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An automatic pipeline for the design of irreversible derivatives identifies a potent SARS-CoV-2 M\textsuperscript{pro} inhibitor

Graphical Abstract

Covalentizer

In brief

Designing covalent inhibitors is a challenging task. Covalentizer is a computational pipeline for identifying irreversible inhibitors based on known non-covalent binders. Within the PDB, Zaidman and Gehrtz et al. identified >1,553 structures with covalent predictions, leading to the discovery of covalent kinase inhibitors and an inhibitor of SARS-CoV-2 M\textsuperscript{pro}.

Highlights

- Covalentizer was developed to suggest covalent analogs of non-covalent binders
- Experimentally validated against kinase targets as well as SARS-CoV-2 M\textsuperscript{pro}
- Application against the entire PDB uncovered numerous covalent opportunities
- A web server, as well as all the current predictions, are publicly available

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An automatic pipeline for the design of irreversible derivatives identifies a potent SARS-CoV-2 M\textsuperscript{pro} inhibitor

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SUMMARY

Designing covalent inhibitors is increasingly important, although it remains challenging. Here, we present covalentizer, a computational pipeline for identifying irreversible inhibitors based on structures of targets with non-covalent binders. Through covalent docking of tailored focused libraries, we identify candidates that can bind covalently to a nearby cysteine while preserving the interactions of the original molecule. We found ~11,000 cysteines proximal to a ligand across 8,386 complexes in the PDB. Of these, the protocol identified 1,553 structures with covalent predictions. In a prospective evaluation, five out of nine predicted covalent kinase inhibitors showed half-maximal inhibitory concentration (IC\textsubscript{50}) values between 155 nM and 4.5 \mu M. Application against an existing SARS-CoV M\textsuperscript{pro} reversible inhibitor led to an acrylamide inhibitor series with low micromolar IC\textsubscript{50} values against SARS-CoV-2 M\textsuperscript{pro}. The docking was validated by 12 co-crystal structures. Together these examples hint at the vast number of covalent inhibitors accessible through our protocol.

INTRODUCTION

Covalent irreversible inhibitors have become increasingly popular over the last decade as chemical probes and drugs. Most often these inhibitors target a cysteine residue to form the covalent bond. Several rationally designed irreversible inhibitors targeting cysteines were approved by the FDA in recent years, with notable examples, such as ibrutinib (Burger and Buggy, 2013), afatinib (Sequist et al., 2013), and osimertinib (Oxnard et al., 2016). Irreversible inhibitors offer a variety of advantages over non-covalent ones. These include: (1) prolonged residence time (Bradshaw et al., 2015), (2) an ability to compete with high-affinity natural substrates (Lonsdale and Ward, 2018; Michalczewski et al., 2008; Yun et al., 2008), (3) their improved selectivity when targeting non-conserved cysteine residues (Cohen et al., 2005; Ghosh et al., 2019), (4) targeting shallow binding sites (Sutanto et al., 2020), and (5) covalent binding can enable targeting of especially challenging targets such as the G12C oncogenic K-Ras mutation (Canon et al., 2019; Nnadi et al., 2018; Zeng et al., 2017).

Historically, most covalent inhibitors were designed by the addition of an electrophile to an already known reversible inhibitor that suitably binds next to a cysteine residue (Angst et al., 2020; Dubiella et al., 2015; Hagel et al., 2015; Vazquez-Rodriguez and Wright, 2019; Ward et al., 2015; Weisner et al., 2015). More recently, covalent inhibitors are also being discovered by empirical screening of covalent fragment libraries (Backus et al., 2016; Craven et al., 2018; Johansson et al., 2019; Kathman et al., 2014, 2015; Parker et al., 2017; Resnick et al., 2019) and by covalent virtual screening (Bensinger et al., 2019; Chowdhury et al., 2019; London et al., 2014; Nnadi et al., 2018; Rachman et al., 2019; Scarpino et al., 2018; Shraga et al., 2019; Toledo Warshaviak et al., 2014). While covalent fragment and virtual screening can potentially discover new scaffolds, the binding affinity of primary hits may be relatively low, and often require laborious medicinal chemistry to reach suitable potency.
Covalent derivatization of an already known reversible binder, can endow the compound with the added benefits of irreversible binding mentioned above, while retaining at least part of the binding energy from the reversible recognition, and possibly even have improved potency. Still, this approach is far from trivial. Three crucial questions have to be answered: (1) Which electrophilic moiety to use? (2) What is the optimal vector on the scaffold to attach through? (3) What linker, if any, would optimize the placement of the electrophile with respect to the binding mode of the scaffold and the position of the target residue? There are numerous possible answers for these questions. Furthermore, the covalentized (derivative containing the electrophile) version of the non-covalent inhibitor should be synthetically accessible. Therefore, tools that would be able to address this design problem algorithmically, would significantly simplify covalent inhibitor design and has the potential to discover many potent covalent binders for a large variety of targets.

Computational approaches to address this challenge are scarce. DUckCov (Rachman et al., 2019), a covalent virtual screening method, begins with non-covalent docking of a library of covalent compounds, while using pharmacophoric constraints for hydrogen bonds, as well as for the covalent warhead. This is followed by covalent docking of the ligands with the strongest non-covalent affinities. CovaDOTS (Hoffer et al., 2019) uses a set of synthetic schemes and available building blocks to create covalent analogs of existing non-covalent ligands, but was only assessed retrospectively. Cov_FB3D (Wei et al., 2020) constructs de novo covalent ligands and was retrospectively assessed on recapitulation on known covalent inhibitors.

Here, we present a computational pipeline to identify potential existing non-covalent binders for covalentization (creation of a covalent analog). Given a complex structure or model of a ligand in the vicinity of a cysteine residue, we elaborate the ligand or its substructures with various electrophiles. This ad hoc library of covalent analogs is covalently docked to the target protein and the original (non-covalent) structure is used as a filter to identify high-confidence covalent candidates. We applied this protocol—covalentizer—to the entire PDB to identify thousands of potential candidates amenable for irreversible inhibition, and made both the protocol and the database of pre-computed candidates publicly available to the community (https://covalentizer.weizmann.ac.il). We have prospectively synthesized and tested several predictions of various covalent kinase inhibitors proposed by the protocol and succeeded in five out of nine designs with half-maximal inhibitory concentration (IC50) values of between 155 nM and 4.2 µM.

In early February 2020, the COVID-19 pandemic started to spread globally (Wu et al., 2020; Zhu et al., 2020). We turned to the pre-compiled database of covalentizer results to look for possible candidate inhibitors for SARS-CoV-2 proteins. The search found a reversible small-molecule inhibitor designed against the main protease of the SARS-CoV virus (PD: 3V3M; Jacobs et al., 2013), which has 96% sequence identity to the main protease of SARS-CoV-2, with a promising covalent prediction. We synthesized the prediction and validated irreversible binding to the SARS-CoV-2 main protease (Mpro). We further optimized the non-covalent affinity of the compound, resulting in improved analogs. Co-crystal structures confirmed the computational model. This example highlights the strength of our method—the design was already available, and enabled very rapid development. The database suggests that hundreds more such examples await testing.

RESULTS

The covalentizer pipeline

For a given complex structure with a non-covalent ligand in the vicinity of a target cysteine residue, the pipeline (Figure 1) comprises four consecutive steps: fragmentation, electrophile diversification, covalent docking, and root-mean-square deviation (RMSD) filtering.

Fragmentation

In this step, the ligand is broken down and divided into two parts via synthetically accessible bonds (Lewell et al., 1998). Doing this recursively, results in a list of substructures (Figure 1A). For each substructure, we augment the list with its corresponding Murcko scaffold (Bemis and Murcko, 1996), which is the naked ring system, without any decoration, to allow more exit vectors from which the electrophile can be added next. The motivation for this fragmentation step is 3-fold. First, as mentioned, fragmenting the molecule exposes new vectors on which to install the electrophile (see Figure 1C, example 2). Second, the additional constraint of forming the covalent bond might cause a slight shift to the molecule’s binding mode from the original crystal structure. Such a shift may propagate and cause a steric clash between the protein and a ligand moiety distal to the electrophile. Since adding the covalent bond is expected to increase the overall potency, we “sacrifice” parts of the molecule to enable the addition of an electrophile. The final ranking of candidate covalent analogs relies on covalent docking, which is sensitive to sub-Å shifts. Hence, occasionally, a truncated version of the ligand will dock well, while the full ligand will not. Thus, including the substructures and their scaffolds maximizes the number of candidates. Since covalent docking accuracy was shown to deteriorate with ligand size and number of rotatable bonds (London et al., 2014), we filter the final list of substructures, to those with 8–25 heavy atoms and up to 5 rotatable bonds. We should note that, after successful docking of a fragment, additional parts of the reversible scaffold that do not interfere with electrophile installation can be added to the design.

Electrophile diversification

For each substructure or scaffold, we generate a library of potential electrophilic analogs, typically resulting in a few hundred analogs (Figure 1B). We consider four kinds of electrophiles ranging in reactivity—vinyl sulfonamides, chloroacetamides, acrylamides, and propynamides—that can all be installed in one step onto a free amine. We add these electrophiles to the substructures using simple connection rules, which, however, do not guarantee synthetic accessibility (see the STAR Methods for more details). We also consider various linkers between the fragment and the electrophile. In our application below, we considered either a methylene linker or various di-amine linkers (Figure S1).

Covalent docking

The structure of the complex is prepared for docking, using all available cysteine rotamers. We use DOCKvalent (London et al., 2014) to dock the electrophile library we described above against the protein (after removing the crystallographic ligand).
RMSD filtering
Compounds that are able to form a covalent bond with the target cysteine while still maintaining the same binding mode of the original reversible ligand are likely candidates for covalent analogs. To assess this, we evaluate the RMSD between each docking prediction and the crystallographic ligand. Due to the fact that RMSD is calculated between two sets of matching atoms, and the reversible ligand is different from the irreversible one, it was calculated based on the maximal common substructure (MCS) between the two molecules. Figure 1C exemplifies predictions with varying RMSDs. For a PDB-wide application of the pipeline, we focused on covalent analogs with a docking position of <1.5 Å RMSD from the crystallographic pose. These results suggest that covalentizer is able to rediscover a substantial portion of known covalent inhibitors, but at the same time emphasize the fact that the success of our protocol is dependent on the performance of the docking program, which is limited.

Covalentizing the PDB
Encouraged by the results in recapitulating known covalent kinase inhibitors we aimed to apply our protocol to the entire PDB. We started from the set of all the protein-small molecule X-ray structures (<3.0 Å resolution) that contained a small molecule with a molecular weight greater than 300 Da, and no DNA/RNA chains. We did not apply a maximum molecular weight cutoff, since the fragmentation process will address larger ligands. As of the date of the search (July 4, 2019) this resulted in 44,990 structures. We filtered these to structures in which a ligand has one of its atoms within 6 Å of the sulfur atom of a free cysteine residue. Disulfides or covalently modified cysteines were excluded. After applying this filter, we ended up with 8,386 such structures, and ~11,000 cysteines. These structures, which constitute the target space for our protocol, contain substantial redundancy. Clustering them with a threshold of 90% sequence identity, results in 2,227 representatives; 38% of the structures are of human proteins and the rest span many other organisms, including rodents, bacteria, and yeast. They also span seven different enzyme classes, with the most prevalent being transferases (41.4%). A total of 928 structures (11.1% of the entire dataset) are kinases. These ~8,400 proteins contain 3,673 different ligands, each binding next to a cysteine (Data S1B). The ligand that is most abundant in this
Covalentizer successfully recapitulates known covalent kinase inhibitors

Examples of covalent kinase inhibitors (green) for which covalentizer was able to find a substructure match (magenta) under the 1.5 Å threshold. (A) ERK2, PDB: 4ZZO (Ward et al., 2015). (B) EphB3, PDB: 5L6P (Kung et al., 2016). (C) EGFR (7290M), PDB: 4I24 (Gajiwala et al., 2013). (D) JAK3, PDB: 5TOZ (Telliez et al., 2016). The electrophiles span acrylamides (A, D), a substituted acrylamide (C) and chloroacetamide (B).

Exploring additional linkers

As mentioned above, the entire database was processed using direct attachments of the electrophiles to atoms of the substructures, as well as with a methylene linker. The use of longer and more diverse linkers for the addition of an electrophile would allow the targeting of cysteines further from the ligand, thus increasing the available target space, as well as diversifying the introduced chemistry. To investigate this further, we searched the covalent inhibitor discovery literature (Caldwell et al., 2019; Engel et al., 2015; Liang et al., 2017; Shi et al., 2019) for the most common di-amine linkers used in the last decade, which led to the selection of 9 aromatic linkers and 17 aliphatic linkers (Figure S1). Since including all of these linkers increases the computational demands of the pipeline, we restricted its application to the subset of liganded kinase structures in the PDB. Since these linkers can enable ligands to reach further cysteines at extended distances, the search criteria was extended to a distance of up to 10 Å from the ligand (instead of 6 Å, previously).

The final subset includes 1,880 PDB structures that contain a Cys residue of up to 10 Å away from one of 1,398 various ligands. The size of the custom-made libraries of electrophilic analogs for a particular reversible ligand, containing these linkers, now extends to a few thousand compounds. Overall, we generated in silico over 3 million electrophilic compounds with di-amine linkers for the kinase subset. The results show candidates of <1.5 Å MCS-RMSD between the original reversible ligand and the electrophilic candidate for 411 protein structures. A total of 186 of these structures (45%) was not found in the previous run, showing the potential of using these more sophisticated linkers to reach farther cysteines and to covalentize more ligands. Out of these 186, some target new cysteines that are further than 6 Å from the ligand, while others are due to better docking of the fragments to the same cysteine made possible by the new linkers.

Covalent inhibitors found for various kinases

Kinase inhibitors comprise 22% of the covalentizer results. We selected a subset of these for prospective validation. We chose the candidates based on three features: (1) low RMSD relative to the parent reversible ligand; (2) the addition of the electrophile is not predicted to interfere with the kinase hinge binding region; and (3) ease of synthesis. This required manual inspection of pre-selected low RMSD results. Overall, we made and tested nine compounds (Figure 4) targeting five different kinases. In some cases, addition of the electrophile required removal of large parts of the parent reversible ligand (Figure S2). The compounds were each tested in a kinase activity assay against the target kinase in the structure from which the covalentizer result was derived. The assay was performed at ATP concentration equal to the Km of the kinase in question, with a 2 h pre-incubation of the inhibitor at 25°C.

Four of the nine compounds did not show inhibition under the assay conditions ($IC_{50} > 10 \mu M$). Three compounds targeting ERK2 showed IC50 values of 2.9–4.52 μM. The remaining covalentizer hits included a 2.01 μM inhibitor (3) of FGFR4 derived from the non-selective kinase inhibitor ponatinib, and a 155 nM inhibitor (4) of GSK3β.

We focused on two of the ERK2 inhibitors (1 and 2) for further characterization. We assessed their irreversible binding to ERK2 by intact protein mass spectrometry (10 μM ERK2, 100 μM compound, 3 h incubation at room temperature). The expected protein–compound adducts were detected (100% and 85% labeling, respectively; peak-to-peak Δm 265–270 Da for both compounds; Figure 4B) with no additional adducts derived from multiple reactions, highlighting the moderate reactivity of
Figure 3. PDB-wide application of covalentizer identifies candidate irreversible inhibitors for more than 1,500 structures

(A) We filtered the PDB for structures that had only protein chains (no DNA/RNA), and contained a small molecule of at least 300 Da. This threshold was set to ensure some minimal initial fit/binding affinity to the target, as well as to filter out non-ligand small molecules, such as crystallization reagents. We used a PyMOL-based script to filter only the structures in which at least one ligand atom is <6 Å away from the sulfur atom of a cysteine residue. This cysteine also has to be free (no disulfide or other covalent modifications). After running the protocol and filtering only for results with <1.5 Å RMSD of the MCS between the reversible ligand and the covalent analog generated by covalentizer, there were 1,553 structures for which at least one such prediction was obtained.

(B) The top 1% of results have an RMSD under 0.5 Å; 23% are between 0.5 and 1 Å, and 76% are between 1 and 1.5 Å.

(C) The distribution of the four electrophiles used is balanced, with 29% between the reversible ligand and the covalent prediction was easily accessible by replacing a PyMOL-based script to filter only the structures in which at least one ligand atom is <6 Å away from the sulfur atom of a cysteine residue. This cysteine also has to be free (no disulfide or other covalent modifications). After running the protocol and filtering only for results with <1.5 Å RMSD of the MCS between the reversible ligand and the covalent analog generated by covalentizer, there were 1,553 structures for which at least one such prediction was obtained.

Beyond further optimization of the S1 and S2 pocket binding it appeared that relief of steric strain around the amide nitrogen played a part, since change to a methyl amide (in the Ugi reaction while keeping the acrylamide fixed). We explored a variety of replacements for the initial p-tert-butyphenyl motif protruding into the S2 pocket (Figure 5), most of them did not result in improved potency (Figure S2C). Similarly, independent optimization of binding to the S1 pocket only led to the identification of one beneficial change (23, IC₅₀ = 65.58 µM), with a meta chloro-substitution of the pyridine (Figure S2A). Other substituents (-Br, -OEt, -CF₂CH₃) led to inactive compounds.

To optimize 10, we have made and tested close to 140 analogs (Data S3; Figures 5 and S5), exploring all three components of the Ugi reaction while keeping the acrylamide fixed. We explored a variety of replacements for the initial p-tert-butyphenyl motif protruding into the S2 pocket (Figure 5), most of them did not result in improved potency (Figure S2C). Similarly, independent optimization of binding to the S1 pocket only led to the identification of one beneficial change (23, IC₅₀ = 65.58 µM), with a meta chloro-substitution of the pyridine (Figure S2A). Other substituents (-Br, -OEt, -CF₂CH₃) led to inactive compounds.

Beyond further optimization of the S1 and S2 pocket binding it was clear that extension of the ligand toward the S3 and S4 pockets should prove fruitful. For example, a reversible covalent α-ketoamide inhibitor (Zhang et al., 2020) (biocatalytic IC₅₀ = 0.67 ± 0.18 µM) probes the S3/4 region with an additional hydrogen bond to the backbone of Glu166. In a large-scale fragment screen, numerous fragments were able to bind in these pockets (Dauangamath et al., 2020). In this case, we exhaustively synthesized analogs of 10, using 34 available isocyanides. Starting from 10, simple alkyl chain extension resulted in compounds with improved potency (Figure 5D). Namely compound 11, harboring a phenethylamide motif, was particularly potent with an IC₅₀ of 2.95 µM (Figure 5G) and Kᵦ/Kᵦᵦ of 18.4 M⁻¹ s⁻¹ (Figure S4C). After chiral chromatography (see Methods S1) of the molecule, one enantiomer showed slightly better inhibition (IC₅₀ = 2.86 µM), while the other enantiomer was inactive (IC₅₀ = 86.32 µM; Figure S4D). A co-crystal structure of 11 with Mₚro showed that the fluorophenyl binds through hydrophobic interactions with Met165 and Gln189, explaining its improved potency (Figure 5C; Data S3).

It appears that relief of steric strain around the amide nitrogen also plays a part, since change to a methyl amide (in 12, relative to 10) also resulted in increased potency. Opposed to our initial assumption of independent optimization of S1–3 pocket binding,
the combination of beneficial structural motifs in a third generation of Ugi products led to inhibitors with diminished potency compared with 11 (see Figure S5B; Data S3). One explanation for this behavior is the high plasticity of Mpro, leading to induced fit effects.

Removal of the furanyl in ML188 and replacement with an electrophile in 10 initially led to a loss in potency, which in this case was overcome by optimization of the non-covalent affinity in the S3 region to give compound 11. Re-installing the furanyl ring in combination with the S3-optimized phenethylamide motif led to compound 17 with a similar IC50 (2.72 μM; Figure S4B), suggesting that the marked improvement of this side chain is particular to the covalently bound conformation.

In conclusion, we successfully executed a mode of action change toward irreversible targeting of the catalytic cysteine residue in Mpro, which may have improved activity in cells as well as long-term strategic benefits to safeguard against viral evolution.

**DISCUSSION**

Designing new covalent inhibitors is challenging. Here, we leveraged the subset of protein targets for which a structure of a known binder is available, to computationally enumerate and evaluate exhaustive sets of covalent derivatives. Automating the protocol allowed us to apply it to the entire PDB and assess the applicability of this approach. Prospective testing against six real-world targets demonstrated that irreversible ligands can be reached with little synthesis, and structures validated the binding-pose prediction.

A main advantage of our work is the wide exploration of X-ray structures, which produced an extensive list of candidates waiting to be explored. This allowed us to quickly find a promising lead series against SARS-CoV-2 Mpro. This prediction, which was based on a historic non-covalent SARS-CoV Mpro inhibitor (Jacobs et al., 2013), was pre-calculated and ready for synthesis at a moment’s notice. We have made thousands of such predictions available through a public website (https://covalentizer.weizmann.ac.il) that updates weekly with the release of new structures to the PDB. It also allows covalentizing of user-uploaded structures. The covalentizer was also integrated into PDBe-KB (Protein Data Bank in Europe-Knowledge Base) (PDBe-KB consortium, 2020), a community-driven resource collating functional annotations and predictions for structure data in the PDB archive. PDBe-KB allows understanding covalentizer data in the context of results from a diverse group of related bioinformatics resources and research teams. We believe these would enable wide application and experimental testing of new covalent inhibitors.

Despite the success of our protocol, several caveats remain. First is the fact that currently the protocol does not take into account the synthetic feasibility of the proposed designs. When selecting candidates for prospective evaluation, we found that some of the molecules required complicated synthesis. Incorporating into our pipeline a strategy, such as DOTS (Hoffer et al., 2018, 2019), other retrosynthesis algorithms (Delepine et al., 2018; Law et al., 2009; Watson et al., 2019) or even the use of synthetic feasibility scores (Ertl and Schuffenhauer, 2009; Huang et al., 2011; Podolyan et al., 2010), can improve the quality of proposed candidates in the future.

Another point for improvement is the relatively weak potency of our prospective designs in comparison with their parent compounds (Figure S2). One likely explanation for these lower
affinities is the removal of non-covalent affinity elements, which are not sufficiently compensated by the gains from covalent bond formation. For example, in compound 2 (derived from PDB: 4QTA) more than 350 Da of the original compound (Chai-kuad et al., 2014) is removed (Figure S2), resulting in three orders of magnitude loss in potency. However, the remaining covalent fragment, despite its very small size, still shows substantial inhibition of ERK2. Another example is compound 1, its parent compound (PDB: 4QP9) has an IC50 of 71 nM (Burdick et al., 2015); however, the propyl-pyrazole group we have omitted to accommodate the electrophile (Figure S2) improved the parent reversible binder by more than 150-fold. Even omitting only the propyl moiety from the parent compound already decreased its potency by \( \frac{1}{24} \) fold. A third example is compound 8, which did not show inhibition, while its parent compound (PDB: 5IHA) has an IC50 of 9 nM (Toure´ et al., 2016). Although keeping the hinge binding motif, we omitted groups from both ends, one due to clash with the added electrophile, and the other due to synthetic difficulty. While the same fragment was not tested by the authors of the parent compound paper, they did test a matched pair, showing that a single N to C substitution in the pyridine ring that we removed caused a decrease of 50-fold in potency, underscoring pyridine’s importance.

The acetyl versions of 1 and 2 we have made and tested corroborate this hypothesis. Both non-covalent compounds showed marked reduction in potency compared with the parent compound. 2-Ac became inactive and 1-Ac lost more than 40-fold potency, suggesting that the decrease in potency compared with the parent compound originated from the trimming of the propyl-pyrazole group rather than from the covalentization process itself. Finally, the loss of a hydrogen bond between the MPro backbone NH of Gly143 and the furanyl oxygen of ML188
(PDB: 3V3M), due to the replacement with the acrylamide, decreased potency, under our assay conditions of 15 min pre-incubation, by more than 30-fold. These results suggest that a careful examination of the binding energy contribution is required for the parts that are omitted to accommodate the new electrophile.

However, as we saw for both ERK2 (1 versus 5) and Mpro (10 versus 11), improving reversible recognition is able to improve potency, and even to surpass the parent compound in the case of 11. For this series we believe that further structure-based optimization of binding to the S3 pocket, including H bonding to Glu166, can pave the way for sub-micromolar Ugi-type covalent Mpro inhibitors. Thus, in many cases where irreversible binding is needed, to avoid resistance, for instance, or improve selectivity, our protocol can provide a promising starting point for optimization.

Another possible explanation for the relatively low affinity of the irreversible binders are slight inaccuracies in the covalent warhead positioning, which results in sub-optimal covalent bond formation. Perhaps due to the fact that the docking program does not take into account the actual formation of the covalent bond, and ignores, for instance, the transition state energy of the rate-determining step of the covalent bond formation, but rather evaluates the binding energy of the adduct. The performance of covalentizer is limited by the performance of the docking program that it uses. Having said that, the fragmentation technique does allow to find solutions that might not have been found using the full molecules (Data S1A). An obvious way to improve covalentizer is to improve the covalent docking program, or to replace it with a better performing one. Several programs have been reported retrospectively to perform better than DOCKvalent (Scarpino et al., 2018) and thus may be used to replace it in the protocol. We should also note that, while some successful covalent virtual screening prospective results are reported, these were typically a result of testing several predictions per target, whereas here we tested a very limited number of predictions per protein.

For the above reasons, we suspect that only a subset of the results that passed the 1.5 Å threshold are worth pursuing, depending on the position on which the electrophile is installed and whether the addition is mutually exclusive with an important part of the reversible recognition element of the original molecule. For example, while selecting kinase inhibitors, we only considered predictions for which the electrophile can be added without removing the hinge binding region, which is known to be crucial for binding (Xing et al., 2015). Another important aspect in selecting candidates for testing is the complementation of the docked fragment to include as much as possible of the reversible scaffold that is not mutually exclusive with the electrophile. The importance of this is demonstrated by the cases of 5 and 1, as well as in the case of 10, for which only a fragment of the molecule docked well but we complemented it to the full compound. In the future such considerations may also be automated.

It is interesting to note the similar activity of a methylene-chloroacetamide (1), compared with its acrylamide analog (6). Such electrophile replacements can be very useful in rational design of irreversible inhibitors, especially if they prove to work across various scaffolds. Geometrically, the additional methylene before the chloroacetamide makes the distance from the ring to the thiol similar to that of the acrylamide (Figure S2). In terms of reactivity, however, the acrylamide, conjugated to the azaindole is activated (Flanagan et al., 2014) and thus is likely closer in reactivity to the chloroacetamide. Indeed, a methylene linker would be the minimal linker element required to insulate against π-conjugation, allowing easier prediction of intrinsic reactivity. No-linker designs connected to extended π-systems, such as heteroarenes, often exhibit a range of intrinsic reactivities (Flanagan et al., 2014; Resnick et al., 2019), which remain challenging to predict computationally (Lonsdale et al., 2017; Palazzesi et al., 2019, 2020) and thus require careful evaluation.

Many additional designs remain to be discovered beyond the ones we made available through the covalentizer server. New electrophiles and linkers, which will enable new geometric trajectories between the cysteine and the molecule, can considerably expand the design space. We tested this idea computationally using a library of linkers curated from the literature (Figure S1), on a subset of kinases from our database, showing an increase in the number of structures that can be covalentized. Recent covalent “warheads,” including reversible covalent warheads, such as cyanoacrylamides (Bandyopadhyay and Gao, 2016), and chlorofluorooacetamides (Shindo et al., 2019), became available, both for cysteine residues (Backus, 2019), but also for other amino acids (Martin-Gago and Olsen, 2019; Ray and Murkin, 2019; Shannon and Weerapana, 2015). These can be incorporated with little effort into the covalentizer pipeline. Since cysteine is one of the least-abundant natural amino acids, additional covalent chemistries will substantially expand the number of ligands that can be potentially addressed.

In summary, we show that, using covalent docking, we were able to make irreversible analogs of ligands for which a complex structure is available. We made our discoveries public in the form of a database of the results we obtained by running our protocol on the entire PDB, which is automatically updated weekly with newly released entries, as well as a web-tool for applying the protocol on new targets given by users. Using the protocol, we discovered several covalent kinase inhibitors and optimized a potent covalent COVID-19 protease inhibitor, with a low-cost, modular, and fast synthesis. We hope our results will encourage researchers to apply covalent inhibitors for a wide range of targets.

**SIGNIFICANCE**

Covalent inhibitors have increased in popularity over the past decade, owing to several advantages over non-covalent ligands. However, their discovery and rational design is still challenging. By developing the covalentizer we show that many non-covalent small-molecule binders for which a structure is available, with a variety of proteins, have the potential to be transformed into covalent binders. We prospectively validated our protocol against various kinases, as well as in a rapid drug discovery campaign against the SARS-CoV-2 main protease. These results point to many other potential targets that may be targeted by a covalent analog of a known ligand, potentially shortening the time and effort in finding future covalent inhibitors. Our web server [https://covalentizer.weizmann.ac.il/](https://covalentizer.weizmann.ac.il/) allows users with little to no computational background to use our predictions.
for covalent analogs, as well as using the *covalentizer* on their own models. We hope that our results and server will facilitate a broad use of covalent inhibitors.

**CONSORTIA**

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**STAR METHODS**

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**SUPPLEMENTAL INFORMATION**

Supplemental information can be found online at [https://doi.org/10.1016/j.chembiol.2021.05.018](https://doi.org/10.1016/j.chembiol.2021.05.018).

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**AUTHOR CONTRIBUTIONS**

D.Z. and N.L. designed the algorithm. D.Z. implemented the algorithm, performed all computational experiments as well as ERK2 LC/MS experiments. P.G. led all organic syntheses, with help from M.F., D.F., and A.D., as well as ERK2 LC/MS experiments. F.J.D. performed MPro crystallography. R.G. performed ERK2 LC/MS and LC/MS/MS experiments. E.R. performed MPro LC/MS studies. J.P. implemented the webserver. D.O., C.S.-D., P.L., and M.A.W. took care of MPro protein production. The Covid-Moonshot Consortium contributed MPro-related reagents and protocols. D.Z., P.G., and N.L. wrote the manuscript. All authors contributed to the final form of the manuscript.

**DECLARATION OF INTERESTS**

N.L. is a member of the SAB of Tutas medicines, Monte Rosa Therapeutics and MetaboMed.
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## STAR METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Bacterial and Virus Strains** | | |
| E. Coli BL21(DE3)-R3-pRARE | Walsh Lab | N/A |
| **Chemicals, peptides, and recombinant proteins** | | |
| pGEX-6P-1-Mpro plasmid | Walsh Lab | N/A |
| **Deposited Data** | | |
| SARS-CoV-2 Mpro with LON-WEI-b8d98729-23 (Mpro-x2540) | This paper | PDB: 5RG7 |
| SARS-CoV-2 Mpro with LON-WEI-b8d98729-43 (Mpro-x2694) | This paper | PDB: 5RH5 |
| SARS-CoV-2 Mpro with LON-WEI-b8d98729-18 (Mpro-x2703) | This paper | PDB: 5RH6 |
| SARS-CoV-2 Mpro with LON-WEI-b8d98729-20 (Mpro-x2705) | This paper | PDB: 5RH7 |
| SARS-CoV-2 Mpro with LON-WEI-b8d98729-8 (Mpro-x2776) | This paper | PDB: 5RH9 |
| SARS-CoV-2 Mpro with LON-WEI-adc59df6-2 (Mpro-x3110) | This paper | PDB: 5RL0 |
| SARS-CoV-2 Mpro with LON-WEI-adc59df6-27 (Mpro-x3113) | This paper | PDB: 5RL1 |
| SARS-CoV-2 Mpro with LON-WEI-adc59df6-26 (Mpro-x3115) | This paper | PDB: 5RL2 |
| SARS-CoV-2 Mpro with LON-WEI-adc59df6-39 (Mpro-x3117) | This paper | PDB: 5RL3 |
| SARS-CoV-2 Mpro with LON-WEI-adc59df6-3 (Mpro-x3124) | This paper | PDB: 5RL4 |
| SARS-CoV-2 Mpro with LON-WEI-adc59df6-30 (Mpro-x3359) | This paper | PDB: 5RL5 |
| SARS-CoV-2 Mpro with LON-WEI-adc59df6-47 (Mpro-P0047) | This paper | PDB: 7NW2 |
| SARS-CoV Mpro with ML188 | (Jacobs et al., 2013) | PDB: 3V3M |
| Nitrate reductase from Ulva prolifera | (You et al., 2018) | PDB: SYLY |
| Human mineralocorticoid receptor | (Lotesta et al., 2016) | PDB: SHCV |
| Human progesterone receptor | (Williams and Sigler 1998) | PDB: 1A28 |
| ERK2 | (Ward et al., 2015) | PDB: 4ZZO |
| EphB3 | (Kung et al., 2016) | PDB: 5L6P |
| EGFR (T790M) | (Gajiwala et al., 2013) | PDB: 4I24 |
| JAK3 | (Telliez et al., 2016) | PDB: STQZ |

### Software and Algorithms

| Resource | Source | Link |
|----------|--------|------|
| Covalentizer | London Lab | https://covalentizer.weizmann.ac.il/ https://github.com/LondonLab/Covalentizer |
| PyMOL | Schrödinger | https://pymol.org/ |
| Prism | GraphPad | https://www.graphpad.com/scientific-software/prism/ |
| DOCKvalent 3.7 | (London et al., 2014) | http://dock.compbio.ucsf.edu/DOCK3.7/ |
| Illustrator | Adobe | http://adobe.com/ |
| OpenBabel | OpenBabel | http://openbabel.org/wiki/Main_Page |
| RDKit version 2018.09.3 | RDKit | RDKit.org |

(Continued on next page)
RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Nir London (nir.london@weizmann.ac.il)

Materials availability
Small molecules synthesized in this study can be custom synthesized following the synthetic procedures in the supplemental information. This study did not generate other new unique reagents.

Data and code availability
The code generated in this study is available at GitHub: https://covalentizer.weizmann.ac.il/
https://github.com/LondonLab/Covalentizer.
The Covalentizer database and webserver is available online: http://covalentizer.weizmann.ac.il/

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Microbe strains
The pGEX-6P-1-Mpro plasmid was transformed into a competent E. coli expression cell line based on a multi-phage resistant BL21(DE3) derivative, containing the pRARE2 plasmid for improved expression of proteins that utilise rare E. coli tRNA genes (BL21(DE3)-R3-pRARE). Multiple transformant colonies were used to inoculate a starter culture supplemented with 100 mg/ml Carbenicillin. The culture was then grown to log phase for ~8 h. Ten millilitres of the starter culture was used to inoculate one litre of auto induction medium supplemented with 10 ml of glycerol and 100 μg/ml carbenicillin. The cultures were grown at 37°C, 200 rpm for 5 h then switched to 18°C, 200 rpm for 10 h.

METHOD DETAILS

Programs and libraries
RDKit was used for 2D molecular handling, conformation generation and RMSD calculation. RDKit: Open-source cheminformatics; version 2018.09.3; RDKit.org. Marvin was used in the process of preparing the molecules for docking, Marvin 17.21.0, ChemAxon (https://www.chemaxon.com). OpenBabel (http://openbabel.org/wiki/Main_Page) was used to switch between molecular file formats. DOCKvalent (London et al., 2014) was used for virtual covalent docking. The Covalentizer code is available at https://github.com/LondonLab/Covalentizer.

Enumerating substructures for covalentization
Fragmentation and scaffold extraction was done using RDKit’s implementation of the Recap algorithm (Lewell et al., 1998) and the Murcko Scaffold (Bemis and Murcko, 1996) functionality respectively.

Covalentizing a substructure
For each substructure or scaffold, we generated a library of potential electrophilic analogs using SMARTS based reactions. The reaction rules were: 1. Adding an electrophile (including the nitrogen) to any non-substituted aromatic carbon, as well as all aliphatic...
carbon with one or two bonded atoms, excluding carbons which are already connected to nitrogen. 2. Adding an electrophile to a free amine, either primary or secondary. In this case the nitrogen is completed with the rest of the electrophile. The first rule will usually require more complicated synthesis, whereas the second rule, will allow to use the same ligand as a starting material for a nucleophilic substitution of the acyl form of the electrophile with the free amine.

Docking and RMSD calculation
RDKit and Marvin were used to create 250 conformations for each electrophilic analog. Covalent docking was done using DOCKovalent—a virtual screening program. We docked the appropriate analog library for each target, while saving 10 structures for each analog to increase the number of final candidates. When docking the larger linker based libraries we only used the top scoring structure for each analog, due to the large number of structures for analysis. Alternative rotamers for the cysteine residue were generated with pymol based scripts. DOCKovalent does not allow for protein flexibility, however, sampling the cysteine’s rotamers before docking does introduce some flexibility. Water molecules as well as any co-crystallised ligand were not taken into account during docking. We used RDKit to filter only for results with a MCS that has an RMSD of less than 1.5 Å to the original ligand. For this we used the FindMCS function with both FindMCSringMatchesRingOnly and completeRingsOnly attributes set to True.

Hardware and run time
Running Covalentizer on all suitable structures from the PDB was done using a PBS scheduler, on a computer cluster with 528 cores. The overall runtime was several days. Running Covalentizer on a single structure on a similar sized cluster takes only a few hours, and many structures can run in parallel.

Computational optimisation of the Mpro inhibitor
We used the RDKit reaction functionalities, as well as OpenBabel (http://openbabel.org/wiki/Main_Page) to prepare virtual libraries of analogs of compound 10. The Ugi reaction has three reactants: amine, isocyanide, aldehyde and carboxylic acid. The carboxylic acid is set constant to acrylic acid, since we didn’t want to change the electrophilic component. In the virtual libraries, we left it as the reversible furan moiety for convenience in modeling. We thus created three such libraries, each one by replacing one of the three other Ugi reactants with commercially available building blocks. Using RDKit, we generated up to 100 constrained conformations of each molecule, by fixing the conformation of three components as in the crystal structure, and changing only the conformation of the variable part. We then used the Rosetta modeling suite in order to choose the best conformation for each compound, when bound to the protease. For each molecule, we then defined this set of constrained conformations as an extra residue for Rosetta, and used Rosetta Packer (Kuhlman and Baker, 2000) to choose the best conformation, while allowing side-chain flexibility. Eventually, we chose analogs only for the amine and the isocyanide components, as the aldehyde component was highly optimised already. We chose 9 isocyanide replacements and 14 amine replacements (one of them was not based on docking). Most combinations of these components were made by Enamine and tested as part of the Covid-Moonshot effort (Chodera et al., 2020; Douangamath et al., 2020).

Intact protein LC/MS
The LC/MS runs were performed on a Waters ACQUITY UPLC class H instrument, in positive ion mode using electrospray ionization. UPLC separation used a C4 column (300 Å, 1.7 μm, 21 mm × 100 mm). The column was held at 40 °C and the autosampler at 10 °C. The mass data were collected on a Waters SQD2 detector with an m/z range of 2–3071.98. The desolvation temperature was 500 °C with a flow rate of 1000 L/h. The voltages used were 0.69 kV for the capillary and 46 V for the cone. Raw data was processed using openLYNX and deconvoluted using MaxEnt.

2 μM recombinant Mpro was incubated with 2 - 200 μM for 90 minutes in 50 mM Tris pH 8 300 mM NaCl in room temperature. 2 μM recombinant ERK2 was incubated with 100 μM for 3 hours in 50 mM Tris pH 8 300 mM NaCl in room temperature. 2 μl sample was diluted with 22 μl buffer + 6 μl 2.4% formic acid, and 10 μl were injected with the following gradient at 0.4 ml/min: 20% acetonitrile + 0.1% formic acid for 2 minutes, followed by a rise to 80% acetonitrile over 2.5 minutes, 0.5 minutes at 80% acetonitrile, fall back to 20% acetonitrile over 0.2 minutes, and 0.8 minutes at 20% acetonitrile. The mass data were collected at a range of 750-1750 m/z. Raw data was processed deconvoluted using a 37000-51000 kDa window, 1 Da/channel resolution.

LC/MS/MS
50 μl of 10 μM Recombinant ERK2 was incubated in Tris 50 mM pH = 8, 0.3 M NaCl with 100 μM compounds or DMSO. The compounds were then removed by methanol-chloroform precipitation of the protein. The dry pellet was dissolved in 40 μl of 50 mM Tris pH = 8 + 5% SDS and heated to 95 °C for 6 minutes. The concentration of the protein was estimated using BCA assay (using purified ERK2 heated in 50 mM Tris pH = 8 + 5% SDS as the standard). 50 pmol of each sample (~2 μg) were diluted to 20 μl with Tris 50 mM pH = 8 + 5% SDS, reduced with DTT (1 μl of 0.1 M in 5% SDS/Tris 50 mM pH = 8, 45 minutes 65 °C), cooled to room temperature, then
alkylated with 1 μl of 0.2 M iodoacetamide in water (30 minutes room temperature in the dark). The protein was then isolated and trypsinized on s-traps (Protifi) according to the manufacturer’s instructions. Triplicates were prepared for each molecule.

**LC/MS/MS running**
ULC/MS grade solvents were used for all chromatographic steps. Each sample was loaded using split-less nano-Ultra Performance Liquid Chromatography (10 kpsi nanoAcquity; Waters, Milford, MA, USA). The mobile phase was: A) H2O + 0.1% formic acid and B) acetonitrile + 0.1% formic acid. Desalting of the samples was performed online using a reversed-phase Symmetry C18 trapping column (180 μm internal diameter, 20 mm length, 5 μm particle size; Waters). The peptides were then separated using a T3 HSS nanocolumn (75 μm internal diameter, 250 mm length, 1.8 μm particle size; Waters) at 0.35 μL/min. Peptides were eluted from the column into the mass spectrometer using the following gradient: 4% to 30%B in 155 min, 35% to 90%B in 5 min, maintained at 90% for 5 min and then back to initial conditions. The nanoUPLC was coupled online through a nanoESI emitter (10 μm tip; New Objective; Woburn, MA, USA) to a quadrupole orbitrap mass spectrometer (Q Exactive HFX, Thermo Scientific) using a FlexIon nanospray apparatus (Proxeon). Data was acquired in data dependent acquisition (DDA) mode, using a Top10 method. MS1 resolution was set to 120,000 (at 200 m/z), mass range of 375-1650 m/z, AGC of 3e6 and maximum injection time was set to 60 msec. MS2 resolution was set to 15,000, quadrupole isolation 1.7 m/z, AGC of 1e6, dynamic exclusion of 45 sec and maximum injection time of 60 msec.

**LC/MS/MS analysis**
Analysis was done using MaxQuant 1.6.3.4. The sequence of ERK2 was used for the analysis. The digestion enzyme was set to Trypsin/P with a maximum number of missed cleavages of 2. Carbamidomethyl and the modification by the molecules were included as variable modifications on cysteine. The “Re-quantify” option was enabled. Contaminants were included. Peptides were searched with a minimum peptide length of 7 and a maximum peptide mass of 4,500 Da. “Second peptides” were enabled and “Dependent peptides” were disabled. The option “Match between runs” was enabled with a Match time window of 0.7 min and an alignment window of 20 min. An FDR of 0.01 was used for Protein FDR, PSM FDR and XPSM FDR. The triplicate measured for each compound (or for DMSO-treated protein) was analyzed separately. Following MaxQuant analysis, only fully cleaved peptides were quantified and cysteine-containing peptides that were not modified by either iodoacetamide or compound were ignored. The intensity for each peptide was calculated as the average of the three triplicates. If the intensity was zero for one of the replicates, the average of the two others was used, and if two replicates showed zero intensity the peptide was ignored. The intensities for the non-cysteine containing peptides were averaged for each data set and used to normalize the intensity of cysteine containing peptides.

**Biochemical assays**
While covalent inhibitors display time-dependent inhibition, determination of IC₅₀’s at a specific incubation time allows their evaluation. For kinase inhibition we determined all IC₅₀’s at 2h pre-incubation. For Mpro inhibition we determined all IC₅₀’s at 15 min. pre-incubation.

**Kinase activity assays**
Biochemical Kinase inhibition assays for compounds 1-9, 1-Ac and 2-Ac, were carried out at Nanosyn, Santa Clara. Test compounds were diluted in 100% DMSO using 3-fold dilution steps. Final compound concentration in assay ranged from 10 μM to 0.0565 nM. Compounds were tested in a single well for each concentration and the final concentration of DMSO in all assays was kept at 1%. Reference compound, Staurosporine, was tested in an identical manner. Compounds were preincubated in 25°C for 2 hours before the measurements, and the kinase reactions were then performed for an additional 3 hours. For ERK2, the kinase concentration was 0.25-0.35 nM, the ATP concentration was 25 μM. For MELK, the kinase concentration was 0.06 nM, the ATP concentration was 30 μM. For VEGFR2, the kinase concentration was 0.25 nM, the ATP concentration was 80 μM. For GSK3β, the kinase concentration was 0.09 nM, the ATP concentration was 10 μM. For FGFR4, the kinase concentration was 0.17 nM, the ATP concentration was 250 μM.

**Biochemical Mpro inhibition assay**
All UGI Compounds and in particular compounds 10-40 were seeded into assay-ready plates (Greiner 384 low volume 784900) using an Echo 555 acoustic dispenser, and DMSO was back-filled for a uniform concentration in assay plates (maximum 1%). Reagents for Mpro assay were dispensed into the assay plate in 10 μl volumes for a final of 20 μl. Final reaction concentrations were 20 mM HEPES pH=7.3, 1mM TCEP, 50 mM NaCl, 0.01% Tween-20, 10% glycerol, 5 mM Mpro, 375 mM fluorogenic peptide substrate ([5-FAM]-AVLQSGFR-[Lys(Dabcyl)]-K-amide). Mpro was pre-incubated for 15 minutes at room temperature with compound before addition of substrate. Protease reaction was measured continuously in a BMG PheraStar FS with a 480/520 ex/em filter set. Data was mapped and normalized in Genedata Screener.

Mpro CryoCrystallography Mpro protein was expressed and purified as discussed previously (Douangamath et al., 2020). The pGEX-6P-1-Mpro plasmid was transformed into a competent E. coli expression cell line based on a multi-phase resistant BL21(DE3) derivative, containing the pRARE2 plasmid. Transformant colonies were used to inoculate a starter culture supplemented with 100 μg/ml Carbenicillin. The culture was then grown to log phase for ~8 h. Ten milliliters of the starter culture was used to inoculate one litre of auto induction medium supplemented with 10 ml of glycerol and 100 μg/ml carbenicillin. The cultures were grown at 37°C, 200 rpm for 5 h then switched to 18°C, 200 rpm for 10 h. The cells were harvested by centrifugation and stored at ~80°C. For purification, cells
were resuspended in 50 mM Tris pH 8, 300 mM NaCl, 10 mM Imidazole, 0.03 μg/ml Benzonase. The cells were disrupted on a high-pressure homogeniser (3 passes, 30 kpsi, 4 °C). The lysate was clarified by centrifugation at 50,000 x g. The supernatant was then applied to a Nickel-NTA gravity column and washed and eluted with 50 mM Tris pH 8, 300 mM NaCl, and 25–500 mM imidazole pH 8. N-terminal His-tagged HRV 3 C Protease was then added to the eluted protein at 1:10 w/w ratio. The mixture was then dialysed overnight at 4 °C against 50 mM Tris pH 8, 300 mM NaCl, 1 mM TCEP. The following day, the HRV 3 C protease and other impurities were removed from the cleaved target protein by reverse Nickel-NTA. The relevant fractions were concentrated and applied to an S200 16/60 gel filtration column equilibrated in 20 mM Hepes pH 7.5, 50 mM NaCl buffer. The protein was concentrated to 30 mg/ml using a 10 kDa MWCO centrifugal filter device.

Apo Mpro crystals were grown using the sitting drop vapour diffusion method at 20 °C by adding 150 nl of protein (5 mg/ml in 20 mM Hepes pH 7.5, 50 mM NaCl) to 300 nl of crystallisation solution (11% PEG 4K, 6% DMSO, 0.1M MES pH 6.7) and 50 nl of seed stock prepared from initial crystal hits. 55 nl of a 100 mM compound stock solution in DMSO was added directly to the crystallisation drops using an ECHO liquid handler (final concentration 10% DMSO) and drops were incubated for approximately 1 hour prior to mounting and flash freezing in liquid nitrogen. Data were collected at Diamond Light Source on beamline I04-1 at 100K and processed using XDS (Kabsch, 2010) and either xia2 (Winter et al., 2013), autoPROC (Vonrhein et al., 2011) or DIALS (Winter et al., 2018). Further analysis was performed with XChemExplorer (Krojer et al., 2017): electron density maps were generated with Dimple (Keegan et al., 2015); ligand-binding events were identified using PanDDA (Pearce et al., 2017) (both the released version 0.2 and the pre-release development version (https://github.com/ConorFWild/pandda)); ligands were modelled into PanDDA-calculated event maps using Coot (Emsley et al., 2010); restraints were calculated with GRADE (Long et al., 2017); and structures were refined with BUSTER (Bricegne et al., 2011). Coordinates, structure factors and PanDDA event maps for all data sets are deposited in the Protein Data Bank under PDB IDs 7NW2, 5RG5, 5R55, 5RH5, 5RH6, 5RH7, 5RH9, 5RL0, 5RL1, 5RL2, 5RL3, 5RL4 and 5RL5. Data collection and refinement statistics are summarised in Data S3E. The ground-state structure and all corresponding datasets are deposited under PDB ID 5R8T.

QUANTIFICATION AND STATISTICAL ANALYSIS

No statistical analysis was performed. Experimental results are presented as means and with error bars indicating standard deviation over n=2 or 3 as indicated in the figure legends.