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Interleukin-2 druggability is modulated by global conformational transitions controlled by a helical capping switch

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Interleukin-2 (IL-2) is a small α-helical cytokine that regulates immune cell homeostasis through its recruitment to a high-affinity heterotrimeric receptor complex (IL-2Rα/IL-2RβγC). IL-2 has been shown to have therapeutic efficacy for immune diseases by preferentially expanding distinct T cell compartments, and several regimens have been developed for combination therapies. The conformational plasticity of IL-2 plays an important role in its biological actions by modulating the strength of receptor and drug interactions. Through an NMR analysis of milliseconds-timescale dynamics of free mouse IL-2 (mIL-2), we identify a global transition to a sparse conformation which is controlled by a helical capping switch ("switch") at the loop between the A and B helices (AB loop). Binding to either an anti-mouse IL-2 monoclonal antibody (mAb) or a small molecule inhibitor near the loop induces a measurable response at the core of the structure, while locking the switch to a single conformation through a designed point mutation leads to a global quenching of core dynamics accompanied by a pronounced effect in mAb binding. By elucidating key details of the long-range allosteric communication between the receptor binding surfaces and the core of the IL-2 structure, our results offer a direct blueprint for designing precision therapeutics targeting a continuum of conformational states.

IL-2 | immunomodulation | protein dynamics | NMR | drug design

An extensive body of work has revealed that the conformational plasticity of IL-2 plays an important role in targeting the IL-2 receptor signaling axis in both agonist and antagonist mode; however, a more detailed understanding of such processes is needed for actionable value in drug development. Using mouse IL-2 as a model, our current body of work combines methyl-based chemical exchange NMR spectroscopy (CPMG and CEST) with thermodynamic measurements and immunoassays to characterize a global conformational transition to a sparsely populated autoinhibited conformational state, together with its functional significance. Targeting this minor state through mutations and small molecules can be exploited to fine tune the affinity and selectivity for different IL-2 binding partners, toward engineering novel cytokine functions.

Significance

An extensive body of work has revealed that the conformational plasticity of IL-2 plays an important role in targeting the IL-2 receptor signaling axis in both agonist and antagonist mode; however, a more detailed understanding of such processes is needed for actionable value in drug development. Using mouse IL-2 as a model, our current body of work combines methyl-based chemical exchange NMR spectroscopy (CPMG and CEST) with thermodynamic measurements and immunoassays to characterize a global conformational transition to a sparsely populated autoinhibited conformational state, together with its functional significance. Targeting this minor state through mutations and small molecules can be exploited to fine tune the affinity and selectivity for different IL-2 binding partners, toward engineering novel cytokine functions.

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Data deposition: Nuclear magnetic resonance assignments for the free wild-type (WT) mouse IL-2 (mIL-2), mIL-2 in complex with JES6-1 single-chain Fv (scFv) antibody, mIL-2 in complex with IL-2 receptor (IL-2R), and R52A mIL-2 mutant have been deposited in the Biological Magnetic Resonance Data Bank (BMRB), http://www.bmrb.wisc.edu, under accession numbers 27969, 27976, 27971, and 27974, respectively.

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IL-2Rα displaces the JES6-1 antibody and releases the cytokine to signal through the high-affinity heterotrimeric receptor on IL-2Rα^{high} T_{reg} cells (Fig. 1B). Of particular relevance is the mobility of the AB loop (the loop between α-helices A and B), which has been shown to undergo a large conformational change upon JES6-1 binding. The AB loop is the main binding site for JES6-1 and the IL-2Rα subunit, preventing signaling of IL-2Rα^{low} effecter cells (right). However, an exchange mechanism between JES6-1 and the IL-2Rα subunit allows a preferential release of IL-2 for exclusive signaling on IL-2Rα^{high} T_{reg} cells, biasing toward an immunosuppressive response (left). (C) Overlay of 1H,13C-HMQC spectra of selectively labeled mIL-2 at the I_{101} and I_{137} positions, recorded in the free (black) or as a stoichiometric complex with JES6-1 scFv (cyan) or IL-2 receptor (magenta), acquired at 800 MHz, 25 °C. The arrows highlight major chemical shift effects. Selected methyl assignments are shown. (D) Mapping of methyl chemical shift changes on the crystal structure of mIL-2/JES6-1 complex (PDB ID 4YQ0) and (E) on the overlaid IL-2Rα subunit from the homologous IL-2 quaternary complex structure (PDB ID 2B31). The mIL-2 residues with CSPs > 0.05 ppm are shown with cyan spheres in the schematic representation of JES6-1-bound mIL-2 (blue) and with magenta spheres in IL-2Rα-bound mIL-2 (green).

**Results**

**Long-Range Effects on mIL-2 Core Dynamics upon Recognition of Its Binding Partners.** Due to their hydrophobic character, methyl groups are well-suited NMR probes for monitoring the dynamics of the structural core of proteins (21, 22). The chemical shifts of methyl groups report on side chain rotameric states and local magnetic environment, while their NMR measurement is less influenced by the size of the system under investigation due to favorable relaxation properties (23). To gain insight into conformational
changes on mIL-2 induced by its binding partners, we prepared selective \(^1\)H\(^{13}\)C MILV (Met, Ile, Leu and Val)-methyl labeled samples on a uniform \(^1\)C, perdeuterated background. We obtained NMR assignments of mIL-2 backbone and methyl groups using a combination of transverse relaxation–optimization spectroscopy (TROSY)-based triple-resonance and three-dimensional (3D) out-and-back side chain transfer experiments. Under the conditions of our NMR experiments, mIL-2 is strictly monomeric, as characterized by size exclusion chromatography coupled to multiance laser light scattering (SEC-MALS) (SI Appendix, Fig. S1). To obtain stereospecific resonance assignments of methyl groups and to reduce spectral overlap observed in the free mIL-2 methyl spectrum, which is typical of α-helical proteins, we also prepared a selectively \(1\text{H}_6^{13}\text{C}_5\text{H}_4\), proS L\(^{1\text{I}}\)\(^{13}\)C\(^1\)H\(_2\) labeled protein sample (24). This approach allowed us to obtain unambiguous side chain resonance assignments of Met \(\delta_1\), Ile \(\delta_1\), Leu \(\delta_1\), and Val \(\gamma_1\) (60 methyl groups in total) (SI Appendix, Fig. S1) (25), distributed throughout the structure, enabling us to comprehensively map the cytokine’s surface and hydrophobic core in the IL-2Ra and JES6-1-bound states (26, 27). Fig. 1C shows \(^1\)C-\(^1\)H band-selective optimized flip angle short transient (SOFAST) heteronuclear multiple quantum coherence (HMQC) of (L\(^{1\text{V}}\)proS)-methyl labeled mIL-2 unbound (Fig. 1C, black) and as a stoichiometric complex with JES6-1 single-chain Fv (scFv) (Fig. 1C, cyan) or IL-2RaX (Fig. 1C, magenta). The weighted chemical exchange spectra (CSPs) are presented along the mIL-2 sequence in SI Appendix, Fig. S2. In agreement with previous surface plasmon resonance (SPR) measurements (10), under our NMR sample conditions, mIL-2 forms tight complexes with either JES6-1 or IL-2RaX. Exchange between the free and bound states is slow on the chemical shift timescale, as shown by detailed NMR titrations where we observe a unique set of peaks corresponding to the free and bound states with large CSPs (up to 0.6 parts per million [ppm]) between them.

Notably, either receptor or antibody binding leads to measurable conformational changes at the corresponding primary binding sites (for example, L\(486\), L\(50\), and L\(546\) in the AB loop) but also induces long-range effects at remote sites distributed throughout the structure, including L\(35\)α, L\(35\)δ, in the A helix, L\(75\)α, L\(80\)α, and L\(80\)δ in the B helix, I\(101\)δ, in the C helix, and V\(129\)γ2, V\(130\)β2, L\(336\)α, and L\(376\)δ in the D helix. Mapping the most affected methyl groups on the mIL-2 structure (Fig. 1D and E) highlights a contiguous path connecting the AB loop to the hydrophobic core of the structure. Overall, the methyl CSPs reveal a global response of the mIL-2 structure, indicative of allosteric communication between the primary binding site at the AB loop region and core residues of the cytokine (SI Appendix, Fig. S2). Given that the four α-helical bundle core of apo-IL-2 is maintained in the complexed forms with the IL-2R receptor or JES6-1 antibody (backbone heavy atom rmsd of 0.8 Å and 2.3 Å, respectively), our results suggest that binding of either JES6-1 or IL-2RaX at the AB loop region induces a remodeling of core side chains. Specifically, the observed methyl chemical shift changes can arise from either changes in the lowest energy rotamer state or through perturbations in ensembles of rotamers sampled by the free and bound forms.

**Free mIL-2 Samples a Global Transition to an Excited-State Conformation.** Based on our observation that the \(^1\)H–\(^{13}\)C transverse relaxation–optimization spectroscopy (TROSY) spectrum of free mIL-2 was of marginal quality (SI Appendix, Fig. S1) with a significant fraction of completely broadened amide resonances (33%), we hypothesized that, in solution, mIL-2 undergoes exchange between different conformational states. The loss of signal for amide resonances corresponding to residues in the AB loop due to \(^1\)N line broadening suggests that the loop is switching between multiple backbone conformations with different chemical environments on an intermediate (microseconds to milliseconds [μs–ms]) timescale. This hypothesis is consistent with a recent molecular dynamics (MD) simulation-based model in which free mIL-2 samples distinct conformations of the AB loop akin to the crystallographically observed structures in complexes with the JES6-1 antibody and IL-2Ra receptor (10, 15).

To characterize the conformational landscape sampled by free mIL-2 in solution, we performed a series of \(^1\)H chemical exchange saturation transfer (CEST) (28) and \(^1\)C single-quantum methyl Carr–Purcell–Meiboom–Gill (CPMG) relaxation dispersion experiments (29). Briefly, in CPMG experiments, the effective transverse relaxation rate (\(R_{2eff}\)) is measured as a function of refocusing pulse frequency (vCPMG), which quenches the effects of conformational exchange, producing so-called dispersion profiles. Such profiles can be fit to extract the rate of exchange between the ground and excited conformational states, \(k_{ex}\). the population of the excited state, \(p_e\), and the chemical shift differences between the two exchanging states \(\Delta \omega_0\) (ppm). In CEST, the intensities of protein resonances are measured in the presence of a weak off-resonance radiofrequency (\(B_1^*\)) field, typically between 5 and 50 Hz, where a series of two-dimensional (2D) experiments are acquired with a varying offset of the \(B_1^*\) field. When the field offset coincides with the resonance of an exchanging minor state, saturation transfer occurs during a fixed period, leading to an attenuation of the resonance of the otherwise “invisible,” minor state. While, in principle, both experiments can be used to quantify conformational exchange processes in biomolecules (30), CEST is sensitive to motions ranging from <50 to 500 s\(^{-1}\) whereas CPMG on the order of 100 to 3,000 s\(^{-1}\). As a result, CEST experiments are typically recorded at a reduced temperature (4 to 10 °C), in order to slow the exchange to a measurable regime.

Due to poor quality of the amide spectra of free mIL-2, we turned to methyl spectra as a readout in both CPMG and CEST experiments. Conformational exchange was observed for the resonances of several methyl groups distributed throughout the mIL-2 structure, as illustrated by representative residues on the B helix (L\(80\)α, C helix (I\(101\)β), and D helix (L\(133\)β) (Fig. 2). The resonances of 20 methyls exhibiting well-defined CPMG dispersion curves (SI Appendix, Fig. S3 and Table S1) were used in a quantitative analysis. The remaining mIL-2 resonances exhibited low signal-to-noise (S/N) dispersion curves (at 18.8 T in a quantitative analysis. The remaining mIL-2 resonances exhibited low signal-to-noise (S/N) dispersion curves (at 18.8 T magnetic field) or high spectral overlap. Dispersion curves were fitted globally using a two-site exchange model, yielding a \(k_{ex}\) of 1,000 ± 72 s\(^{-1}\) and an excited state population of 8.0 ± 0.4% (SI Appendix, Fig. S4). Additionally, \(^1\)C-CEST profiles for the same residues (L\(80\)δ, I\(101\)β, and L\(133\)β) revealed significant dips in the \(^1\)C dimension, characteristic of an excited state (Fig. 2B and SI Appendix, Fig. S5). To quantify the extent of structural adaptations upon formation of the excited state for each methyl site participating in the exchange process, we extracted values of the corresponding \(^1\)C chemical shift changes, \(\Delta \omega_0\), from independent fits of the CPMG relaxation dispersion and CEST datasets. We observed good correlation between the resulting \(\Delta \omega_0\) values for most methyl probes (Fig. 2C), suggesting that the two datasets are likely reporting on a similar exchange process, with the caveat that the CEST data were recorded at a lower temperature (10 °C relative to the standard temperature of 25 °C used for CPMG data acquisition). Thus, the two observed outliers in the data (L\(133\)β, L\(80\)α) could arise from the temperature dependence of the chemical shifts of the major and minor states.

Methyl groups exhibiting CPMG dispersion curves are plotted on the homology-based model of free mIL-2 in Fig. 2D and color-coded according to the magnitude of the fitted \(\Delta \omega_0\) values, which report on differences in the local magnetic environment between the major and minor conformations. Large \(\Delta \omega_0\) values were observed for methyls at the AB loop (L\(486\)δ, L\(546\)δ), in addition to the C terminus of the B helix facing toward the loop.
(L80δ2, V83γ1, L84δ1) and throughout the hydrophobic core of the structure (I101δ1, V130γ2, L133δ1, and I137δ1). Thus, our CPMG data suggest that free mIL-2 samples a global, cooperative transition to an excited state, which involves a conformational “switch” of the AB loop, coupled to a cooperative change affecting the core methyl groups (Fig. 2E). Notably, a pronounced structural change relative to the free form can be observed in the X-ray structure of JES6-1–bound mIL-2 (10), in which the AB loop (residues Y45 to T55) undergoes an ∼38° rigid-body rotation to adopt a conformation that is optimal for interactions with the antibody surface. Taken together, conformational plasticity of the AB loop is a key component of the solution dynamics of free mIL-2, relevant for the formation of a high-affinity immunomodulatory complex.

To characterize the major conformation sampled by the AB loop in solution, we analyzed methyl nuclear Overhauser effect (NOE) intensities recorded in a 3D C_{αα}C_{αα}H_{αα} SOFAST nuclear Overhauser effect spectroscopy (NOESY) experiment (Fig. 2F), relative to the corresponding distances observed in 1) a model of free mIL-2 built using the human IL-2 apo-structure as a template (Protein Data bank [PDB] ID code 1M47), 2) the cocrystal structure of mIL-2 in complex with the JES6-1 antibody (Ab)
A Conserved Hydrogen Bond Network Coupled to the Structural Core Stabilizes the Excited mIL-2 Conformation. We next sought to characterize the structural features of the excited-state conformation identified by our CPMG and CEST experiments. Excited states play important roles in protein function, including catalysis (31-33) and complex formation via conformational selection or induced fit-type mechanisms (34, 35). However, their de novo structure modeling is challenged by the ambiguity in interpreting methyl chemical shifts. A qualitative comparison of the placement of residues undergoing conformational exchange in free mIL-2 (Fig. 2D) with those perturbed upon formation of the IL-2Rα and JES6-1-bound states (Fig. 1D and E) showed a significant overlap, both for residues that are within the AB loop and other binding surfaces (L80, M53, and L86) and also at the core of the structure (L133 and L137), suggesting that the same sites that participate in the formation of the excited state also undergo structural changes upon formation of the mIL-2 tertiary complexes.

Our NOE-based analysis has shown that the backbone conformation of the AB loop in the major, “closed” state of mIL-2 is inconsistent with the JES6-1-bound structure. Close inspection of the structure reveals that, in the JES6-1 complex, the AB loop instead participates in a network of C-capping interactions with the A helix (Fig. 3A and D). Specifically, the side chain guanidino group of R52 forms multiple hydrogen bonds with the backbone carbonyl oxygens of M42, Y45 located at the C-terminal end of A helix, and with R46, L48 at the N-terminal part of the AB loop (Fig. 3B and C). Conversely, in the free structure of the homologous human IL-2 (PDB ID code 1M47, termed “uncapped” state), the R52 side chain is solvent-exposed, and M53 is buried at the hydrophobic interface between the A and B helices. The “closed/uncapped” conformation is optimal for interactions with coreceptors, as exemplified by the IL-2Ra complex where R52 forms a salt bridge with a conserved aspartic acid on the receptor surface (PDB ID code 2BS1). JES6-1 binding therefore induces a transition from a “closed” to “open” AB loop conformation, which exposes the hydrophobic side chains of L50, M53, and L54 toward the surface of the molecule while the side chain of R52 becomes buried. In this “capped” state, the loss of hydrophobic packing contacts is compensated by the formation of multiple C-capping hydrogen bonds with the A helix, and the backbone of the loop itself (Fig. 3A and C).

According to our NOE-based analysis, the placement of M53 at the hydrophobic interface between the AB loop and the core of the structure promotes the “closed” loop conformation, which is the major solution state. We sought to determine whether the excited state sampled by free mIL-2 encompasses the C-capping features of the “open” state. If the dynamics observed by NMR correspond to a concerted process, such as an open-to-closed transition of the AB loop, then mutations which destabilize the open state should alter the observed exchange parameters in a uniform manner. To perturb the “open” state conformation, we used the mutant R52A, which eliminates all C-capping interactions (Fig. 3C and SI Appendix, Fig. S6) (36), and compared the resulting CPMG relaxation dispersion and CEST data performed under identical conditions to our established results for the WT. Remarkably, Ala mutation of R52 suppressed the CPMG relaxation dispersion profiles of all methyl probes that were undergoing conformational exchange in the WT form (Fig. 3E and F and SI Appendix, Fig. S3B). Due to the significantly reduced dispersion range observed for methyl resonances of the R52A mIL-2 mutant, the χ2 surface obtained from a global fit of the CPMG data does not show a well-defined minimum in terms of the estimated population and exchange rate, kex. Consistently, in CEST profiles recorded using the R52A mutant, the second dip was barely observed (I1016v) or notably absent (L708, I1336, I1336, L1336) (Fig. 3E and F and SI Appendix, Fig. S5), indicating that the population of the excited state (“open” state) was below the detection limits of the experiment. Both observations are consistent with a significant “dampening” of µs-ms dynamics. The R52A mutation quenched relaxation not only at the side chain methyl but also the backbone amide groups, alleviating resonance broadening and leading to a significant improvement in the quality of the 1H-1H TROSY spectrum (SI Appendix, Fig. S6A). These observations imply the presence of a global conformational exchange process on a µs-ms timescale, involving a switching of the AB loop coupled to a transient twisting or “breathing” of the α-helices. Notably, the conserved C-capping sequence motif at the AB loop is critical for the formation of the excited state.

To determine how destabilization of the excited state affects mIL-2 binding to JES6-1, we used isothermal titration calorimetry (ITC) and measured the binding free energy (ΔG), and its enthalpic (ΔH) and entropic (–TΔS) components both for WT and R52A mIL-2 (Fig. 3G and H). Consistently with previous SPR measurements (10) and our observation of a tight complex in slow exchange by NMR, WT mIL-2 binds JES6-1 with a high affinity of 2 nM. The interaction is largely entropy-driven, with a positive net enthalpy change of 12.7 kcal·mol−1 and a favorable entropic contribution of −24.3 kcal·mol−1 (293 K, Fig. 3H and SI Appendix, Table S3). Notably, R52A binds to JES6-1 with three orders of magnitude lower affinity than WT mIL-2, due to less favorable binding entropy (Fig. 3G and H). Given that the side chain of R52 forms a salt bridge with E60 of JES6-1 in the X-ray structure of the complex (Fig. 3J), the observed 1,000-fold decrease in binding affinity reflects the combined effects of 1) loss of binding enthalpy and 2) increase in free energy of the “closed” state, which is competent for binding. Thus, our ITC results show that perturbations of the capping hydrogen-bond network coupled to global changes in core dynamics abrogate JES6-1 binding by destabilizing the open conformation, consistently with our NMR measurements showing a shift of the conformational equilibrium toward the closed state.

To determine the functional activity of the R52A mutant, we probed JES6-1 antibody effects on mIL-2-induced signaling in IL-2Ra+ subpopulations of mouse CTLL-2 cells as a surrogate for its effects on IL-2Ra+ immune cell subsets (Fig. 3H). Consistent with our NMR and ITC results, JES6-1 showed a fivefold reduced WT mIL-2-mediated signal transducers and activators of transcription 5 (STAT5) activation. However, STAT5 signaling potency by R52A was independent of JES6-1, as demonstrated by the comparable effective concentration, 50% (EC50) values, suggesting that the weaker affinity to JES6-1 was primarily responsible for the behavior of the R52A mutant (Fig. 3G).
**Figure 3.** Effects of a conserved helical capping motif on IL-2 dynamics and function. (A) Crystal structure of mIL-2 (schematic) bound to JES6-1 (blue surface) with a detailed view of the mIL-2/JES6-1 interface on the AB loop and B helix of the cytokine (PDB ID 4YQX). C-capping hydrogen bonds involving the R52 side chain are shown as black lines. (B) The AB loop in mIL-2 adopts either an uncapped “closed” conformation (apo state) or a capped “open” conformation (JES6-1-bound state). The open conformation is stabilized by an intramolecular C-capping interaction between the side chain of R52 and the backbone of M42-Y45-R46. Black dashes indicate hydrogen bonds. The R52 side chain does not contribute to intramolecular stabilizing interactions in the closed conformation. Black dashed arrow shows the direction of loop movement between the closed and open states. (C) Close-up view of the AB loop in the closed and open conformations. The R52A mutation eliminates the C-capping interactions. (D) Sequence conservation pattern in the AB loop region in various IL-2 orthologs. (E) Comparison of $^{13}$C-CPMG relaxation dispersion profiles for L806, I1016, and L1336 of WT mIL-2 (purple) and R52A (green), at 600 MHz and (F) $^{13}$C-CEST profiles for WT mIL-2 (red) and R52A (green). Conformational exchange profiles throughout the mIL-2 structure are quenched by the R52A mutation. Experimental errors in CPMG and CEST data are determined as described in Fig. 1. (G) Thermodynamic fingerprints of the interactions between WT and R52A mIL-2 with JES6-1. Isothermal titration calorimetry thermographs and curve fits for titrations. The R52A mutation leads to a three-order-of-magnitude reduction in binding affinity. To minimize enthalpy of solvation effects, all experiments were performed in 20 mM phosphate buffer, pH 7.2, 150 mM NaCl. Data shown are representative of duplicate experiments. (H) STAT5 phosphorylation ($p$STAT5) response to WT and R52A mIL-2 in the free state or JES6-1 immunocomplexes treatment in IL-2Rα+ mouse CTLL-2 cells. R52A shows attenuated immunomodulation by the JES6-1 Ab while signaling via IL-2Rα+ is at similar levels to WT. μcal, median fluorescence intensity.

**Allosteric Communication in mIL-2 through a Remodeling of Side Chain Rotamers.** Our methyl-based NMR probes revealed a global response of the mIL-2 structure upon binding to JES6-1, alongside the crystallographically observed conformational changes of the AB loop (Fig. 1 C and D). Consistently, our NMR relaxation measurements suggest a coupling of dynamic motions between the loop and core methyls. We sought to identify a plausible mechanism linking AB loop movement to changes in core side chain packing during transitions between “closed/uncapped” and “open/capped” conformations. To enumerate all possible side chain rotamers that can be adopted by each residue, we performed a global analysis of compatible rotamer pairs using a satisfiability-based approach in Rosetta (Materials and Methods and SI Appendix, Fig. S8) and mapped our results on the free and JES6-1-bound mIL-2 structures (Fig. 4 and SI Appendix, Fig. S9). Using the backbone conformations of the “open” and “closed” states as inputs, our analysis highlights differences in rotamer sets that can be accessed by the free (closed) and JES6-1-bound (open) states. We identified a large set of residues (L28, L39, M42, L48, M53, F58, F132, L133, W136, and F139) (Fig. 4 A and B) spanning the AB loop, A Helix and D Helix, and part of the hydrophobic core. For these residues, the space of rotamers was significantly different between the closed and open states, indicating a plausible remodeling of packing interactions. Specifically, a 10-residue segment (V129 to
F139) forming the hydrophobic face of the amphipathic D helix exhibits expansions and contractions in allowed rotamer sets as IL-2 transitions between the two states (SI Appendix, Fig. S8). A concerted motion of the D helix upon binding of IL-2Rα has been highlighted as having a functional role in the formation of the high-affinity heterotrimeric signaling complex (37). These results are consistent with our observed NMR chemical shift mapping of the IL-2Rα and Ab-bound states (Fig. 1 C and D), suggesting that the conformational transition between the free and complexed states can lead to a redistribution of the rotameric states at the hydrophobic core. A plausible allosteric communication network starts at the AB loop on one end of the structure, traverses through the inner core of mIL-2, and ends at the N terminus of the A and D helices adjacent to the binding site of the IL-2Rα receptor (Fig. 4C). The hydrophobic core residues exhibiting differences in rotamer sets also include sites with significant chemical exchange contributions in our CPMG data, indicating the presence of dynamics at the μ–ms timescale (Fig. 4C, orange circles). Taken together, our results highlight a plausible allosteric communication network in the IL-2 structure mediated via sequential remodeling of side chain packing interactions.

Skewing the Dynamic Landscape of mIL-2 by Ligand Binding. The R52A mutation characterized here destabilizes the open mIL-2 conformation by perturbing the C-capping hydrophobic pocket network between the AB loop and B helix, leading to quenching of conformational exchange throughout the core of the structure. We hypothesized that a small molecule binding preferentially to the closed AB loop conformation would impact the dynamic landscape of mIL-2 in a similar manner. We used a known compound targeting human IL-2 (hIL-2) (Ro 26-4550), to compete with IL-2Rα binding (7, 38). The coocrystal structure (PDB ID code 1M48) shows that Ro 26-4550 is nestled in a hydrophobic pocket at the interface between the AB loop and B helix, with the hIL-2 side chains of M39, V69, and L72 (mIL-2 residues M53, V83, and L86) packing against the terminal aromatic ring (7). We performed an NMR titration of increasing molar ratios of Ro 26-4550 on isoleucine, leucine, and valine-methyl-labeled WT mIL-2 using 1H-13C HMQC spectra as a readout (Fig. 5A). Under our NMR conditions, we observed the formation of a moderate-affinity (μM range dissociation constant [Kd]) complex in fast exchange with the free mIL-2 form where residues in close proximity to the AB loop (L4861, L5461, and L6081) experience the largest effects (SI Appendix, Fig. S7). However, CSPs extend beyond the AB loop, to sites in the hydrophobic core (L8061, V129γ2, V130γ2, L133β2, and I137ε1) at distances >12 Å from the inhibitor binding site in the coocrystal structure (Fig. 5C). A quantitative NMR line shape analysis of the resonances of V129γ2 and L133β2 yielded an equilibrium dissociation constant of 39.4 ± 5.5 μM (Fig. 5B), suggesting a two-state cooperative transition impacting all sites in the ligand-bound form. To further examine whether Ro 26-4550 binding affects the μ–ms dynamics of mIL-2, we repeated 13C-CPMG relaxation dispersion experiments, under saturation binding, identical protein concentration and NMR sample conditions, and compared with our established data recorded for the free form (Fig. 5D). Our results indicate that Ro 26-4550 binding strongly dampens relaxation of methyl groups that are undergoing μ–ms conformational exchange throughout the mIL-2 structure. Thus, in agreement with our allosteric network model of mIL-2, stabilization of the closed/uncapped conformation through ligand binding reduces dynamic exchange at the hydrophobic core of mIL-2 as the conformational equilibrium is skewed toward the closed state. The observation of residual exchange at the CPMG timescale suggests that sampling of the open state is still permitted by the presence of the ligand.

Discussion
The unique conformational plasticity of IL-2 appears to play an important role in targeting the IL-2 receptor signaling axis in both agonist and antagonist mode. However, this plasticity also presents an opportunity to target or otherwise manipulate the conformational landscape of IL-2 for drug discovery, but more detailed insights are needed for actionable value. Evidence for a functional role of such conformational plasticity was provided in previous studies focusing on both human and mouse IL-2. These studies employed small molecule binding (6, 7), antibody engineering (9, 12), or directed mutagenesis (8) to induce minor perturbations in the cytokine’s core α-helical fold, leading to distinct immunomodulatory functions. Specifically, elucidating a link between
specific allosteric effects on the IL-2 structure, induced by the JES6-1 and S4B6 antibodies, with the targeting of opposing immune cells types expressing different levels of receptor subunits in vivo has provided a solid foundation for modulation through the stabilization of different IL-2 conformational states. Conversely, the high-throughput screening of hIL-2 libraries in yeast led to the identification of a variant, termed “super-2,” comprising a helix that is locked in a conformation which is primed for interactions with the IL-2R subunit, thereby leading to preferential expansion of cells lacking IL-2Rα (8). These results were leveraged to discover a human T456-based Ab, akin to JES6-1, of significant therapeutic potential (11). Thus, given the sequence and structure conservation of the IL-2 fold across species, the results established for the mouse IL-2 paradigm directly relate to human IL-2, whose use as a multifaceted protein therapeutic (Proleukin) has been hampered by an incomplete understanding of its precise functional properties. Therefore, a detailed characterization of the relevant conformational states and their dynamic interconversion process in solution would provide a road map for drugging IL-2, through a range of approaches.

Our methyl-based NMR measurements from two complementary methodologies, CPMG and CEST, provide strong support that free mIL-2 is highly dynamic at the μs-ms timescale and samples an excited state via a concerted conformational transition of the AB loop and core of the structure. Functionally relevant excited states have been previously described for a range of protein and nucleic acid systems (30) and are typically separated by relatively low energy barriers from the ground state. The inherent plasticity of the AB loop and allosteric cross-talk with the hydrophobic core suggested an avenue for biasing the conformational equilibrium by perturbing key features present in the excited state conformation. Based on our analysis of crystallographically observed snapshots of IL-2, we hypothesized that the excited state might share similar features with the JES6-1 antibody-bound structure, with respect to a stabilizing C-capping motif, and tested this hypothesis through a detailed dynamic, binding, and functional characterization of a designed Ala mutant. In fact, our data demonstrate that the structural basis for such functional plasticity of the AB loop and allosteric cross-talk with the hydrophobic core engaged in a cooperative motion on the preequilibrium between closed (major form, 92%) and open (minor form, 8%), with the AB loop and the hydrophobic core engaged in a cooperative motion on the preequilibrium between closed (major form, 92%) and open (minor form, 8%).

Fig. 5. Small molecule binding at the AB loop quenches dynamics of mIL-2. (A) The 2D 1H,13C-HMQC spectra of ILV-methyl-labeled mIL-2 in the free state (red), and with increasing concentrations of Ro 26-4550 inhibitor (1:4 molar ratio, shown in black). Data were recorded at 800 MHz, 25 °C. Fast-exchange chemical shift changes are highlighted with arrows for select methyl resonances. (B) NMR line shape analysis for the V129y2 and L133s2 methyl resonances using TITAN (Materials and Methods), with indicated equilibrium dissociation constant and errors propagated from the spectral S/N. Recorded NMR spectra are shown in blue, with simulated line shapes in red. Ratios of mIL-2 to inhibitor are indicated in each panel. (C) Two views showing the mapping of residues undergoing significant chemical shift perturbations (CSPs) onto the ribbon representation of mIL-2. Methyl groups with marked CSPs (>0.05 ppm) are shown as black spheres. The inhibitor is shown as a blue ball-and-stick diagram on the overlaid hIL-2 complex structure (PDB ID code 1M48). (D) The 13C single quantum CPMG relaxation dispersion profiles in the absence (purple) and presence of Ro 26-4550 at saturating concentration (black) for selected residues in the AB loop (L54s1), B Helix (L80s1), C Helix (I101s1), and D Helix (V130y2, L133s1, I137s1). CPMG experiments were performed at a 1H field of 600 MHz and 25 °C.
timescale. In the closed/uncapped (productive) state, the AB loop is in a conformation that is primed to interact with the IL-2Rα receptor. Formation of a C-capping hydrogen network locks the AB loop in an open (autoinhibited) state, in which the loop adopts a conformation that is optimal for interactions with the JES6-1 antibody (Fig. 6, left). Disruption of the C-capping hydrogen bond network by mutation or small molecule binding shifts the equilibrium toward the closed state and induces global changes in core dynamics which ultimately lead to a reduction in Ab binding by three orders of magnitude (Fig. 6, right). While the “closed” and “open” conformations mediate high-affinity complex formation with the receptor and Ab, respectively, the binding process is likely achieved via the formation of an initial encounter complex where additional induced structural adaptations follow the selection of preexisting conformational states. This model is consistent with the majority of protein–ligand binding examples where both processes contribute to complex formation (34).

A consequence of allosteric cross-talk in the IL-2 fold is that the protein becomes globally sensitive to mutations. Point mutations can induce long-range effects, which may explain the high level of sequence conservation of IL-2 residues removed from the primary receptor recognition sites. Further examples of such long-range modulations are provided by engineered IL-2 agonists. IL-2 superkine (also denoted as “super-2”) functions independently of IL-2Rα and has enhanced binding to IL-2Rβγ, despite five of the six mutations being clustered on the BC loop and within the C helix core, removed from the IL-2Rβγ primary binding site. Instead, a cluster of hydrophobic substitutions (F80, V85, and V86) induce a subtle change in C helix orientation, leading to an increased affinity for IL-2Rβγ. Super-2 exerts a more potent expansion of tumor-specific cytotoxic T cell populations, and a reduced expansion of Treg cells (8). Moreover, Super-2 can be used as a platform for additional mutations that inhibit binding to the γc receptor and, consequently, heterodimerization of IL-2Rβγ and γc, thereby acting as a potent antagonist (13). Conversely, to bypass allosteric effects present in the IL-2 fold, a de novo design approach was used to generate a mimetic of IL-2/IL-15 showing an α-helical fold with a wellpacked hydrophobic core, denoted Neoleukin-2/15 (Neo-2/15) (16). This synthetic molecule retains the binding surfaces for the IL-2Rβγ/γc receptors while also alleviating the requirement of IL-2Rα/IL-15Rα for formation of a productive signaling complex. Neo-2/15 exhibited enhanced antitumor activity in mouse models, was more stable, and lacked detectable immunogenicity (16). These studies highlight the potential for modulating IL-2 function through different design approaches, showing varying levels of responsiveness to regulatory molecules and conditions at the vicinity of the membrane signaling complex.

In summary, our results highlight the role of conformational plasticity at conserved features of the IL-2 fold, acting locally and allosterically to form an “excited” state with discrete functional properties. Our NMR-based characterization offers a window to access otherwise hidden states of the energy landscape that can be then exploited to fine-tune the affinity and selectivity for different IL-2 binding partners toward engineering novel cytokine functions. For example, our established allosteric coupling between the receptor binding sites to the hydrophobic core of the structure presents an opportunity to alter IL-2 function through
the introduction of core mutations and induction of allosteric effects without directly affecting the receptor/Ab binding surfaces. Indeed, the allosteric pathway elucidated in detail here using mouse IL-2 as a model system has been recently exploited for human IL-2 to develop Tα-biased antibodies that appear to work by a similar mechanism (11). Given that the IL-2 fold presents a limited surface area with overlapping binding epitopes, our findings have important practical utility from an engineering and synthetic biology perspective.

Materials and Methods

Specific details about mIL-2 NMR sample preparation and stereospecific isotopic labeling, backbone and methyl resonance assignments, relaxation dispersion and 13C CEST experiments, SEC MALS and ITC experiments, side chain rotamer analysis, and STAT5 signaling are outlined in detail in SI Appendix.

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