Insulin in motion: The A6-A11 disulfide bond allosterically modulates structural transitions required for insulin activity

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The structural transitions required for insulin to activate its receptor and initiate regulation of glucose homeostasis are only partly understood. Here, using ring-closing metathesis, we substitute the A6-A11 disulfide bond of insulin with a rigid, non-reducible dicarba linkage, yielding two distinct stereo-isomers (cis and trans). Remarkably, only the cis isomer displays full insulin potency, rapidly lowering blood glucose in mice (even under insulin-resistant conditions). It also possesses reduced mitogenic activity in vitro. Further biophysical, crystallographic and molecular-dynamics analyses reveal that the A6-A11 bond configuration directly affects the conformational flexibility of insulin A-chain N-terminal helix, dictating insulin’s ability to engage its receptor. We reveal that in native insulin, contraction of the $C_{\alpha}$-$C_{\alpha}$ distance of the flexible A6-A11 cystine allows the A-chain N-terminal helix to unwind to a conformation that allows receptor engagement. This motion is also permitted in the cis isomer, with its shorter $C_{\alpha}$-$C_{\alpha}$ distance, but prevented in the extended trans analogue. These findings thus illuminate for the first time the allosteric role of the A6-A11 bond in mediating the transition of the hormone to an active conformation, significantly advancing our understanding of insulin action and opening up new avenues for the design of improved therapeutic analogues.

Insulin is fundamental to the physiological regulation of blood glucose concentration1. A deficiency in insulin results in diabetes, a major economic and primary health care burden across both developed and developing countries. Insulin therapy is essential in both type 1 diabetes and late-stage type 2 diabetes, with current therapeutic insulins being designed to restore the normal biphasic insulin response to food intake2,3. While such therapeutic analogues are largely successful in controlling blood glucose levels, their means of administration and their pharmacokinetic and pharmacodynamic profiles are far from ideal, putting patients at risk of both hyper- and hypoglycaemia. Notably, none of the currently available therapeutic insulin analogues have employed in their design an atomic-level understanding of how insulin engages its receptor, as such detail has only recently begun to emerge4,5. A thorough understanding of the conformational changes involved in insulin/insulin receptor interaction therefore has the potential to lead to a new generation of insulin analogues with improved pharmacological properties.

Insulin is a two-chain polypeptide, comprising an A chain of 21 residues that includes two $\alpha$ helices (residues A1 to A8 and A12 to A18, respectively), and a B chain of 30 residues that includes a single $\alpha$ helix (residues B9 to B19) (Fig. 1a). Integral to insulin’s structure are its three disulfide bonds — one intra-chain (CysA6-CysA11) and two inter-chain (CysA7-CysB7 and CysA20-CysB19) (Fig. 1a). Formation of these disulfide linkages ensures both the
or cis and adopts either a trans configuration. Introduction of a dicarba bond into a number of small polypeptides leads to a new understanding of the critical interplay between A-chain conformational flexibility and restraint that is allosterically regulated by the dynamics and biological characterization of these dicarba insulin isomers. Taken together with a re-analysis of extant T-state insulin crystal structures, our structural, molecular analogy of human insulin. Here, the highly hydrophobic N-terminus of the insulin A-chain necessitated the C-terminal asparagine residue on resin cleavage, Fmoc-L-Asp-O\textsubscript{t}-Bu was loaded onto Rink amide resin via its side chain. Microwave-accelerated SPPS in combination with HATU–DIPEA activation and Fmoc-protected amino acids were used to generate the truncated peptide sequence (Fig. 2), carrying through each intermediate without purification and characterization. Two strategically placed L-allylglycine residues were incorporated into the primary sequence to facilitate formation of the intra-chain dicarba bridge, and cysteine residues were orthogonally protected to later aid regioselective dicarba oxidation and tethering of the location of the three \(\alpha\) helices and the three disulfide bonds. (b) Schematic diagram of native cystine and isomeric cis- and transdicarba bridges.

**Figure 1.** Insulin sequence and structure. (a) Primary sequence (top) of the A (blue) and B (grey) chains of human insulin, highlighting disulfide bonds (yellow), site 1-binding residues (underlined) and site 2-binding residues (bold)\textsuperscript{14}. Ribbon diagram of insulin (2Zn-coordinated \(\alpha\)\(\beta\) conformation\textsuperscript{16} PDB entry 1MSO) showing the location of the three \(\alpha\) helices and the three disulfide bonds. (b) Schematic diagram of native cystine and isomeric cis- and transdicarba bridges.

**Results**

**Chemical synthesis.** While dicarba bonds have been previously introduced into the analogous bond of insulin-like peptides INSIL3 and INSIL7\textsuperscript{21–23}, no strategy exists in the literature to generate A6-A11 intra-chain dicarba analogues of human insulin. Here, the highly hydrophobic N-terminus of the insulin A-chain necessitated the development of an interrupted solid phase peptide synthesis (SPPS)-catalysis approach\textsuperscript{24} to overcome deleterious aggregation and achieve quantitative ring closing metathesis (see Supplementary Methods). Additionally, to ensure exclusive generation of the required C-terminal asparagine residue on resin cleavage, Fmoc-L-Asp-O\textsubscript{t}-Bu was loaded onto Rink amide resin via its side chain. Microwave-accelerated SPPS in combination with HATU–DIPEA activation and Fmoc-protected amino acids were used to generate the truncated peptide sequence (Fig. 2), carrying through each intermediate without purification and characterization. Two strategically placed L-allylglycine residues were incorporated into the primary sequence to facilitate formation of the intra-chain dicarba bridge, and cysteine residues were orthogonally protected to later aid regioselective dicarba oxidation and tethering of the location of the three \(\alpha\) helices and the three disulfide bonds. (b) Schematic diagram of native cystine and isomeric cis- and transdicarba bridges.
of the B chain (Fig. 2). It was critically important to perform the catalysis without the five N-terminal residues; performing the ring closure on the full A-chain sequence (21 mer), unlike other insulin-super family molecules, gave only poor conversion. Hence, ring-closing metathesis (RCM) of the fully protected, truncated resin-tethered peptide 1 (16 mer) was performed in the presence of 20 mol% second-generation Grubb’s catalyst in DCM with 0.4 M w/v LiCl in DMF. Under these conditions, microwave irradiation of the peptidyl-resin at 100 °C for 2 h resulted in near quantitative conversion to the desired carbocycle 2. Continued microwave-accelerated SPPS was then performed and the remaining five residues (GIVEQ) were appended to the N-terminus to deliver the complete dicarba insulin A chain. Mass spectral analysis of the 21-mer gave the required molecular ions for carbocycle 3 and the RP-HPLC trace showed the formation of two geometric isomers (E-3 and Z-3) in a 3:1 ratio (see Fig. S1c). Following resin cleavage, crude peptide 3 was exposed to an acidic cleavage mixture containing 2,2′-dipyridyl disulfide to facilitate concerted tert-butyldiprotection and pyridinyl-reprotection of residue CysA7. Each of the resultant isomeric dicarba insulin peptides 4 were then purified before being subjected to regioselective chain coupling. Construction of the complementary insulin B chain 5 was achieved through microwave-accelerated SPPS, in combination with HBTU/HOBt–DIPEA activation and Fmoc-protected amino acids, on preloaded Fmoc-Thr(tBu)-PEG-PS resin. During chain elongation, orthogonally protected Cys(Trt) and Cys(Acm) residues were strategically incorporated into the primary sequence in positions B7 and B19 respectively. Preparative RP-HPLC gave the required insulin B chain 5 in 30% yield and 90% purity. The monocyclic A-B conjugates were prepared by combination of the dicarba insulin A chains (E-4, Z-4) with the insulin B chain 5 under basic conditions. In all cases, oxidation was complete within minutes giving the required 51 amino acid peptides (Fig. 2, E-6 and Z-6). Mass spectral analysis of the isolated solids supported formation of the covalent A-B dimers. The final disulfide bridge in the c[Δ4A6,11]-dicarba human insulins (Fig. 2, E-7 and Z-7) was formed on exposure of each of the monocyclic A-B conjugates (E-6, Z-6) to iodine under acidic conditions. Removal of the acetamidomethyl (Acm) protecting groups at positions A20 and B19 resulted in spontaneous oxidation of the liberated free thiol groups to give the two target isomeric trans- and cis-dicarba insulin peptides (Fig. 2, peptides E-7 and Z-7, respectively), which were then purified by RP-HPLC and independently subjected to biological testing and structural analysis.

**Stereochemical assignment of the dicarba bridge.** The Cα chemical shifts of peptides comprising Δ4-diaminosuberic acid (Δ4Sub) residues show appreciable differences between the cis-(Z) and trans-(E) configurations25. These features were used to identify the stereochemistry of the insulin A-chain A6-A11 dicarba
bridge without the need for structural calculations. Hence, TOCSY and $^{13}$C-HSQC spectra were acquired on each of the dicarba insulin A-chain peptides (E-4 and Z-4). $\Delta^{15}$Sub H$^\beta$ resonances for each E- and Z-dicarba isomer were readily identifiable at $\sim$5.6 ppm, which facilitated the assignment of associated $\Delta^{15}$Sub H$^\beta$ resonances as well as $\Delta^{15}$ Sub C$^\gamma$ resonances in the $^{13}$C HSQC. As with other dicarba peptide sequences, only minor differences in carbon chemical shifts were observed between the two isomers, with the major difference occurring at the C$^\gamma$ atoms of the dicarba bridge (Fig. S2)$^{29}$. Consistent with the previously reported model for stereochromical assignment, the upfield C$^\gamma$ shifts for the cis isomer presented at $\delta^{30.9}$ and $\delta^{32.8}$ and those for the trans isomer appeared at $\delta^{36.4}$ and $\delta^{36.9}$. The stereochemistry of the trans isomer E-7 was also confirmed by X-ray crystallography. Additionally, an independent, stereoselective synthesis of cis-[A6–11]-dicarba insulin A chain (Z-3) (Fig. SII) was achieved using a preformed, orthogonally protected Z-configured diaminobutyric acid residue (cis-S1)$^{25}$ and SPPS (Fig. S3) and found to be identical to material obtained via RCM (Fig. S1c).

**Receptor binding and activation.** The affinities of the two dicarba insulin isomers for the human insulin receptor isoforms IR-A and IR-B and the human type 1 insulin-like growth factor receptor (IGF-1R) were determined using ligand competition binding assays$^{19}$. The cis isomer Z-7 was found to bind IR-B and IR-A with similar affinity to native insulin, whereas the trans isomer E-7 had a $\sim$50-fold lower affinity for IR-B and IR-A than native insulin (Fig. 3a, Table S1 and Fig. S4a). The cis isomer Z-7 bound IGF-1R with similar affinity to insulin (Table S1 and Fig. S4c), whereas trans isomer E-7 exhibited negligible affinity to IGF-1R.

In receptor phosphorylation assays, the cis isomer was equipotent in insulin activation of IR-B (Fig. 3b) and IR-A (Fig. S4b), but slightly poorer than insulin in activation of IGF-1R (Fig. S4d), whereas the trans isomer was $\sim$1000 fold less potent than insulin in activating IR-B (Fig. 3b). The trans isomer had very low binding affinity for the IR-A and IGF-1R and its activity at these receptors was left undetermined.

In summary, these data indicate that (i) only one of the two stereochromical isomers of A6-A11 dicarba insulin (viz., the cis isomer) is a potent analogue, and (ii) the equipotency of the cis analogue to native insulin is maintained despite the increase in rigidity across the A6-A11 connection afforded by the unsaturated dicarba bond.

**Metabolic and mitogenic activity.** The cis isomer was equipotent to native insulin in promoting glucose uptake by NIH3T3-L1 adipocytes (Fig. 3c), aligning with its equipotency in the above receptor binding and receptor activation assays. However, the cis isomer was 5–10 fold less potent than native insulin in promoting DNA synthesis, despite its equal affinity for IR and IGF-1R (Fig. 3d). The trans isomer was unable to stimulate DNA synthesis significantly above basal levels, correlating with its poor receptor binding ability.

**Insulin tolerance test.** The cis isomer lowered blood glucose more effectively than native insulin in an insulin tolerance test in mice (Fig. 3e,f) measured as described$^{27}$. This was also evident in insulin-resistant mice by the root-mean-square deviations (RMSDs) of the bond angle and bond lengths from ideality; Table 1). The cis isomer structure the A1-A10 segment is not involved in crystal contacts (or at least not in the putative conformation in which it has been modelled).

**Crystal structure determination.** The X-ray crystal structures of the cis- and trans isomers were determined using diffraction data to 2.70 Å and 1.55 Å resolution, respectively, with the crystals being grown under similar conditions (see Methods). Data processing and refinement statistics are presented in Table 1. Both crystal structures exhibit a T-state-like conformation in that the B chain N-terminal segment adopted an extended conformation folded back against the interface between the B chain helix and the polypeptide linker between the two A chain helices (Fig. 5a,b). Both crystal structures also contain dimers formed in a fashion closely similar to that of the classical insulin T-state dimer. In the crystal structure of the cis isomer, resides A1 to A10 appear largely disordered, with the A chain N-terminal helix being represented by no more than a relatively featureless “blob” of difference electron density (Fig. 5d). The dimensions of this blob loosely approximate that of the polypeptide core of the 8-mer $\alpha$-helix of the native hormone and likely reflect a crystallographic superposition of the helix in a variety of azimuthal and axial orientations. Various unsuccessful attempts were made to obtain alternative crystals of the cis isomer with higher structural definition of the A-chain N-terminal helix. Nevertheless, we were able to build a tentative model of the cis isomer into the available electron density and to refine this model crystallographically to acceptable $R_{\text{meas}}$/$R_{\text{free}}$ statistics, though with poor overall stereochemistry (as evidenced by the root-mean-square deviations (RMSDs) of the bond angle and bond lengths from ideality; Table 1). The Ramachandran plot statistics for the two structures are: trans isomer: 99% in the favoured region, none in the disallowed region; cis isomer: 72% in the favoured region, 11% in the disallowed region (the latter values aligning with the poor stereochemical nature of the model). We note that, in the trans isomer structure, the A1-A10 segment is involved in crystal contacts whereas in the cis isomer structure the A1-A10 segment is not involved in crystal contacts (or at least not in the putative conformation in which it has been modelled).
Figure 3. Insulin receptor binding, activation and biological activities of cis- and trans isomers. (a) Competition binding of insulin (squares), cis- (triangles) and trans- (circles) isomers with europium-labelled insulin. Results are expressed as a percentage of binding in the absence of competing ligand (%B/B0). (b) Activation of IR-B by increasing concentrations of dicarba insulins (10 min stimulation) is expressed as receptor phosphorylation as a percentage of the maximal phosphorylation induced by insulin. Insulin vs cis isomer (non significant); insulin vs trans isomer ****(P ≤ 0.0001) (2-way ANOVA; Dunnett’s multiple comparison) (c) Glucose uptake stimulated by increasing concentrations of insulin or cis isomer is expressed as fold glucose uptake (pmol/min/mg) above basal. Insulin vs cis isomer (ns) (paired T-test). (d) DNA synthesis in response to increasing concentrations of dicarba insulins is shown as percentage incorporation of ³H-thymidine (³H-Thy) above basal. All data in (a–d) are the mean ± S.E.M. n = at least 3 independent experiments. (e) Insulin tolerance test in mice fed on a normal diet (chow), or (f) on a high fat diet were administered through intraperitoneal injection (ip) with 0.75 IU/kg insulin (squares; solid lines) or cis isomer (triangles; dotted lines) under non-fasting conditions and tail vein blood glucose was measured via glucose meter at indicated times. n = 5–6 per group. Blood glucose levels are expressed as change over basal levels (mmol/L). Chow diet, insulin vs cis isomer ***(P ≤ 0.01); high fat diet, insulin vs cis isomer ***(P ≤ 0.01) (paired T-test).
In order to compare the three-dimensional structure of the trans isomer determined here with those of native insulin, we began by analysing the conformations of residues A1-A11 in extant insulin crystal structures in the Protein Data Bank (PDB; see Table S3). Inspection of T-state structures of (receptor-free) insulin monomers, including monomers from T2, T6, T3R3 and T3Rf3 assemblies by alignment of the respective B-chain helices (residues B8-B20) revealed that they could be partitioned into two classes on the basis of the conformation of the A-chain N-terminal helix: Class 1 (30 structures), in which residues A1 to A9 exhibit the classical (i, i+4) α-helical hydrogen-bonding pattern; and Class 2 (93 structures), wherein residues A1 to A5 form a single α-helical turn and residues A3 to A9 adopt a wider helix conformation, approximating an (i, i+5) π-helix (Fig. 5c,f). The distinction between these two classes is most apparent in the hydrogen bonding exhibited by the backbone amide of Thr A8: in Class 1 structures the amide forms a canonical α-helical (i, i+4) hydrogen bond with the backbone carbonyl oxygen of GluA4; however, in the Class 2 structures, the amide hydrogen bonds to the backbone carbonyl of ValA3. In many cases, these two classes occur within the same crystallographic asymmetric unit structure, with 27 out of 30 Class 1 structures being from crystals that also contain within their asymmetric unit an insulin of Class 2 conformation.

Concomitant with these differences in helical conformation are differences in the relative azimuthal positioning of residues A1-A5 about the helix axis (Fig. 5f and Table S4). In addition, the corresponding mean CαA6-to-CαA11 distance of the Class 1 insulins (4.78 ± 0.13 Å) is slightly longer than the mean CαA6-to-CαA11 distance of the Class 2 insulins (4.55 ± 0.37 Å). Concomitantly, the mean CαA7-to-CαB7 distance of the Class 1 insulins (4.62 ± 0.12 Å) is slightly shorter than the mean CαA7-to-CαB7 distance of the Class 2 insulins (4.76 ± 0.11 Å) (Table S5).

The crystal structure of the trans isomer corresponds to a Class 1 insulin conformation. Residues A1-A9 exhibit a classical α-helical geometry, with the azimutal positioning of residues A1-A5 being closely similar to that of native insulins within Class 1 (Fig. 6a). In addition, the CαA6-to-CαA11 distance (5.17 Å) in the trans isomer is more similar to the corresponding average CαA6-to-CαA11 distance of the Class 2 insulins than the average CαA6-to-CαA11 distance of the Class 2 and IR-bound native insulin (PDB entries 4OGA, 3W12, 3W13) (Table S5).

Our analyses are consistent with the report of Kaarlsholm et al. of two forms of insulin that differ at residues A1-A5 through a rotation of 32° about the Cys46 Cα-NH bond in the crystal structure of 2Zn porcine insulin. The Class 2 conformation of the A chain N-terminal helix is also observed in insulin analogues synthetically engineered to reposition the B chain C-terminal segment away from the hormone core, e.g., des[23–30]-insulin (PDB...
Our study is, however, the first to recognize the allosteric role played by the A6-A11 disulfide bond in this conformational switch without movement of the B chain C-terminal segment.

As far as we can ascertain from the structures of the insulin/IR Site 1-complexes (PDB entries 4OGA, 3W12 and 3W13, at resolution (3.5–3.9 Å)), the A-chain N-terminal helix of insulin (and insulin analogues) exhibits a Class 2-like structure when bound to IR (see Fig. 6). This is apparent in the larger-diameter A-chain N-terminal helix and shorter CαA6-CαA11 distance than those observed for Class 1 insulins, and in a similar azimuthal positioning of the constituent Cα atoms to those within Class 2 structures. The ability of the insulin A-chain N-terminal helix to adopt these two classes of structure indicates both (i) a degree of conformational flexibility in the three covalent bonds that connect that helix to the remainder of the hormone (viz., the A6-A11 and A7-B7 disulfide bonds and the downstream peptide bond), and (ii) rotameric plasticity of the side chains of the residues that form the interface between that helix and the remainder of the hormone (Fig. 5c,f).

Molecular dynamics simulations. Molecular dynamics (MD) simulations were conducted to explore the significance of conformational flexibility in the A6-A11 linkage and the A1-A9 helix — in particular, the distinction between the Class 1 and Class 2 conformations. Three high-resolution X-ray crystal structures were selected, each having a different conformation of the A6-A11 disulfide linkage (PDB entry 1G7B chains E and F, Class 2; PDB entry 1MSO chains A and B, Class 1; and PDB entry 3I3Z chains A and B, Class 2). Each structure was also modified to contain cis and trans isomers, resulting in nine starting structures. Multiple 200 ns MD simulations were then conducted based on each starting structure. In all simulations, it was evident that the A-chain N-terminal helix was highly mobile and, in many cases, interchange between classes was also observed. Indeed, although each starting structure belonged initially to a particular Class, that distinction appeared not to survive the pre-optimization phase, and hence did not bias the simulations.

Root mean square deviations (RMSDs) with respect to a representative Class 2 structure were calculated for all atoms of residues A1 to A4 for each frame of each MD simulation. From the RMSD plots (Fig. 5g–i), it is seen that both insulin and the cis isomer can approach the Class 2 conformation (RMSD < ~1.5 Å), albeit rarely (Fig. 5g,i). In contrast, the trans isomer almost never adopted a Class 2 conformation, with RMSD values being generally > 2 Å and very rarely < 1.5 Å (two disparate frames out of 12,000 had an RMSD < 1.5 Å) (Fig. 5h).

Interestingly, three different hydrogen-bonding partners for the backbone amide of Thr A8 were observed in the structures generated in the MD simulations: Val A3 (i, i + 5; π-helix), Glu A4 (i, i + 4; α-helix) and Gln A5 (i, i + 3; 310-helix) (Table S6). For both insulin and the two dicarba isomers, Glu4 and Gln5 were the dominant hydrogen bonding partners. A backbone hydrogen bond between ThrA8 and ValA3 was seen only in ~4% of simulation frames for insulin and the cis isomer and never in any of the trans isomer MD runs. Cluster analysis suggests

**Table 1.** X-ray diffraction data processing and refinement statistics. “Diffraction data are from a single crystal in both instances. Resolution limits were set based on the CC1/2 correlation statistic being assessed significant at the P = 0.001 level of probability. Free set comprised 5% of the reflections.

|                      | cis isomer | trans isomer |
|----------------------|------------|--------------|
| **Data collection**  |            |              |
| Space group          | 12, 3      | P2, 3        |
| Cell dimensions a, b, c (Å) | 79.78, 79.78, 79.78 | 77.28, 77.28, 77.28 |
| Resolution (Å)       | 28.20–2.70 | 30.0–1.55    |
| Rmerge               | 0.066 (1.653) | 0.144 (5.03) |
| Rwork (I)            | 16.19 (1.13) | 11.72 (0.38) |
| CC1/2                | 0.999 (0.280) | 0.999 (1.07) |
| Completeness (%)     | 98.8 (98.9) | 99.9 (1.00)  |
| Redundancy           | 6.0 (6.2)  | 10.8 (10.4)  |
| Molecules/asymmetric unit | 1          | 2            |
| **Refinement**       |            |              |
| Resolution (Å)       | 28.20–2.70 | 30.0–1.55    |
| No. reflections      | 2403       | 22449        |
| Rmerge/Rwork         | 0.223/0.282^2 | 0.195/0.215^2 |
| No. atoms            | 404        | 827          |
| Protein              | 120.       | 33.4         |
| Ligand/ion           | 0          | 0            |
| Water                | 0          | 97           |
| B-factors            |            |              |
| Protein (Å^2)        | n.a.       | 46.7         |
| Water                | n.a.       | 46.7         |
| R.m.s. deviations    |            |              |
| Bond lengths (Å)     | 0.011      | 0.006        |
| Bond angles (°)      | 1.5        | 0.8          |

1DEI) and NMeAla-B26-D1 insulin analogue (PDB 2WRX). Our study is, however, the first to recognize the allosteric role played by the A6-A11 disulfide bond in this conformational switch without movement of the B chain C-terminal segment.

As far as we can ascertain from the structures of the insulin/IR Site 1-complexes (PDB entries 4OGA, 3W12 and 3W13, at resolution (3.5–3.9 Å)), the A-chain N-terminal helix of insulin (and insulin analogues) exhibits a Class 2-like structure when bound to IR (see Fig. 6). This is apparent in the larger-diameter A-chain N-terminal helix and shorter CαA6-CαA11 distance than those observed for Class 1 insulins, and in a similar azimuthal positioning of the constituent Cα atoms to those within Class 2 structures. The ability of the insulin A-chain N-terminal helix to adopt these two classes of structure indicates both (i) a degree of conformational flexibility in the three covalent bonds that connect that helix to the remainder of the hormone (viz., the A6-A11 and A7-B7 disulfide bonds and the downstream peptide bond), and (ii) rotameric plasticity of the side chains of the residues that form the interface between that helix and the remainder of the hormone (Fig. 5c,f).
that structures with Thr\textsuperscript{A8}-Glu\textsuperscript{A4} and Thr\textsuperscript{A8}-Gln\textsuperscript{A5} backbone hydrogen bonds are not distinct conformations and transitions between them are part of a natural “breathing” mode of the helix, possibly explaining why Thr\textsuperscript{A8}-Gln\textsuperscript{A5} backbone hydrogen bonds have not been found in extant crystal structures. The average C\textalpha{\textsuperscript{A6}-to-C\textalpha{\textsuperscript{A11}} distance for Class 1 insulin and for cis isomers were effectively the same within the MD trajectories and the X-ray structures (Table S5). Curiously, the average C\textalpha{\textsuperscript{A6}-to-C\textalpha{\textsuperscript{A11}} distances for Class 2 structures in the MD simulations (4.05 Å and 3.97 Å for the cis isomer and insulin, respectively) were found to be shorter than seen in the crystal structures (4.50 Å and 4.55 Å for the cis isomer and insulin, respectively), suggesting that the distance is possibly distorted from its optimal value in the crystalline phase, or that there are differences in the respective stereochemical libraries that underpin MD and crystallographic refinement methodologies.

It is important to note that the variations in the RMSDs through the time course of the simulations (Fig. 5g–i) are larger for the cis isomer ($\sigma = 1.3$ Å) than for either the trans isomer ($\sigma = 1.0$ Å) or insulin itself ($\sigma = 0.9$ Å).
This indicates that the N-terminus of the cis isomer is more conformationally flexible than that of the other analogues, suggesting an explanation for the disorder of this helix in the X-ray crystallographic maps. Although the cis isomer is seen to access the Class 2 conformation in MD simulations and the trans isomer is not, their average RMSDs from the Class 2 conformation across all simulation frames are similar (3.4 and 3.5 Å, respectively) and are higher than for insulin (3.0 Å). These increased RMSDs for the dicarba insulin analogues correlate with statistically significant decreases in the helicity of the A-chain isomer, such reduced helical content in solution is consistent with the disorder of the cis isomer. In the case of the trans isomer, CD measurement indicates that the N-terminal helix observed in its crystal structure. In the case of the cis isomer, both molecules within the crystallographic asymmetric unit of the trans isomer show an ordered A chain N-terminal helix. Unlike in the structure of the cis isomer, the respective A-chain N-terminal helices of the two trans isomers are involved in crystal contacts, suggesting their crystallographic order may be brought about by these contacts: in solution, the trans isomer is not, their average RMSDs from the Class 2 conformation across all simulation frames are similar (3.4 and 3.5 Å, respectively) and are higher than for insulin (3.0 Å). These increased RMSDs for the dicarba insulin analogues correlate with statistically significant decreases in the helicity of the A-chain isomer, such reduced helical content in solution is consistent with the disorder of the cis isomer. In the case of the trans isomer, CD measurement indicates that the N-terminal helix observed in its crystal structure. In the case of the cis isomer, both molecules within the crystallographic asymmetric unit of the trans isomer show an ordered A chain N-terminal helix. Unlike in the structure of the cis isomer, the respective A-chain N-terminal helices of the two trans isomers are involved in crystal contacts, suggesting their crystallographic order may be brought about by these contacts: in solution, the trans isomer A chain is likely disordered. In support, our preliminary atomic force microscopy analysis (to be published elsewhere) shows an increased rate of fibrillation of both dicarba isomers compared to that of insulin. A key feature of fibril formation is proposed to be loss of helical structure as the A-chain N-terminal helix transitions to a β sheet structure. Consistent with our CD and MD data, earlier studies show that the A-chain N-terminal helix is susceptible to perturbation through removal of the A6-A11 disulfide bond or residue substitution elsewhere within the helix, and that it can undergo conformational fluctuation. An explanation for the cis isomer being active and the trans isomer being inactive arises from our analysis of the β-sheet insulin structures. The existence of the two Classes (1 and 2) of T-state insulin structures suggests that the hormone is capable of transition between the two forms. As far as can be ascertained at the resolution of the IR-site-1-complexed structures of insulin, the IR-bound conformation of the hormone corresponds to Class...
2. The crystal structure of the trans isomer indicates that it has a Class 1 conformation, and its inactivity suggests that it is incapable of transitioning to a Class 2 structure. This hypothesis is supported by the MD simulations, which show that, although the trans isomer is capable of transiently adopting a range of structures, the Class 2 conformation appears to be precluded. An overlay of trans isomer and Class 1 insulin X-ray structures onto those of Class 2 and receptor-bound (PDB entry 4OGA) insulins shows a distinction between the two sets of structures (Fig. 6a), with the N-terminal end of the helix axis adopting different rotational configurations with respect to the hormone core (highlighted in Supplementary Information Movie 1). The overlay also indicates that there would be a significant steric clash between side chain of Glu^{44} of the Class 1/trans isomer structures and the side chains of Asn^{711} within the receptor αCT helix (Fig. 6b).

Hence, introduction of the cis and trans dicarba linkages is seen to affect the structural dynamics of the A chain N-terminal helix in three ways: (i) increased short-range motion: both cis and trans linkages disrupt the residue backbone conformations (i.e., helicity), as evidenced in both the CD spectroscopy and the MD simulations—this possibly due to the increased rigidity of the dicarba linkages transmitting more vibrational energy from the rest of the molecule into the helix in comparison with the less rigid disulfide bond of insulin; (ii) increased long-range motion in the cis isomer, evidenced by the increased range of RMSDs (Fig. 5g) and the corresponding increased standard deviation seen in the MD simulations, as well as in the disorder apparent in the cis isomer X-ray structure; and (iii) decreased conformational flexibility in the trans isomer: the longer trans-dicarba bond (Table S5) changes the conformation of the A6-A11 loop, preventing the formation of a Thr^{A8}-Val^{A3} hydrogen bond and hence the adoption of a Class 2 conformation. The inability of the trans isomer to adopt a Class 2 conformation (as seen within the MD simulations) then precludes its engagement with IR site 1 and results in its observed biological inactivity.

The activity of the cis isomer demonstrates that reduction-oxidation of the native A6-A11 disulfide bond (if it occurs) does not play an obligatory role in IR binding and activation. Rather, we suggest that the A6-A11 disulfide bond is able to modulate the insulin conformation through allostery, key elements of which are revealed by our study. The ability of the A6-A11 cystine bridge to adopt a range of disulfide conformations (Table S3) allows switching of the N-terminal region of the insulin A chain between active (Class 2) and inactive (Class 1) conformations. The active conformation requires a short A6-A11 Cα-Cα distance and a concomitant lengthening of the distance between the Cα atoms of A7 and B7 (Table S5). The short A6-A11 Cα-Cα distance of the Class 2 structures appears to result in a “pulling” of the N-terminal end of A1-A8 helix away from the volume that would be occupied by IR αCT in a putative receptor complex (a conformation also adopted by mutation of B26 to NMeAla or NMeTyr^{25}). In contrast, the longer A6-A11 linkage seen in Class 1 structures results in the base of the A chain N-terminal helix being closer to the B chain as a direct consequence of the A7-B7 disulfide bond length (Table S5). Critically, the short A6-A11 Cα-Cα distance seen in the active Class 2 conformations appears to be necessary to allow a precise rotation of the A-chain N-terminal helix, which in turn positions the side chains of residues A2 to A4 in an orientation compatible with receptor binding. In insulin, it is the adoption of a particular A6-A11 disulfide configuration that gives the required Cα-Cα distance and hence access to the active conformation. While both cis and trans A6-A11 dicarba bonds promote flexibility in the A1-A8 helix, it is only the cis dicarba linkage that can allow the A6 and A11 Cα atoms to come into the close proximity required to adopt a Class 2 structure. The increased interchain length resulting from the presence of a trans carbon-carbon double bond means that the active conformation is not accessible to the trans isomer.

Conclusions and Outlook

Through detailed analysis of the structure and function of the two dicarba insulin isomers generated through novel chemistry, our investigation provides the first description of the key role that the evolutionarily-conserved A6-A11 cystine bridge plays as an allosteric switch of insulin activity. We reveal that both the configuration and flexibility of the A6-A11 disulfide linkage are essential elements in insulin's adoption of an active conformation. Additionally, we conclude that the A6-A11 dicarba linkage of the cis isomer can also adopt a configuration that enables insulin receptor binding. This leads to effective lowering of blood glucose levels in mice and a significantly reduced ability to promote mitogenic signaling, a highly desirable biological property. Thus, the cis isomer represents an analogue with promising clinical potential. In conclusion, we have revealed a novel mechanism underlying insulin receptor engagement that dictates downstream biological outcomes. Using these findings as a basis we can now design improved insulin analogues for the treatment of diabetes.

Methods

Materials. Actrapid Insulin was purchased from Lyppard Australia Pty Ltd. Hybridoma cells expressing antibodies specific for the IR α subunit (83-7) and the IGF-1R α subunit (24-31) were a gift from Prof. K Siddle^{41–43}. [3H]-Thymidine was purchased from PerkinElmer Life Sciences. hIR-A and hIR-B overexpressing R fibroblast cells (derived from IGF-1R knockout mouse embryonic fibroblasts, a gift from Prof. R. Baserga (Philadelphia, USA)^44), were produced according to Denley et al.^{45} hIR-A over-expressing L6 myoblasts were provided by Dr. B.F. Hansen (Novo Nordisk A/S, Denmark). P6 cells (BALB/c3T3 cells overexpressing the human IGF-1R) were from Prof. R. Baserga^{45}.

Synthesis of dicarba insulins. An interrupted solid-phase peptide synthesis (SPPS)-catalysis approach^{44} was developed to overcome deleterious aggregation and achieve quantitative ring-closing metathesis of the dicarba insulin A chain. Construction of the complementary insulin B chain was achieved through microwave-accelerated SPPS. The monocyclic A-B conjugates were prepared by combination of the dicarba insulin A chains with the insulin B chain under basic conditions resulting in spontaneous oxidation of the liberated free thiol groups to give the two target isomeric trans and cis dicarba peptides. The details of this method are provided in the Supplemental Methods.
Receptor competition binding assays and kinase receptor activation assays (KIRA). Human IR-A, IR-B and IGF-1R binding affinities and receptor phosphorylation were measured as described by Denley et al.\textsuperscript{26} Assays were performed in triplicate in at least three independent experiments.

DNA synthesis assay. DNA synthesis was carried out as described in Gaugin et al.\textsuperscript{46} Briefly, L6 rat skeletal myoblasts overexpressing human IR-A, were plated in a 96-well flat bottom plate (1.5 × 10^4 cells/well) and grown overnight at 37°C, 5% CO_2. Cells were starved in serum-free medium for 4 h before treatment with increasing ligand concentrations for 18 h in Dulbecco’s minimal essential medium with 1% bovine serum albumin. The cells were incubated with 0.13 μCi/well [^3]H]-thymidine for 4 h, shaken for 2 h with 50 μL disrupting buffer (40 mM Tris pH 7.5/10 mM EDTA/150 mM NaCl) and then harvested onto glass fibre filters (Millipore\textsuperscript{®}) using a MICRO 96\textsuperscript{TM} Skatron harvester (Molecular Devices). The filters were counted in a Wallac MicroBeta counter (PerkinElmer Life Sciences). Assays were performed in triplicate in at least three independent experiments.

Glucose uptake assay. Briefly, NIH3T3-L1 myoblasts (up to passage 20) grown in DMEM supplemented with 10% newborn calf serum, 2 mM L-glutamine, 100 U/mL penicillin, 100 μg/mL streptomycin at 37°C were seeded into 24-well plates at 5 × 10^4 cells/well and grown for 8 days to confluence and were then differentiated into adipocytes as described\textsuperscript{57}. Glucose uptake in response to insulin and the cis isomer was measured essentially as described\textsuperscript{48}. Briefly, 3T3-L1 adipocytes were serum starved in serum free DMEM/1% BSA for 4 h, washed twice with Krebs-Ringer phosphate buffer (KRP, 12.5 mM HEPES, 120 mM NaCl, 6 mM KCl, 1.2 mM MgSO_4, 1 mM CaCl_2, 0.4 mM Na_2HPO_4, 0.6 mM Na_2HPO_4 (pH 7.4)) containing 1% BSA and incubated for 15 min at 37°C. Insulin or the cis isomer was added at decreasing concentrations (100-0.3 nM) for 30 min at 37°C. For the final 10 min, 2-deoxyglucose (DOG) uptake was initiated by the addition of 50 μM cold deoxyglucose and 1 μCi \[^3^H]\)-deoxyglucose per well. The assay was terminated by rapidly washing the cells three times with ice-cold KRP buffer. Cells were solubilized in 0.5 M NaOH/0.1% SDS and \[^3^H]\) content was determined by scintillation counting. Nonspecific 2-DOG uptake was determined in the presence of 50 μM cytochalasin B.

Insulin tolerance test. Eight-week-old C57BL6 male mice were fed either a standard rodent chow diet containing (wt/wt) 77% carbohydrate, 20% protein, and 3% fat from Ridley AgriProducts (Pakenham, Victoria, Australia) or a high fat diet (HFD) containing (wt/wt) 57% carbohydrate, 19% protein and 15% fat from Specialty Feeds (SP08-044, Glen Forrest, Western Australia, Australia) for 12 weeks. Mice (5 or 6 mice per group) were injected ip with 0.75 U/kg insulin or cis isomer under non-fasting conditions and tail vein blood glucose was measured via glucometer at indicated times\textsuperscript{47}. Experimental procedures were carried out in accordance to the protocols approved by the Austin Health Animal Ethics Committee (AEC 2011/04936).

Circular Dichroism (CD). CD was carried out as previously described\textsuperscript{29,49}. Briefly, CD spectra were recorded on a Jasco J-815 CD spectrometer; spectra were from 260 to 190 nm with a 1.0 nm step size using a 1.0 s response time and 1.0 nm bandwidth in a quartz cuvette with a 0.1 cm path length. Insulin and dicarba insulin analogues were diluted in 10 mM phosphate buffer (pH 7.4) to a concentration of 20 μM using a 1.0 s response time and 1.0 nm bandwidth in a quartz cuvette with a 0.1 cm path length. Insulin and dicarba insulin analogues were diluted in 10 mM phosphate buffer (pH 7.4) to a concentration of 20 μM using a 1.0 s response time and 1.0 nm bandwidth in a quartz cuvette with a 0.1 cm path length. Spectra were background-corrected by subtraction of the spectrum of buffer alone. Temperature denaturation was achieved by automated thermal control increasing by 2°C/min at 1° intervals. Samples were diluted to 10 μM for equilibrium denaturation studies in guanidine hydrochloride (1.0–8.0 M).

Bioinformatics. A dataset of insulin structures was constructed from all medium- and high-resolution X-ray crystal structures (better than 2.8 Å) in PDB Archive Version 4.0 (Jul 2011). 275 unique insulin structures were found in 105 PDB files; these included human, bovine and porcine insulins and synthetic mutants. 152 structures were found to exhibit the T (or R) state (data not shown) and 123 the T state. (Note: structures with A6-A11 S->S bond lengths outside the range of 1.95 to 2.05 Å (indicative of poor modelling of the disulfide), as well as structures involving insulin in complex with the insulin degrading enzyme, were not included in the dataset.) Analyses were performed only on T-state insulins (i.e., T-state insulins from monomeric, T_3, T_6, T_R, or T_R 3 structures). Of the T-state structures, 30 exhibited a Class I conformation and 93 a Class 2 conformation. Custom programs were used to extract the C_GA6->C_GA17 and C_GA7->C_GA17 distances and A6-A11 disulfide conformations, as well as the hydrogen bonding pairs, from each structure (see Supplementary Information). The approach used to determine the relative azimuthal positioning of residues within the segment A1-A10 of each insulin is also described in the Supplementary Information.

Molecular Dynamics (MD). Molecular dynamics simulations were performed using the AMBER14 program package\textsuperscript{50}. Amber ff14SB force field parameters\textsuperscript{51} were used for all standard amino acid residues. RESP charges\textsuperscript{52} and force-field parameters for the dicarba linkages were determined using the PyRED program\textsuperscript{53,54} (see Supplementary Information). Initial atomic coordinates were taken from three high-resolution T-state insulin PDB entries: 1G7B, 1MSO A,B, and 3I3Z A,B. Each of these structures has a different conformation for the A6-A11 disulfide bond. cis and trans isomers were created from each entry, giving a total of nine different starting structures. After minimization and equilibration (see Supplementary Information), the system was heated to 400 K and the dynamics simulated for 10 ns. Each high-temperature simulation was analysed to identify distinct conformations of the insulin analogue. Four structures (or 8 in the case of the cis isomer, see Supplementary Information) were chosen from each run to use as starting geometries for subsequent 200 ns room-temperature MD simulations. Full details of simulation protocols are provided in the Supplementary Information. Processing of simulations and cluster analysis was performed using CPPTRAJ\textsuperscript{55}. The same custom programs as described above under Bioinformatics were used to assess the backbone hydrogen-bonding patterns, interatomic distances and A6-A11 disulfide conformations in each MD simulation frame. DSSP\textsuperscript{56} was also used to identify residue backbone conformations.
Cryostalization.  **trans isomer:** An initial sparse-matrix sitting-drop vapor-diffusion crystallization screen was conducted at the CSIRO Collaborative Crystallisation Centre (CSIRO C3, Parkville, Australia). Based on hit conditions determined, a ca 50 μm crystal was then grown at 20 °C in a 24-well Linbro plate by vapor diffusion from a hanging drop of 1 μL of 1 mg/ml protein dissolved in 10 mM HCl mixed with 2 μL crystallant solution (0.9 M potassium sodium tartrate, 0.1 M Tris HCl (pH 8.5), 0.5% PEG 5000 MME) on a siliconized coverslip (Hampton Research) placed over 0.4 mL crystallant. The crystal was placed in a second hanging drop vapor diffusion condition, a 2 μL drop of 1.0 M potassium sodium tartrate, 0.1 M Tris HCl pH 8.5 and equilibrated against 0.4 mL of 1.5 M potassium sodium tartrate overnight at 20 °C, after which the well solution was exchanged for saturated potassium sodium tartrate and incubated for 24 h. The crystal was cryo-cooled by direct plunging into liquid nitrogen (without addition of a cryo-stabilizing solution).  **cis isomer:** Following an initial sparse-matrix screen identical to that described above for the **trans** isomer, a ca 25 μm crystal was subsequently grown at CSIRO C3 in a 96-well additive screen using a SWISSCII plate in vapor diffusion format at 20 °C using a sitting drop of 150 nL of 1 mg/ml protein dissolved in 10 mM HCl mixed with 150 nL 0.8 M potassium sodium tartrate, 0.1 M Tris-HCl pH 8.5, 0.5% PEG 5000 MME, 4% acetonitrile. This crystal was briefly dipped in paraffin oil (Hampton Research) and cryo–cooled by direct plunging into liquid nitrogen.

**Diffraction data collection and processing.** Diffraction data for both crystals were collected at the MX2 beamline at Australian Synchrotron2 at ~100 K and at λ = 0.9537 Å. Diffraction data were processed and merged using XDS15, statistics are presented in Table 1.

**Crystallographic structure solution and refinement.** Molecular replacement solutions for both isomers were obtained using PHASER83, with the starting model in both cases being a porcine insulin monomer obtained from PDB entry 1B2486. Structure refinement was performed with PHENIX59 iterated with manual model building within COOT61. In the case of the **cis** isomer, the density remained exceptionally poor for the A chain N-terminal helix and for residues in vicinity of the dicarba bond. Attempts were made to refine the structure as an ensemble62, but without success. Final refinement statistics are presented in Table 1.

**Statistical Analyses.** Statistical analysis of receptor binding, receptor activation and DNA synthesis assays were performed using a 2-way ANOVA with a Dunnett’s multiple comparison. Insulin tolerance test and glucose uptake assay data were analysed with a paired *T*-test. Significance was accepted at P < 0.05.

**Data Availability Statement.** All data generated or analyzed during this study are included in this published article (and its Supplementary Information files). Protein Data Bank (PDB) coordinates for the crystal structure of **trans** isomer have been deposited with accession code PDB 5T7R. The model coordinates and associated crystallographic structure factors of the **cis** isomer are included as **cis** _isomer_.cif; the unreliability of the model structure as an ensemble62, but without success. Final refinement statistics are presented in Table 1.

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Author Contributions
B.V.L. and A.B. conducted chemical syntheses, S.C.O. performed in vitro assays, biophysical analyses with technical support by C.D. S.A. performed insulin tolerance test, N.L.H. conducted MD analyses, J.G.M. performed all crystallography experiments and analyses. N.L.H., M.C.L., A.J.R. and B.E.F. designed studies, analyzed results and wrote the paper.

Additional Information
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