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Insulin Fibrillation and Protein Design: Topological Resistance of Single-Chain Analogs to Thermal Degradation with Application to a Pump Reservoir

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Abstract

Insulin is susceptible to thermal fibrillation, a misfolding process that leads to nonnative cross-β assembly analogous to pathological amyloid deposition. Pharmaceutical formulations are ordinarily protected from such degradation by sequestration of the susceptible monomer within native protein assemblies. With respect to the safety and efficacy of insulin pumps, however, this strategy imposes an intrinsic trade-off between pharmacokinetic goals (rapid absorption and clearance) and the requisite physical properties of a formulation (prolonged shelf life and stability within the reservoir). Available rapid-acting formulations are suboptimal in both respects; susceptibility to fibrillation is exacerbated even as absorption is delayed relative to the ideal specifications of a closed-loop system. To circumvent this molecular trade-off, we exploited structural models of insulin fibrils and amyloidogenic intermediates to define an alternative protective mechanism. Single-chain insulin (SCI) analogs were shown to be refractory to thermal fibrillation with maintenance of biological activity for more than 3 months under conditions that promote the rapid fibrillation and inactivation of insulin. The essential idea exploits an intrinsic incompatibility between SCI topology and the geometry of cross-β assembly. A peptide tether was thus interposed between the A- and B-chains whose length was (a) sufficiently long to provide the "play" needed for induced fit of the hormone on receptor binding and yet (b) sufficiently short to impose a topological barrier to fibrillation. Our findings suggest that ultrastable monomeric SCI analogs may be formulated without protective self-assembly and so permit simultaneous optimization of pharmacokinetics and reservoir life.

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Abbreviations: (BrdU) bromodeoxyuridine, (CD) circular dichroism, (Gu) guanidine, (HI) human insulin, (HPI) human proinsulin, (HPLC) high-performance liquid chromatography, (IGF-1R) insulin-like growth factor-1 receptor, (IGF-I) insulin-like growth factor-I, (IR-B) insulin receptor isoform B, (MALDI-TOF MS) matrix-assisted laser desorption/ionization-time of flight mass spectrometry, (MS) mass spectrometry, (NaCl) sodium chloride, (NMR) nuclear magnetic resonance, (NPH) neutral protamine Hagedorn, (PBS) phosphate-buffered saline, (PDB) Protein Data Bank, (rp-HPLC) reverse-phase high-performance liquid chromatography, (SCI) single-chain insulin, (ThT) thioflavin T

Keywords: amyloid, closed-loop system, insulin pump, intraperitoneal pump, pump reservoir

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Introduction

nsulin is susceptible to physical and chemical degradation. Physical degradation refers to an irreversible change in physical state without change in covalent structure; the latter is designated chemical degradation. The major barrier to the storage and use of insulin above room temperature (such as in an intraperitoneal pump) is fibrillation, a universal property of polypeptides leading to cross-β linear polymers.¹ Studies of fibrillation provide a foundation for the design of ultrastable and ultrafast insulin analogs with potential application to the artificial pancreas project.²,3

Insulin formulations are shipped and stored at 4-8 °C; shelf life is typically 2 years. Degradation accelerates rapidly with increasing temperature. Vials, once used, must be kept below 30 °C and discarded after 30 days (15 days in the case of diluted solutions of Humalog® or Novolog®). Mechanisms of chemical degradation (such as scission of disulfide bonds leading to covalent polymers) are well characterized.4-9 Studies of insulin analogs suggest that flexible regions are particularly susceptible to chemical degradation, and so stabilizing substitutions protect from such processes.¹⁰ By contrast, physical degradation is not well understood.^{1,11} Despite the importance of fibrillation in biotechnology and medicine, how globular proteins form fibrils remains enigmatic. Although molecular models have been proposed, 12,13 the structure of an insulin fibril has not been elucidated at atomic resolution.

The present study exploited general structural features of an insulin fibril as a basis for rational design. It has been established, for example, that it is the monomeric polypeptide (as distinct from native self-assemblies) that mediates aggregation-coupled misfolding. Accordingly, conditions that favor (or disfavor) native self-assembly are associated with delayed (or accelerated) fibrillation.9 Insulin fibrillation is thus promoted by factors that destabilize its classical pathway of assembly, presumably by increasing the availability of the monomer.¹⁴ Fibrillation is further accelerated by partial unfolding as induced by agitation or elevated temperatures.¹⁵ The zinc insulin hexamer (depicted in Figure 1) has long provided a vehicle for pharmaceutical formulation due to its stability. Sequestration of the insulin monomer by native selfassembly forestalls partial unfolding and nonnative aggregation (Figure 2).

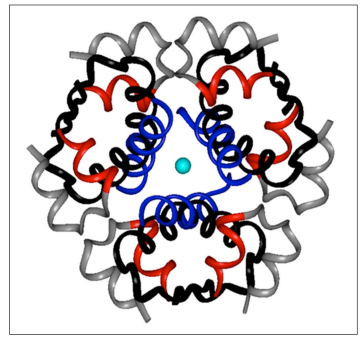


Figure 1. Structure of insulin as an R_6 hexamer [Protein Data Bank (PDB) entry 1 TRZ]. The two axial zinc ions (center) are shown as overlying light blue spheres. The A-chains are shown in red (residues A1–A8) and gray (A9–A21); the B-chains are shown in blue (R-specific B1–B8 α -helix) and black (B9–B30). Phenolic ligands are not shown.

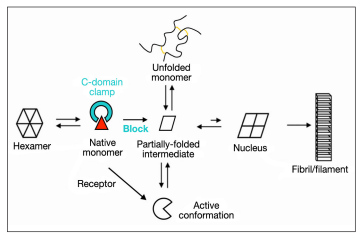


Figure 2. Mechanism of insulin fibrillation via partial unfolding of monomer. The native hexamer is protected by self-assembly (far left). Disassembly leads to an equilibrium between native and partially folded monomers (triangle and rhombus, respectively). An engineered linker between A- and B-chains (electric blue) impedes formation of this partial fold, which may otherwise either (i) unfold completely as an off-pathway event (top) or (ii) aggregate to form a nucleus en route to a protofilament (far right). Sufficient play must be present in the linker to permit induced fit on receptor binding (bottom). Figure adapted from Nielsen and colleagues. Page 18.

Use of insulin analogs in pumps (such as Humalog or Novolog) was based on the principle that mutational destabilization of subunit interfaces within the hexamer would facilitate absorption.^{7,16–20} Such analogs have indeed enabled better glycemic control with reduced risk of hypoglycemia.^{21–23} Although advantageous for pump delivery,²¹ this strategy enhances rates of misfolding and fibrillation.^{24,25} Case reports have documented occasional pump occlusion due to fibrillation,^{25,26} posing a risk of metabolic decompensation. Although ultrarapid formulations would enhance the safety and efficacy of closed-loop systems, extension of this mutational strategy encounters a molecular trade-off: accelerated disassembly ordinarily promotes degradation.

We sought to circumvent the trade-off of conventional prandial insulin design through the use of single-chain insulin (SCI) analogs, also designated miniproinsulins.²⁷ Although SCI analogs are refractory to fibrillation,9,28 their investigation has been discouraged by low activities.²⁹ An extreme combination of these properties is exemplified by SCI-50, a single-chain analog whose crystal structure (Figure 3B) is essentially identical to that of wild-type insulin (Figure 3A).²⁹ Whereas SCI-50 contains a single peptide bond between Lys^{B29} and Gly^{A1}, in wild-type insulin, the C-terminus of the B-chain adjoins (and is often saltbridged to) the N-terminus of the A-chain. Uniquely among insulin analogs, SCI-50 is both refractory to fibrillation³⁰ and devoid of biological activity.²⁹ The tether between chains presumably constrains conformational changes required in concert for nonnative cross-β assembly and receptor binding.31,32 This approach thus seemed only to replace the trade-off of prandial analog design with another and more severe limitation.

Our interest in the potential therapeutic use of SCI analogs was motivated by two findings. On the one hand, molecular models of insulin fibrils in general posit parallel extension of A- and B-chains with retention of native disulfide pairing;¹² in particular, cross-β assembly requires splaying of the C-terminus of the B-chain from the N-terminus of the A-chain (despite their proximity in native structures) by distances greater than 30 Å.30 On the other hand, systematic studies of cross-linked insulin analogs (i.e., containing chemical tethers) have suggested that these termini are between 16 and 24 Å apart in a hormone-receptor complex.³³ Together, these findings suggested a reciprocal modulation of fibrillation rate and biological activity as a function of connectingpeptide length: whereas tight tethers (as in SCI-50) would preclude fibrillation at the expense of activity, loose tethers (as in proinsulin) would, at least in part, restore

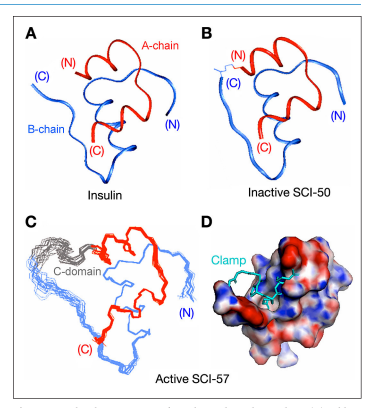


Figure 3. Molecular structures of insulin and insulin analogs. **(A)** Ribbon model of an isolated protomer (T-state conformation; PDB entry 4INS). The A-chain is shown in red, and B-chain in blue; disulfide brides are not depicted. **(B)** Ribbon model of *des*-Thr^{B30}-SCI-50, an inactive 50-residue single-chain analog in which a peptide bond links Lys^{B29} to Gly^{A1,29} The structure represents the T-state protomer of a T₃R^f₃ zinc hexamer (PDB entry 1PID). **(C)** NMR-derived ensemble of SCI-57 as an engineered monomer in solution.³⁴ The A-chains are shown in red, B-chains in blue, and linkers (sequence GGGPRR) in gray. Coordinates were aligned according to the main-chain atoms of residues B4–B28 and A2–A20. **(D)** A rotated view of SCI-57 highlights the linker (sticks; electric blue) as a tether between A- and B-domains (space-filling representation). The surface is color-coded according to electrostatic potential (red, electronegative; purple, electropositive; and white, neutral). Images in panels C and D were obtained from PDB entry 2JZQ.

activity at the cost of fibrillation.³⁰ Might there be an optimal intermediate length that is both protective and compatible with native activity? We have described the chemical synthesis and nuclear magnetic resonance (NMR) structure of an SCI of 57 residues (SCI-57) whose connecting peptide lies within the receptor-active distance range (six residues; ~16-24 Å in length) and whose insulin moiety has been optimized with respect to stability and receptor binding.34 The structure of SCI-57 (Figure 3C) is remarkable for the damping of conformational fluctuations by its tether, packed as a clamp within an interchain crevice (Figure 3D). Here, we demonstrate that SCI-57 retains native potency in a diabetic rat model, exhibits native mitogenicity in a human cancer cell line, and displays extraordinary resistance to chemical and physical degradation as a monomer at an elevated temperature. Together, our

findings highlight the promise of intermediate-length SCI analogs as ideal pump insulin analogs.

Methods

Elements of Protein Design

SCI-57 was designed to achieve the following objectives:³⁴

Optimization of α-Helical Stability

Because physical degradation involves a change in conformation from α -helix to β -sheet, we sought to optimize intrinsic helical propensity. Similarly, because rates of chemical degradation are inversely correlated with their thermodynamic stabilities, we sought to maximize stability. 34

Engineered Monomer

Because insulin absorption is slowed by native assembly, we sought an analog lacking functional dimer-, trimer-, or hexamer-forming surfaces.³⁵

Electrostatic Balance

Because an optimal analog must be stored at high concentration, we sought to maintain a native-like isoelectric point.

Potency

To control glycemia, we sought an analog with potency similar to that of human insulin.

Resistance to Fibrillation

Because cross- β assembly requires splaying of A- and B-chains, 12,30,36 we sought an SCI framework as a topological barrier to fibrillation.

<u>Mitogenicity</u>

To reduce the risk of tumorigenesis,³⁷ we sought an analog whose mitogenicity was similar to that of human insulin.

Progress toward the first three objectives was documented in our initial study;³⁴ see also *Appendix*.

Materials

Insulin and proinsulin were provided by Eli Lilly and Co. (Indianapolis, IN). Insulin-like growth factor-I (IGF-I) was obtained from GroPep, Ltd. (Adelaide, Australia) and thioflavin T (ThT) from Sigma (St. Louis, MO).

Chemical Protein Synthesis

SCI analogs were prepared by native fragment ligation as described.³⁴ [His^{A8}, Asp^{B10}, Asp^{B28}, Pro^{B29}]-insulin was

prepared by chain combination following solid-phase synthesis of the A- and B-chain analogs.³² Predicted molecular masses were verified by mass spectrometry (MS). Purity was >98% as assessed by reverse-phase high-performance liquid chromatography (rp-HPLC).

Mass Spectrometry

Matrix-assisted laser desorption/ionization-time of flight mass spectrometry (MALDI-TOF MS) employed a pulsed nitrogen laser source ($\lambda=337$ nm) and acceleration voltage of 28 kV operated in the linear mode (Voyager Biospectrometry Workstation, PerSeptive Biosystems, Framingham, MA); 100–200 spectra were obtained per sample.

Thermodynamic Modeling

Guanidine denaturation was monitored by circular dichroism (CD) at 222 nm; data were fitted by nonlinear least squares to a two-state model as described. 30,38

Preparation of Fibrils

Proteins were made $60~\mu\text{M}$ in degassed phosphate-buffered saline [PBS; 10~mM phosphate, 140~mM sodium chloride (NaCl), pH 7.4], incubated in presterilized glass vials with airtight sealed caps (Allergy Labs, Oklahoma City, OK), and rocked on a Becton-Dickinson nutator (BD Diagnostic Systems, Franklin Lakes, NJ) at $\sim 60~\text{rpm}$. Aliquots, withdrawn with a sterilized single-use syringe, were added to ThT solution for fluorescence assays of fibrillation.

Degradation Assays

Aliquots obtained at indicated time points (10 µl) were mixed with 3 ml ThT assay buffer [5 mM ThT in 50 mM tris(hydroxymethyl)aminomethane, hydrochloride (pH 7.5) and 100 mM NaCl]. Fluorescence emission spectra were collected from 470–500 nm following excitation at 450 nm; ThT in buffer without protein was used as baseline. Changes in covalent structure were evaluated by rp-HPLC and MS.

Modeling of Protofilament

A model of a two-layered insulin protofilament was constructed using InsightII (Accelrys, Inc., San Diego, CA) as described³⁰ based on earlier cryo-electron microscopy studies.¹²

Receptor Binding

Assays employed lectin-purified and detergent-solubilized insulin receptor [insulin receptor isoform B (IR-B)] or insulin-like growth factor-1 receptor (IGF-1R) as described.^{39,40} Respective dissociation constants were determined

by competitive displacement of ¹²⁵I-[Tyr^{A14}]-insulin or ¹²⁵I- [Tyr³¹]-IGF-I (kindly provided by Novo-Nordisk, A/S, Bagsværd, Denmark). Each determination was performed with at least three replicates. The percentage of tracer bound in the absence of competing ligand was <15% to avoid ligand-depletion artifacts.

Mitogenicity

The human breast cancer cell line MCF-7 was maintained in culture essentially as described. Medium was changed every 2 days and cells were subcultured when they reached 90% of confluence. Cell proliferation was determined by the incorporation of bromodeoxyuridine (BrdU) using a kit (Roche Diagnostic, Indianapolis, IN). Approximately 3.5×10^3 cells were seeded into each well of a 24-well plate and cultured for 24 h; medium was then replaced with fresh serum-free medium containing 0.1% bovine serum albumin. After 24 h, insulin analogs or IGF-I were added at various concentrations in fresh medium; media were changed each day for 5 days. BrdU incorporation was then determined according to the manufacturer's directions.

Biological Potency

Male Lewis rats (~250 g body weight) were rendered diabetic with streptozotocin as described.⁴¹ Human insulin and analogs (SCI-57 and a two-chain version lacking the six-residue linker) were purified by HPLC, lyophilized, and dissolved in insulin-related sterile diluent (Eli Lilly and Co., Indianapolis, IN). Rats were injected subcutaneously at time 0 with 1.5 U/kg body weight in 100 µl of diluent. Blood was obtained from clipped tip of the tail at time 0 and every 10 min up to 90 min. Blood glucose was measured using a Hypoguard Advance Micro-Draw (Hypoguard USA, Inc., Minneapolis, MN) meter.

Results

Our study focused on properties pertinent to implantable pumps and closed-loop systems in real-world environments. The thermodynamic stability of SCI-57 at 37 °C (ΔG_u 4.3 kcal/mol), as probed by chemical denaturation studies (**Figure 4A**), is significantly higher than the stabilities of wild-type insulin, KP-insulin, or Asp^{B28}-insulin (ΔG_u 2.4 kcal/mol; **Figure 4B**); such stabilities (column two of **Table 1**) were obtained under zincfree conditions predictive of relative susceptibility to chemical degradation.⁶ The net gain in stability ($\Delta \Delta G_u$ 1.9 \pm 0.2 kcal/mol) presumably reflects the combined effects of stabilizing (His^{A8}, Asp^{B10}, and linker contacts) and destabilizing modifications (Asp^{B28} and Pro^{B29}).

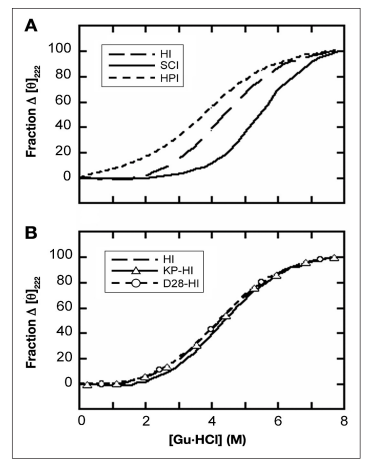


Figure 4. Guanidine (Gu) denaturation studies. **(A)** Comparison of human insulin (HI), human proinsulin (HPI), and 57-residue singlechain insulin (SCI). **(B)** Comparison of HI, [Lys^{B28}-Pro^{B29}]-insulin (KP-HI; the active component of Humalog), and Asp^{B28}-insulin (D28-HI; the active component of Novolog). Studies were conducted at 37 °C. Thermodynamic parameters are given in **Table 1**; unfolding was monitored by CD at helix-sensitive wavelength 222 nm.

Loose tethering, as probed by comparison of insulin and proinsulin, resulted in itself in only a small gain in stability ($\Delta\Delta G_u$ 0.4 \pm 0.2 kcal/mol). In accordance with inferred free-energy values, 50% unfolding of SCI-57 required a higher concentration of guanidine (5.5 \pm 0.1 M; column three of **Table 1**) than did 50% unfolding of proinsulin (4.0 \pm 0.2 M), wild-type insulin or prandial insulin analogs (4.1 M).

Degradation was monitored on gentle agitation in glass vials at 37 °C (**Figure 5** and column five of **Table 1**). Whereas wild-type insulin or prandial analogs were inactivated by fibrillation within 5 days and proinsulin within 15 ± 2.5 days, SCI-57 exhibited no signs of fibrillation up to 180 days (at which point the study was terminated). Analysis of chemical-degradation products by rp-HPLC and MS revealed only a *des*-amido species, consistent with conversion of Asn→Asp. Despite the harsh conditions employed (elevated temperature and

Table 1.	
Thermodynamic Stabilities ^a and Fibrillation Lag Times ^b (p	pH 7.4 and 37 °C)

	0 1		
ΔG_u (kcal/mol)	C _{mid} (M)	m (kcal/mol/M)	Fibrillation lag time (days)
2.4 ± 0.1	4.1 ± 0.2	0.57 ± 0.02	3.5 ± 0.6
2.4 ± 0.1	4.1 ± 0.1	0.60 ± 0.01	1.7 ± 0.3
2.4 ± 0.1	4.1 ± 0.1	0.59 ± 0.01	2.6 ± 0.3
2.8 ± 0.1	4.0 ± 0.2	0.71 ± 0.03	15 ± 2.5
4.3 ± 0.1	5.5 ± 0.1	0.79 ± 0.02	>180
	2.4 ± 0.1 2.4 ± 0.1 2.4 ± 0.1 2.8 ± 0.1	ΔG_u (kcal/mol) C_{mid} (M) 2.4 ± 0.1 4.1 ± 0.2 2.4 ± 0.1 4.1 ± 0.1 2.4 ± 0.1 4.1 ± 0.1 4.1 ± 0.1 4.0 ± 0.2	ΔG_{u} (kcal/mol) C_{mid} (M) m (kcal/mol/M) 2.4 ± 0.1 4.1 ± 0.2 0.57 ± 0.02 2.4 ± 0.1 4.1 ± 0.1 0.60 ± 0.01 2.4 ± 0.1 4.1 ± 0.1 0.59 ± 0.01 2.8 ± 0.1 4.0 ± 0.2 0.71 ± 0.03

^a Thermodynamic stabilities were inferred from CD-detected guanidine denaturation studies by application of a two-state model. ³⁴ Data are obtained in part from Table 1 of Hua and colleagues. ³⁴ Stability (kcal/mol) is defined as the free energy of unfolding (ΔG_{ii}) at 37 °C as extrapolated to zero denaturant concentration; ³⁸ uncertainties are ±0.1 kcal/mol. C_{mid} is defined as that molar concentration of guanidine-HCl associated with 50% protein unfolding. The m value (kcal/mol/M) is defined as the slope in plotting the unfolding free energy versus molar concentration of denaturant; this slope is often found to be proportional to the protein surface area exposed on unfolding.

^b Fibrillation lag times were determined by ThT fluorescence following gentle agitation in the presence of an air-liquid interface as described. ³⁰ Proteins were made in 60 μM PBS (pH 7.4) in the absence of zinc ions and gently rocked at 37 °C in a glass vial in the presence of an air-water interface. ⁴²

^c This analog, the active component of Novolog, contains the amino-acid substitution Pro^{B28}→Asp.

^d This analog, the active component of Humalog, contains amino-acid substitutions Pro^{B28}→Lys and Lys^{B29}→Pro.

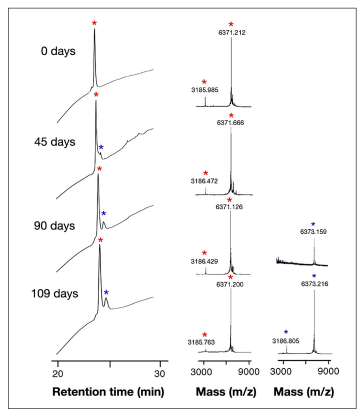


Figure 5. Chemical and physical stability of SCI-57: time course of rp-HPLC chromatogram (left) and MALDI-TOF mass spectrum (right) from 0–109 days. The SCI-57 solution remained clear and without visible precipitation during incubation (see main text). At indicated times (0–109 days), aliquots were applied onto a reverse-phase C8 HPLC column; components were eluted with a methanol gradient. Molecular masses were determined by MS using vendor-provided software (Perkin-Elmer, Wellesley, MA). The difference of 1–2 mass units between the native analog and degradation products represents deamidation (likely to be conversion of Asn^{A21} to Asp). Respective red and blue asterisks represent major peak (native analog) and minor degradation product. Mass 3186 Da represents the SCI with two charges.

gentle agitation against a nonsilanized glass surface in the presence of an air-liquid interface), no chain cleavage or formation of covalent polymers was detected. Further, no reductions were observed in receptor-binding activity as evaluated at successive time points during the test; representative data are shown in **Figure 6A**. By contrast, residual binding of the degraded solutions of insulin or prandial analogs was reduced by more than 100-fold.

Because of the presence of Asp^{B10} in SCI-57 and the association of this substitution with increased mitogenicity and carcinogenicity in rats,37 cross-binding to IGF-1R was evaluated in relation to IGF-1, wild-type insulin, Asp^{B10}-insulin, and a corresponding two-chain insulin analog lacking a connecting domain ([His^{A8}, Asp^{B10}, Asp^{B28}, Pro^{B29}]-insulin) (**Figure 6B**). In accordance with earlier studies, 40 baseline affinities (dissociation constants K_d) of IGF-1 and wild-type insulin were 0.041 \pm 0.005 and 9.57 ± 0.31 nM, respectively. As expected, Asp^{B10}-insulin and [His^{A8}, Asp^{B10}, Asp^{B28}, Pro^{B29}]-insulin exhibited augmented binding to IGF-1R (K_d 2.7 \pm 0.003 and 4.31 ± 0.006 nM, respectively). Remarkably, interposition of the six-residue linker reduced IGF-1R binding to a level lower than that of wild-type insulin (K_d 9.89 \pm 0.035 nM). Mitogenicity was directly evaluated in cell culture employing human breast-cancer cell line MCF-7. Whereas mitogenicities of Asp^{B10}-insulin and [His^{A8}, Asp^{B10}, Asp^{B28}, Pro^{B29}]-insulin were intermediate between those of wildtype insulin and IGF-1, as probed by both cell number and DNA content, no significant differences were observed between SCI-57, wild-type insulin, and KP-insulin (data not shown). Thus, the linker may be regarded as a tuning element that downregulates potential mitogenicity.

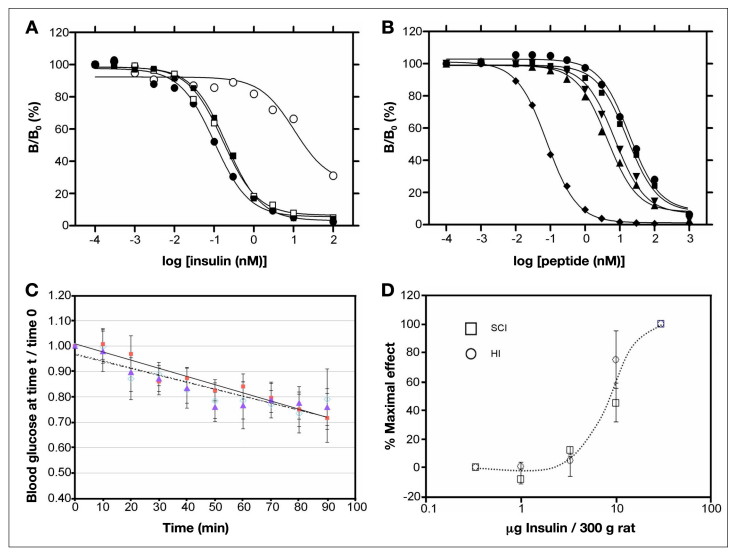


Figure 6. Receptor-binding studies and biological activity in rats. (A) Stability of the receptor-binding activity of SCI-57 and wild-type insulin on prolonged gentle agitation at 37 °C. The following representative competitive-displacement curves are shown: assays conducted at the start of the incubation (SCI-57 ■; insulin •), assay performed after 42 days of agitation (HI \circ), or assay performed after 90 days (SCI-57 \square). Results are expressed as the ratio (B/B₀) of ¹²⁵I-[Tyr^{A14}]-insulin specifically bound at a given concentration of cold insulin relative to bound ¹²⁵I-[Tyr^{A14}]-insulin specifically bound in absence of competing unlabeled insulin. (B) Competitive binding of insulin analogs to IGF-1R: SCI-57 (■), wild-type insulin (•), [His^{A8}, Asp^{B10}, Asp^{B29}, Pro^{B29}]-insulin (♠), Asp^{B10}-insulin (♥), and human IGF-I (•). (C) Decrease in blood glucose concentration with time (in minutes): regular human insulin (⟨⟩), the 57-residue single-chain analog (SCI; ■), and its two-chain derivative lacking the linker (♠). Initial rates and maximal extents of glucose-lowering activity were indistinguishable. (D) A log plot of concentration-dependent efficacy in the rat shows no difference relative to wild-type insulin.

The biological potency of SCI-57 was assessed in a streptozotocin-induced rat model of diabetes mellitus. No significance differences were observed between SCI-57 and human insulin with respect to initial rate of fall in blood glucose concentration (**Figure 6C**) and dose-response relationship (**Figure 6D**). In this model, Humulin®-R U-100 (Eli Lilly and Co.) insulin and Humalog exhibit indistinguishable pharmacodynamics (presumably due to differences between rat skin and human skin and to differences in subcutaneous depot sizes) but each is markedly more rapid than Lantus® (Sanofi, Paris, France) or neutral protamine Hagedorn (NPH) insulin (Humulin N; Eli Lilly and Co.).

Discussion

Insulin functions in the bloodstream as a monomer, and yet it is the monomer that is susceptible to fibrillation and most forms of chemical degradation. Degradation in vials and delivery devices has long been delayed through protective self-assembly. Use of Zn^{2+} -coordinated insulin hexamers as a formulation vehicle recapitulates the logic of the pancreatic β -cell whose secretory granules contain microcrystalline arrays of zinc insulin hexamers. X-ray structures of such hexamers are of exquisite beauty (**Figure 1**). Multiple crystal forms have been described *in vitro*, defining three structural families (T_6 , $T_3R_3^f$, and R_6).

The T-state (**Figure 3A**) resembles the structure of an insulin monomer in solution.^{31,44,45}

Absorption of regular insulin is limited by the kinetic lifetime of the Zn-insulin hexamer as disassembly to smaller dimers and monomers enables capillary absorption.35 The essential idea underlying design of Humalog and Novolog was thus to accelerate disassembly. This was accomplished by destabilization of the classical dimer-forming surface (the C-terminal antiparallel β-sheet). Humalog contains substitutions Pro^{B28}→Lys and Lys^{B29}→Pro, an inversion that mimics the sequence of IGF-I; Novolog contains substitution Pro^{B28}→Asp.^{46,47} Although the substitutions impair dimerization, the analogs are competent for assembly of a phenol-stabilized zinc hexamer. 47 This assembly protects the analog from fibrillation in the vial but following injection, the hexamer rapidly dissociates. The instability of these analogs on dilution underlies their reduced shelf life after opening, especially on dilution. In addition to effects of reduced selfassembly, the present studies have shown that zinc-free monomers of Asp^{B28}-insulin and KP-insulin at 37 °C are intrinsically more susceptible to fibrillation than is wildtype insulin. Because of the importance of formulation stability, a molecular trade-off is thus posed by the use of destabilizing mutations to enhance rates of absorption.

A NMR study of an engineered insulin monomer demonstrated that the N-terminal A-chain segment (residues A1–A8), although a canonical α -helix in crystal structures (red in Figure 1), exhibits anomalous ¹³C chemical shifts in solution.⁴¹ These findings provided evidence that, in the absence of self-assembly, this segment undergoes significant conformational fluctuations. This interpretation was supported by analysis of amide-proton exchange rates in D₂O: protection factors in this segment were observed to be negligible.⁴⁸ Such anomalous NMR features suggest that the distorted monomeric conformation mediating formation of an amyloidogenic nucleus (trapezoid in Figure 2) contains a partial fold in which the N-terminal segment of the A-chain is poorly ordered (central panel of Figure 7A). Solid-state NMR studies have shown that in a fibril, this segment (like most of the A- and B-chains) forms a β-sheet (**Figure 7B**). Because retention of native disulfide pairing imposes a parallel orientation between chains, 12 this class of models requires splaying of the two chains with distances between the C-terminus of the B-chain and N-terminus of the A-chain greater than 30 Å (Figure 7C).30 It may therefore be of critical importance that the length of the six-residue linker of SCI-57, even at maximal extension, is constrained by covalent structure to be less than 24 Å.

The present study suggests that monomeric SCI analogs may be engineered to exhibit a combination of favorable features and thereby circumvent the trade-off of conventional prandial insulin analog design. We envisage that the foreshortened connecting domain acts as a clamp to stabilize the α-helical conformation of the N-terminal A-chain segment and to disallow splaying of the chains as required for cross-β assembly. Potential advantageous features promise to combine native potency and mitogenicity with marked resistance to chemical and physical degradation as encountered in the reservoir of an insulin pump under real-world conditions. Although such extensive modification of the insulin molecule may in principle be precluded by immunogenicity in patients on long-term use (an open question), we envisage that an engineered ultrastable monomeric SCI analog may yield ultrafast rates of absorption, enhancing the efficacy and safety of closed-loop systems.⁵⁰ Further studies of

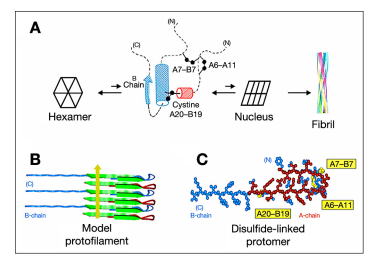


Figure 7. Structural model of insulin protofibrils. (A) Structures at successive stages of insulin fibrillation. Hexameric insulin is shown in cartoon form at far left. Putative amyloidogenic intermediate (cylinder model)30 and unknown nucleus (cartoon) are shown at center. Model of an insulin fibril¹² is shown at far right. In the partial fold, cylinders and the arrow indicate molten supersecondary structure (residues B9-B26 and A16-A20). Dashed lines indicate disordered regions; disulfide bridges are indicated by black balls (sulfur atoms). (B) Schematic representation of stacked insulin main chains in a protofilament as proposed by Jimenez and colleagues.¹² The cross-β unit consists of alternatively stacked A- and B-chains (red and blue, respectively) to form in-register parallel β-sheets (segments A1-A6, A12-A21, B1-B6, and B12-B30). The yellow arrow indicates direction of long axis of fibril. (C) An insulin protomer in a protofilament (as viewed down its long axis) is predicted to exhibit an extended polypeptide conformation with central bend; disulfide bridges are in yellow. Models were constructed wherein two β-strand regions of the A-chain (A1–A5 and A12–A21) were built with dihedral angles φ and ψ set at -140 and 140 °, respectively; corresponding B-chain segments (B1-B5 and B12-B30) were likewise constructed, and the two chains connected by disulfide bridges (A7-B7 and A20-B19). Interchain β -sheet-related hydrogen bonds were then imposed by time-averaged simulated-annealing.4

the pharmacokinetic and immunogenic properties of SCI analogs in large animal models and healthy human volunteers will be required to test these ideas in an effort to reduce this molecular technology to practice.

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Appendix

Summary of SCI-57 Design and Prior Characterization

SCI-57 contains two known helix-stabilizing substitutions. One (Thr^{A8} \rightarrow His)¹ enhances the C-terminal residue (C-cap)² of the N-terminal A-chain α -helix;³⁴ the other (His^{B10} \rightarrow Asp)⁵ enhances the N-terminal residue (N-cap)² of the central B-chain α -helix;³⁴ the other (His^{B10} \rightarrow Asp)⁵ enhances the N-terminal residue (N-cap)² of the central B-chain α -helix;³⁴ the other (His^{B10} \rightarrow Asp)⁵ enhances the N-terminal residue (N-cap)² of the central B-chain α -helix;³⁴ the other (His^{B10} \rightarrow Asp)⁵ enhances the N-terminal residue (N-cap)² of the central B-chain α -helix;³⁴ the other (His^{B10} \rightarrow Asp)⁵ enhances the N-terminal residue (N-cap)² of the central B-chain α -helix;³⁴ the other (His^{B10} \rightarrow Asp)⁵ enhances the N-terminal residue (N-cap)² of the central B-chain α -helix;³⁴ the other (His^{B10} \rightarrow Asp)⁵ enhances the N-terminal residue (N-cap)² of the central B-chain α -helix;³⁴ the other (His^{B10} \rightarrow Asp)⁵ enhances the N-terminal residue (N-cap)² of the central B-chain α -helix;³⁴ the other (His^{B10} \rightarrow Asp)⁵ enhances the N-terminal residue (N-cap)² of the central B-chain α -helix;³⁴ the other (His^{B10} \rightarrow Asp)⁵ in the context of insulin analogs for the insulin receptor.³⁴ In the context of otherwise offsetting modifications, i.e., a gain of two basic residues (in linker sequence GGGPRR) offset by two acidic residues (Asp^{B10} \rightarrow and Asp^{B28} \rightarrow , gain of a titratable imidazolic group (His^{A8} \rightarrow) offset by loss of His^{B10} \rightarrow , and the balanced elimination of opposing chain termini (A1 α -amino group and B30 α -carboxylate). Isoelectric focusing studies revealed an isoelectric point of 5.3, indeed lower than that of wild-type insulin (pI 5.8).

Table A1. Mitogenicity Assay ^a		
A. Relative to IGF-I		
IGF-I:Asp ^{B10} -insulin	p = .040	
IGF-I:KP-insulin	p = .021	
IGF-I:SCI-57	p = .028	
IGF-I:two-chain control ^b	p = .041	
IGF-I:insulin	p = .031	
B. Relative to wild-type insulin		
Insulin:Asp ^{B10} -insulin	p = .051	
Insulin:KP-insulin	p = .068	
Insulin:SCI-57	p = .091	
Insulin:two-chain control ^b	p = .049	

 ^a Statistical analysis of BrdU studies as depicted in Figure A1.
 ^b Two-chain control designates [His^{A8}, Asp^{B10}, Asp^{B28}, Pro^{B29}]-

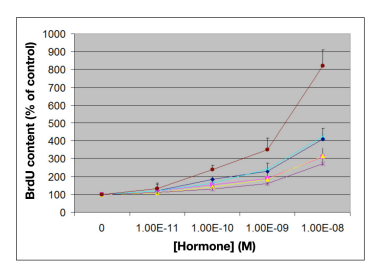


Figure A1. Mitogenicity studies of MCF-7 human breast-cancer-derived cell line. Relative changes in BrdU content after 5 days in culture (vertical axis) are shown as a function of hormone concentration (horizontal axis). From top to bottom: (maroon) human IGF-I; (light blue) Asp^{B10}-insulin; (dark blue) [His^{A8}, Asp^{B10}, Asp^{B28}, Pro^{B29}]-insulin; (yellow) SCI-57; (pink) KP-insulin; and (purple) wild-type insulin. Error bars indicate standard deviation of triplicate determinations. Statistical p values are provided in Table A1. [His^{A8}, Asp^{B10}, Asp^{B28}, Pro^{B29}]-insulin corresponds to the insulin core of SCI-57 and so provides a two-chain control. KP-insulin designates [Lys^{B28}, Pro^{B29}]-insulin, the active component of Humalog.

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