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Introduction

A structural/functional overview

The mechanistic target of rapamycin (mTOR) plays a crucial role in cell growth and homeostasis.¹⁻³ It belongs to the PI3K-related protein kinase (PIKK) family which shares a complex domain organization.^{4,5} mTOR consists primarily of four parts: the Nterminal HEAT (N-HEAT) (Huntingtin, EF3A, ATM, TOR) and middle HEAT (M-HEAT) repeats, the FAT domain (coined after FRAP, ATM, and TRRAP), and the kinase domain (Fig. S1a[†]).⁶⁻⁹ Together with specific proteins, mTOR assembles into two functional complexes (mTORCs), mTORC1 and mTORC2 (Fig. S1b[†]).¹⁰⁻¹⁶ The complexes share an accessory protein, mammalian lethal with SEC13 protein 8 (mLST8), but differ in other components.¹⁷⁻¹⁹ Regulatory-associated protein of mTOR

The allosteric mechanism of mTOR activation can inform bitopic inhibitor optimization[†]

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mTOR serine/threonine kinase is a cornerstone in the PI3K/AKT/mTOR pathway. Yet, the detailed mechanism of activation of its catalytic core is still unresolved, likely due to mTOR complexes' complexity. Its dysregulation was implicated in cancer and neurodevelopmental disorders. Using extensive molecular dynamics (MD) simulations and compiled published experimental data, we determine exactly how mTOR's inherent motifs can control the conformational changes in the kinase domain, thus kinase activity. We also chronicle the critical regulation by the unstructured negative regulator domain (NRD). When positioned inside the catalytic cleft (NRD IN state), mTOR tends to adopt a deep and closed catalytic cleft. This is primarily due to the direct interaction with the FKBP-rapamycin binding (FRB) domain which restricts it, preventing substrate access. Conversely, when outside the catalytic cleft (NRD OUT state), mTOR favors an open conformation, exposing the substrate-binding site on the FRB domain. We further show how an oncogenic mutation (L2427R) promotes shifting the mTOR ensemble toward the catalysis-favored state. Collectively, we extend mTOR's "active-site restriction" mechanism and clarify mutation action. In particular, our mechanism suggests that RMC-5552 (RMC-6272) bitopic inhibitors may benefit from adjustment of the (PEG₈) linker length when targeting certain mTOR variants. In the cryo-EM mTOR/RMC-5552 structure, the distance between the allosteric and orthosteric inhibitors is ~22.7 Å. With a closed catalytic cleft, this linker bridges the sites. However, in our activation mechanism, in the open cleft it expands to ~24.7 Å, offering what we believe to be the first direct example of how discovering an activation mechanism can potentially increase the affinity of inhibitors targeting mutants.

> (Raptor) is unique to mTORC1, while Raptor-independent companion of mTOR (Rictor) to mTORC2. mTORC1 is a dimer of heterotrimers containing three basic proteins, mTOR, Raptor, and mLST8, while mTORC2 is a dimer of heterotetramers with four conserved subunits, mTOR, Rictor, mLST8, and mammalian stress-activated map kinase interacting protein 1 (mSIN1).20 mTORCs' activation can be triggered through the PI3K/AKT/mTOR pathway.1,21-23 When growth factors such as insulin bind to receptor tyrosine kinases (RTKs), PI3K is activated, converting lipid phosphatidylinositol 4,5bisphosphate (PIP₂) to phosphatidylinositol-3,4,5-triphosphate (PIP₃).²⁴ PIP₃-mediated activation of PDK1 stimulates AKT phosphorylation, which subsequently inhibits a negative regulator, the tuberous sclerosis complex (TSC).²⁵⁻²⁸ In signaling, TSC acts as the GTPase activating protein (GAP) and inhibits GTP-bound Ras homolog enriched in brain protein (RHEB, a member of the Ras superfamily).^{29,30} Thus, AKT-mediated inhibition of TSC promotes RHEB activation. Active RHEB binds to the FAT domain of mTOR in mTORC1, allosterically changing the alignment of the active-site residues in the kinase domain in favor of catalysis. Full activation of mTORC1 requires the uptake of amino acids, such as Arg, Leu, and Glu, which regulate the recruitment of mTORC1 to lysosomes and



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endosomes through the Rag GTPases-mediated pathway.^{1,19,31-33} mTORC2 is predominately mediated by growth factor recruitment. Binding of mSIN1 to PIP₃ can release mTORC2 inhibition. mTORC2 activity can be tuned by the negative feedback of mTORC1. Ribosomal protein S6 kinase 1 (S6K1), a substrate of mTORC1, can phosphorylate insulin receptor substrate 1 (IRS1), which then attenuates the PI3K/AKT signaling, thus mTORC2 activity.³⁴

The classes and status of mTOR inhibitors

Studies associated constitutive mTORCs' activation with cancer.^{22,35-40} Alterations of mTOR regulators in the PI3K/AKT/ mTOR pathway, as well as mTOR itself, can stimulate signaling, triggering cell growth and proliferation,7,41-47 impacting neurodevelopmental disorders,48 neurodegenerative diseases, and type 2 diabetes.14,34 The activation of the PI3K/ AKT/mTOR pathway imparts non-single residue mutational resistance in acute lymphoblastic leukemia (ALL) linked to the oncoprotein Bcr-Abl1, which is encoded by the fusion BCR-ABL1 gene in the Philadelphia chromosome.49 Suppressing aberrant mTOR can extend life and healthspan.3,16,50,51 mTOR inhibitors can be classified into three generations. The first is rapalogs, the variants of rapamycin. Notable successes of rapalogs include temsirolimus (Torisel) by Wyeth, everolimus (Afinitor) by Novartis, and ridaforolimus by Merck, with a bulkier group replacing the hydroxyl group at C40 of rapamycin. The U.S. Food and Drug Administration (FDA) has approved temsirolimus for treating renal cell cancer and everolimus for treating breast cancer, renal cell carcinoma, and neuroendocrine carcinoma. Ridaforolimus' effectiveness is under evaluation in clinical trials. These drugs bind to the interfacial pocket between the FKBP12-rapamycin-binding (FRB) domain of mTOR and the 12 kDa FK506-binding protein (FKBP12), which appears to cause steric hindrance and obstruct the active-site cleft.4,52 The drug prefers binding to mTORC1 over mTORC2.53,54 In mTORC2, the rapamycin/FKBP12 binding site is occluded by Rictor's Cterminal domain, making it insensitive to rapamycin and rapalogs. However, under prolonged treatment, rapamycin can inhibit mTORC2.21 Side effects limit rapalogs' broad clinical applications.⁵⁵ In addition to rapalogs, there are non-rapalog allosteric inhibitors of mTOR. These inhibitors are specifically engineered to target the same binding site as rapamycin or rapalogs, but they exhibit distinctive interaction patterns.56 Like other kinase inhibitors, the second generation binds to the ATPbinding pocket of the kinase domain in mTORC1 and mTORC2,57 but often lack selectivity. Examples encompass sapanisertib (MLN0128, INK-128 or TAK-128), vistusertib (AZD2014), AZD8055, torkinib (PP 242), torin-1, torin-2, OSI-027 (ASP7486), and PP30. The dual PI3K/mTOR inhibitors effectively inhibit both PI3K and mTOR due to the catalytic domains' structural similarities.58 Illustrative instances include omipalisib (GSK2126458), dactolisib (NVP-BEZ235), gedatolisib (PF-05212384), apitolisib (GDC-0980), bimiralisib (PQR309), voxtalisib (SAR245409), panulisib (P7170), and PI-103. The combination regimen involving both allosteric and orthosteric drugs has been recognized as having increased potential in effectively

suppressing mTOR activity, making it a valuable approach for overcoming drug resistance.^{59–61} The third generation, known as rapalinks, are bitopic inhibitors that combine features of the first and second generations. Rapalinks can simultaneously occupy the cavity between the mTOR FRB domain and FKBP12, and the ATP-binding site.^{34,61–65} This dual binding mechanism enhances the affinity and selectivity of the inhibitors towards mTOR.

mTOR suppresses catalytic activity by restricting substrate access to the active site

The obligate cellular localization and substrate recruitment of mTORCs rely on their components.9,66 X-ray and cryo-EM structures of mTORC1 and mTORC2 provided structural insights into their regulation,2,4,14,53,67-69 and studies largely focused on mTORCs' ability to sense substrate signals. For both complexes, mTOR is the catalytic core. A recent crystal structure featured a complex of N-terminal truncated mTOR (Δ NmTOR) with mLST8, a subunit of both mTORC1 and mTORC2 (Δ NmTOR/mLST8).⁴ mLST8 is a scaffolding protein for mTORC2 cofactors Rictor and mSIN1 but apparently not critical for mTORC1.70 The crystal structure suggested that mTOR suppressed the catalytic activity by restricting substrate access to the active site.4 This mechanism resembles those of PI3K and SHP2.24,71 Two factors contribute to the restricted access. First, the extensions of the N-lobe by the mTOR FRB domain and the C-lobe by mLST8 generate a deep "V-shape" catalytic cleft. Second, the ka9b-helix (residues 2425–2436) and the negative regulator domain (NRD, residues 2430-2450) block the active site. The static crystal structure could not capture the conformational dynamics required to precisely discern how these domains and motifs orchestrate the conformational changes of mTOR for its kinase activity, in the wild type and the mutant.72,73

Here we compiled cellular and medicinal experimental data and performed extensive molecular dynamics (MD) simulations to establish and depict the mechanism of the intrinsic activation of mTOR. We then apply this mechanism to describe how oncogenic mutations of mTOR can promote its activation, and the drug actions. Building upon this mechanism, we suggest a drug strategy that is promising to effectively target the mTOR variants.

Results

Description of Δ NmTOR/mLST8, and comparison of mTOR with PI3K

The crystal structure (PDB ID: 4JSP) of Δ NmTOR/mLST8 has a compact architecture (Fig. 1a).⁴ Δ NmTOR (residues 1376– 2549) consists of the FAT domain with most α -helical repeats forming a "C-shape" α -solenoid conformation and the kinase domain with a typical bilobate conformation. The FAT domain winds around half of the kinase domain. The mTOR kinase domain has approximately 300 residues more than those of canonical kinases (Fig. 1b). The kinase domain contains three inserted domains: the N-lobe inserted FRB (residues 2021–2118) domain, the C-lobe inserted Lst8 binding element (LBE,

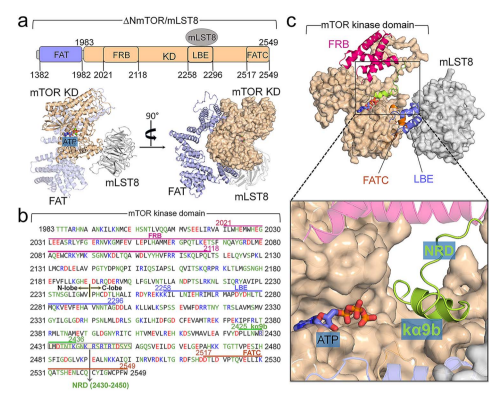


Fig. 1 The mTOR domain, sequence, and structure. (a) Domains and structure of Δ NmTOR/mLST8. (b) The sequence of the kinase domain of mTOR. There are several important inserted domains or motifs within the kinase domain of mTOR: the FRB domain (residues 2021–2118) in the N-lobe, and the LBE domain (residues 2258–2296), the k α 9b-helix (residues 2425–2436), the NRD (residues 2430–2450), and the FATC domain (residues 2517–2549) in the C-lobe. (c) Conformation of the catalytic cleft of mTOR with the k α 9b-helix and NRD inside the catalytic cleft, blocking the active site.

residues 2258-2296) domain, and the C-lobe inserted FAT Cterminal (FATC, residues 2517-2549) domain.9 The FRB domain may operate as a gatekeeper, providing a specific binding site for both rapamycin and substrates (Fig. 1c).^{2,4} The LBE domain is bound to mLST8. The FATC domain appears to be indispensable for C-lobe integration, packing beside the activation loop (A-loop). In addition to these three inserted domains (FRB, LBE, and FATC), the mTOR kinase domain has other inserted motifs. Within the N-lobe, there are insertions of a kinked helix (k α 1) immediately before FRB (Fig. S2a[†]), and a β strand (k β 0) and two short helices (k α 2 and k α 2b) right after FRB (Fig. S2b[†]). The ka1-helix is a conserved motif in other PIKKs and PI3K.^{4,74,75} In PI3K, the ka1-helix is followed by the ka2. The position of ka2 in PI3K is taken by k β 0, ka2, and ka2b in mTOR, likely as an anchor for the FRB domain onto the Nlobe. Within the C-lobe of the mTOR kinase domain, ka9b is also embedded, together with the LBE and FATC domains forming an A-loop-centered interaction spine in the vicinity of the active site (Fig. S2c[†]). Analogous to class III PI3KC, the Aloop displays an ordered helix (kaAL). Unlike the A-loop in the inactive conformation of those canonical kinases, kaAL within the A-loop of mTOR does not block the active site. The four inserted helices, ka3, ka9, ka9b, and ka10, form a compact structure centered on ka9b (Fig. S2d[†]).

The α C-helix of mTOR in both mTORC1 and mTORC2 is in the "IN" conformation,^{4,14,69,76} characteristic of active kinases.

The crystal structure of $\Delta NmTOR/mLST8$ shows that mTOR inhibition is primarily due to the blocked catalytic cleft,4,76,77 resembling PI3K, whose active site is blocked by the iSH2 domain. PI3K activation is initiated with nSH2 release, followed by the rotation of iSH2 and the exposure of the active site.⁷¹ In mTOR, the ka9b-helix and NRD inside the catalytic cleft can obstruct substrate access, suppressing its kinase activity. Experimental data showed that NRD removal greatly enhanced the basal activity of mTOR in vitro and in vivo,78-80 supporting the critical role of NRD in the regulation. The C-terminal of the $k\alpha$ 9b-helix partially overlaps the N-terminal of NRD (Fig. 1b). In this work, we aim to unravel the activation mechanism of the mTOR core, not considering mTOR effectors, such as Raptor, Rictor, and mSIN1. Experimental data indicated that the ΔNmTOR/mLST8 complex has kinase activity comparable to mTORC1,⁴ indicating that it is a functional complex. Thus, we constructed the simulation models based on the crystal structure of the $\Delta NmTOR/mLST8$ complex. Since mTORC1 and mTORC2 share the same binding posture between mTOR and mLST8 (Fig. S3[†]),^{14,69} we expect the elucidated mechanism to explain the basic activation of mTOR for both complexes.

Activation mechanism of mTOR by NRD release

To explore how NRD affects mTOR activity, we constructed two Δ NmTOR/mLST8 complex models, Δ NmTOR^{WT}/mLST8 with NRD in the catalytic cleft (termed IN NRD) and Δ NmTOR^{Δ NRD}/

mLST8 in the absence of NRD and performed MD simulations (Table S1[†]). During the simulations, we observed that mTOR in both systems underwent conformational changes in the kinase domain, mainly in the "V-shape" catalytic cleft. However, their changes differed. To characterize these changes, we measured the distance of FRB-LBE $(d_{\text{FRB-LBE}})$ (Fig. S4a[†]) and the angle of FRB-ATP-LBE ($\theta_{\text{FRB-ATP-LBE}}$) (Fig. S4b[†]) along the trajectories for these two systems. In $\Delta NmTOR^{WT}/mLST8$, $d_{FRB-LBE}$ and $\theta_{\text{FRB-ATP-LBE}}$ consistently decrease, ultimately populated at ~ 23 Å (Fig. 2a) and \sim 52° (Fig. 2b), respectively, which are much smaller than those (\sim 31 Å and \sim 67°) in the crystal structure, eventually resulting in a deeper and narrower catalytic cleft (Fig. 2c, left). In sharp contrast, the catalytic cleft of mTOR in the $\Delta NmTOR^{\Delta NRD}/mLST8$ complex was more prone to open (Fig. 2c, right). The removal of NRD resulted in $d_{\text{FRB-LBE}}$ and $\theta_{\text{FRB-ATP-LBE}}$ peaking at ~36 Å and ~73°, respectively, much larger than for the $\Delta NmTOR^{WT}/mLST8$ complex with NRD inside the catalytic cleft. The FRB domain harbors the second substrate binding site.⁴ This conformational change may favor the recruitment of the substrate at the FRB site (Fig. S5[†]).

These observations were consistent with the principal component analysis (PCA). The first two principal components (PCs) of the PCA for the kinase domains of mTOR in both Δ NmTOR^{WT}/mLST8 and Δ NmTOR^{Δ NRD}/mLST8 were clustered according to three timeframes, 0-500 ns, 500-1000 ns, and 1000-1500 ns (Fig. 3a), which reflect the motion patterns of the kinase domains during different periods. Overall, the kinase domain of mTOR in Δ NmTOR^{WT}/mLST8 exhibits relatively light variations, as evidenced by the considerable overlap of the three clusters of the first two PCs. For $\Delta NmTOR^{WT}/mLST8$, in the first 500 ns, the main change was the loss of half of the helical component of the ka9b-helix, which couples with NRD approaching the FRB domain (Fig. 3b). This leads to the formation of some salt bridges between the positively charged residues (K2440, R2241, R2443) in NRD and the negatively charged residues (E2032 and E2033) in the FRB helix (residues 2022-2040) (Fig. S6[†]). Such interactions favor the FRB domain getting closer to the LBE domain (or mLST8), giving rise to a closed catalytic cleft. This conformation persisted for 500-1500 ns. In sharp contrast, the $\Delta NmTOR^{\Delta NRD}/mLST8$ complex

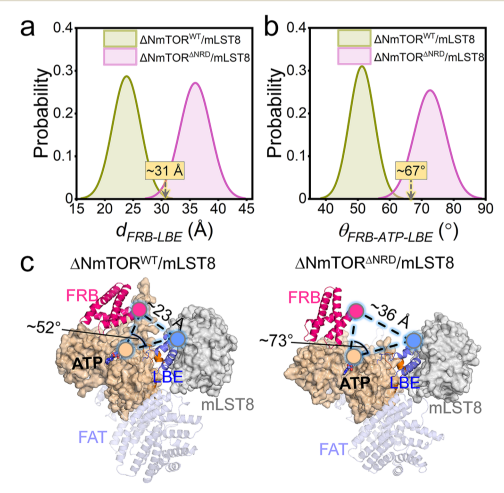


Fig. 2 NRD regulation of the mTOR kinase domain. Probability distributions of (a) the distance between the FRB and LBE domains ($d_{FRB-LBE}$) and (b) the angle of FRB-ATP-LBE ($\theta_{FRB-ATP-LBE}$), and (c) final snapshots to represent $d_{FRB-LBE}$ and $\theta_{FRB-ATP-LBE}$ of mTOR in Δ NmTOR^{WT}/mLST8 and Δ NmTOR^{Δ NRD}/mLST8. $d_{FRB-LBE}$ is the distance between the C α atom of residue M2039 in the FRB domain and the C α atom of residue M2277 in the LBE domain. $\theta_{FRB-LBE}$ is the angle between the vector from the phosphorus atom of the γ -phosphate of ATP to the C α atom of residue M2039 in the FRB domain and the vector from the phosphorus atom of the γ -phosphate of ATP to the C α atom of residue M2277 in the LBE domain.

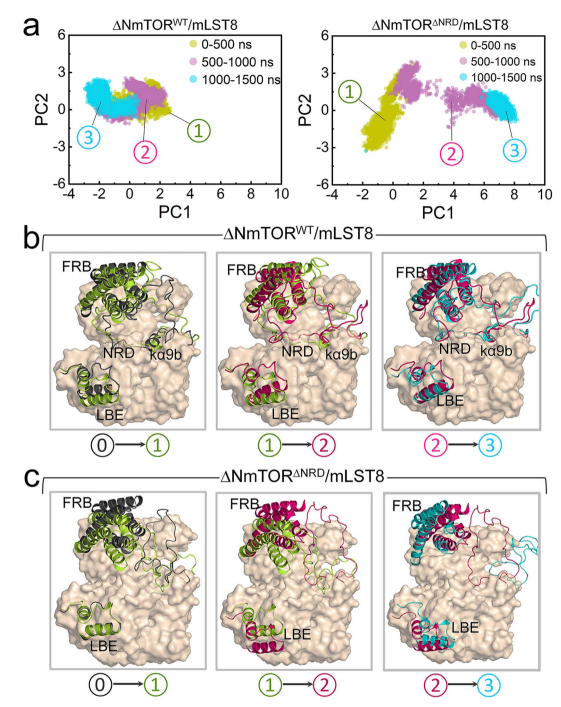


Fig. 3 PCA for the mTOR kinase domain in Δ NmTOR^{WT}/mLST8 and Δ NmTOR^{Δ NRD}/mLST8. (a) The projections of the first and second principal components (PC1 and PC2) from the PCA, and (b) structural alignment for the mTOR kinase domain in the three different simulation time frames (0–500 ns, 500–1000 ns, and 1000–1500 ns) for the Δ NmTOR^{WT}/mLST8 and Δ NmTOR^{Δ NRD}/mLST8 systems. In (b) and (c), structure (\odot) represents the initial configuration of the mTOR kinase domain, and structures (\odot), (\odot), and (\odot) are the representative configurations of the mTOR kinase domain in the 0–500 ns, 500–1000 ns, and 1000–1500 ns trajectories, respectively.

displayed a remarkably sparse distribution of the first two PCs, indicating that NRD release results in a remarkable variation in the mTOR kinase domain (Fig. 3a). In Δ NmTOR^{Δ NRD}/mLST8, the FRB domain initially underwent a weak rotation. Without restriction from NRD, the FRB domain moved away from the LBE domain (or mLST8), significantly enlarging the entrance of the catalytic cleft and shallowing it (Fig. 3c). Simultaneously, the

helical components of $k\alpha$ 9b entirely disappeared, and the region after NRD shifted out and moved away from the catalytic cleft. These observations support the essential role of NRD in regulating mTOR activity. Importantly, the IN NRD not only directly blocks the active site but also serves as the determinant for maintaining a deep and closed catalytic cleft through its restriction of the FRB domain, thereby prohibiting substrate

access to the active site. Upon NRD release, the FRB domain will move, giving rise to an open catalytic cleft and making the secondary substrate-binding site in the FRB domain more accessible, increasing the probability of substrate recruitment and access to the active site.

Oncogenic activation of mTOR

mTOR mutations are frequently detected in cancer patients. The activating mutations upregulate signaling through the PI3K/AKT/mTOR pathway, driving cancer.⁸¹ Most mTOR kinase domain mutations occur in the ka3-, ka9-, ka9b-, and ka10helices (Fig S2d†),⁴ disturbing the ka9b-centered packed structure. The top-3 most frequent mutating sites are S2215 at the interface between the N- and C-lobe, I2500 in the ka10-helix, and L2427 in the ka9b-helix (Fig S7[†]).² Here, we chose L2427. The three cancer databases, TCGA, GENIE, and COSMIC cataloged 26, 19, and 7 individuals harboring the L2427R, L2427Q, and L2427P mutations, respectively. We verified L2427R pathogenicity,82 modeled an L2427R mutated complex $(\Delta NmTOR^{L2427R}/mLST8)$ and conducted MD simulations (Table S1[†]). As expected, this mutation significantly affected the conformation and mobility of the ka9b-helix and NRD, with the helical components of the helix entirely disappearing (Fig. 4a). Compared to $\Delta NmTOR^{WT}/mLST8$, the averaged interaction energy between NRD and FRB dramatically increases from \sim -198.0 kcal mol⁻¹ to \sim -85.7 kcal mol⁻¹ (Fig. 4b), indicating that the conformational change weakens their interactions. In turn, this resulted in increasing the mobilities of FRB, $k\alpha$ 9b, and NRD, as well as in the unstructured region after NRD (Fig. 4c).

The sparse distribution of the first two components observed in the PCA further confirms the significant conformational change in the Δ NmTOR^{L2427R}/mLST8 kinase domain (Fig. 5a), with the L2427R mutation driving fast unfolding of the k α 9bhelix (Fig. 5b). Without the strong NRD–FRB interaction, NRD and the unstructured region after it move out from the catalytic cleft (termed OUT NRD) and extend. The structural alignment of the mTOR kinase domains in the 0–500 ns, 500–1000 ns, and 1000–1500 ns trajectories further confirms the large movement and large fluctuation of FRB, k α 9b, and NRD. Their instabilities facilitate the opening of the catalytic cleft, supporting the premise that some mTOR-activating mutations may loosen the structural framework around the k α 9b-helix,⁴ promoting the shift of the conformational ensemble of mTOR toward the catalysis-favored state.

mTOR drug discovery

mTOR is a compelling drug target, with recent advances garnering increasing attention.^{34,64,83-86} Beyond fundamental therapeutic indices such as solubility, affinity, and broad kinase toxicity with off-target effects, ensuring high selectivity for mTORC1 over mTORC2 is critical. mTORC2 inhibition has been

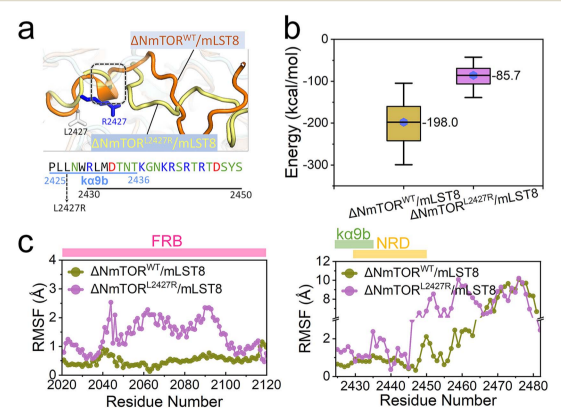


Fig. 4 Oncogenic L2427R mutation increases the high dynamics of k α 9b, NRD, and FRB. (a) Comparison of the conformations of the k α 9b-helix and NRD, (b) interaction energy between NRD and FRB, and (c) root mean square fluctuations (RMSF) of the FRB domain (left), and the k α 9b-helix, NRD, and the unstructured region after NRD (right) for the Δ NmTOR^{WT}/mLST8 and Δ NmTOR^{Δ NRD}/mLST8 systems.

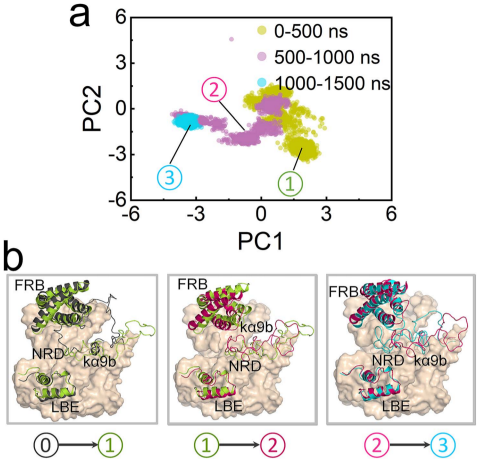


Fig. 5 PCA for the mTOR kinase domain in Δ NmTOR^{L2427R}/mLST8. (a) The projections of the first and second principal components (PC1 and PC2) from the PCA, and (b) structural alignment for the mTOR kinase domain in the three different simulation time frames (0–500 ns, 500–1000 ns, and 1000–1500 ns) for the Δ NmTOR^{L2427R}/mLST8 system. In (b), structure (\odot) represents the initial configuration of the mTOR kinase domain, and structures (\odot , (\odot), and (\odot) are the representative configurations of the mTOR kinase domain in the 0–500 ns, 500–1000 ns, and 1000–1500 ns trajectories, respectively.

associated with compromised insulin sensitivity and glucose intolerance.55 A noteworthy advancement in mTOR drug discovery is the emergence of novel bitopic inhibitors. Shokat et al. devised Rapalink-1, a compound connecting an allosteric inhibitor (rapamycin) and an orthosteric inhibitor (MLN0128) using a PEG-rich linker (Fig. 6a).62,63 The linked chemotypes in the bitopic inhibitor trigger a mutually beneficial interaction, which can cooperatively enhance the binding of both components, augmenting the affinity and selectivity for the target protein. Some residues in the FRB and kinase domains contribute to the allosteric and orthosteric drug binding. Their mutations promote resistance to the monovalent drug interventions. Patients harboring the A2034V and F2108L mutations within the mTOR FRB domain demonstrate resistance to the allosteric drug (everolimus) and the M2327I mutation is able to disrupt the binding of the orthosteric drug (AZD8055).61,87 With the cooperation of rapamycin and MLN0128, Rapalink-1 can effectively overcome the resistance.⁶¹ Recently, Revolution Medicines Inc. created two exceptionally potent bitopic inhibitors, RMC-5552 and RMC-6272 with a linker comprised of eight EG units (PEG₈) to connect a rapalog with "rigid MLN" and

XL388, respectively. These inhibitors exhibited elevated selectivity for mTORC1 and potent anti-tumor effects, positioning them as promising contenders for targeted therapeutic approaches.⁶⁵ RMC-5552 is currently undergoing clinical trials (NCT ID: NCT05557292 and NCT04774952) to evaluate its effectiveness in treating patients with recurrent glioblastoma and relapsed/refractory solid tumors.

Our simulations indicate that mTOR adopts an open catalytic cleft when activated. In its wild-type form, mTOR maintains a closed catalytic cleft. However, oncogenic mutations, such as L2427R, result in a high population of the open catalytic cleft, accompanied by the FRB domain displacement. A cryo-EM structure of the mTOR/RMC-5552 complex (PDB ID: 8ERA) reveals a closed catalytic cleft of mTOR (Fig. 6b), suggesting that RMC-5552 exhibits potency against wild-type mTOR. However, its effectiveness against oncogenic mTOR remains uncertain. The distance between the allosteric and orthosteric inhibitors is ~22.7 Å in the cryo-EM mTOR/RMC-5552 structure. The length of the linker in bitopic inhibitors is critical for high binding affinity.^{62,63,88,89} In mTOR with a closed catalytic cleft, RMC-5552 and RMC-6272, with the PEG₈ linker, aptly occupy the space

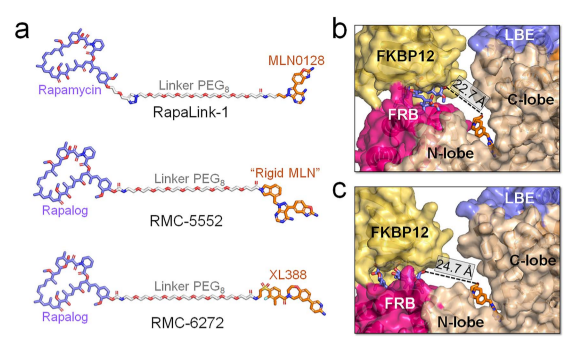


Fig. 6 Structural insights into the binding of the bitopic inhibitors and mTOR. (a) Structures of mTOR's bitopic inhibitors, RapaLink-1, RMC-5552, and RMC-6272. Binding modes between RMC-5552 and mTOR with the (b) closed and (c) open catalytic cleft. The distances between allosteric (blue sticks) and orthosteric (orange sticks) inhibitors in (b) and (c) were measured. The RMC-5552/mTOR complex in (b) is extracted from the cryo-EM structure (PDB ID: 8ERA), in which the mTOR adopts the closed catalytic cleft. In (c), the RMC-5552/mTOR complex was modeled by integrating RMC-5552 with a representative configuration of mTOR extracted from the trajectories of the Δ NmTOR^{Δ NRD}/mLST8 system, in which mTOR populates the open catalytic cleft.

between mTOR's allosteric and orthosteric sites. In our modeled RMC-5552/mTOR complex with an open catalytic cleft the distance between the allosteric and orthosteric inhibitors expands to ~24.7 Å (Fig. 6c). This indicates that bitopic inhibitors with PEG₈ linker may encounter reduced affinity to mTOR with open cleft, possibly diminishing their effectiveness against oncogenic mTOR variants. At the same time, long linkers are more flexible, and depending on their chemistry, can be allergenic. Collectively, our analysis suggests that it might be advantageous to consider adjusting the linker length and/or chemistry of the bitopic inhibitors when targeting a certain mTOR variant. In experiments, it would be valuable to have the compounds with shorter and longer linkers to determine if the linker length is indeed responsible for the observed changes in distance.

Discussion

The allosteric mechanism underlying mTOR activation

Here we provide a detailed account of the allosteric mechanisms underlying the activation of the wild-type and mutated mTOR at the atomic level. Our findings are consistent with a broad range of experimental data and support the previously proposed concept of "active-site restriction".^{4,78-80} We demonstrate that two partially overlapping motifs, the k α 9b-helix and NRD, play a crucial role in regulating the conformation and activity of the mTOR kinase domain. The FRB domain and mLST8 extending from the N- and C-lobes of the mTOR kinase domain, respectively, create a deep "V-shape" catalytic cleft that

partially restricts substrate access to the active site. The NRD can switch between two states, inside (IN) and outside (OUT) the catalytic cleft (Fig. 7 & Movie S1[†]). As proposed earlier,⁴ when the NRD and the ka9b-helix are in the IN state, they can obstruct the catalytic cleft, effectively preventing substrate access. Our simulations reveal that the NRD can also directly interact with the FRB domain, further restricting its movement and stabilizing the deep and closed "V-shape" cleft. This interaction strengthens the restriction imposed on substrate access. Conversely, when the NRD is in the OUT state, these restrictions are relieved, leading to increased mobility of the FRB domain. The relaxed FRB domain becomes more likely to move away from the mTOR LBE domain (or mLST8), resulting in a shallow and open catalytic cleft, with substrates able to more easily access the active site. The movement of the FRB domain also exposes its secondary substrate-binding site, facilitating substrate recruitment. These structural changes create a seemingly optimal environment for efficient catalysis. Our results also shed light on why the deletion of NRD (residues 2430-2450), rather than the partially overlapping/adjoining unstructured segment of 2443-2486, can significantly increase the kinase activity of mTOR.76

mTOR's autoinhibition and its release

Wild-type mTOR predominantly adopts an autoinhibited state characterized by a closed catalytic cleft (Fig. 7). Activating physiological interactions allosterically release mTOR from this state to assume an active conformation. In mTORC1, these include the interaction between RHEB and the mTOR FAT

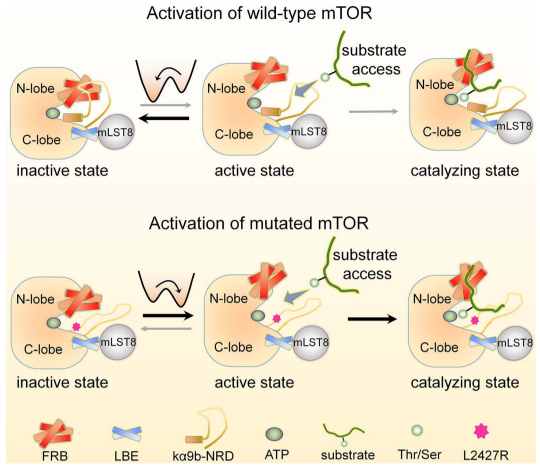


Fig. 7 A schematic illustration of mTOR's activation mechanism. The release of NRD from the catalytic cleft of mTOR leads to the opening of the catalytic cleft and the substrate-binding site on FRB making them more accessible, which allows for catalysis. Oncogenic mutations, such as L2427R, can promote this activation process, shifting the conformational ensemble of the mTOR kinase domain toward the catalysis-preferred state.

domain. For mTORC2, they include the binding of the mSIN1 PH domain to the membrane. The interaction between RHEB and the mTOR FAT domain is known to allosterically promote activation of mTORC1 by releasing the FAT domain autoinhibition,² with RHEB's C-terminal farnesylation promoting its lysosomal membrane interactions and activation,⁹⁰ while the binding of the pleckstrin homology (PH) domain of mSIN1 to the membrane facilitates activation of mTORC2. RHEB also stimulates a conformational change that allosterically rearranges mTORC1 active-site residues, accelerating catalysis.²

The k α 9b-helix plays a critical role in regulating the activity of mTOR. The movement of this helix is closely linked to the mobility of the NRD. When the NRD exhibits higher mobility, it is more likely to shift out from the catalytic cleft (Fig. 7). Several oncogenic mutations were identified within the k α 3-, k α 9-, and k α 10-helices of mTOR, which are neighboring motifs that surround the k α 9b-helix. These mutations have the potential to disrupt packing around k α 9b, leading to its, and the NRD's, increased mobility, ultimately promoting a shift of the mTOR ensemble towards a catalytically favored state. The L2427R mutation, which stimulated unfolding of the k α 9b-helix, provides an example. The mutation enhanced the mobility of

both the ka9b-helix and the associated NRD and FRB. Our mTOR^{L2427R} simulations observed that NRD tends to adopt an OUT conformation, favoring catalytic activity. We expect that other oncogenic mutations at the 2427-site, such as L2427R and L2427Q, or mutations at positions adjacent to the 2427-site, like S2215 and I2500, may similarly enhance mTOR activity through such a mechanism. Increased dynamics of the FRB domain can also destabilize the IN conformation of the NRD, leading to a preference for the OUT state. Oncogenic mutations in the FRB domain (e.g., E2029K, E2032A, E2033G) may weaken the interactions with the NRD, resulting in a shallower and more open catalytic cleft, which, in conjunction with the OUT orientation of the NRD, further promotes mTOR activity. This also suggests the presence of allosteric communication between these mutations and the orthosteric ATP-binding site.91,92 As recorded in the ClinVar database, pathogenic mTOR variants manifest as somatic and germline mutations. Somatic and germline alterations can drive cancer, while germline mutations are also commonly detected in disorder diseases. These allele origins of mTOR variants can share identical mutations, suggesting that individuals with disorder diseases may have an elevated risk of developing cancer.93,94

8

Different than other protein kinases, mTOR may regulate its activity through dynamic opening and closing of its deep catalytic cleft

mTOR is a protein kinase, but its mechanism differs. Kinases' catalytic activity is governed by the energy barrier of the conformational change toward the active state (Ka) and substrate accessibility to the active site (Km).⁹⁵ Typically, activation involves an OUT-to-IN movement of the aC-helix and collapsed-to-extended transition of the A-loop. These features can be used in kinase drug discovery strategies.96 In contrast, the mTOR *a*C-helix maintains the IN conformation, suggesting that it is constitutively active. mTOR appears to have evolved to regulate its activity through the dynamic opening and closing of its catalytic cleft, as observed in our simulations. The depth of mTOR's active site may restrict phosphorylation sites adjacent to bulky substrate structures. Common mTORC1 and mTORC2 substrates include the AGC kinase family, e.g., AKT, S6K1, SGK1, PKN1, and PKCs. They primarily phosphorylate serine or threonine within hydrophobic motifs (HM), TOR interacting motifs (TIM), and turn motifs (TM), rather than those within the A-loop of the bulk kinase domain.20 mTOR protein kinase populates heteromeric mTOR complexes, unlike most protein kinases in human cells that exist as monomers or homomeric assemblies. Other mTORCs' subunits play obligatory roles in cellular localization and regulation, primarily through substrate recruitment (Km).9 Raptor is responsible for recruiting mTORC1 substrates through direct binding to the TOR signaling (TOS) motifs within the substrate, whereas for mTORC2, mSIN1, rather than Rictor, appears to be the substrate recruiter. Anchoring mSIN1 onto the complex appears the only explored function of Rictor.²⁰ Experimental data have also suggested mSIN1 inhibition of mTORC277 through interaction of mSIN1's PH domain with mTOR kinase domain. Competitive binding of mSIN1 PH domain to PIP₃ relieves kinase domain inhibition. Binding of mSIN1 Ras binding domain (RBD) to H-Ras and K-Ras can dampen MAPK signaling.66,97,98 DEP domaincontaining mTOR interacting protein (DEPTOR) operates as an endogenous mTOR inhibitor. DEPTOR binding to the mTOR FAT domain induces an inactive conformation, partially inhibiting mTORC1 and mTORC2.99 Proline-rich AKT substrate 40 kDa (PRAS40) inhibits mTORC1 by blocking the first substrate-binding site on Raptor and the second on the mTOR FRB domain.² These interactions can compete with substrates, such as S6K1 and eukaryotic translation initiation factor 4Ebinding protein 1 (4E-BP1), binding to mTORC1. RHEB's regulation of mTORC1 does not relate to substrate recruitment. It can associate with the FAT domain of mTORC1 promoting allosteric changes that favor catalysis by altering the alignment of active site residues.² mLST8 is a common component in both mTOR complexes but functions differently in mTORC1 and mTORC2.70 Since mLST8 is indispensable for maintaining the Rictor-mTOR interaction but apparently not Raptor-mTOR, it is required for phosphorylation of AKT and protein kinase C alpha (PKCa) by mTORC2 but not S6K1 by mTORC1.50,69

Our work elucidates exactly how the domains and inherent motifs of mTOR regulate conformational changes in the mTOR

kinase domain, thereby controlling kinase activity. Our mechanism focuses on conformational sampling, thereby elucidating activation principles of both mTORCs and helps explain how mutations can drive mTOR transition to the active state. In particular, our proposed allosteric activation mechanism can suggest how to optimize bitopic inhibitors informing selective targeting of the mTOR mutants. Finally, it connects protein conformational ensembles to function, mechanisms of activation and drug discovery, in line with theory.^{100,101}

Conclusions

In this work, we decipher the mechanism of intrinsic and oncogenic activation of mTOR at the atomic level. Our findings highlight the significance of the unstructured N-terminal regulatory domain (NRD) (residues 2430-2450) as a key regulator and show that NRD dynamics are closely associated with conformational changes in the kinase domain. When NRD is located inside the catalytic cleft along with ka9b, it obstructs the active site, effectively blocking mTOR activity. NRD interacts directly with the mTOR FRB domain, thereby constraining its motion. This interaction contributes to maintaining a deep and closed catalytic cleft, which restricts substrate access. Upon NRD release from the catalytic cleft, the mTOR FRB domain moves away from the mTOR LBE domain (or mLST8), resulting in an open conformation of the catalytic cleft and a more accessible substrate-binding site on FRB, which is favorable for substrate access and recruitment. The motion of NRD is affected by the dynamics of the adjacent ka9b-helix. The oncogenic mutation L2427R promotes unfolding of the ka9bhelix. This results in increasing the mobilities of NRD, $k\alpha 9b$, and FRB, which reduces the interaction between NRD and FRB, accelerating mTOR activation by shifting the equilibrium toward the catalysis-favored state. These scenarios lead us to expect that mutations that can break the ka9b-helix-centered packed structure to promote mTOR activation follow this activation principle. Our proposed mechanism aligns with previous experimental observations, expands the concept of the activesite restriction mechanism of mTOR, and clarifies how oncogenic mutations can drive mTOR activation. Our simulations may help in more potent bitopic inhibitors that could effectively target mTOR mutants. This study advances our understanding of mTOR regulation and offers insights for therapeutic intervention.

Materials and methods

Modeling of different Δ NmTOR/mLST8 complex systems

We modeled three different Δ NmTOR/mLST8 complex systems based on the crystal structure of N-terminal truncated mTOR complexing with mLST8 (PDB ID: 4JSP). They are Δ NmTOR^{WT}/ mLST8, Δ NmTOR^{Δ NRD}/mLST8, and Δ NmTOR^{L2427R}/mLST8 (Table S1†). The missing regions in the crystal structure were modeled using the AlphaFold2 package.^{102,103} Based on the active-site restriction mechanism that the k α 9b-helix and NRD block the active site of mTOR, we manually adjusted NRD, leading it to approach the active site. The resulting complex was used as $\Delta NmTOR^{WT}/mLST8$. Based on $\Delta NmTOR^{WT}/mLST8$, the $\Delta NmTOR^{\Delta NRD}/mLST8$ and $\Delta NmTOR^{L2427R}/mLST8$ systems were constructed by deleting the 2430–2450 segment, and substituting L2427 with Arg, respectively. An ATP molecule and two magnesium ions (Mg²⁺) were placed in the ATP-binding pocket of mTOR in the three complexes. The resulting systems were solvated using the TIP3P solvent model. The Na⁺ and Cl⁻ ions were added into the solvated systems to neutralize the systems and to generate a physiological salt concentration of ~0.15 mol L⁻¹.

MD simulation protocol

The all-atom MD simulations were performed for the three systems using the NAMD 2.13 package¹⁰⁴ with the CHARMM¹⁰⁵ all-atom additive force field (version C36).106,107 The simulation protocol closely followed the methodology employed in our previous studies.71,108-111 Prior to MD simulation, we performed several cycles of minimization and dynamics to eliminate atom contacts within the systems. Each system underwent 1.5 µs MD simulations under the NPT ensemble with 3D periodic boundary conditions. To assess the consistency of outcomes, we executed three parallel trajectories concurrently for each individual system. The results derived from these three parallel trajectories exhibited similarities and were found to be comparable. Throughout the simulations, pressure was maintained at 1 atm using the Langevin piston control algorithm, while the temperature was kept at 310 K using the Langevin thermostat method with a damping coefficient of 1 ps^{-1} . The SHAKE algorithm was used to constrain covalent bonds, including hydrogen atoms. A 2 fs time-step was used in the simulations. The long-range electrostatic and short-range van der Waals (vdW) potential energies were calculated using the particle mesh Ewald method and switching functions, respectively. The analysis used the FORTRAN script in the CHARMM package (version c45b1) and the TCL script in the VMD package.

Data availability

Data is available upon request.

Author contributions

YL, MZ, HJ, and RN conceived and designed the study. YL and MZ conducted the simulations. YL also analyzed the results and drafted the manuscript. YL, HJ, and RN validated, reviewed, and edited the manuscript. RN supervised the project.

Conflicts of interest

The authors declare no potential conflicts of interest.

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