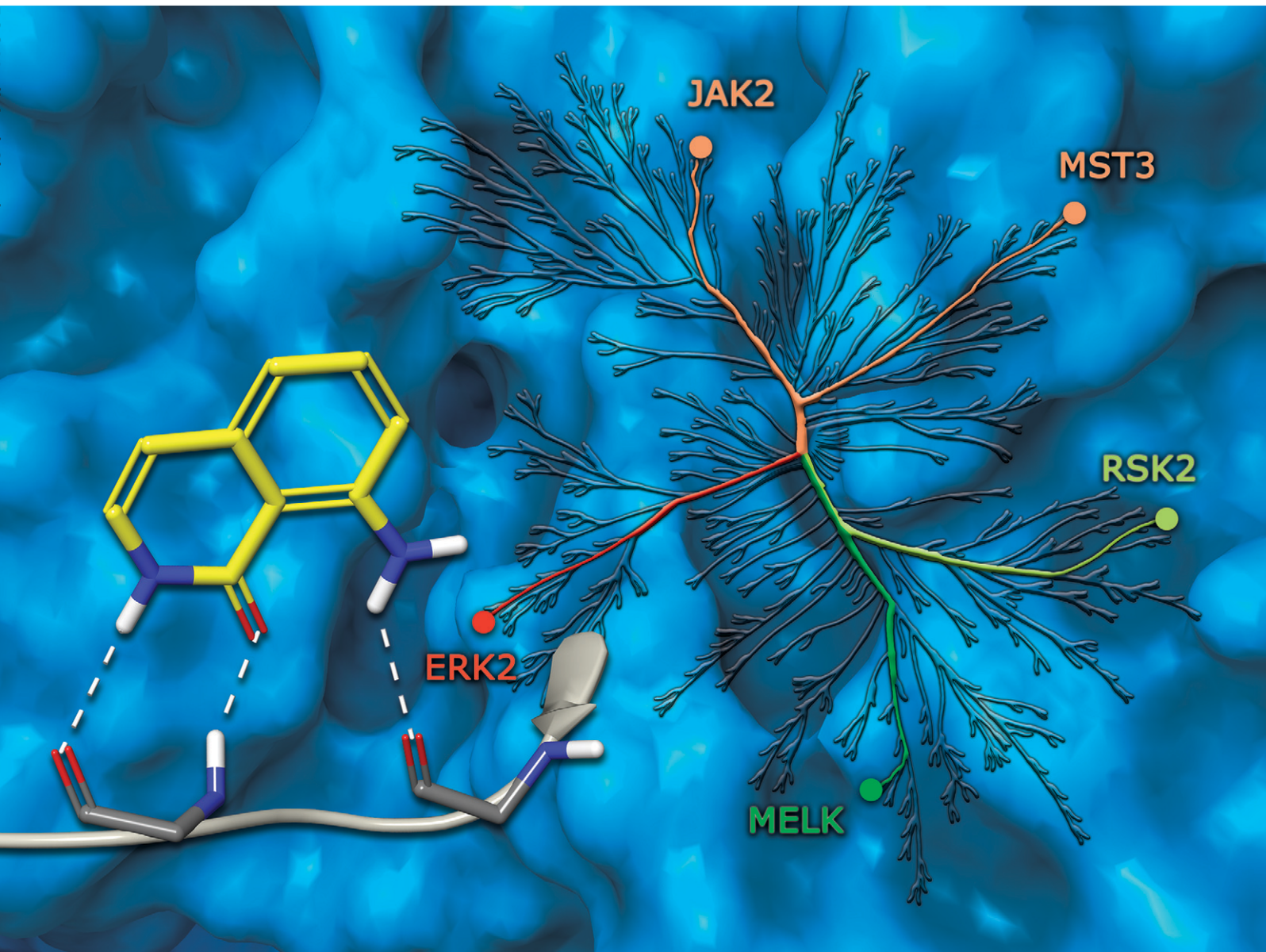


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Discovery of a novel kinase hinge binder fragment by dynamic undocking

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One of the key motifs of type I kinase inhibitors is their interactions with the hinge region of ATP binding sites. These interactions contribute significantly to the potency of the inhibitors; however, only a tiny fraction of the available chemical space has been explored with kinase inhibitors reported in the last twenty years. This paper describes a workflow utilizing docking with rDock and dynamic undocking (DUck) for the virtual screening of fragment libraries in order to identify fragments that bind to the kinase hinge region. We have identified 8-amino-2*H*-isoquinolin-1-one (**MR1**), a novel and potent hinge binding fragment, which was experimentally tested on a diverse set of kinases, and is hereby suggested for future fragment growing or merging efforts against various kinases, particularly MELK. Direct binding of **MR1** to MELK was confirmed by STD-NMR, and its binding to the ATP-pocket was confirmed by a new competitive binding assay based on microscale thermophoresis.

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Introduction

Protein kinases are a major class of drug targets, with over 30 marketed drugs and 250 drug candidates undergoing clinical studies, mainly but not only in various oncology indications.¹ The majority of kinase inhibitors target ATP binding sites (type I), which contain a conserved hinge region. For this reason, one of the main challenges in the development of kinase inhibitors is obtaining selectivity. Another major challenge is avoiding an already congested IP space, as the diversity of hinge binder scaffolds used in type I kinase inhibitors is relatively limited.² Consequently, these chemotypes have barely been sampled, which prompts the thorough exploitation of their chemical space.^{3,4} For example, it has been shown recently that only 1% of potential hinge binders are present in known kinase inhibitors.⁵

Fragment-based drug discovery (FBDD) has already been used in many kinase programs⁴ and has the potential to

identify novel fragment-sized hinge binders and specifically “evolve” the fragment for the targeted kinase.^{6,7} In fact, the first FBDD derived drug on the market, vemurafenib, targets the oncogenic V600E mutant of the B-Raf kinase.⁸ Structure-based virtual screening approaches have been used effectively for both the identification of novel fragment-sized hinge binders and their optimization. In one case study, Kolb and colleagues screened 730 000 compounds and discovered two ligands with different hinge binder moieties. The initial set was first filtered by kinase hinge binding pharmacophore restraints and the resulting 21 418 compounds were docked to the ATP site.⁹ In another study, Urich and colleagues extracted core fragments from 2.3M commercially available compounds. The resulting unique fragments were filtered for kinase hinge pharmacophores, and were subsequently docked into a panel of protein kinases. This strategy identified a number of hinge binder fragments with no previously reported activity against the investigated kinases.¹⁰

In this work, we used a novel screening strategy to find potent hinge binding fragments. Our dynamic undocking¹¹ based approach identified **MR1** (Fig. 1A) that has been experimentally validated against five kinase targets (Table 1). Future fragment growing or merging efforts toward these targets could avoid a congested druggable chemical space (e.g. for JAK2) and could provide a suitable chemical starting point to unmet medical needs (e.g. for MELK). JAK2 belongs to the non-receptor tyrosine kinase family, for which the V617F mutation is known to be implicated in myeloproliferative disorders.^{12,13} MST3 belongs to the Ste20

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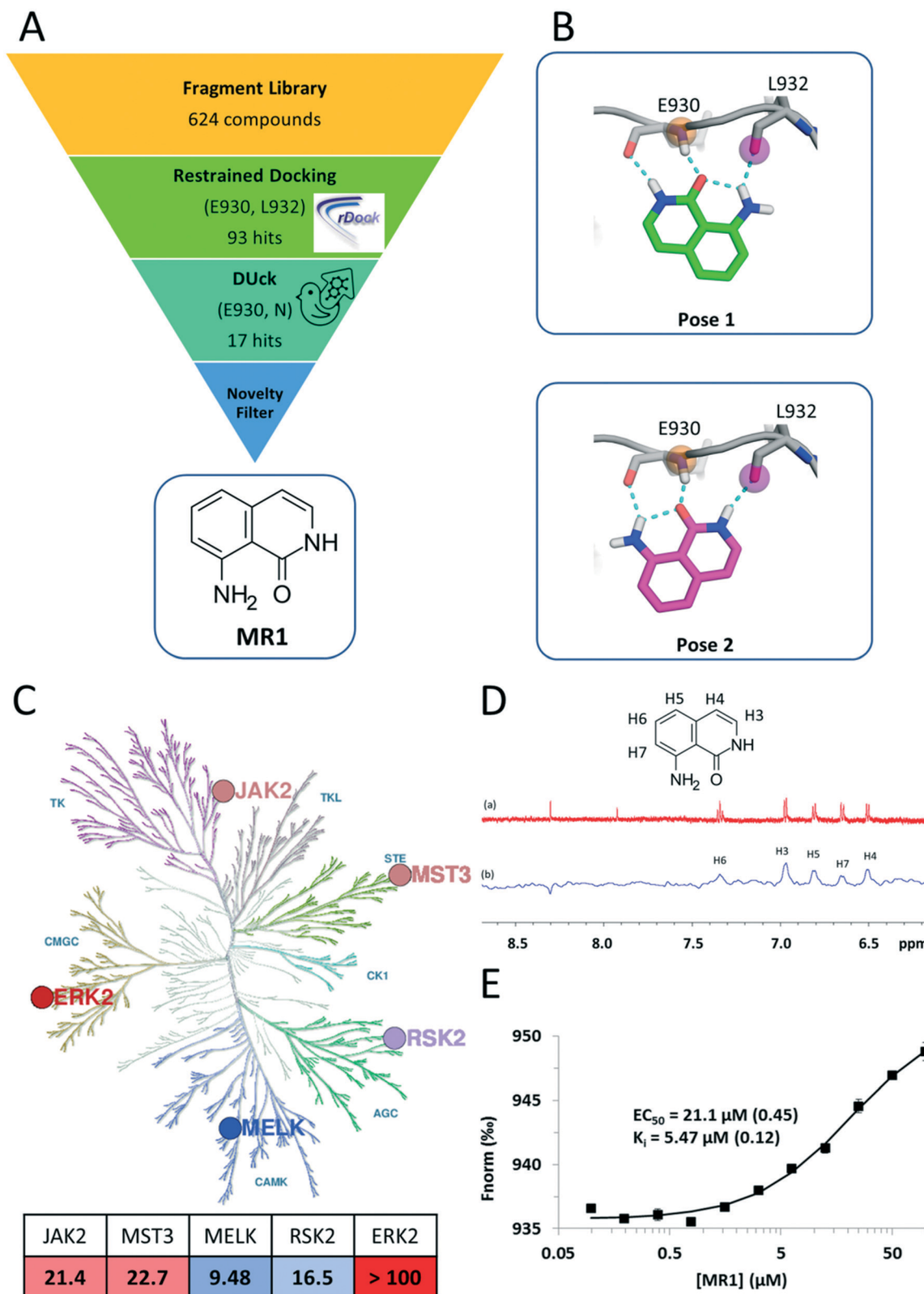


Fig. 1 A) Summary of the virtual screening workflow. B) Predicted binding modes of MR1 in the JAK2 ATP pocket. Both binding modes utilize tridentate H-bond interactions with the backbone carbonyl and amide groups of the hinge residues E930 and L932. C) Kinase inhibitory profile of MR1 overlaid on the kinase phylogenetic tree²⁸ (coloured blue to red, from most active to least active). The table summarizes the respective IC₅₀ values in μM units, with the same colouring scheme (illustration reproduced courtesy of Cell Signaling Technology, Inc., www.cellsignal.com). D) ¹H NMR spectrum of MR1 (a) and saturation transfer difference NMR spectrum of MR1 in the presence of MELK (b). E) In the competitive, MST-based binding assay, titration of MELK with MR1 reveals concentration-dependent binding with a K_i value of 5.47 μM and confirms the ATP-pocket as the location of binding.



Table 1 Summary of the kinase targets investigated in this study

Branch	Kinase	Uniprot ID	Clinical indications
AGC	RSK2	P51812	Oncogenesis and leukemia ¹⁵
CAMK	MELK	Q14680	Oncogenesis and cancer treatment resistance ¹⁶
CMGC	ERK2	P28482	Cervical ¹⁷ and colorectal ¹⁸ cancer
STE	MST3	Q9Y6E0	Breast cancer ¹⁴
TK	JAK2	O60674	Myeloproliferative neoplasms (MPNs) ^{12,13}

serine/threonine protein kinase family, which has been shown to promote proliferation and tumorigenicity.¹⁴ Maternal embryonic leucine zipper kinase (MELK) is a serine/threonine kinase belonging to the CAMK family, and was initially found to be expressed in a wide range of early embryonic cellular stages.

More recently, MELK has been identified to be present in several human cancers and stem cell populations with a unique spatial and temporal pattern, which suggests a prominent role in cell cycle control, cell proliferation, apoptosis, cell migration, cell renewal, embryogenesis, oncogenesis, and cancer treatment resistance and recurrence.¹⁶ ERK2 is part of the Ras/Raf/MEK/ERK pathway, which is often overactivated in a very wide range of cancers, while RSK2 is a downstream effector of this pathway and phosphorylates substrates involved in transcription, translation, cell cycle regulation, and cell survival.¹⁵

Results and discussion

Here, we present a novel and potent hinge binding fragment, **MR1** (Fig. 1A), that was discovered through virtual screening (VS) of our in-house fragment library¹⁹ using kinase hinge pharmacophore information in a hierarchical VS strategy (Fig. 1A). Our initial screening has been performed on JAK2 as a prototypic protein kinase with a large number of type I inhibitors. It is well known that the kinase hinge region exhibits a pronounced H-bond pattern with ATP or ATP-site competitors and that these interactions are imperative for type I kinase inhibitors to bind. As such, a typical VS strategy would incorporate such information by ensuring that the retained ligands maintain these features through the use of *e.g.* pharmacophoric restraints. Here, we employ an additional filter that emphasizes the importance of the defined H-bonds through dynamic undocking (DUck¹¹), which estimates the robustness of H-bonds. DUck is an orthogonal method to approaches meant to predict binding free energies. Instead, it assesses structural stability through evaluation of the robustness of H-bonds. The VS strategy consisted of docking with rDock²⁰ using pharmacophoric restraints to ensure that the features necessary for H-bonding with the hinge were present, then using DUck to estimate the resistance to rupture of these H-bonds. This strategy led to the discovery of **MR1**, which was tested on five kinases from the major branches of the kinase phylogenetic tree (Fig. 1C), each with several relevant indications (Table 1). MST3 and MELK, in particular, have only few known potent inhibitors.

MR1 was found to inhibit four of the target kinases with IC₅₀ values in the low to mid-micromolar range (corresponding to ligand efficiencies between 0.54–0.59),²¹ which is in line with its small size and other hinge binders of this complexity.⁷ ERK2 was the only kinase that was not inhibited; one possible reason behind this is the higher ATP concentration (100 μM) in the ERK2 inhibition assay, as compared to the rest of the kinases (10–50 μM), resulting in a stronger competition of ATP toward the binding site. The assessment of the binding mode of **MR1** was based on the docking and DUck scores. Two plausible solutions were generated (Fig. 1B), both of which adopt a tridentate interaction with the hinge. This was verified also for MELK: here, pose 2 is clearly preferred (*W*_{QB} value of 7.7 kcal mol⁻¹, *vs.* 1.6 kcal mol⁻¹ for pose 1). The ability of fragments to bind simultaneously in multiple orientations has been reported in numerous occasions, and can even be favourable from an entropic perspective.²² Depending on how the fragment is then elaborated, it freezes into one binding mode or the other,^{23,24} which will likely stay conserved during the elaboration process²⁵ (this also means that the mentioned entropic gain will be lost, but ideally, this is compensated by a larger favourable enthalpy decrease due to additional secondary interactions between the protein and the elaborated ligand). The tridentate interaction is not often observed within kinases (a bidentate interaction being the most common), but – as supported by the inhibition data – a third interaction allows for high versatility.²⁶ This can be constrained at the optimization phase for introducing family-specific substituents. In fact, it is not the hinge binder but rather its decoration that determines kinase selectivity. Consequently, selective kinase inhibitors can stem from an unselective fragment.^{7,27}

In addition to identifying **MR1**, we have observed a stepwise enrichment of known hinge binders along the workflow: while the whole fragment library contained known hinge binders in 14.4% of the compounds (90 out of 624), this ratio was 64.7% for the virtual hits after the DUck calculations (11 out of 17). This can be considered as a retrospective validation of the presented virtual screening workflow.

Next, we aimed to confirm the direct and ATP-competitive binding of **MR1** experimentally. For this purpose, we have selected MELK, as **MR1** displayed the strongest inhibitory activity against this kinase. Direct binding to MELK was confirmed by an STD-NMR measurement (Fig. 1D), while the binding site was validated with a competitive MST-based (microscale thermophoresis) assay developed in our lab (Fig. 1E).



The assay is based on the displacement of a fluorescently labelled type I reference ligand (see the Experimental section) and it has additionally enabled us to quantify the dissociation constant of **MR1** from MELK as 5.47 μM . Its high ligand efficiency ($\text{LE} = 0.67$) nominates **MR1** as a viable starting point for optimizing ATP-site kinase inhibitors.

Experimental

Virtual screening

The structures were prepared using MOE 2016,²⁹ by removing water and cofactors, capping the termini and gaps, and for protonation with default settings. For the virtual screening, the JAK2 PDB structure, 3E64 (chain A) was used.³⁰ For docking, the cavity was defined in the prepared structure by the reference ligand method, using the crystallized ligand as reference. The fragment library consisted of 624 compounds and was prepared with LigPrep,³¹ so that ligands above 300 Da would be ignored, at most eight stereoisomers, six tautomers and eight ring conformers would be generated and lastly, probable ionization states within the pH range of six to eight would be generated.

The prepared library was docked with pharmacophoric restraints at the hinge region, namely, Glu930, O and Leu932, N. The pharmacophore was defined as a 2 Ångström radius around the mentioned receptor atoms. If the feature did not adhere to the positional constraints, rDock would assign a positive (unfavourable) pharmacophore restraint score, for which the cutoff was set to 0.5. Furthermore, a high-throughput VS (HTVS) protocol was implemented, which consisted of three stages, for which at every stage the number of docking runs increases, and the rDock "SCORE.INTER" filter becomes stricter. The filter was adapted to scores expected for fragment-sized molecules which resulted in 93 compounds in total. DUCK was performed on this set of compounds, pulling from Leu932, N (pulling from Glu930, O did not result in any additional filtering). The first step for a DUCK simulation is the definition of the chunk (a part of the protein structure) that represents the local environment surrounding the residue interacting with the ligand. When selecting residues for the chunk, the following guidelines were considered: I) selecting as few residues as possible to reduce computational time, II) residues were not selected if they would block the ligand from exiting the pocket during the simulations based on the directionality of the H-bond, III) residues were not removed if this would lead to the possibility of solvent entering the pocket from areas other than where the ligand is exiting, and lastly IV) preserving the local environment of the interacting atoms in the already prepared structures. The sequence gaps created during the process of selecting the chunk residues were capped. For this, each section of residues was split into separate chains, and the termini of each chain were acetylated or methylated. Lastly, the chunk was checked for clashes possibly created during the capping of the chains. The chunk included the

following residues: 853–859, 861–865, 879–882, 898, 902, 911, 912, 927, 929–941, 976, 978, 980–984 and 993–996. After production of the chunk, DUCK performs I) automatic ligand parameterization in MOE, II) minimization, III) equilibration, and IV) two SMD simulations (at two different temperatures, 300 K and 325 K), in which the distance between the interacting atoms in the ligand and protein is increased from 2.5 to 5.0 Å, and V) if the W_{QB} value (work necessary to break the H-bond) in the previous step reaches a pre-defined threshold, then the system is sampled by a short unbiased MD simulation, after which the resulting new structures are fed into steps IV) and V); the last two steps being repeated in a finite number of cycles (replicas).¹¹ Steps II) to V) were performed with GPU-based pmemd.cuda in AMBER. Here, up to five replicas of steps IV) and V) were performed, during which a W_{QB} threshold of 6 kcal mol⁻¹ (work necessary to break the H-bond) was used, so that the simulations were discontinued if the measured W_{QB} in any replica was below the threshold. If five runs were completed, the lowest obtained W_{QB} value was used, which resulted in 17 compounds that surpassed the threshold. Finally, we checked the novelty of these fragments against known kinase inhibitors by substructure searches, which ultimately led to the selection of compound **MR1** for experimental validation.

To suggest a binding pose for MELK, **MR1** was docked in all liganded MELK structures available. Both poses in Fig. 1B were identified and submitted to a DUCK simulation, pulling from the NH atom in the hinge (Cys 89 N). Chunks were derived by selecting residues within 10 Å from Cys89 N.

To check for known hinge binders in the fragment library, all kinase inhibitors found in the ATP binding site were downloaded from the KLIFS database.³² The cores (5 Ångstrom from the hinge NH group) were extracted from known binders from KLIFS and from the compounds in the fragment library. A substructure search was performed to find which of the fragment cores in the library were present in the known hinge binder cores.

Kinase inhibition assay

MR1 (purity 97% by LC/MS) was tested against the target kinases with the Z'-LYTE kinase inhibition assay (Life Technologies). The assay employs a fluorescence-based format and is based on the different sensitivities of phosphorylated and nonphosphorylated peptides to proteolytic cleavage. A suitable peptide substrate is labelled with two fluorophores, forming a FRET pair. After incubating the kinase + peptide + test compound mixture for an hour, a development reaction is carried out. Any peptide that was not phosphorylated by the kinase is cleaved, disrupting the resonance energy transfer between the FRET pair. The reaction progress is quantified based on the ratio of the detected emission at 445 nm (coumarin) and 520 nm (fluorescein), *i.e.* the ratio of cleaved *vs.* intact peptides. A more detailed description of the assay is available on the website of Life Technologies.³³ IC₅₀ values were determined



from 10-pt titration measurements (with duplicate datapoints) using the SelectScreen™ Biochemical Kinase Profiling Service available at Life Technologies.

NMR measurements

The human MELK protein (Q14680) DNA was obtained from OriGene and the kinase-uba domain (1–337) sequence was cloned into a modified pET vector encoding an N-terminal His-tag. The recombinant protein was expressed in *E. coli* BL21 pLysS cells, harvested by centrifugation and purified by immobilized metal affinity chromatography and anion exchange chromatography. ¹H and STD-NMR measurements were performed using a 600 MHz Bruker Avance III spectrometer equipped with a 5 mm cryo-TXI (¹H, ¹³C, ¹⁵N) probe with z-gradient at 298 K. The MELK protein and **MR1** were dissolved in a 10% (v/v) D₂O and H₂O mixture containing 20 mM Tris buffer (pH 8.0), 260 mM NaCl, 1 mM TCEP and 5% glycerol. Spectra were acquired with water suppression using excitation sculpting with the pulsed gradient scheme. For the ¹H and STD measurements, the MELK and **MR1** concentrations were 2.14 and 50 μM, respectively. As a reference, STD experiments were also performed without the target, containing the ligand species alone.

STD-NMR spectra were acquired using a series of 40 equally spaced 50 ms Gaussian-shaped pulses for selective saturation of the protein, with a total saturation time of 2 s and a 50 ms spinlock to suppress protein signals. The frequency of the on-resonance saturation was set at 1.0 ppm and the off-resonance saturation frequency was set at 40.0 ppm. A total of 2000 scans were collected for each pseudo-2D experiment.

Microscale thermophoresis binding assay

Microscale thermophoresis measurements were conducted on a NanoTemper Monolith NT.115 device, using the red fluorescence channel. MELK was stored and applied in a buffer consisting of 20 mM Tris, 260 mM NaCl, 1 mM TCEP and 5% glycerol, at a pH of 8.0. The **MR1** stock was prepared in DMSO and diluted into the protein buffer, with the final DMSO concentrations not exceeding 1%. We have applied the ligand displacement assay principle to detect the thermophoresis of the fluorescently labeled MELK reference ligand Kinase Tracer 236 (ThermoFisher, cat. no. PV5592) at increasing concentrations of **MR1**. A titration curve was acquired with serial 1:1 dilutions starting from 100 μM ligand concentration (with 0.56 μM protein, 50 nM Kinase Tracer 236, 60% LED power, 20% MST power), with 11 datapoints. Each datapoint was acquired in triplicate. The primary result was a relative EC₅₀ value of **MR1** against MELK (21.1 μM), which was converted to a K_i value of 5.47 μM, using the following formula:

$$K_i = \frac{EC_{50}}{\frac{[T]_{50}}{K_d} + \frac{[P]_0}{K_d} + 1} \quad (1)$$

In formula (1), [P]₀ is the protein concentration at 0% binding, [T]₅₀ is the concentration of free tracer at 50% and K_d is the dissociation constant of the tracer and protein, which was evaluated to be 208 nM under the assay conditions in a direct binding measurement.

Conclusions

Here, we report the discovery of a novel hinge binder using a new computational screening protocol based on dynamic undocking. 8-Amino-2*H*-isoquinolin-1-one (**MR1**) was found to inhibit kinases from four major branches of the kinase phylogenetic tree, with low to mid-micromolar potencies (translating to ligand efficiencies between 0.54–0.59). All of these kinases are relevant clinical targets, mostly for oncological indications. In particular, MST3 and MELK have few known potent inhibitors, presenting the opportunity to create novel chemical matter for these kinases with a less congested IP space. In addition to its versatile inhibitory profile, direct binding of **MR1** to MELK was confirmed by STD-NMR, and its binding to the ATP site was validated by an MST-based (microscale thermophoresis) ligand displacement assay developed as part of this work. Together, these findings nominate **MR1** as a viable fragment starting point for a range of type I kinase inhibitor programs.

Abbreviations

AGC	PKA, PKG and PKC containing families
ATP	Adenosine triphosphate
CAMK	Ca ²⁺ /calmodulin-dependent protein kinases
CMGC	CDK, MAPK, GSK3 and CLK containing families
DMSO	Dimethyl sulfoxide
Duck	Dynamic undocking
ERK2	Common alternative name for mitogen-activated protein kinase 1 (MAPK1)
JAK2	Janus kinase 2
LC/MS	Liquid chromatography/mass spectrometry
MD	Molecular dynamics
MELK	Maternal embryonic leucine zipper kinase
MST3	Common alternative name for serine/threonine-protein kinase 24 (STK24)
NMR	Nuclear magnetic resonance
RSK2	Ribosomal protein S6 kinase alpha-3
SMD	Steered molecular dynamics
STD	Saturation transfer difference
STE	Yeast sterile 7-, 11- and 20-homologue kinases
TK	Tyrosine kinases

Conflicts of interest

There are no conflicts to declare.



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