

Structure of the active core of human stem cell factor and analysis of binding to its receptor Kit

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Stem cell factor (SCF) is an early-acting hematopoietic cytokine that elicits multiple biological effects. SCF is dimeric and occurs in soluble and membrane-bound forms. It transduces signals by ligand-mediated dimerization of its receptor, Kit, which is a receptor tyrosine kinase related to the receptors for platelet-derived growth factor (PDGF), macrophage colony-stimulating factor, Flt-3 ligand and vascular endothelial growth factor (VEGF). All of these have extracellular ligand-binding portions composed of immunoglobulin-like repeats. We have determined the crystal structure of selenomethionyl soluble human SCF at 2.2 Å resolution by multiwavelength anomalous diffraction phasing. SCF has the characteristic helical cytokine topology, but the structure is unique apart from core portions. The SCF dimer has a symmetric 'head-to-head' association. Using various prior observations, we have located potential Kit-binding sites on the SCF dimer. A superimposition of this dimer onto VEGF in its complex with the receptor Flt-1 places the binding sites on SCF in positions of topographical and electrostatic complementarity with the Kit counterparts of Flt-1, and a similar model can be made for the complex of PDGF with its receptor.

Keywords: crystal structure/helical cytokine/hematopoiesis/multiwavelength anomalous diffraction (MAD)/stem cell factor

Introduction

Stem cell factor (SCF) is an early-acting hematopoietic cytokine that binds at the cell surface to its receptor, Kit, whereby it produces other biological effects in addition to those on hematopoiesis (see reviews by Galli *et al.*, 1994; Lev *et al.*, 1994; Besmer, 1997; Broudy, 1997). SCF, which is produced by various fibroblast-type cells

including bone marrow stromal cells, has also been called Kit ligand (KL), mast cell growth factor (MGF) and steel factor. The biochemistry and molecular biology that identified SCF and Kit as a ligand–receptor pair were preceded by an array of elegant animal biology studies that anticipated the underlying molecular mechanisms responsible for the genetics (Russell, 1979). Mice with mutations in the steel (*Sl*) locus (gene for SCF) or in the dominant-spotting (*W*) locus (*c-kit*, the gene for Kit) show complex phenotypes that include varying degrees of macrocytic anemia, sterility, lack of coat pigmentation and mast cell deficiency. Kit mutations in man are responsible for the autosomal dominant congenital pigmentation disorder, piebaldism. Consistent with these phenotypes, in the last 10 years a host of *in vitro* and *in vivo* experiments have clearly established Kit-mediated roles for SCF in early stages of hematopoiesis, in gametogenesis, in melanocyte proliferation and function and in mast cell proliferation, maturation and activation (Galli *et al.*, 1994; Lev *et al.*, 1994; Besmer, 1997; Broudy, 1997). Potential therapeutic applications of SCF include the treatment of anemias, boosting the mobilization of hematopoietic stem/progenitor cells to the peripheral blood for harvest and transplantation, and increasing the efficiency of gene transduction for gene therapy (Galli *et al.*, 1994; McNiece and Briddell, 1995; Glaspy, 1996; Broudy, 1997).

SCF is expressed as membrane-associated forms of either 248 or 220 amino acids (Galli *et al.*, 1994; Lev *et al.*, 1994; Besmer, 1997; Broudy, 1997). The two forms are a consequence of alternative mRNA splicing that includes or excludes exon 6. Exon 6 encodes a proteolytic cleavage site such that soluble SCF^{1–165} is released from the 248 amino acid precursor. Residues 166–189 represent a tether to the membrane, residues 190–221 represent a hydrophobic transmembrane segment and residues 222–248 represent a cytoplasmic domain. The 220 residue form lacks the cleavage site and tends to remain membrane bound. Soluble SCF exists as a non-covalently associated homodimer (Arakawa *et al.*, 1991). Each SCF monomer contains two intra-chain disulfide bridges, Cys4–Cys89 and Cys43–Cys138 (Langley *et al.*, 1992). The N-terminal 141 residues of SCF have been identified as a functional core, SCF^{1–141}, which includes the dimer interface and portions that bind and activate the receptor Kit (Langley *et al.*, 1994).

It has been proposed that SCF is a member of the helical cytokine structural superfamily characterized by a double-crossover four-helix bundle topology (Bazan, 1991). Three-dimensional structures are known for many of the family members and, from a comparison of the structures and sequences, the members have been classified into three subgroups (Sprang and Bazan, 1993): short-chain, long-chain and interferon-like.

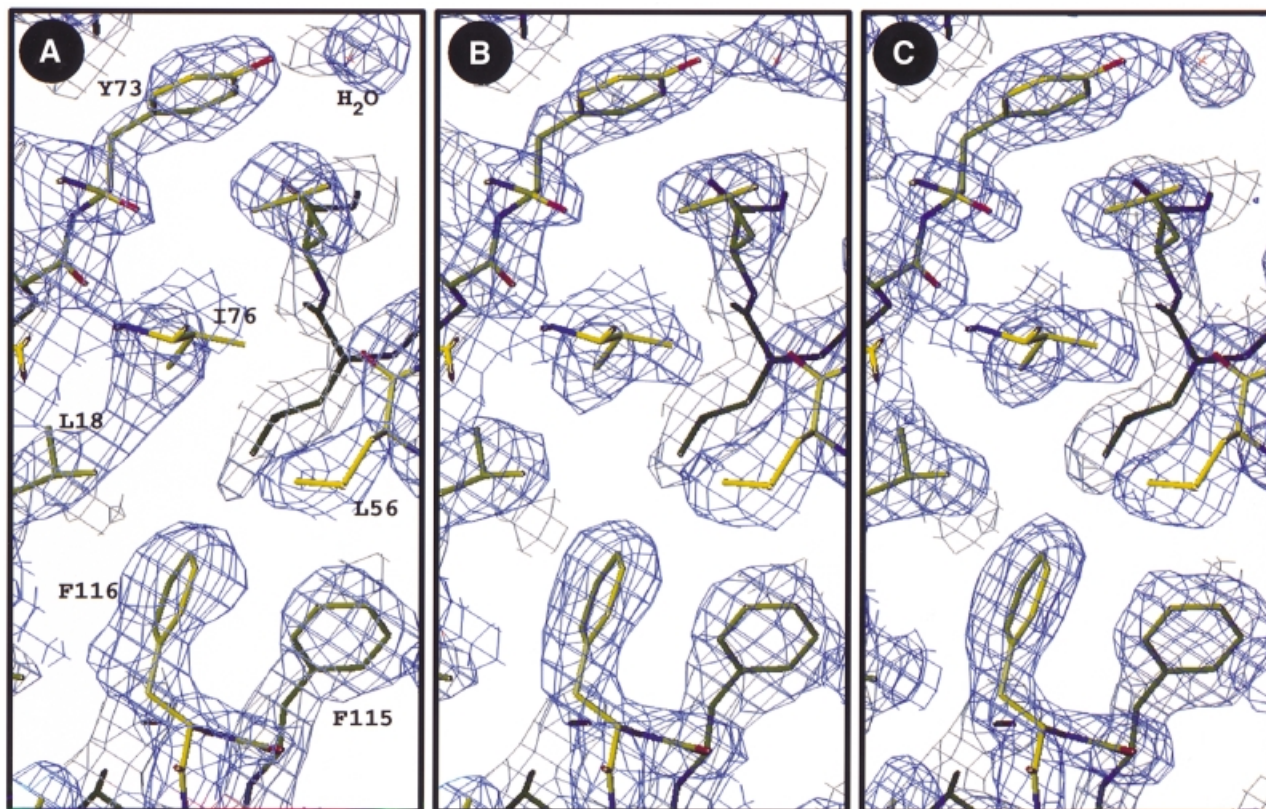


Fig. 1. Representative electron density distributions. (A) MAD-phased experimental map calculated at 2.3 Å resolution. (B) The experimental map after 4-fold averaging. (C) The current $2F_o - F_c$ map superimposed with the model refined at 2.2 Å resolution. Each map is contoured at 1.0σ . Figures were drawn by O (Jones *et al.*, 1991).

Most helical cytokines signal through members of the hematopoietic cytokine receptor superfamily, which are without intrinsic kinase activity (Heldin, 1995). SCF, in contrast, signals through a class III receptor tyrosine kinase (i.e. Kit). This class of kinases also includes the receptors for platelet-derived growth factor (PDGF), macrophage colony-stimulating factor (M-CSF) and Flt-3 ligand, and it is related to the class V receptor tyrosine kinases (Flt-1, Flk-1/KDR and Flt-4) for vascular endothelial growth factors (VEGFs) and less closely to the class IV receptors (FGFRs) for fibroblast growth factors (FGFs) (Fantl *et al.*, 1993; Heldin, 1995; Rousset *et al.*, 1995). The receptors in these classes have ‘split’ kinase domains intracellularly and multiple immunoglobulin (Ig)-like domains extracellularly.

The structures of PDGF (Oefner *et al.*, 1992), M-CSF (Pandit *et al.*, 1992) and VEGF (Muller *et al.*, 1997) have all been determined by X-ray crystallography, as has the complex of VEGF with domain 2 of its receptor, Flt-1 (Wiesmann *et al.*, 1997). Recently, instructive structures have also been reported for FGF2 bound to a two-domain fragment of FGFR1 (Plotnikov *et al.*, 1999) and for the comparable complex between FGF1 and FGFR2 (Stauber *et al.*, 2000). The ligands for the class III and class V receptors are all dimeric, whereas the class IV receptors use both FGF monomers and heparin as ligands. SCF, like the other dimeric ligands, initiates signal transduction by direct dimerization of its receptor, Kit, and the two juxtaposed receptors undergo tyrosine autophosphory-

lation (Heldin, 1995; Broudy, 1997), which initiates downstream intracellular signaling.

Here we report the crystal structure of the core fragment of recombinant human stem cell factor, SCF¹⁻¹⁴¹, as determined at 2.2 Å resolution from multiwavelength anomalous diffraction (MAD) measurements. Incorporating data from mutagenesis and other structure–function studies, we locate putative receptor-binding sites on the surface of the symmetric SCF dimer. From a comparison of these results with structural and functional data for the related ligand–receptor systems, we model the complex of SCF with the receptor Kit and suggest a similar mode of association between other class III and class V receptors and their ligands.

Results and discussion

Structure determination

Both native and selenomethionyl (SeMet) human SCF¹⁻¹⁴¹ were expressed as recombinant proteins in *Escherichia coli* (Langley *et al.*, 1994). Crystals grew in space group $P2_12_12_1$ with four SCF subunits and 39% solvent in the asymmetric unit. Our attempts to solve the structure of SCF¹⁻¹⁴¹ by molecular replacement from other cytokine structures gave good rotation solutions, but no significant translation function peaks. We then evaluated experimental phases in a MAD experiment. Four-wavelength data were measured from a single, frozen SeMetSCF¹⁻¹⁴¹ crystal and analyzed with MADSYS (Hendrickson,

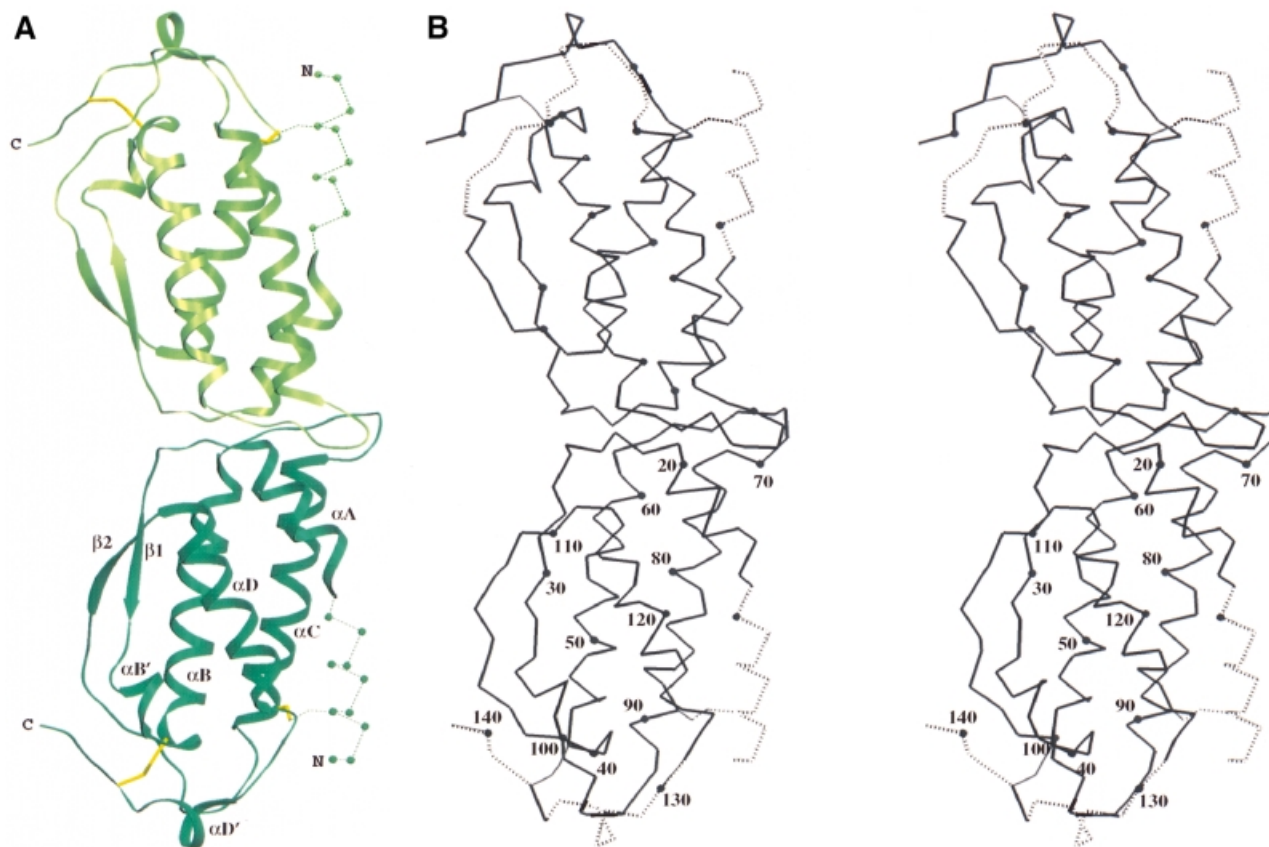


Fig. 2. Overall structure of an SCF dimer. (A) Ribbon diagram. (B) C_{α} stereodiagram of the AB dimer. Figures were drawn by SETOR (Evans, 1993).

1985). Twelve selenium sites were found in four congruent sets associated with the respective SCF subunits in the crystal. A MAD-phased electron density map was calculated at 2.3 Å resolution (Figure 1A) and improved by molecular averaging (Figure 1B) and refinement (Figure 1C).

An atomic model was fitted to the experimental maps and refined at 2.2 Å resolution to an R -value of 0.199 ($|F| > 2\sigma$) with stereochemical ideality typified by the r.m.s. deviation from bond ideality of 0.016 Å. There are no residues in energetically disallowed regions of the Ramachandran plot. This model for SeMetSCF¹⁻¹⁴¹ has 3804 non-hydrogen atoms from 448 amino acid residues, 264 water molecules, three Ca²⁺ ions and one polyethylene glycol (PEG) moiety. All four polypeptide chains (designated A, B, C and D) are sufficiently disordered before residue 11 to preclude modeling of this portion, and none of them is fully ordered through to the end. Specifically, A92–103, B130–136, B139–141, C92–103, C127–141, D91–103 and D128–141 are all disordered. This disorder is such that, of the eight disulfide bridges, only two are seen. To test whether the reducing agent used to crystallize SeMetSCF¹⁻¹⁴¹ (see Materials and methods) might have broken these bonds and caused the disorder, we also refined the structure of native SCF¹⁻¹⁴¹ crystallized without reducing agent. The two crystals were nearly isomorphous, and the two structures showed the same pattern of order–disorder.

Structure of SCF

The four independent SCF subunits in the crystal are similar but distinctive, and identification of the AB and CD pairs as molecular dimers is unmistakable. None of the SCF subunit copies is complete, but each flexible portion except for the N-terminus is stabilized by lattice contacts to another subunit. Thus, through the combination of chains A and B, there are images for all residues except 1–10, and the position of Cys89, to which Cys4 must bridge, determines the approximate course of this disordered segment. The overall structure of this composite SCF dimer is shown in Figure 2A, and the C_{α} backbone for the actual AB dimer is drawn in stereo in Figure 2B. Topologically, the SCF structure is similar to that of other short-chain helical cytokines (Rozwarski *et al.*, 1994), with a core of four helices (αA , αB , αC and αD) and two β -strands, $\beta 1$ between αA and αB and $\beta 2$ between αC and αD . Apart from the tight $\beta 2$ – αD connection, however, the segments outside these core elements are unique in conformation if not in length. In particular, there is an additional one-turn helix, $\alpha B'$, between $\beta 1$ and αB , there is an exceptional hairpin loop between αB and αC at the dimer interface, and there is another extra one-turn helix, $\alpha D'$, in the C-terminal extension. The bounds of secondary structure elements are given in Figure 3.

The core SCF dimer has its subunits arranged in a head-to-head manner with the opposed four-helix bundle axes nearly coincident (Figure 2). This gives the molecule

an elongated shape, $\sim 85 \times 30 \times 20$ Å. Approximately 855 Å² of surface area from each protomer is buried into the dimer interface. The interface is dominated by contacts from the C-terminal end of αA and the αA - $\beta 1$ connection of one monomer to the αB - αC loop of the other monomer (Figure 2), and the reciprocal pair is related by an approximate dyad axis of symmetry. The actual symmetry operators have rotational and translational components of 176.3° and 0.33 Å, respectively, for the AB dimer and 177.4° and 0.04 Å, respectively, for the CD dimer. The two dimers thereby deviate significantly and similarly (with A matched to C and B to D) from true 2-fold symmetry. Nevertheless, since interatomic contacts at the interface are symmetric, we presume that these deviations reflect flexibility rather than inherent asymmetry.

The crystal structure is compatible with solution biochemistry. Consistent with the relative rates of *in vitro* oxidation of methionyl residues (Hsu *et al.*, 1996), Met36 and Met48 are buried in the hydrophobic core whereas Met27 is solvent accessible. Furthermore, as predicted on the basis of fluorescence spectroscopy studies (Arakawa *et al.*, 1991), Trp41 is buried within the hydrophobic core.

Natural SCF and Chinese hamster ovary (CHO) cell-expressed recombinant SCF are heavily glycosylated, with both N- and O-linked carbohydrates. All four potential N-linked sites of human SCF¹⁻¹⁶⁵ are in the SCF¹⁻¹⁴¹ portion that we have crystallized (Langley *et al.*, 1992; Lu *et al.*, 1992). Although the recombinant proteins expressed in bacteria are non-glycosylated, both human and rat SCF expressed in *E. coli* and then refolded *in vitro* have native structures, as judged by biophysical methods and *in vitro* biopotency assays (Arakawa *et al.*, 1991; Langley *et al.*, 1992). The crystal structure of our recombinant SCF is compatible with the glycosylation pattern found for SCF from mammalian cells. Thus, the potential site at Asn72, which is unglycosylated in both human and rat SCF expressed from mammalian cells, is buried in the dimer interface, whereas the site at Asn120, which is fully glycosylated in both species, is solvent accessible. Other sites (Asn65 in both human and rat, human Asn93 and rat Asn109) are glycosylated in some molecules but not others. These sites are also accessible in the atomic model. Asn93 is located in the highly flexible region between αC and $\beta 2$, and its side chain is disordered.

Although natural SCF is a non-covalently associated dimer, recombinant human SCF produced in *E. coli* can fold alternatively *in vitro* into a covalently linked dimer. These dimers have Cys4-Cys89' and Cys43-Cys138' intermolecular disulfide bonds (Lu *et al.*, 1996). The disulfide-linked SCF dimer and native non-covalently associated SCF dimer are similar with regard to biochemical and biophysical properties, biopotency and receptor-binding affinity. It was proposed that the disulfide-linked dimer arises from a double swap of αA and αD helices between the monomers (Lu *et al.*, 1996). The crystal structure of SCF, however, suggests that a single swap at the αB - αC loop near residue 68 is more likely.

Comparison with other short-chain helical cytokines

Although SCF has the characteristic features of the short-chain helical cytokines, as among other members, both sequence and structure are highly divergent. If anything,

SCF resembles the others less than they resemble one another (Table I). Our comparison of SCF with other short-chain helical cytokine structures [granulocyte-macrophage colony-stimulating factor (GM-CSF) (Diederichs *et al.*, 1991), M-CSF (Pandit *et al.*, 1992), interleukin (IL)-2 (McKay, 1992), IL-4 (Wlodaver *et al.*, 1992) and IL-5 (Milburn *et al.*, 1993)] shows greatest structural similarity with M-CSF or IL-4, but even here fewer than half of the residues can be superimposed (Table I). Sequence similarities are essentially random. Our structure-based sequence alignment (Figure 3) has pairwise identities ranging from 6.7 to 18.8% (Table I) and not even a single residue in SCF is conserved in all of the others. Nevertheless, the core elements are remarkably similar in structure.

Core portions aside, SCF differs markedly from other short-chain helical cytokines, as indeed they differ from one another (Figure 3; Rozwarski *et al.*, 1994). First, helix αA of SCF is unusually shortened at its N-terminus. Its disordered extension must deviate toward αC , as in M-CSF but not in the others, by virtue of the Cys4-Cys89 disulfide bridge in common with M-CSF. Secondly, the conformation of the αA - $\beta 1$ connection is distinctive as required for the dimer interface, and the $\beta 1$ - αB connection uniquely has $\alpha B'$. Again at the dimer interface, the αB - αC hairpin loop extends out distinctively along the dyad axis. Thirdly, the unusually long αC - $\beta 2$ loop of SCF is both highly flexible (only one ordered copy) and with a unique path when ordered. Finally, the C-terminal extension after αD compares only to that of M-CSF, and then only in its general direction of exit out past αB and the β -strands.

SCF and M-CSF are similar in gene structure, alternative splicing, proteolytic maturation, disulfide bridging, dimer assembly and receptor type (these similarities also extend to the Flt-3 ligand; Lyman and Jacobsen, 1998). Despite negligible sequence identity, an alignment and secondary structure prediction prompted by these relationships (Bazan, 1991) fits the actual structure amazingly well, except for shifts in αB and in the αC - $\beta 2$ loop. Here reality confounds logic; unexpectedly, comparable glycosylation sites (Asn120 in SCF and Asn122 in M-CSF) are displaced by one helical turn, and comparable disulfide bridges (Cys43-Cys138 in SCF and Cys48-Cys139 in M-CSF) are not superimposable structurally (Figure 4).

Comparison with other cytokine dimers

Helical cytokines dimerize in various ways (Sprang and Bazan, 1993). Among the five dimeric helical cytokines for which crystal structures have been described previously [M-CSF, IL-5, ciliary neurotrophic factor (CNTF), interferon- γ (IFN- γ) and IL-10], only IFN- γ and IL-10 are similar dimers. These latter two have a 'tip-to-tip' packing with helix axes approximately perpendicular. Otherwise, the only salient feature in common is having the subunits oriented with bundle axes aligned in parallel and helix dipoles positioned to compensate. There is 'head-to-head' packing of the four-helix bundles in M-CSF, 'tail-to-tail' packing in IL-5 and 'side-to-side' packing in CNTF. Moreover, IFN- γ , IL-10 and IL-5 are all interdigitated dimers with helices swapped between subunits.

SCF, in keeping with its relationship to M-CSF, is a non-interdigitated 'head-to-head' dimer (Figure 4). The

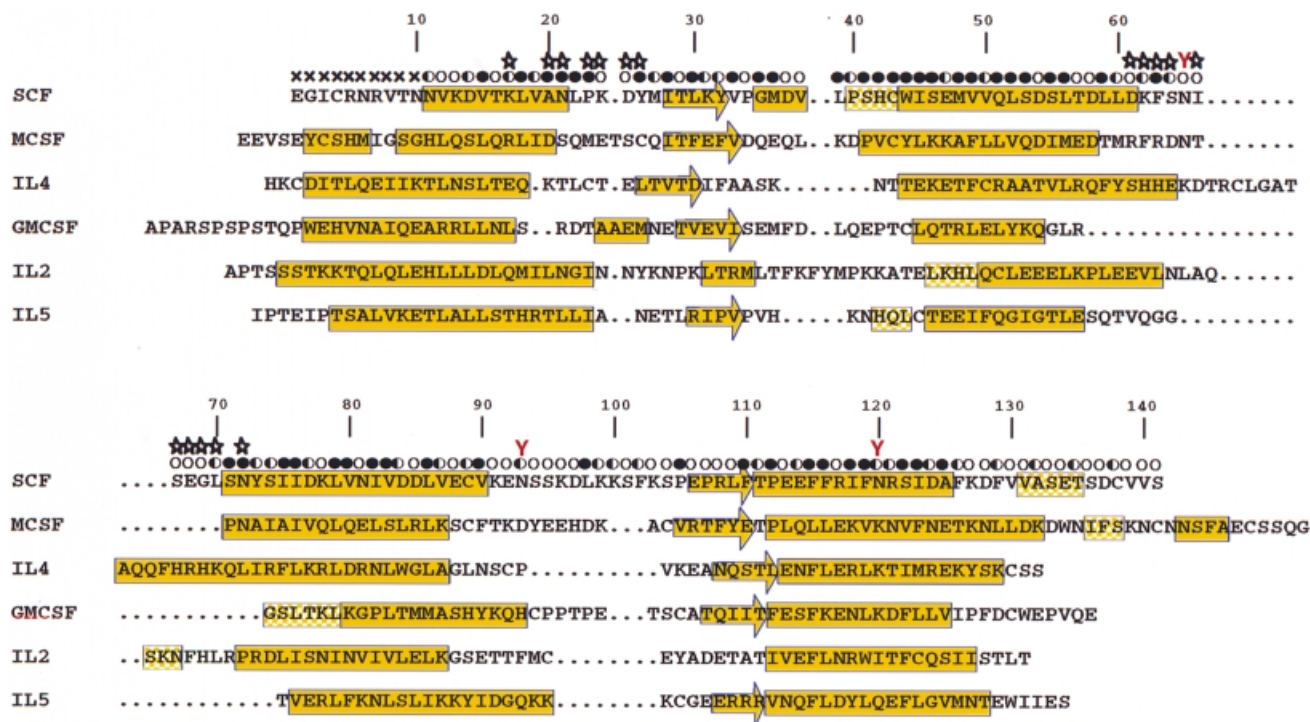


Fig. 3. Structure-based sequence alignment of SCF with other short-chain helical cytokines of human species. Dots denote gaps. The secondary structure elements were assigned according to the output of the PROCHECK program (Laskowski *et al.*, 1993) except the helix assignment for residues 35–38, which was identified by inspection of the hydrogen bond pattern. Secondary structures are shown in yellow with filled boxes referring to α -helices, half-filled boxes to 3_{10} -helices and arrows to β -strands. The solvent accessibility of the SCF dimer is indicated for each residue by an open circle if the fractional solvent accessibility is >0.4 , a half-filled circle if it is 0.1 – 0.4 , and a filled circle if it is <0.1 . Residues at the SCF dimer interface are identified by stars, and the N-linked glycosylation sites by red Ys above the Asn residues.

Table I. Structural and sequence comparisons of short-chain helical cytokines

	SCF	M-CSF	IL-4	GM-CSF	IL-2	IL-5
SCF		14.1 (13.0)	12.7 (12.3)	12.5 (23.5)	18.8 (16.4)	6.7 (21.1)
M-CSF	64 (1.76)		14.8 (18.9)	13.8 (18.3)	17.5 (17.1)	10.5 (18.6)
IL-4	63 (1.58)	54 (1.82)		26.6 (25.0)	14.5 (22.2)	18.9 (18.9)
GM-CSF	48 (1.63)	58 (1.81)	64 (1.56)		9.8 (26.0)	20.4 (14.7)
IL-2	48 (1.70)	57 (1.58)	69 (1.33)	61 (1.48)		14.5 (22.2)
IL-5	45 (1.70)	38 (1.72)	53 (1.32)	49 (1.33)	62 (1.37)	

Structural comparisons and sequence comparisons between the short-chain helical cytokines are given in the lower left and upper right triangles, respectively. Structural comparisons are given as the maximum number of equivalent α -carbon atoms between two short-chain helical cytokines, and the r.m.s. deviation (\AA) (in parentheses). Sequence comparisons are given as the percentage of sequence identity from sequence alignment based on structural superimposition, and that based on the sequence alignment from the BESTFIT program of the GCG package (in parentheses). The latter alignment is based only on maximizing the percentage of identity, similarities and length of the matching sequences, and the sequences submitted to the BESTFIT program were restricted within the region as defined in the PDB files, including the disordered residues. With the advantage of the relatively large number of independent data points (15 pairs), we analyzed the correlation between sequence identity and structural deviation. Without any restriction of structural alignment, the correlation coefficient (C) between structural deviation and sequence identity is -0.21 and the Student's t probability (P) is 0.44 , suggesting little correlation between a specific sequence and the tertiary fold. With the restriction of structural alignment, however, C is -0.30 and P is 0.28 , indicating that the structure-based sequence identity and structural deviation are weakly connected (as also observed in another highly diverged protein family, hemoglobin; Aronson *et al.*, 1994).

interfaces between protomers are completely different, however. One αA – $\beta 1$ loop of M-CSF is situated between the αA – $\beta 1$ and αB – αC loops of the other protomer, whereas in SCF each αA – $\beta 1$ loop interacts only with the αB – αC loop of the partner. This staggered mode of M-CSF dimerization (Figure 4B) is dictated by the position of the Cys31–Cys31' intermolecular disulfide bond in M-CSF. The dyad axes are oriented similarly in

the two cases (perpendicular to the bundle axis and parallel to the $\alpha A\alpha D$ and $\alpha B\alpha C$ helix planes), but whereas the dyad axis in SCF nearly intersects the bundle axis, that in M-CSF is offset toward the $\alpha A\alpha D$ helix pair (Figure 4). Thus, when one protomer of an SCF dimer is superimposed onto a protomer from M-CSF, the superimposition of the two mates requires a translation of 3.8 \AA but a rotation of only 4.7° .

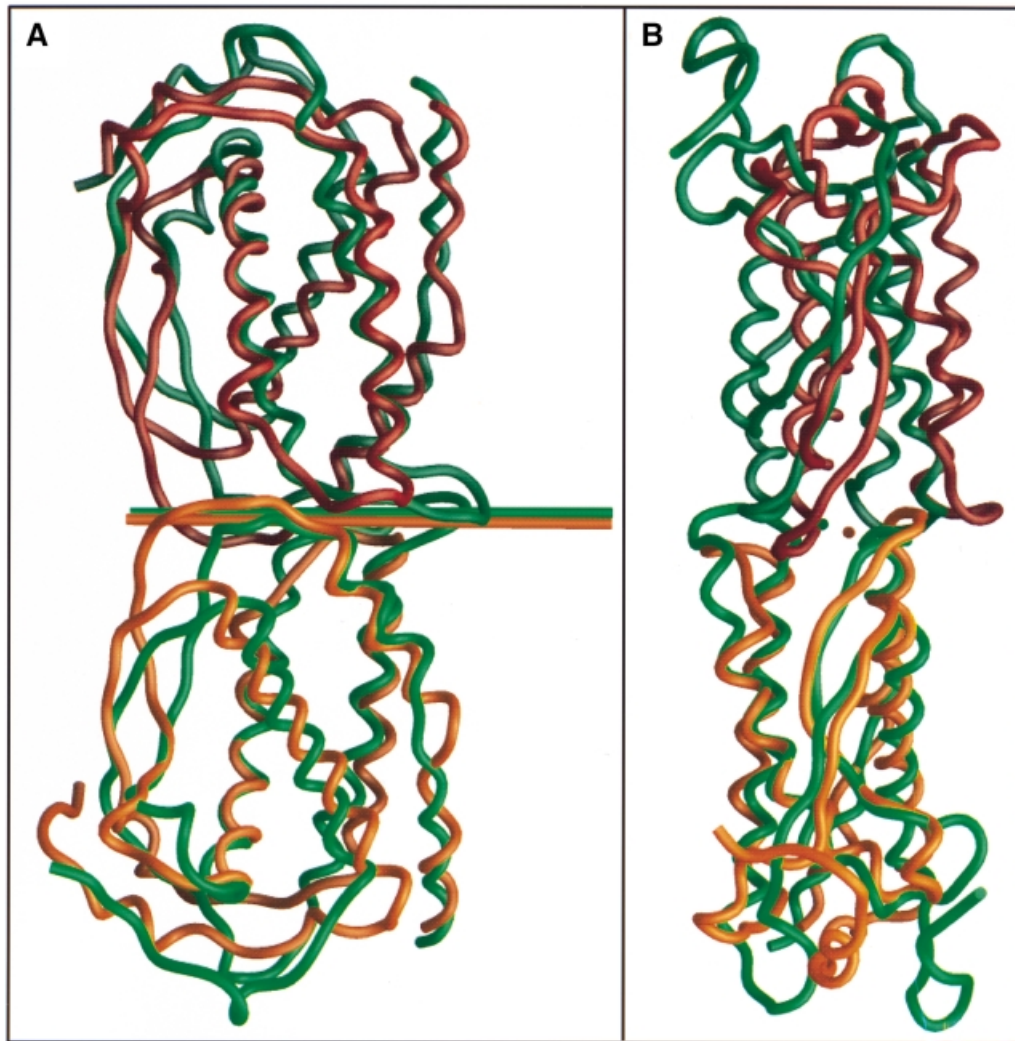


Fig. 4. Comparison of SCF dimer (shades of green) and M-CSF dimer (shades of brown). (A) View as in Figure 2. (B) View perpendicular to that in (A), along the dyad axis of M-CSF. Symmetry axes are shown as lines in (A) and dots in (B). Figures were drawn by GRASP (Nicholls *et al.*, 1991).

Location of the binding site for the receptor Kit

SCF binds with high affinity (nM range) to its receptor (Philo *et al.*, 1996; Broudy, 1997). Various structure–function studies and analyses help to define residues of SCF that may be involved in this binding. These studies include mutagenesis, immunochemical mapping, comparative analyses of inter-species ligand–receptor interactions and analyses of glycosylation.

From studies of truncation and point mutants, Langley *et al.* (1994) demonstrated that the N-terminal residues 1–4 and 1–10 and the Cys4–Cys89 disulfide bond are required for receptor binding and bioactivity, and that the Cys43–Cys138 disulfide bond and C-terminal residues past 127 are not required for receptor binding but may have some roles in bioactivity. Moreover, alterations at Asn10 and Asn11 brought about by chemical isomerization or by mutagenesis have positive or negative effects on bioactivity, depending on the substitution (Hsu *et al.*, 1998). A quadruple mutant of SCF (Arg121Asn, Asp124Asn, Lys127Asp and Asp128Lys) was found to be defective in bioactivity (Matous *et al.*, 1996). The molecular cause of this

deficiency may be specific to Lys127 or due to indirect electrostatic effects. Arg121 and Asp124 are adjacent to the main N-linked glycosylation site, which is not involved in binding (see below), and Asp128 is absent in the 1–127 truncation mutant that retains full receptor-binding activity (Langley *et al.*, 1994). Moreover, a study of human–murine SCF chimeras narrowed the important receptor recognition epitopes to within residues 1–35 and 79–97 (Matous *et al.*, 1996), and the epitope of a neutralizing monoclonal antibody was mapped to the region of residues 60–95 (Mendiaz *et al.*, 1996) and 79–97 (Matous *et al.*, 1996).

Although SCF molecules from different mammalian species are very similar (>75% identity), there are substantial differences in inter-species receptor activation. Human SCF activates mouse Kit very poorly, rodent SCF has only slightly lower potency than human SCF in binding/activating human Kit (Martin *et al.*, 1990; Lev *et al.*, 1992), and dog SCF activates human Kit slightly better than does human SCF itself (K.E.Langley, unpublished data). It is likely that the receptor-binding regions involve residues that are different between man and mouse



Fig. 5. Sequence alignments of SCF from human, mouse, rat and dog (Anderson *et al.*, 1990; Huang *et al.*, 1990; Martin *et al.*, 1990; Shull *et al.*, 1992). The residues that are conserved in human and dog but different from rat and mouse are shadowed in yellow. Five regions of divergent sequence are identified (Roman numerals). Dots denote gaps, and dashes indicate residues identical to the human residues.

but conserved between man and dog. These residues can be classified into five groups in the sequence (Figure 5). Most residues in group III (45–58) are buried and those in group II (24–34) are close to the dimer interface. The residues in groups I (1–15), IV (80–117) and V (130–140) are more likely to be involved in direct receptor binding.

Human SCF expressed in CHO cells is ~30% carbohydrate by weight (Arakawa *et al.*, 1991). The main glycosylation site is at Asn120 (Langley *et al.*, 1992; Lu *et al.*, 1992). Glycosylation at this site, which is near the center of the α D helix, does not appear to influence biological activity; therefore, the area around this residue cannot be involved in receptor binding. Glycosylation of human SCF at either Asn65 or Asn93 lowers the biological activity ~10-fold; therefore, these residues may be near but not directly at the binding site.

Taken together, these observations indicate that the receptor-binding site may include residues from the first few N-terminal residues, the 79–95 region (mainly located on the α C helix) and the C-terminal end of α D (around 127). These regions are contiguous on the SCF surface in our atomic model. The putative receptor-binding site of M-CSF was mapped to a similar region (Taylor *et al.*, 1994).

Structural characteristics of SCF-Kit and related ligand-receptor complexes

Class III and class V receptor tyrosine kinases are distinguished from one another by the number of Ig repeats in their extracellular ligand-binding portions (five for class III and seven for class V) and by the length of kinase insert. These Ig-like receptors share similar signal transduction pathways, chromosomal localization and gene organization (Rousset *et al.*, 1995), but their ligands

come with completely unrelated topologies as typified by PDGF and VEGF (cystine knot) on the one hand, versus M-CSF, SCF and Flt-3 ligand (helical cytokine) on the other. Even receptors of the same class have unrelated ligands; thus, both SCF and PDGF use class III receptors and VEGF and Flt-3 ligand use class V receptors. The amino acid sequences of the ligands are extremely dissimilar even when the fold is the same, as for PDGF versus VEGF (25% identity) and M-CSF versus SCF (14% identity).

Although Ig-like receptors have very similar kinase portions (~70% amino acid sequence identity between III and V), their Ig-like domains are dissimilar in sequence both between repeats within a molecule and also at comparable positions between different receptors (Rousset *et al.*, 1995). Nevertheless, there are features of the receptor–ligand interaction that the class III and class V receptors have in common. First, for every studied example, the ligand-binding function has been localized to the first three Ig-like domains and, where defined further, to domains D2 and D3 specifically (Heidaran *et al.*, 1990; Blechman *et al.*, 1993; Lev *et al.*, 1993; Wang *et al.*, 1993; Davis-Smyth *et al.*, 1996; Barleon *et al.*, 1997). Secondly, the ligands for all of these receptors are functional as dimers; M-CSF, VEGF and PDGF are covalently linked dimers, while SCF and Flt-3 ligand are non-covalently associated dimers. In each case, signaling occurs through ligand-mediated receptor oligomerization (Heldin, 1995). For SCF-Kit, it has been shown directly by biophysical methods that complexes containing two SCF subunits and two Kit extracellular domain molecules can form in solution (Philo *et al.*, 1996).

The structure of domain D2 of receptor Flt-1 in complex with VEGF (Wiesmann *et al.*, 1997) provides a template

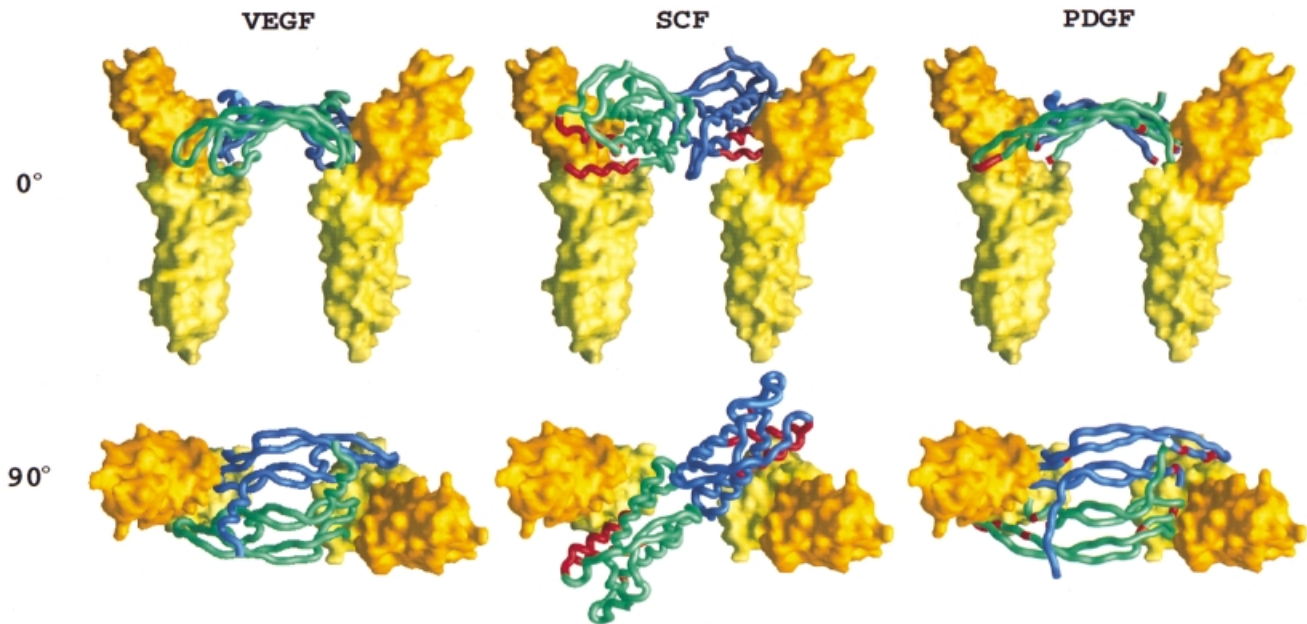


Fig. 6. Ligand (worm structures)–receptor (yellow structures) models of: VEGF–Flt-1, SCF–Kit and PDGF–PDGF receptor. The VCAM-1 structure (PDB code: 1vca) was used, without any modification, to approximate the receptor models. Receptor models are presented as yellow surfaces. Background portions of ligands are colored light blue for one protomer and green for the other; receptor-interacting residues identified from site-directed mutagenesis experiments [VEGF (Muller *et al.*, 1997), PDGF (Fenstermaker *et al.*, 1993)] and other experimental data (SCF; see text) are colored magenta. Figures were drawn by GRASP (Nicholls *et al.*, 1991).

for ligand interactions with PDGF-related receptors. Wiesmann *et al.* (1997) modeled the interaction of VEGF with D1D2D3D4(Flt-1) and discussed the likelihood that other ligand complexes with class III and class V receptors may be similar. In light of the structure of SCF and the identified location of receptor-binding sites, we have modeled the SCF–Kit complex by analogy.

The D2(Flt-1) domain is similar in structure to telokin, as predicted (Harpaz and Chothia, 1994), and thereby also to both domains in the structure of vascular cell adhesion molecule (VCAM)-1 (Jones *et al.*, 1995). To test the validity of VCAM-1 as a model for D2D3(Flt-1) and D2D3(Kit), we used a prediction-based threading program (Fisher and Eisenberg, 1996) to thread the sequences of the Ig-like domains of Flt-1 and Kit into the telokin and VCAM-1 structures. Fits were achieved with moderate to very high confidence of similarity. The resulting structure-based sequence alignment of D2D3(Kit) with the VCAM-1 template (five gaps) has a continuous domain boundary, and residues Cys151 and Cys183 in D2(Kit) are positioned properly to make an additional disulfide bridge between strands C and F.

We next constructed a model of the VEGF–D2D3(Flt-1) receptor complex from a rigid-body superimposition of VEGF (Muller *et al.*, 1997) and VCAM-1 so as to mimic the reported VEGF–D2(Flt-1) structure (Wiesmann *et al.*, 1997). Then, keeping the dyad symmetric receptor pair fixed, we successively replaced VEGF with other Ig-like receptor ligands of known three-dimensional structure: PDGF (Oefner *et al.*, 1992), M-CSF (Pandit *et al.*, 1992) and SCF (this work). Each was placed on the dyad axis and positioned to optimize contacts between the VEGF-binding site on the receptor and the putative receptor-binding regions of the ligands. Remarkably, these disparate dimeric ligands have similar spacings between binding

sites and a satisfactory fit is possible for each (Figure 6). We also constructed simple homology models of the various receptors with changes in the backbone only to accommodate insertions and deletions. The model for SCF complexed with D2D3(Kit) shows a striking electrostatic complementarity between a highly negative binding surface on SCF and a positive surface on Kit (Figures 7A and B). The glycosylation sites on both molecules are also compatible with unimpeded interaction.

The receptor Kit is activated by both soluble and membrane-bound forms of SCF, and signaling from the membrane-bound form appears to have *in vivo* roles (see review by Lyman and Jacobsen, 1998). Moreover, as in the case of D4(Flt-1) (Barleon *et al.*, 1997), D4(Kit) may be involved in inter-receptor contacts in the signaling dimer (Blechman *et al.*, 1995) [although this proposal for Kit has been questioned (Philo *et al.*, 1996; Lemmon *et al.*, 1997)]. The model that we have constructed for the SCF–Kit complex is compatible with these properties (Figure 7). The C-termini of the SCF dimer are directed oppositely from those of Kit, as would be appropriate for a cell–cell contact, and the receptor units would cross naturally at D4. It is noteworthy that the ligands of two other Ig-like class III receptors also have membrane-bound forms (M-CSF and Flt-3 ligand) (Kawasaki and Ladner, 1990; Lyman and Jacobsen, 1998).

Materials and methods

SCF expression, purification and analyses

Human SCF^{1–141} was expressed recombinantly in *E. coli* as described previously (Langley *et al.*, 1994). For SeMetSCF^{1–141}, the expression vector was transfected into the methionine auxotrophic *E. coli* strain FM5. Fermentation was carried out at 30°C in 8 l of minimal medium consisting of ammonium sulfate (10 g/l), glucose (5 g/l), methionine (0.125 g/l), phosphate salts, magnesium, citric acid, trace metals and vitamins. When

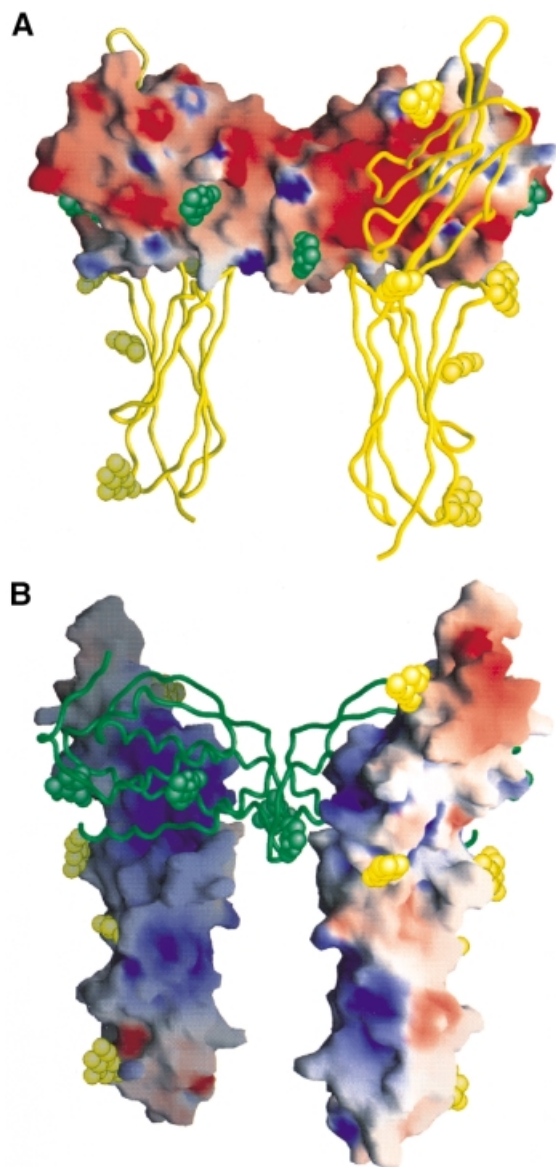


Fig. 7. Electrostatic and carbohydrate surfaces of SCF and homology-modeled receptor Kit. (A) Electrostatic surface of SCF and worm of D2D3(Kit). (B) Electrostatic surface of the Kit fragment and worm of SCF. Negative potential is colored red and positive potential blue, with greatest saturations at -10 and $+10$ kT, respectively. Carbohydrate moieties are represented by CPK models of β -D-N-acetylglucose (green for SCF moieties and yellow for potential Kit moieties). Figures were drawn by GRASP (Nicholls *et al.*, 1991).

an OD_{600} of 3–5 was reached, a feed medium was added that consisted of the following components in a total volume of 1 l: 100 g of ammonium sulfate, 450 g of glucose, 2 g of methionine, magnesium, trace metals and vitamins. At an OD_{600} of 12.4, induction medium (1 l containing 100 g of ammonium sulfate, 300 g of glucose and 1 g of SeMet) was added and fermentation proceeded at 30°C . Five hours later (at an OD_{600} of ~ 16), the temperature was raised to 42°C to induce SCF expression and additional SeMet (1 g) was added. Cells were harvested 4 h after the temperature shift (OD_{600} of ~ 16). SeMetSCF¹⁻¹⁴¹ expression was estimated as 0.5 g/l. Both SCF¹⁻¹⁴¹ and SeMetSCF¹⁻¹⁴¹ were purified with minor modifications to previously described procedures (Langley *et al.*, 1992, 1994). Both retain the initiating methionine (or SeMet) residue [position (-1)] (Langley *et al.*, 1994). N-terminal amino acid sequencing was performed as described (Lu *et al.*, 1991). About 90% SeMet was present in SeMetSCF¹⁻¹⁴¹ at each of the Met positions, based on amino acid analysis and N-terminal sequencing results (i.e. lack of recovery of Met residues for SeMetSCF¹⁻¹⁴¹ in comparison with SCF¹⁻¹⁴¹; data not shown).

Crystallization

Crystals were obtained by use of the hanging drop vapor diffusion method under aerobic conditions. The initial crystals were grown by mixing 1 μl of protein solution (44 mg/ml SCF¹⁻¹⁴¹ or 38 mg/ml SeMetSCF¹⁻¹⁴¹ in 10 mM sodium phosphate pH 6.5, 80 mM NaCl) with 1 μl of crystallization reservoir solution. The crystallization reservoir solution included 25% (w/w) PEG400, 240 mM CaCl₂ and 100 mM HEPES pH 7.4 for SCF¹⁻¹⁴¹, and 22% (w/w) PEG400, 220 mM CaCl₂, 100 mM HEPES pH 7.2 and 5–10 mM dithiothreitol (DTT) for SeMetSCF¹⁻¹⁴¹. Crystallization trays were incubated at 20°C and crystals reached full size in ~ 3 days with typical dimensions of $0.5 \times 0.2 \times 0.2$ mm. Microseeding and lower concentrations of DTT solution (~ 2 mM) were needed to reproduce SeMetSCF¹⁻¹⁴¹ crystals subsequently. An extant SeMetSCF¹⁻¹⁴¹ crystal was washed with its reservoir solution and then crushed to produce microseeds, which were stored in 50 μl of a stabilizing solution of 32% (w/w) PEG400, 260 mM CaCl₂, 100 mM HEPES pH 7.4 at room temperature. For microseeding experiments, the seed stock was diluted by 10- to 10 000-fold with crystallization reservoir solution. A 1 μl aliquot of this prepared precipitant was mixed with 1 μl of the protein solution to make the droplet. The crystal for MAD phasing was grown from a crystallization reservoir solution containing 2 mM DTT.

Diffraction measurements

X-ray diffraction data from SCF¹⁻¹⁴¹ crystals were recorded on two Hamlin–Xuong area detectors at 293K at a home source. The data were integrated using the UCSD software package and scaled using AGROVATA and ROTAVATA as implemented in the CCP4 suite (CCP4, 1994). The MAD experiments for SeMetSCF¹⁻¹⁴¹ were conducted at the X4A synchrotron beam line of Brookhaven National Laboratory using Fuji image plates. A single crystal was frozen at 110K using paratone-N (Exxon) as a cryoprotectant. The MAD data were collected at four wavelengths (before the edge, at the SeK edge, at the peak and after the peak) in oscillations of 1.3 – 1.5° without overlap. The SeMetSCF¹⁻¹⁴¹ crystal was oriented such that the *b*-axis was parallel to the oscillation axis, and a mirror geometry was used during data collection. The MAD data were processed using DENZO and Scalepack (Otwinowski, 1993; Gewirth, 1995) (Table II).

Molecular replacement attempts

Structure determination by the molecular replacement method was attempted for the home source data set. The MERLOT (Fitzgerald, 1988) and AMoRe (CCP4, 1994) programs were used with various four-helix bundle structures as search models, and a good rotation solution was obtained. The rotation solution agreed well with the orientation of helical bundles (approximately along the *b*-axis of the unit cell) that was deduced from native Patterson maps. Dissimilarities among the helical cytokines and the multiplicity of subunits (four) hampered detection of any significant translation function peaks.

Phase evaluation

The processed MAD data were passed through the MADSYS programs (Hendrickson, 1985). Algebraic and probabilistic MAD phasing procedures (Hendrickson, 1985; Pähler *et al.*, 1990) were applied for phase determination (Table II). Selenium sites were located using the HASSP program (CCP4, 1994) in F_A Patterson and difference Fourier maps and refined by MADSYS programs. The choice of enantiomer was determined by comparison of the electron density maps computed from the two enantiomeric selenium structures to maximum Bragg spacings of 2.6 Å. The phases were improved by 4-fold non-crystallographic symmetry (NCS) averaging. The rotation–translation matrices of the NCS axes were determined by TOSS (Hendrickson, 1979) from the selenium sites and subsequently refined by LSQRHO (W.A.Hendrickson, unpublished) and RAVE (Kleywegt and Jones, 1994), and the averaging procedure by DM (CCP4, 1994).

Model building and refinement

The initial model of SeMetSCF¹⁻¹⁴¹ was built into the averaged map at 2.3 Å by using program O (Jones *et al.*, 1991). The model includes 98 core residues for each of the four molecules in an asymmetric unit. The remote wavelength after the SeK peak was used for the refinement with the Bijvoet difference applied to Se scattering factors. The *R*-value for this model, before any refinement, was 42.1% in the resolution range 10.0–2.3 Å. NCS restraints were applied during the initial rounds of refinement. After several iterations of least square and simulated annealing refinement with X-PLOR (Brünger *et al.*, 1987) and manual rebuilding against SIGMAA (Read, 1986) and $2|F_o| - |F_c|$ maps, the crystallographic *R*-value is 19.9% for the current model (Table III). The

Table II. MAD data collection and phasing statistics

Data collection (25–2.0 Å) ^a						
Wavelength (Å)	Unique reflections		Completeness (%)	Signal (I/σ)	R_{sym} (%)	
$\lambda_1 = 0.9919$ (pre-edge)	65 810		95.1	18.4	6.7	
$\lambda_2 = 0.9793$ (inflection)	65 759		95.0	16.7	5.8	
$\lambda_3 = 0.9791$ (peak)	65 665		94.9	15.2	6.7	
$\lambda_4 = 0.9686$ (remote)	65 689		94.9	16.0	5.6	
Anomalous diffraction ratios (20–2.6 Å) ^b						
	λ_1	λ_2	λ_3	λ_4	f' (e)	f'' (e)
λ_1	0.035 (0.030)	0.051	0.042	0.035	−4.0	−0.5
λ_2		0.052 (0.029)	0.033	0.051	−10.3	3.8
λ_3			0.070 (0.031)	0.041	−8.1	5.6
λ_4				0.055 (0.030)	−3.9	3.8
MAD phasing (25–2.6 Å) ^c						
$R(\circ F_{\text{T}}) = 0.044$	$R(\circ F_{\text{A}}) = 0.39$		$\langle \Delta(\Delta\Phi) \rangle = 41.6^\circ$		$\langle \sigma(\Delta\Phi) \rangle = 18.7^\circ$	

^aUnique reflections are determined by point group 222 (not mmm) to distinguish Bijvoet-related reflections. $R_{\text{sym}} = 100 \times \sum_{\text{hkl}} \sum_i |I_i - \langle I \rangle| / \sum_{\text{hkl}} \sum_i I_i$, where I_i is the i th measurement of reflection hkl and $\langle I \rangle$ is the weighted mean of all measurements of I .

^bAnomalous diffraction ratios = $\langle \Delta|F|^2 \rangle^{1/2} / \langle |F|^2 \rangle^{1/2}$, where $\Delta|F|^2$ is the absolute value of the Bijvoet (diagonal elements) or dispersive difference (off-diagonal elements), respectively. Values in parentheses are for centric data.

^c $R = \sum_{\text{hkl}} \sum_i |F_i| - \langle F \rangle / \sum_i |F_i|$. $\circ F_{\text{T}}$ is the structure factor due to normal scattering from all the atoms. $\circ F_{\text{A}}$ is the structure factor due to normal scattering from the anomalous scatterers only, and $\Delta\Phi$ is the phase difference between $\circ F_{\text{T}}$ and $\circ F_{\text{A}}$. $\Delta(\Delta\Phi)$ is the difference between two independent determinations of $\Delta\Phi$.

Table III. Lattice and refinement statistics

	SeMetSCF ¹⁻¹⁴¹ (λ_4)	SCF ¹⁻¹⁴¹
Lattice		
space group	$P2_12_12_1$	$P2_12_12_1$
cell constants (a, b, c) (Å)	71.8, 82.6, 88.2	73.0, 84.7, 88.8
Z_{a} ^a	4	4
Refinement ^b		
resolution range (Å)	20.0–2.2	8.0–3.3
completeness (%)	96.6	98.6
unique reflections ^c	49 851	7990
R -value ^d ($ F > 2\sigma$) (%)	19.9	20.8
R_{free} ^e (%)	24.2	27.3
R_{sym} ^f (%)	5.6	15.2
Model parameter		
total non-H atoms	3804	3502
total residues	448	447
total water molecules	264	0
total metal ions	3	0
r.m.s. bond length/angle	0.016/2.5°	0.017/3.0°
average B -factor (Å ²)	32.1	18.7
main-chain r.m.s. B (bond/angle) (Å ²)	1.2/1.6	1.9/2.2
side-chain r.m.s. B (bond/angle) (Å ²)	2.1/2.4	3.0/3.3

^a Z_{a} , number of molecules in the asymmetric unit.

^bThe reflection data higher than the resolution range were not included in the refinement due to poor R_{sym} in these resolution shells.

^cUnique reflections are determined by point group 222 for the SeMetSCF¹⁻¹⁴¹ data set to distinguish Bijvoet-related reflections and by point group mmm for the SCF¹⁻¹⁴¹ data set.

^d R -value = $\sum_{\text{hkl}} | |F_{\text{o}}| - |F_{\text{c}}| | / \sum_{\text{hkl}} |F_{\text{o}}|$

^eA subset of the data (6%) was excluded from the refinement and used for the free R -value calculation.

^f R_{sym} for the SeMetSCF¹⁻¹⁴¹ data set was calculated in the resolution range of 25–2.2 Å and the for SCF¹⁻¹⁴¹ data set in the resolution range 13–3.3 Å.

sites of Ca²⁺ ions, a component of the crystallization medium, were located from a Bijvoet difference Patterson map at the remote wavelength before the SeK edge. The SCF¹⁻¹⁴¹ model was obtained by subjecting the refined SeMetSCF¹⁻¹⁴¹ model to refinement against the area-detector data set from the SCF¹⁻¹⁴¹ crystal using the X-PLOR program (Brünger *et al.*, 1987). The atomic coordinates have been deposited in the Brookhaven Protein Data Bank with accession code of 1scf.

Structure analysis

Solvent accessibilities were defined compared with the corresponding Gly-X-Gly peptide (Shrake and Rupley, 1973), as calculated by X-PLOR (Brünger *et al.*, 1987). Structural superimpositions were based on α -carbon atoms alone, with r.m.s. deviations calculated only from atom pairs identified as equivalent. The coordinates were taken from the Brookhaven Protein Data Bank with entry codes: M-CSF, 1hmc (Pandit

et al., 1992); IL-4, 1rcb (Wlodaver *et al.*, 1992); GM-CSF, 1gmf (Diederichs *et al.*, 1991); IL-2, 3ink (McKay, 1992); and IL-5, 1hul (Milburn *et al.*, 1993). Initial segments of equivalence between two structures were defined according to equivalent secondary structure elements. These structures were then superimposed using the program TOSS (Hendrickson, 1979), and the number of equivalent atoms was extended using the Lsq_imp command in program O (Jones *et al.*, 1991) with the criteria of a cut-off distance of 3.0 Å and a minimum fragment length of three consecutive residues. Different initial equivalent segments did give different results in the structural alignment, as Rozwarski *et al.* (1994) observed in their study. We tried several initial sets of equivalent segments for each alignment and retained the one that generated the greatest number of equivalent atoms after the Lsq_imp extension.

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