A Thing of Beauty: Structure and Function of Insulin’s “Aromatic Triplet”

Michael A. Weiss\(^1\) and Michael C. Lawrence\(^{2,3}\)

\(^1\)Department of Biochemistry & Molecular Biology, Indiana University School of Medicine, 635 Barnhill Drive, Indianapolis, IN 46202 USA

\(^2\)The Walter and Eliza Hall Institute of Medical Research, 1G Royal Parade, Parkville, Victoria 3052, AUSTRALIA

\(^3\)Department of Medical Biology, University of Melbourne, Parkville, Victoria 3010, AUSTRALIA

Abstract

The classical crystal structure of insulin was determined in 1969 by D.C. Hodgkin and colleagues following a 35-year program of research. This structure depicted a hexamer remarkable for its self-assembly as a zinc-coordinated trimer of dimer. Prominent at the dimer interface was an “aromatic triplet” of conserved residues at consecutive positions in the B chain: Phe\(^B24\), Phe\(^B25\) and Tyr\(^B26\). The elegance of this interface inspired the Oxford team to poetry: “A thing of beauty is a joy forever” (John Keats as quoted by Blundell, T.L., et al. Advances in Protein Chemistry 26:279–286 (1972)). Here, we revisit this aromatic triplet in light of recent advances in the structural biology of insulin bound as a monomer to fragments of the insulin receptor. Such co-crystal structures have defined how these side chains pack at the primary hormone-binding surface of the receptor ectodomain. On receptor binding, the B-chain \(\beta\)-strand containing the aromatic triplet (residues B24-B28) detaches from the \(\alpha\)-helical core of the hormone. Whereas Tyr\(^B26\) lies at the periphery of the receptor interface and may functionally be replaced by a diverse set of substitutions, Phe\(^B24\) and Phe\(^B25\) engage invariant elements of receptor domains L1 and \(\alpha\)CT. These critical contacts were anticipated by the discovery of diabetes-associated mutations at these positions by D.F. Steiner and his colleagues at the University of Chicago. Conservation of Phe\(^B24\), Phe\(^B25\) and Tyr\(^B26\) among vertebrate insulins reflects the striking confluence of structure-based evolutionary constraints: foldability, protective self-assembly and hormonal activity.

Keywords

protein structure; protein evolution; protein recognition; structure-activity relationships; hormone-receptor recognition

Michael A. Weiss, Dept. of Biochemistry and Molecular Biology, Indiana University School of Medicine, Indianapolis, IN 46202 USA., weissama@iu.edu. Michael C. Lawrence, Walter and Eliza Hall Institute of Medical Research, Parkville, Victoria 3052, AU., lawrence@wehi.edu.au.

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1 INTRODUCTION

Insulin has provided an enduring model for studies in protein chemistry and cell biology virtually since the hormone’s landmark discovery in Toronto in 1921 [1]. The sequence of bovine insulin, the first protein sequence to be determined [2, 3], established that this class of biological heteropolymers in fact had specific sequences—and so made concrete the concept of a genetic code [4]. Insulin was the second protein to be crystallized (after urease) [5, 6] and provided the first example of protein crystallization as a process that in itself can confer pharmacologic advantage (for review, see [7]). Efforts to determine the three-dimensional (3D) structure of insulin began in 1934 in the graduate studies of D.C. Hodgkin (then at Cambridge University) [6, 8]. These studies came to fruition in 1969 at the University of Oxford with the first crystal structure of a peptide hormone [9, 10]. It is fitting to begin this contribution, written in honor of the late Prof. Donald F. Steiner, at the point of intersection between his laboratory at the University of Chicago with the Hodgkin laboratory.

In an issue dedicated to Prof. Steiner, it is fitting to include a structural perspective as his studies integrated structure with enzymology, cell biology, and pathophysiology [11, 12]. His faculty career began in the 1960s, a decade of transformational discovery in molecular endocrinology. The first half of this decade, for example, witnessed international efforts to develop methods for the total chemical synthesis of insulin [13]. Major programs were led by P.G. Katsoyannis at the University of Pittsburg [14], by H. Zahn in Wool Institute of Aachen, German [15] and by the late Niu Jingyi and his colleagues at the Institute of Biochemistry, Academia Sinica (Shanghai) [16]. The success of these programs highlighted in the breach a seeming paradox posed by the low yields of insulin chain combination. Although the isolated A- and B chains contained sufficient information to direct formation of the three specific disulfide bridges identified by Sanger (cystines A7-B7, A6-A11 and A20-B19) [17], the reaction was dominated by side reactions leading to cyclic chains and peptide aggregates, including what is now understood to be peptide fibrils [18–21]. This critical barrier to chemical synthesis raised the central question of how efficiency of native disulfide pairing might be achieved in the biosynthesis of insulin in the specialized endoplasmic reticulum (ER) of pancreatic β-cells [22–24].

Such was the context for the seminal discovery of a single-chain insulin precursor by Steiner and Philip E. Oyer in 1967 [22, 23]. Designated proinsulin, the precursor contains a connecting peptide between the C-terminal residue of the B chain (ThrB30 in human insulin) and N-terminal residue of the A chain (conserved as GlyA1). The discovery of proinsulin and the C peptide quickly found medical application through development of radioimmunoassays (RIAs) to probe insulin secretory capacity in normal and pathological states, including residual β-cell function in patients with Type 1 diabetes mellitus (T1DM), β-cell failure in long-established Type 2 DM (T2DM) and inappropriate insulin secretion in β-cell neoplasia (insulinoma) (for review, see [25]). Cellular mechanisms ensuring the fidelity of disulfide pairing and their perturbation in a monogenic DM syndrome due to toxic misfolding of variant proinsulins are reviewed in an accompanying article in this issue by M. Liu et al. [26].
The discovery of proinsulin led to exchange of visitors between Chicago and Oxford, stimulated by the determination of the crystal structure of the zinc insulin hexamer in 1969 and at increasing resolution in 1971 [9, 10]. These years preceded the development of computer-graphics systems for visualization of protein structures. Intuition was facilitated through construction of physical models, hand-built in the first instances.

Such was the collegial respect between Professors Hodgkin and Steiner that one of these rare early models of the insulin monomer (now classified as a T-state protomer of the T6 hexamer) was given to the Chicago laboratory, hand-delivered by a student visiting from Oxford. For more than 40 years this model assumed pride of place in Don’s office (Figure 1). Inset at upper right are pictures of Professors Hodgkin and Steiner holding corresponding models.

Since 1969, the structure of insulin has been determined in a variety of crystal forms [27–31], in solution by NMR methods [32–39], and most recently, in complexes with fragments of the insulin receptor (IR) [40, 41]. Our review focuses on a conserved segment of the B chain—the conserved “aromatic triplet” PheB24, PheB25 and TyrB26 [42]—as the thread of a remarkable story. The positions of these residues in the zinc insulin hexamer (T6) are shown in Figure 2A, and their positions in the dimer are shown in Figure 2B. Interest in the contribution of the aromatic triplet to the structure and function of insulin thus began with the first crystal structure of insulin, deepened with the first discovery of mutations in the insulin gene associated with diabetes mellitus [43–46] and continues in ongoing efforts to decipher the structure of the hormone-receptor complexes [47, 48], including by single-particle cryo-electron microscopy [49].

## 2 | CRYSTAL STRUCTURES OF INSULIN

The 1969 crystal structure of insulin (then designated 2-Zn insulin and now classified as a T6 hexamer [50]) was among the first set of protein structures to be determined (for reviews, see [6, 51]). Each of these early structures (spanning globins, ribonucleases and proteases) uncovered a new aspect of protein structure and its relationship to function. These features were to inform general principles of protein folding and assembly in the ensuing decades.

The 2-Zn insulin structure depicted a 32-point-group hexameric self-assembly whose dimer-related interfaces contained a short anti-parallel β-sheet (Fig. 2C) and whose trimer-related interface was stabilized by histidine-mediated 2-zinc-ion coordination [9, 10]. Formation of these interfaces concealed extensive non-polar surfaces that would otherwise be exposed to solvent in an isolated protomer. The specific role of zinc ions in the structure of the insulin hexamer motivated continuing studies of zinc transporters in the cell biology of insulin storage within the secretory granules of pancreatic β-cells [52–54].

Crystal structures of zinc-free insulin dimers by Caspar and colleagues demonstrated that the T2 substructure in the 2-Zn insulin hexamer does not require hexamer assembly or zinc-ion coordination [29, 55].
2.1 | The TR Transition

This original T₆ structure contained a dimer in the crystallographic asymmetric unit, with the hexamer defined by a crystallographic three-fold symmetry axis. The two protomers within the dimer (designated molecules 1 and 2) exhibited subtle differences. More dramatic differences were subsequently observed in the 1976 crystal structure of “4-Zn” insulin [27] in which high concentrations of NaCl induced a conformational change to yield the T₃R₃ hexamer [6, 56]. The T→R transition entailed N-terminal extension of the B-chain α-helix; cylinder models of the constituent TR dimer are shown in Figure 3. Subtle differences were also observed in this transition, including (i) in the conformation of the A7-B7 disulfide bridge, (ii) rotation and reorientation of the A1-A8 α-helix, and (iii) a slight displacement of the B24-B28 β-strand relative to A chain such that an inter-chain hydrogen bond (B25 NH…O=C A19) is broken. The latter feature foreshadowed the frank displacement of the B-chain β-strand on receptor binding as described below (Section 3).

Comparison of the 2-Zn and 4-Zn structures provided a model for analysis of the long-range transmission of conformational change in a protein assembly [56] and inspired speculation regarding which structure (T or R) might best inform structure-activity relationships [6]. One study exploited a key aspect of this transition: a change in the conformation of an invariant glycine at position B8 (purple circles in Figure 3) from the right side of the Ramachandran plot in the T-state β-turn (B7-B10) to the left side to reside within the canonical α-helical island as in the R state [60]. Stabilization of the T state through substitution of D-Ala at B8 impaired binding to the IR by at least 100-fold [60].

The T₆→T₃R₃ transition was “completed” in 1989 through the discovery of R₆ zinc insulin hexamers by G.G. Dodson and colleagues [28]. The conformational plasticity of insulin in the crystalline state (i.e., through the series of structures T₆→T₃R₃→R₆) motivated molecular-dynamics (MD) simulations [61–64] and NMR studies in solution [32–39]. The latter demonstrated that the predominant conformation of the hormone as a monomer in solution (the physiologic state in the bloodstream) closely resembles the crystallographic T state (Figure 4). Such studies exploited engineered insulin monomers to circumvent confounding effects of self-association [59]. In Section 3 we will shall see how this flexibility is exploited to achieve the novel receptor-bound structure of the hormone.

2.2 | The Aromatic Triplet

The sequence of the B chain is notable for the invariance of consecutive phenylalanine residues at positions B24 and B25 (red and green in Figure 5) and for the broad conservation of tyrosine at position B26 (violet). (Divergent rodent sequences may contain ArgB26 or SerB26, substitutions likely to preserve activity [67].) The conserved aromatic triplet is part of a β-strand within a dimer-related anti-parallel β-sheet (B24-B28; Figure 2C). In the T₆→T₃R₃→R₆ series of hexameric structures the side chains of PheB24 and TyrB26 (red and violet in Figure 6) participate in similar structural interactions whereas the position of PheB25 is variable and more peripheral to the hydrophobic core (green). The short-B-chain β-strands follow the B20-B24 β-turn, which contains conserved glycines at B20 and B23 (canonical β-turn positions 1 and 4) [68].
The aromatic side chains of Phe\textsuperscript{B24}, Phe\textsuperscript{B25} and Tyr\textsuperscript{B26} contribute to an extensive non-polar surface of the insulin monomer (Figure 7). This surface is engaged at the dimer interface in crystallographic dimers and hexamers \cite{6, 51}. The side chains of the aromatic triplet thus occupy distinct structural environments in the insulin monomer and are associated with marked differences in structure activity relationships. Interested in these positions was stimulated in the 1980s by the discovery (by Steiner and his colleagues [including Howard S. Tager and Arthur H. Rubenstein] at the University of Chicago) of mutations at B24 and B25 in patients with diabetes mellitus \cite{46}.

**Position B24.**—The aromatic ring of Phe\textsuperscript{B24} lies within a crevice such that one face seals the hydrophobic core and the other exposed to solvent. The ring contacts the methyl groups of Leu\textsuperscript{B15} and the β-protons of Cys\textsuperscript{B19} as a conserved aspect of B-chain super-secondary structure. All natural amino-acid substitutions—with the anomalous exception of Gly—impair receptor binding \cite{69, 70}. Even substitution of Phe\textsuperscript{B24} by its near-isostere Tyr impairs IR binding by at least tenfold \cite{70, 71}. Ser\textsuperscript{B24} is a clinical mutation, known as insulin Los Angeles \cite{72}.

The native activity of Gly\textsuperscript{B24}-insulin may reflect a “register shift” in which Phe\textsuperscript{B25}-Tyr\textsuperscript{B26} take the place of Phe\textsuperscript{B24}-Phe\textsuperscript{B25} with extrusion of a five-residue non-canonical turn (GERGG; residues B20-B24) \cite{39}. An analogous shift may rationalize the enhanced affinity of non-standard insulin analogs containing D-amino-acid substitutions at B24 \cite{38, 39, 73}. In such models the role of Tyr\textsuperscript{B26} is taken by Thr\textsuperscript{B27} in general accordance with divergent rodent sequences (Figure 5 above).

**Position B25.**—The B25 side chain is less well ordered in solution; its aromatic resonances exhibit motional narrowing and fewer inter-residue nuclear Overhauser enhancements (NOEs) contacts relative to those of B24 or B26. Studies by the late H.S. Tager and his then-MD-PhD student Raghu Mirmira (now Director of the Diabetes Center and Professor of Medicine at the Indiana University School of Medicine) established that a broad range of aromatic residues are tolerated by the IR, including β-naphthyl-alanine. A general requirement was established for a trigonal γ-carbon, a shared feature of active aromatic substituents. Leu\textsuperscript{B25}, found as a clinical mutation known as insulin Chicago \cite{74} and containing a tetrahedral γ-carbon, impairs receptor binding by at least 50-fold \cite{75, 76}.

**Position B26.**—Tyr\textsuperscript{B26} lies in part within a crevice between the A- and B chains, contacting the conserved side chains of Val\textsuperscript{B12}, Ile\textsuperscript{A2} and Val\textsuperscript{A3}. This “closed” conformation thus conceals key aspects of the receptor-binding surface as inferred from mutagenesis and visualized in Section 3 through crystallographic studies of model hormone-receptor complexes. Although stabilization of the closed conformation by Tyr\textsuperscript{B26} is not required for biological activity, it protects the free hormone from physical degradation \cite{67}.

Residue-specific photo-cross-linking studies, employing \textit{para}-azido-Phe (Pap) as a photo-activatable probe, implied that residues B24-B26 each pack at the hormone-receptor interface. Domain mapping of photo-products demonstrated that Pap\textsuperscript{B24} and Pap\textsuperscript{B26} photo-cross-linked to the L1 domain of the IR α-subunit \cite{77} whereas Pap\textsuperscript{B25} contacted the C-terminal fragment of the α-subunit \cite{78}. To our knowledge, the use of Pap as a short and
rigid residue-specific photo-probe in insulin was first demonstrated by Steiner in collaboration with Katsoyannis [78]. Rigorous peptide mapping of Pap\textsuperscript{B25}-photo-products by Kurose et al. [78] (exploiting mass-spectrometric methods) provided the first identification of the conserved α\textsuperscript{CT} motif in the IR ectodomain as a critical hormone-binding element; its central structural role is highlighted in Section 3. Although incompletely visualized in electron-density maps, physical evidence that α\textsuperscript{CT} functions in \textit{trans} within the IR (αβ\textsubscript{2})\textsubscript{dimer} was provided by bifunctional Pap derivatives of insulin containing photo-probes at opposite surfaces [41].

Modifications at A2 and A3—sites partially concealed by the C-terminal segment of the B chain in the closed state—are also associated with markedly impaired activities [79, 80]. Even such subtle aliphatic substitutions as Ile\textsuperscript{A2}\textrightarrow allo-Ile (\textit{i.e.}, inversion of β-carbon chirality within the A2 side chain) [81] or substitution Val\textsuperscript{A3}\textrightarrow Leu (insulin \textit{Wakayama} [44]) cause profound defects in IR binding. In the following section we describe how detachment of the aromatic triplet at the receptor surface unmasks these A-chain residues. The latter are critical to receptor engagement via conserved α\textsuperscript{CT} contacts.

3 | INTERACTION OF THE AROMATIC TRIPLET WITH THE RECEPTOR’S PRIMARY INSULIN-BINDING SITE.

3.1 | STRUCTURE OF THE INSULIN RECEPTOR ECTODOMAIN

In order to understand the remarkable role played by the aromatic triplet of insulin in mediating both hormone storage and receptor activation, it is appropriate to provide an overview of the structural biology of the insulin receptor itself. The receptor is a member of the modular receptor tyrosine kinase (RTK) family [82, 83]; two other members belong to the same sub-family of RTKs, namely, the type 1 insulin like growth factor receptor (IGF-1R) [82] and the IR-related receptor (IRR) [84]. These three receptors are distinct from other RTKs in that they are disulfide-linked homodimers, whereas the others are monomers. Conserved modules in the IR are shown in Figure 8.

The sequence of the human IR was determined in 1985 [82, 83]. The mature form of the receptor is a disulfide-linked (αβ\textsubscript{2})\textsubscript{homodimer}, formed by furin processing of disulfide-linked pro-receptor polypeptides into α- and β chains. Within the insulin receptor homodimer, a single disulfide bond links each α chain to its β chain partner and at least two disulfide bonds cross-link the α chains to each other [85].

The receptor is heavily glycosylated: thirteen glycans are N-linked to each α chain and four to each β chain [86], and six glycans are O-linked to each β chain [87]. The human insulin receptor occurs as two splice variants, termed IR-A and IR-B, respectively, with IR-A isoform lacking (and IR-B isoform retaining) the twelve amino-acid protein product of exon 11 [88]. IR-B is the predominant isoform in peripheral tissue and is responsible for metabolic signaling [89]. IR-A is the predominant isoform within fetal tissue and also within the adult brain. IR-A shows enhanced binding to insulin-like growth factors compared to IR-B and is upregulated in certain cancers [89].
The 3D structure of the IR extracellular region (its “ectodomain”) was derived from X-ray analysis of a crystal of the isolated ectodomain in complex with four F\textsubscript{ab} fragments; the latter were attached to the ectodomain fragment in order to overcome the hindrance posed to crystallization by the extensive N-linked glycosylation [90]. Within this structure, the extracellular component of each αβ chain pair was revealed to have the following domain organization [48] (describing from the N terminus of the α chain through to the C terminus of the β chain): a leucine-rich repeat domain (L1), a cysteine-rich region (CR), a second leucine-rich repeat domain (L2), and finally three fibronectin type III domains (FnIII-1, FnIII-2 and FnIII-3; see Figure 8).

Within the holo receptor, the FnIII-3 domain is followed by a transmembrane helical segment (TM), a juxtamembrane region, the tyrosine kinase domain (TK) and a C-terminal tail segment [90]. Domain FnIII-2 contains within its canonical CC’ loop [48] a segment of approximately 110 residues termed the insert domain (ID) that in turn contains the α/β furin processing site (Figure 8). The three-dimensional crystal structure of the ectodomain has a Λ shape, with each “leg” of the Λ containing the L1-CR-L2 module of one receptor monomer juxtaposed in an anti-parallel fashion against the (FnIII-1)-(FnIII-2)-(FnIII-3) module of the opposing receptor monomer [48, 90], placing the sites of membrane entry at ends of the respective “legs” of the assembly (Figure 9).

The “head” of the Λ-shaped assembly contains the [L2-(FnIII-1)]\textsubscript{2} module, the two FnIII-1 domain components of which are disulfide bonded to each other at residue Cys524. The ID domains are mostly disordered and lie within the interior of the Λ-shaped structure, except for a segment (termed αCT, located near the C terminus of the α chain) that assembles as an α helix on the central β sheet (termed L1-β\textsubscript{2}) of the L1 domain of the opposing receptor monomer [48, 91]. A single inter-α-chain disulfide bond connects residues Cys524 (within the CC’ loop of domain FnIII-1) [90, 91], and at least one inter-α-chain disulfide bond occurs within the ID at the triplet Cys682-Cys683-(Ser684)-Cys685 [48, 85]. Each α chain is linked to its β-chain counterpart by a disulfide bond between residues Cys647 (within the α-chain segment of the ID) and Cys860 (within domain FnIII-3; IR-A numbering) [85]. All remaining disulfide bonds are intra-chain and there is a single free cysteine residue within FnIII-3 (Cys872; IR-A numbering) [85]. It should be noted that the crystal contained an (IR-A)-based construct termed IRΔβ, which was engineered to remove the heavily glycosylated segment near the N terminus of the receptor β chain (again to reduce hindrance to crystallization).

The inter-domain interfaces (both intra-monomer and inter-monomer) within the above structure are sparsely packed, with the only exception being that between the L1 domain and the N-terminal region of its immediately downstream CR domain [48, 90]. These loosely-packed interfaces are reminiscent of crystal packing interfaces, suggesting that all or some of them may disassemble upon insulin binding. However, this raises the question as to whether reported apo ectodomain structure has been modulated by F\textsubscript{ab} attachment and/or crystal lattice embedding. Two lines of evidence support the view that the Λ-shaped structure represents that adopted by the ectodomain in the context of a membrane-embedded holo receptor. First, recent negative-stain electron microscopy imaging of holo insulin receptor embedded in lipid nanodiscs show, at least qualitatively, that the Λ-shaped assembly concurs...
with that observed in the nanodisc context [92] (Figure 10). Second, crystallographic analysis of the isolated and growth factor-free IGF-1R ectodomain reveals it to have a similar shape and domain assembly to that of the insulin receptor, albeit that the IGF-1R construct employed included the attachment of a pair of antibody Fv modules [93].

### 3.2 | STRUCTURE OF MODEL HORMONE-RECEPTOR COMPLEXES

Insulin binding to the IR displays complex kinetics, characterized by curvilinear Scatchard plots and negative cooperativity. The latter was demonstrated by accelerated dissociation of labeled insulin upon dilution with cold insulin [94–97]. Further, the accelerated dissociation of insulin is reduced at high cold-insulin concentration (> 1 μM). These data can be explained by a model wherein insulin binding involves the formation of a cross-link between two distinct sites (designated site 1 and site 2’) on the receptor, where ‘ is used here and throughout to denote—within the context being described—entities from the alternate αβ monomer. Site 1 is taken to be the higher-affinity (“primary”) binding site and site 2’ the lower-affinity (“secondary”) binding site. Insulin binding to one site 1 / site 2’ pair is then proposed to reduce the affinity of insulin for the alternate site 1’ / site 2 pair (negative cooperativity). The reduction in accelerated dissociation at high insulin concentrations is explained within this model by postulating that, under such conditions, three insulin molecules can bind to the homodimeric receptor: one cross-linking the site 1 / site 2’ pair with high affinity, one binding to site 1’ and one to site 2. Insulin binding independently to the latter two sites then abrogates both the formation of a high-affinity cross-link between sites 1’ and 2 and the reduction in affinity of the site 1 / site 2’ cross-link.

The complexities of the hormone receptor-binding kinetics taken together with the structural complexities of the receptor itself have confounded attempts to crystallize the ectodomain in complex with insulin. Partial success has thus come via another route, namely, by analysis of insulin bound to fragments of the ectodomain [40, 41]. The smallest receptor fragment known to bind insulin is the isolated two-domain L1-CR construct in the presence of an exogenous peptide segment spanning the receptor αCT helix [98], which together display nanomolar affinity for insulin and arguably reconstitute site 1. This tandem receptor element (L1-CR + αCT) has been termed the insulin “micro-receptor” (μIR) [41], in the same vein of nomenclature as the earlier terms “mini-receptor” (the receptor L1-CR-L2 module plus an exogenous αCT peptide [99, 100]) and “midi-receptor” (the receptor [L1-CR-L2-(FnIII-1)]2 fragment wherein each constituent FnIII-1 domain is extended at its C terminus by the α-chain component of the ID domain [98, 101]).

An X-ray crystallographic structure of a μIR complex consisting of (i) receptor residues 1–310 (an L1-CR fragment), (ii) an αCT peptide fragment spanning residues 704–719 of IR-A and (iii) human insulin, has been determined to a resolution of 3.5 Å [40, 41] (Figure 11).

This structure reveals conformational change both in insulin and in the αCT segment upon their engagement and dissection of the atomic details has led to significant structural insight into the insulin’s classical structure-activity relationships. The details of the insulin / insulin micro-receptor complex are as follow [40, 41].(i) The αCT helix, compared to its conformation within the apo ectodomain structure, has undergone restructuring upon the L1-β2 surface (Figure 12). In particular, its axis is rotated ~30° to lie approximately at right
angles to the direction of the β strands. In addition, it has unwound at its N terminus and wound on at its C terminus; the resultant helix spans residues 705–715 rather than residues 694–710.

(ii) The insulin B-chain helix (observed here to span residues B9-B20) lies on the C-terminal edge of the L1-β2 surface, packing adjacent and anti-parallel to the αCT helix. To allow such accommodation, insulin B-chain residues B24 to B27 fold away from the core of the hormone, with the direction of the polypeptide segment B24-B27 now being approximately perpendicular to the axes of the receptor αCT helix and insulin B-chain helix [41] (Figure 13).

(iii) No crystallographic electron density is apparent in this structure for insulin B-chain residues beyond residue B27, suggesting a lack of specific interaction of these elements with the receptor. (iv) No indication is apparent in this structure that the N-terminal helical extension of the insulin B-chain helix is formed upon microreceptor engagement. (v) The side chains of the αCT residues His 710 and Phe714 engage the exposed hormone core directly, with that of Phe714 effectively occupying the position of the side chain of TyrB26 in receptor-free insulin (Figure 14).

3.3 | PACKING OF THE AROMATIC TRIPLET AT HORMONE-RECEPTOR INTERFACE

Upon displacement of the B-chain C terminus from the hormone core, the side chain of insulin residue PheB24 undergoes a rotameric re-configuration leaving the phenyl ring in effectively the same relative place within the hormone core that it occupies in receptor-free insulin (Figure 15). Such positioning buries insulin residue PheB24 within a large hydrophobic cavity formed by insulin residues TyrA19, CysA20, LeuB15, receptor αCT residue Phe714 and receptor L1 domain residues An15, Leu37 and Phe39 (Figure 16).

In contrast to the positioning of the side chain of PheB24, the side chains of residues PheB25 and TyrB26 are largely exposed within the μIR complex. The side chain of residue PheB25 engages the αCT residues Val715 and Pro718 and is otherwise positioned adjacent to that of insulin residue ThrB27 (Figure 17). The side chain of residue TyrB26 is rotated away from the hormone core, stacking parallel to the side chains of the receptor L1 domain residues Asp12 and Arg14 (Figure 18).

Residue ThrB27 appears weakly ordered in the μIR complexes, with B chain residues beyond B27 almost completely disordered. The latter observation is consistent with the ability of these residues to tolerate significant mutation, chemical modification or deletion [6, 102].

4 | FUTURE PERSPECTIVES

A major challenge is posed by extension of present structures of model hormone-receptor complexes [47, 48] to the holo-receptor. Such progress is likely to require a series of structures, each representing a step in the complex mechanism of conformational change leading to signal propagation. A significant advance was recently obtained through single-particle cryo-electron microscopy analysis of the insulin-ectodomain complex [49]. Because the isolated ectodomain lacks a single high-affinity site and because the present analysis
encompassed only ca. 10% of the images, it is not yet clear how the rearrangement of fibronectin-homology domains observed in the structure [49] relates to the structure of the free ectodomain [47, 48] or μIR complexes [40, 41]. Of particular interest will be understanding the connection between Site 2 as inferred from the cryo-EM structure [49] and Site 2 as defined by kinetic analysis of insulin analogs [96, 97]. We speculate that the aromatic triplet plays a critical role to initiate a series of conformational changes associated with the modular reorganization of the ectodomain on hormone binding.

The aromatic triplet of vertebrate insulins and insulin-like growth factors may not be essential for insulin-like ligands. The recent structure of a venom insulin-like protein from an invertebrate (Conus geographus) provides an example of a “miniature” insulin lacking the C-terminal β-strand of the B chain [103]. It is possible that such minimal insulin scaffolds could provide templates for design of novel therapeutic insulins. The C. geographus ligand contains multiple post-translational modifications whose contributions to stability and activity will require characterization.

A key constraint in the pharmacology of insulin analogs is posed by chemical- and physical degradation [7, 18]. The receptor-bound open conformation of the hormone may be particularly susceptible to such degradation, including formation of amyloid (insulin fibrillation). A promising structure-based route to the engineering of ultra-stable analogs is provided by foreshortened C domains [104, 105]. In such single-chain insulins (SCIs) the aromatic triplet resides in a closed conformation—tethered by a foreshortened connecting peptide—and yet sufficient “play” is provided to enable its detachment on receptor binding. It would be of future interest to investigate crystal- or cryo-EM structures of SCI-receptor complexes as probes of the closed-open transition. In the context of this issue, dedicated to the late Prof. Steiner, it would be fitting if the B-C-A structure of proinsulin [22, 23] should find application in next-generation therapeutic insulin analogs.

We envisage that continuing characterization of an insulin-related superfamily, spanning 500 million years in the natural history of both vertebrates and invertebrates, will provide an increasingly fertile model for studies of protein evolution. How and why particular amino acids are conserved are questions that cross from structural biology to cell biology and physiology. Whereas the structural origins of some evolutionary constraints (such as co-conservation of critical hormone-receptor contacts) may be apparent in structures (e.g., of complexes at higher resolution), deciphering other constraints may require study of off-pathway events, such as toxic protein misfolding [18–21]. Other residues may contribute to on-pathway folding efficiency and yet be dispensable once the native state is achieved.

Extant insulin sequences thus lie at the intersection of multiple explicit and implicit constraints (Figure 19). Many—and indeed perhaps most—residues in insulin are likely to play different roles at distinct stages in the hormone’s complex “life cycle”: dissecting these roles will require high-resolution analysis of conformational change in this remarkable protein from biosynthesis to function and degradation.
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Figure 1.
Office of Prof. D.F. Steiner at the University of Chicago. Shown on table at front in plexiglass box is a plastic model of insulin as a crystallographic protomer. *Inset at top right,* Hodgkin and Steiner with matching molecular models. This figure was originally published in a remembrance of D.F. Steiner [12] and is reprinted here with permission of the authors. The molecular model is now on display as part of an historical collection in the Division of Endocrinology at the Pritzker School of Medicine. One of the authors (MAW) recalls many pleasant and stimulating discussions in this office during which Don references one or more side chains in this model. Often these included the aromatic triplet—Phe$^{B24}$, Phe$^{B25}$ and Tyr$^{B26}$—in light of their special role in the Hodgkin structure.
Figure 2.
Insulin self-assembly as defined in the 1969 crystal structure of insulin [9, 10]. (A) T₆ (or 2-Zn) insulin hexamer, (B) Constituent dimer and (C) anti-parallel C-terminal dimer-related β-sheet (residues B24-b28; arrows) with the side chains of residues B24-B26 shown in red, green and violet, respectively. Coordinates were obtained from Protein databank entry 4INS.
Figure 3.
The two classical conformations of insulin: T and R. TR dimer as extracted from crystal structure of a T₃R₃ zinc insulin hexamer, first observed in the classical “4-Zn” crystal form [27]. The position of B8 is indicated by a purple circle within the B-chain helix (green), which comprises residues B9-B19 in the T state, B1-B19 in the R state (or B3-B19 in the “frayed” R² state).
Figure 4.

$^1$H-NMR spectroscopy defines T-like conformation of an engineered insulin monomer. Solution structure of DKP-insulin highlighting the aromatic triplet (Phe$^{B24}$, Phe$^{B25}$ and Tyr$^{B26}$) in brown relative to Val$^{B12}$, Leu$^{B15}$ and cysteine B19-A20 in blue. In this monomeric analog dimerization was impaired by the paired substitutions Lys$^{B28}$ and Pro$^{B29}$ [57], and the trimer interface was disrupted by Asp$^{B10}$ [58, 59]. The A chain is shown in yellow, and B chain in orange (B1-B19) and brown (B20-B30).
Figure 5.

Conservation of the “Aromatic Triplet”: sequence alignment of residues B20-B30, which encompasses a β-turn (B20-B23), β-strand (B24-B28) and less well-ordered terminal residues (B29 and B30). Phe is invariant among vertebrate insulins at positions B24 (red) and B25 (green) whereas Tyr is highly conserved at position B26 (lilac). The divergent insulin in the rodent O. degu is associated with insulin fibrillation of the islet, leading to β-cell death and senile DM [65, 66].
Figure 6.
Aromatic triplet at dimer interface. Stereo pairs showing Phe$_{B24}^{B24}$ (red), Phe$_{B25}^{B25}$ (green) and Tyr$_{B26}^{B26}$ (violet) in dimer-related β-sheet in (A) T$_6$ hexamer (B) T$_3$R$_3$ hexamer and (C) R$_6$ hexamer. Coordinates were obtained from Protein Databank entries 4INS, 1TRZ and 1ZNJ, respectively. Dotted lines indicate β-sheet-related hydrogen bonds.
Figure 7.
T-state structure of insulin. Aromatic triplet is part of an extensive non-polar surface of the insulin monomer. (A) Stick representation highlighting side chains of Phe$_{\text{B24}}$ (red), Phe$_{\text{B25}}$ (gray; not labeled at right), and Tyr$_{\text{B26}}$ (purple at upper right) relative to $\alpha$-helical residues Val$_{\text{B12}}$ (blue), Leu$_{\text{B15}}$ (green) and Tyr$_{\text{B16}}$ (purple at lower left). The internal disulfide bridge of cystine B19-A20 is shown in yellow. The A chain is shown in dark gray, and B chain otherwise shown in light gray. (B) Corresponding dot-surface representation of the highlighted residues with neighboring residues B20-B23, B25 and B27 in light gray; A20-A21 is dotted in dark gray.
Figure 8.
Primary structure of the insulin receptor (αβ)_2 homodimer. L1: first leucine-rich repeat domain; CR: cysteine-rich region; L2: second leucine-rich repeat domain; FnIII-1, FnIII-2 and FnIII-3: first, second and third fibronectin type III domains; ID: insert domain (IDα: α-chain component; IDβ: β-chain component); αCT: C-terminal segment of the receptor α chain; TM: trans-membrane segment; JM: juxta-membrane segment; TK: tyrosine kinase domain; C-tail: C-terminal tail of the receptor β chain. Inter-chain disulfide bonds are indicated by solid green lines, individual N-linked glycosylation sites by “Y” symbols and the location of the O-linked glycosylation segment by an orange “V” symbol.
Figure 9.
Crystal structure of the insulin receptor (αβ)₂ ectodomain. One αβ monomer is shown in ribbon representation (foreground), the other in molecular surface representation (background). The coloring of domains is as in Panel A. Part of IDα is indicated as a dashed line (foreground monomer only), as it was disordered within the crystal. The schematic is based on PDB entry 4ZXB [48].
Figure 10.
Negative-stain transmission electron microscopy images of holo insulin receptor embedded in nanodiscs [92]. The left-hand panel is an image of the receptor in an insulin-free state, the right-hand panel is an image of the receptor in an insulin-bound state, with the panels underneath each offering an interpretation of the respective images above. The correspondence of the Λ-shaped particle in the left-hand panel to those of that of the crystal structure of the isolated ectodomain is apparent. The right-hand panels show the closing of the receptor legs upon addition of insulin and a likely folding out of the L1-CR-L2 modules. Republished with permission of the Rockefeller University Press, from “Journal of Cell Biology, Gutmann et al., v217, p1643 (2018)”; permission conveyed through Copyright Clearance Center, Inc. [92].
Figure 11.
Interaction of insulin with the receptor’s primary binding site. Schematic shows overall organization of the insulin microreceptor (μIR, the L1-CR module + an αCT peptide) in complex with insulin [40, 41]. The schematic is based on PDB entry 4OGA [41].
Figure 12.
Conformational change in the primary binding site of the IR upon insulin binding. Schematic shows rotation and re-organization of αCT on the surface of domain L1 upon insulin binding. The schematic is based on PDB entries 4ZXB [48] and 4OGA [41].
Figure 13.
Folding away from the hormone core of insulin residues B24-B30 upon engagement with μIR. The receptor-bound insulin B chain is in black and A chain in white, overlaid with the B chain of the free hormone in cyan. The schematic is based on PDB entries 4INS \cite{6} and 4OGA \cite{41}.
Figure 14.
Details of the interaction of αCT residue His710 and Phe714 with the hormone core. The insulin A chain is in white with green side chains, the B chain is in black with light blue side chains. The receptor L1 domain is in tan and the receptor αCT helix is in magenta. The schematic is based on PDB entry 4OGA [41].

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Figure 15. 
Re-positioning of insulin residues B24-B27 upon engagement with μIR. The receptor-bound insulin B chain is in *black* and A chain in *white*, overlaid with the B chain of the free hormone in *cyan*. The schematic is based on PDB entries 4INS [6] and 4OGA [41].
Figure 16.
Atomic environment of Phe\textsuperscript{B24} within the Site 1 \mu R complex. Schematic shows all side chains of all residues within 4 Å of that of insulin PheB24 (unlabeled, orange). Insulin A chain is in \textit{white}, insulin B chain is in \textit{black}, L1 domain of the insulin receptor is in \textit{tan}, αCT segment of the insulin receptor is in \textit{magenta}. Receptor residues have \textit{underlined} labels whereas insulin residues are in \textit{normal} font. The schematic is based on PDB entry 4OGA [41].
Figure 17.
Atomic environment of PheB25 within the Site 1 μIR complex. Schematic shows all side chains of all residues within 4 Å of that of insulin PheB25 (unlabeled, orange). Color and labeling schemes are otherwise as in Figure 12. (q.v.). The schematic is based on PDB entry 4OGA [41].
Figure 18.
Atomic environment of Tyr^{B26} within the Site 1 μIR complex. Schematic shows all side chains of all residues within 4 Å of that of insulin Tyr^{B26} (unlabeled, orange). Color- and labelling schemes are otherwise as in Fig. 12. (q.v.). The schematic is based on PDB entry 4OGA [41].
Figure 19.
Evolutionary constraints and insulin fibrillation: Venn diagram showing intersection of multiple constraints: function, foldability, misfolding and assembly.