification of SK mutants

The native-like recombinant SK (termed nSK) or the various SK mutants were expressed intracellularly in E. coli BL-21 (DE 3) cells after induction with 1 mM IPTG after initial growth until mid-log phase, essentially as described (Nihalani et al., 1998). The bacterial pellets obtained by centrifugation from 1 L shake flask cultures were suspended in 80 mL of 20 mM NaPO\textsubscript{4}, pH 7.2, and subjected to ultrasonication for 10 min with 30-s pulses with equal periods of rest, using a Heat Systems (New York, New York) ultrasonic processor with a medium-sized probe. The lysates were then processed for hydrophobic interaction chromatography (HIC) on phenyl-agarose (Affinity Chromatography Ltd., Isle of Man, U.K.) as follows. After centrifugation at 4°C at 12,000 \( \times g \) for 15 min, the supernatants were made 0.5 M in NaCl, and loaded onto a column of phenyl-agarose (2 \( \times \) 10 cm) in equilibrating buffer, viz., 0.5 M NaCl in 20 mM NaPO\textsubscript{4}, pH 7.2. The column was then washed successively with five bed-volumes each of (in this order): equilibrating buffer, 20 mM NaPO\textsubscript{4} buffer, pH 7.2, distilled water, and finally, 8 M urea. Nearly 90% of the loaded SK activity was recovered by elution with water, and the protein found to be more than 80% pure as judged by SDS-PAGE. This fraction, pooled conservatively, was then loaded onto a Perseptive Biosystem Poros-D anion-exchange column (1 \( \times \) 5 cm) in 20 mM NaPO\textsubscript{4}, pH 7.2, on a Bio-Cad Sprint liquid chromatography work station. The chromatographic elution was carried out (at a flow rate of 1 mL/min) using a linear gradient of NaCl (0 to 1.0 M) in NaPO\textsubscript{4} buffer, pH 7.2 over 20 min. Greater than 75% yield of SK activity, as compared to the preceding step, was obtained, and the SK eluted in and around a molarity of 0.3 NaCl in the gradient. The SK was more than 95% pure at this stage. The major SK-containing fractions were pooled and stored at \(-70^\circ C\) until analyzed. Overall recoveries were consistently 50–65%, yielding between 30–40 mg purified protein/liter of shake-flask grown culture. The purified proteins were subjected to N-terminal sequencing, which showed an extra methionine residue at the N-termini, succeeded by the native sequence of SK.

Characterization of SK mutants

Assays for studying the activation of HPG by SK and SK mutants

A one-stage assay method was used to measure the kinetics of HPG activation by SK or its mutants (Wohl et al., 1980; Shi et al., 1994). Purified SK or mutant SK (0.5–50 nM) were added to a 100 \( \mu \)L quartz assay cuvette containing HPG (1 \( \mu \)M) in assay buffer (50 mM Tris-Cl, pH 7.5) containing 1.0 mM chromogenic substrate (Boehringer-Mannheim). The change in absorbance at 405 nm was then measured as a function of time(t) in a Shimadzu UV-160 model spectrophotometer at 22°C. Appropriate dilutions of standard W.H.O. S. equisimilis streptokinase were used as reference for calibration of international units/mg protein (specific activity) in the unknown preparations (Heath & Gaffney, 1990). The activator activities were obtained from the slopes of the activation progress curves, which were plotted as change in absorbance/t against t (Wohl et al., 1980).

Amidolytic and esterolytic activation of equinolar SK/SK mutant-HPG complexes

HPG (final conc. 5.0 \( \mu \)M) was incubated with 5.5 \( \mu \)M each of SK or mutant SK at 22°C in 50 mM Tris-Cl, pH 7.5, containing 0.5% BSA. The amidolytic activity was determined periodically by transferring a suitable aliquot of the complex (final concentration in reaction, 100 nM) to a 100 \( \mu \)L cuvette containing assay buffer along with 1 mM Chromozyme PL in a final volume of 100 \( \mu \)L. The change in absorbance at 405 nm was monitored continuously to compute the kinetics of amidolytic activation (see Nihalani et al., 1998) by plotting the slopes of the progress curves from the initial periods (2–4 min) as a function of pre-incubation times. The generation of an NPGB-reactive active center by equimolar SK/SK mutant and HPG was monitored at 22°C as described by Wohl et al. (1980). The reaction was initiated by adding 5 \( \mu \)M HPG (final concentration) to an assay cuvette containing buffer (10 mM NaPO\textsubscript{4}, pH 7.2) and SK (5.5 \( \mu \)M) and 100 \( \mu \)M NPGB. The release of p-nitrophenol due to acylation of the active center was detected by continuous monitoring at 410 nm.

Assays for determining the steady-state kinetic constants for HPG activator activity of SK and SK mutants

To determine the kinetic parameters for HPG activation, fixed amounts of SK or SK mutant (1 to 10 nM) were added to a 100 \( \mu \)L assay cuvette containing chromogenic substrate (1 \( \mu \)M) and vary-
as described, in a separate tube (at 22°C) at a final concentration of 0.2 μM in 50 mM Tris-Cl, pH 7.5, containing 0.5% BSA, and suitable aliquots were withdrawn for HPG activator assay using different final substrate HPG concentrations. The kinetic parameters for HPG activation were then calculated from inverse, Lineweaver–Burke plots by standard methods (Wohl et al., 1980).

Assays for determining the steady-state kinetic constants for amidolytic activity of SK and SK mutants

The kinetic parameters of amidolysis by SK or SK mutants complexed with HPN were determined by precomplexing equimolar ratios of SK and HPN (200 nM each) at 22°C in 50 mM Tris-Cl, pH 7.5, containing 0.5% BSA. The amidolytic activity was measured by transferring an aliquot of this mixture to a 100 μL assay cuvette containing buffer (50 mM Tris-Cl, pH 7.5) and varying concentrations of the chromogenic substrate (0.1–2.0 mM) to obtain a final concentration of 100 nM of the complex in the reaction. The reaction was continuously monitored spectrophotometrically at 405 nm for 4–5 min. The kinetic parameters were determined from Lineweaver–Burke plots by standard methods (Wohl et al., 1980). In cases where kinetic constants were determined for SK-HPG complexes, SK/SK mutants were preincubated with equimolar HPG and allowed to mature at 22°C until the generation of maximum amidolytic capability, and then transferred to assay cuvettes as described above for the determination of amidolytic parameters.

Measurement of dissociation constants

Solid phase assays were performed using the procedure described by Reed et al. (1995). Native SK or SK mutants were immobilized onto the wells of polyvinyl microtitre plates (Costar, Cambridge, Massachusetts) [175 μL of protein solution 5.0 μg/mL in 50 mM NaCO₃-NaHCO₃ buffer, pH 9.5 (binding buffer)] after overnight incubation at 4°C. Into each well was then added 200 μL of a 1% (w/v) solution of BSA in PBS, pH 7.4. Plates were then incubated for 1 h at 4°C, following which 175 μL/well of 125I-labeled HPG at different concentrations (5 pM–200 nM) were added and incubated at 4°C for 1 h. The labeling of HPG was done by the Iodogen method (Fraker & Speck, 1978) and the specific activity of the labeled HPG was 3.8 × 10⁶ cpm/mg. After adsorption, the wells were aspirated and then washed twice with 200 μL of PBS containing 0.5% Tween-20. They were then cut and subjected to gamma-counting on a Packard Cobra Autogamma counter.

Binding of SK/SK mutants onto HPG-agarose

The binding characteristics of SK mutants with HPG-agarose was studied by loading ~150 μg of either nSK or a mutant onto the affinity matrix packed in a 1 mL bed volume column equilibrated with 50 mM Tris-Cl, pH 7.5, essentially by the procedure described by Rodriguez et al. (1992). The column was then washed with five bed-volumes of equilibrating buffer. The loading and washing steps were carried out in less than 10 min. Specifically bound protein was eluted with 8 M urea in the same buffer. The chromatography was carried out at 4°C. The protein in each fraction (0.75 mL) was then estimated (Bradford, 1976).

CD measurements

Far-UV CD spectra of the proteins in the wave-length range of 197–250 nm were recorded on a Jobin Yvon Auto Dicrograph Mark-V model using a 0.1 cm path length cell, at protein concentrations of 0.15 mg/mL in PBS, pH 7.4. All samples were centrifuged at 10,000 × g for 10 min, and the clear supernatants used for recording the spectra. Appropriate buffer baselines were subtracted in all cases. A mean residue weight of 114 for SK was used (Radek & Castellino, 1989), and the final spectrum was an average of 10 scans.

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