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Alkynes in covalent enzyme inhibitors: down the kinetic rabbit hole

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Chapter 2



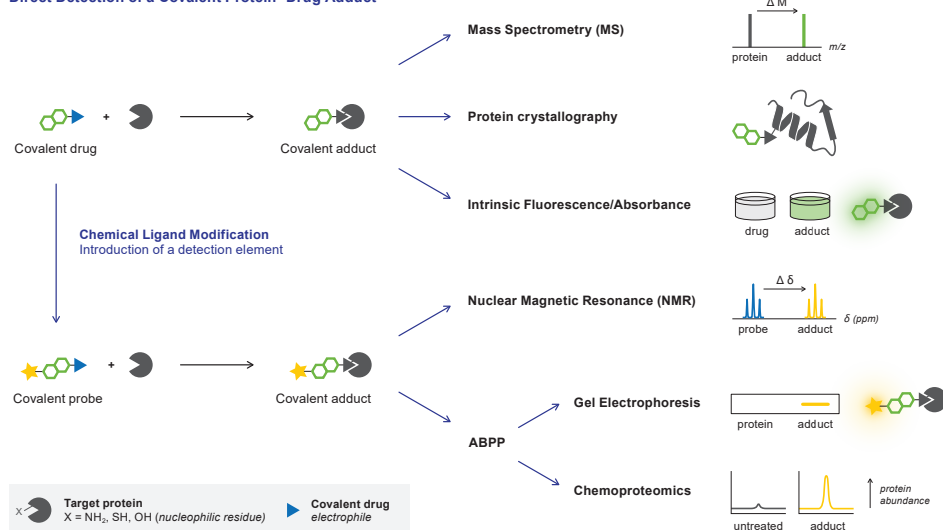
Technologies for Direct Detection of Covalent Protein–Drug Adducts

Adapted from:

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Abstract. In the past two decades, drug candidates with a covalent binding mode have gained interest of medicinal chemists, as several covalent anticancer drugs have successfully reached the clinic. As a covalent binding mode changes the relevant parameters to rank inhibitor potency and investigate structure-activity relationship (SAR), it is important to gather experimental evidence on the existence of a covalent protein–drug adduct. Here, we review established methods and technologies for direct detection of a covalent protein–drug adduct, illustrated with examples from (recent) drug development endeavors. These technologies include subjecting covalent drug candidates to mass spectrometric (MS) analysis, protein crystallography, or monitoring intrinsic spectroscopic properties of the ligand upon covalent adduct formation. Alternatively, chemical modification of the covalent ligand is required to detect covalent adducts by NMR analysis or activity-based protein profiling (ABPP). Some techniques are more informative than others and can also elucidate the modified amino acid residue or bond lay-out. Here, we will discuss the compatibility of these techniques with reversible covalent binding modes, and possibilities to evaluate reversibility or obtain kinetic parameters. Finally, we expand upon current challenges and future applications. Overall, these analytical techniques present an integral part of covalent drug development in this exciting new era of drug discovery.

Direct Detection of a Covalent Protein–Drug Adduct



1. Introduction

Among the most prescribed drugs in the US are successful drugs that were later found to have a covalent binding mode (**Figure 1A**),^{1–2} including established pain killer/anti-inflammatory agent **aspirin**,³ β -lactam antibiotic **penicillin**,⁴ anticoagulant **clopidogrel** (Plavix),⁵ and proton-pump inhibitor (**es**)**omeprazole** (Nexium) for gastroesophageal reflux.⁶ In the past two decades, the paradigm shift from covalent inhibition as an avoided liability toward the development of targeted covalent inhibitors (TCIs) has led to the approval of various drugs with a covalent binding mode (**Figure 1B**).^{1, 7–8} Covalent targeting of noncatalytic cysteine residues at the ATP-binding site of kinases has since proven to be a successful approach to overcome competition by the native substrate,^{9–10} as illustrated by clinically approved covalent Bruton's Tyrosine Kinase (BTK) inhibitors,^{11–13} and covalent (mutant) EGFR inhibitors.^{14–17} Furthermore, a covalent binding mode enabled inhibition of challenging targets for which noncovalent inhibitors could not successfully be developed, as illustrated by two recently approved first-in-class drugs: **sotorasib** (AMG 510) modifies Cys12 in the oncogenic KRAS^{G12C} mutant¹⁸ and **mobocertinib** (TAK788) modifies noncatalytic Cys797 of the EGFR^{ex20ins} mutant.¹⁹ An extensive overview of all FDA-approved drugs (1900–2019) with a known covalent mechanism of action has been compiled by De Vita,²⁰ listing their therapeutic application along with the electrophilic warhead. An update (2020–2022) can be found in **Table S1**.

Typically, a covalent adduct is formed when an electrophilic moiety (or warhead) in the inhibitor is positioned in juxtaposition of a nucleophilic residue in the protein target.^{21–22} Commonly targeted amino acid residues are catalytic cysteine and serine residues as the activated Cys thiolate and Ser hydroxylate are more nucleophilic (low pKa) than their noncatalytic (protonated) counterparts (Cys: pKa = 8–9, Ser: pKa > 13).²³ Popular noncatalytic nucleophilic residues include cysteines, lysines and (N-terminal) threonines.^{23–24} The selection of warhead depends on the identity of the amino acid residue, the nucleophilicity of the targeted amino acid residue, and the desired binding mode (reversible or irreversible).^{10, 24–25} A warhead should have the right balance between intrinsic chemical reactivity and selectivity, quickly forming a covalent adduct with the desired target but not (or much slower) with undesired cellular components.^{26–27} For cysteine-targeting inhibitors, this is typically assessed in indiscriminate thiol reactivity assays with biological thiols such as glutathione (GSH)²⁸ or cysteine.²⁹ The acrylamides and related Michael acceptors – employed in several approved kinase inhibitors – are among the most popular warheads for irreversible covalent targeting of noncatalytic cysteine thiols as the balance of their intrinsic chemical reactivity and selectivity results in a favorable safety profile.^{10, 20} Available warheads, popular as well as upcoming, and their application have been reviewed elsewhere.^{23–24, 30–31} Generally, development of novel TCIs entails introduction of a warhead onto a potent noncovalent scaffold,^{1, 21, 32–34} or high-throughput screening (HTS) of small molecule covalent ligands^{35–37} or covalent fragment libraries,^{31, 37–43} with structure-based lead optimization supported by *in silico* approaches (e.g. covalent docking, virtual screening).^{32, 44–46}

Reversible covalent inhibition is becoming increasingly popular^{47–53} as it combines the high affinity and long residence time of a covalent binding mode with a reduced risk of undesired idiosyncratic toxicity associated with the intrinsic ability to irreversibly modify off-target

proteins.⁴⁹ This approach is especially useful for targets with a relatively short cellular half-life as (proteasomal/proteolytic) degradation of the protein target will induce release of the reversibly bound covalent inhibitor that can engage in inhibition of another target protein. Introduction of an electron-withdrawing cyano group on the α -position of an irreversible covalent acrylamide warhead generates the cyanoacrylamide warhead, which was found to convert the inhibitors into reversible covalent inhibitors with a tunable residence time.⁵⁴⁻⁵⁵ The cyanoacrylamide moiety has gained popularity,⁵⁶⁻⁵⁸ most notably illustrated by reversible covalent BTK inhibitor **rilzabrutinib** (PRN-1008) currently in phase III clinical trials (**Figure 1C**).⁴⁹ Another recent example of the success of reversible covalent inhibition is **nirmatrelvir** (PF-07321332), the principle/novel component of Pfizer's oral antiviral agent Paxlovid that received emergency use authorization in 2021 for treatment of mild-to-moderate coronavirus disease (COVID-19).^{47, 59} **Nirmatrelvir** inhibits SARS-CoV-2 main protease M^{pro} by formation of a reversible covalent thioimide bond between an electrophilic nitrile warhead and the catalytic cysteine thiolate.⁴⁷

As a covalent binding mode changes the relevant parameters to rank drug potency and investigate structure–activity relationship (SAR),^{1, 60-63} it is important to gather experimental evidence on the existence of a covalent protein–drug adduct. Compounds sharing a warhead do not necessarily have the same covalent reactivity, and an electrophile is no guarantee for a covalent protein–drug adduct. Most claims pertaining a covalent binding mode are based on data obtained with the drug itself but there still are examples of clinically approved drugs for which the covalent binding mode is not explicitly demonstrated but assumed based on related compounds or covalent docking (e.g. **remdesivir**).⁶⁴ In this work we review the available methods in the toolbox to validate covalent adduct formation, rather than identification of novel covalent ligands/inhibitors (**Table 1**). Please note that the term inhibitor implies that target binding impairs protein function or blocks a protein–protein interaction, thus not reflecting covalent (partial) agonists⁶⁵⁻⁶⁷ and covalent PROTACs.⁶⁸⁻⁶⁹ We will use the more appropriate neutral term covalent ligand as it describes any covalent modifier without specifying how target engagement affects protein function/binding. We focus on technologies enabling direct detection of the covalent protein–ligand adduct under conditions that distinguish covalent adducts from noncovalent complexes (e.g. an increase in the total mass under denaturing conditions), while indirect covalent adduct detection protocols (e.g. competitive activity-based probe labeling) are occasionally mentioned to exemplify their use as orthogonal validation tools.

Most direct methods only discriminate between noncovalent and covalent protein modification, while others are more informative and provide direct evidence on which amino acid residue is modified. Although it is generally safe to assume that the most nucleophilic (catalytic) amino acid residue will be targeted for covalent modification, TCIs that were unexpectedly found to covalently modify allosteric (less nucleophilic) residues⁷⁰ or even a completely different amino acid⁷¹ illustrate why it is importance to identify the modified amino acid residue. Importantly, covalent adduct formation is not completed instantly upon treating the target protein with an excess covalent ligand.⁶¹ Unless otherwise noted, all procedures involve incubation of protein target and covalent ligand for a sufficient time (ranging from minutes to hours) to allow covalent adduct formation prior to analysis, as it is not possible to detect a covalent adduct

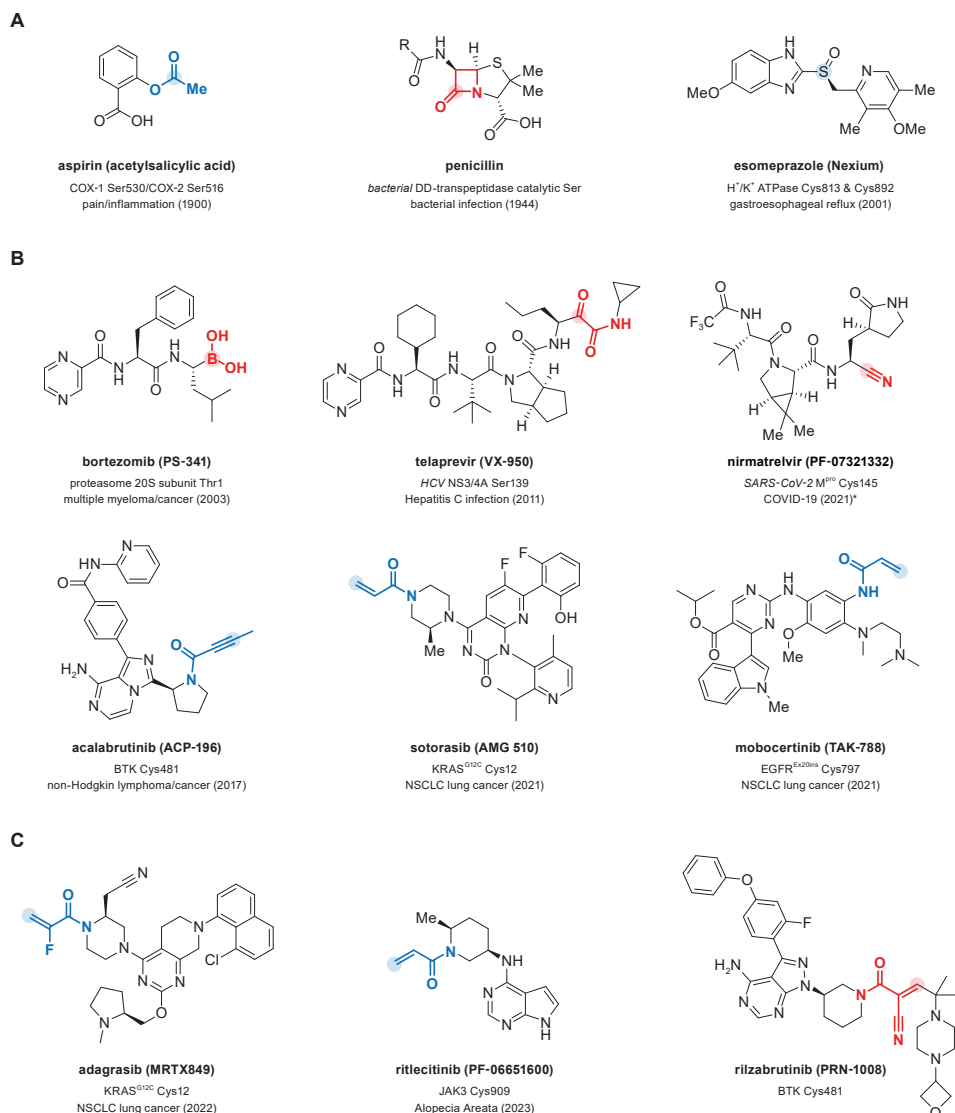


Figure 1 | Drugs with a known covalent binding mode. Structure with reversible warheads in red, irreversible warhead in blue, and the covalently modified atom marked. Compound name, protein target and covalently modified amino acid, therapeutic application and year of first approval. **(A)** Approved covalent inhibitors that were later found to have a covalent binding mode. Warhead not indicated for esomeprazole: metabolic activation of the prodrug precedes formation of a disulfide bond with its target.⁶ **(B)** Approved targeted covalent inhibitors (TCIs) designed to covalently modify their protein target. **(C)** Covalent inhibitors with a novel protein target and/or warhead.

that has not (yet) been formed. Conversion to covalent adduct does not have to be complete but high amounts of unbound protein can complicate detection, especially if unbound protein cannot easily be removed. The focus of this review is on the qualitative detection of covalent

protein–ligand adducts but some of the methods enable quantification of time-dependent covalent occupancy, which might be employed to calculate kinetic rate constants reflecting irreversible covalent inhibitor potency.^{1, 60–61} Details on the kinetic background of covalent adduct formation and potency are beyond the scope of this work,^{61–62} but compatibility with quantification of covalent occupancy will be highlighted. Moreover, special attention will be paid to compatibility with reversible covalent ligands and reversibility assays to assess the (ir)reversible ligand binding mode. Detection of reversible covalent adducts has its unique challenges compared to irreversible covalent adduct detection: detection (and purification) of the reversible covalent protein–ligand adduct is more complicated as the unbound enzyme and covalent adduct are at an equilibrium,⁵⁴ and the covalent occupancy is thus driven by the concentration of (excess) inhibitor.^{48, 61} Furthermore, standard sample preparation conditions (e.g. denaturation, proteolytic digestion, dilution), designed to induce noncovalent inhibitor dissociation, can also induce dissociation of reversible covalent ligands.^{54, 72} Traditional reversibility assays are based on regained enzymatic activity after rapid dilution⁷³ or washout,³⁴ or on detection of released unbound inhibitor upon protein denaturation/digestion⁵⁵ or chasing with a competitive irreversible ABP.⁷⁴ These assays serve to evaluate the reversibility of the adduct formation but irreversible protein modification provides by no means direct evidence of covalency: a covalent drug can have a reversible binding mode, and noncovalent binders can be irreversible.⁶¹

In this work, we will discuss methods for direct detection of covalent protein–ligand adducts (an overview of the methods can be found in **Table 1**). In general, whether the covalent adduct will be detected using a certain technique depends on intrinsic properties of the protein target (e.g. mass, ionizability, crystalline) as well as the inhibitor/ligand (e.g. binding mode, solubility, fluorescence). Each method will be illustrated with examples of advantages and limitations, with specific attention to compatibility with reversible covalent inhibition, identification of the modified amino acid residue and application in (kinetic) evaluation of inhibitor binding mode and/or potency. We start with techniques to detect the covalent protein–drug adduct without chemical modification of the ligand, using the same compound stocks prepared for biochemical *in vitro/in vivo* assays. Predominantly used techniques mass spectrometry (*section 2*) and protein crystallography (*section 3*) will be discussed first, followed by less ubiquitous detection based on the changes in intrinsic spectroscopic properties of the ligand upon covalent adduct formation (*section 4*). Alternatively, ¹³C NMR analysis (*section 5*) and activity-based protein profiling (*section 6*) require chemical modification of the covalent ligand to enable detection of the covalent protein–drug adduct (e.g. introduction of a bioorthogonal handle, reporter tag or isotope labeling). Finally, we expand on current challenges and future applications (*section 7*).

2. Mass Spectrometry (MS)

Initial confirmation of a covalent binding mode is predominantly achieved through mass spectrometric (MS) analysis of the covalent protein–ligand adduct.³⁶ Here, validation of the covalent binding mode is based on the mass increase upon modification of an unbound protein with a covalent ligand, compared to the mass of the unbound protein (**Figure 2**). MS analysis provides confirmation of the biophysical binding event between the protein and ligand but

Table 1 | Technologies for direct detection of covalent protein–drug adducts included in this review.

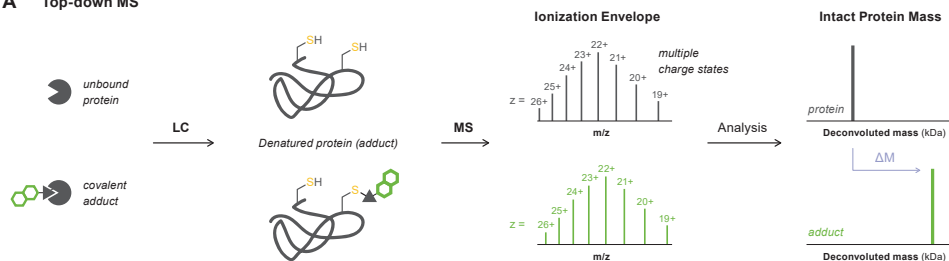
Detection Method	Prerequisites		Compatibility		Structural		Characterization		Notes
	Ligand Resynthesis	Protein Optimization	Reversible Covalent	Whole proteome ^a	Modified Amino Acid ^b	Bond Isoform	Covalent Occupancy	Reversibility	
2. Mass Spectrometry									Relatively fast and easy. Bottom-up MS(/MS) is compatible with large proteins (in mixtures)
2.1. Top-down MS	–	+	+	~ ^c	–	–	+	+	
2.2. Bottom-up MS	–	–	~	+	~ ^d	–	~	–	
2.3. MS/MS	–	–	~	+	+	–	+	–	
3. Protein Crystallography	–	+	++	–	+	+	–	–	Most informative but laborious
4. Intrinsic Fluorescence/Absorbance	– ^e	–	+	–	–	–	~	+	Limited ligand compatibility
5. Nuclear Magnetic Resonance	+	~	++	+	~	++	+	+	Compatible with labile adduct detection in solution
6. Activity-Based Protein Profiling									Detection of modified (off-)target proteins in whole proteomes
6.1. Gel Electrophoresis	+	–	~	++	–	–	+	+	
6.2. Chemoproteomic Platforms	+	–	~	++	+	–	~ ^f	+	
6.3. Homogeneous/plate-based	+	–	+	++	–	–	+	–	

^a Adduct formation in complex mixtures (e.g. lysates, live cells or *in vivo*). ^b Direct detection, not including indirect identification through site-directed mutagenesis of the modified amino acid. ^c Requires enrichment for (modified) protein target. ^d Identification of the peptide containing the modified amino acid residue. ^e Prerequisite for drug ligand class: covalent adduct formation must induce a change in intrinsic spectroscopic properties (e.g. fluorescence, absorbance). ^f Typically assessed in (indirect) competition assays.

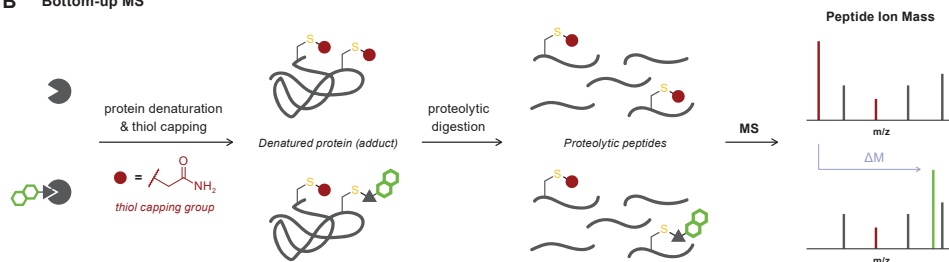
does not elucidate the bond lay-out. MS analysis is generally favored because it consumes a relatively low amount of material and it is compatible with most protein targets. This versatile technique is not only used to validate a covalent binding mode, but also in the discovery of new covalent ligands.^{41–42} Detailed guidelines for mass spectrometric characterization (and quantification) of covalent protein–drug (metabolite) adducts are available elsewhere.^{36, 75–76} Generally, the covalent adduct is formed by incubation of protein with excess inhibitor in an MS-compatible buffer, followed by a purification step such as liquid chromatography (LC) or gel electrophoresis.^{76–77} MS analysis must be performed for the covalent adduct as well as unbound protein, to confirm that the detected mass increase corresponds with covalent ligand modification. For top-down MS analysis (*section 2.1*), the adduct is separated from the unbound protein/inhibitor under denaturing conditions to ensure all noncovalent interactions are disrupted prior to MS analysis of the intact protein–inhibitor adduct (**Figure 2A**). Alternatively, the adduct is submitted to proteolytic digestion with bottom-up MS analysis (*section 2.2*) of the protein-derived proteolytic peptides to identify the peptide sequence modified by an

irreversible covalent ligand (**Figure 2B**). Subsequent peptide ion fragmentation for MS/MS analysis (*section 2.3*) can enable identification of the modified amino acid residue (**Figure 2C**).

A Top-down MS



B Bottom-up MS



C Tandem MS

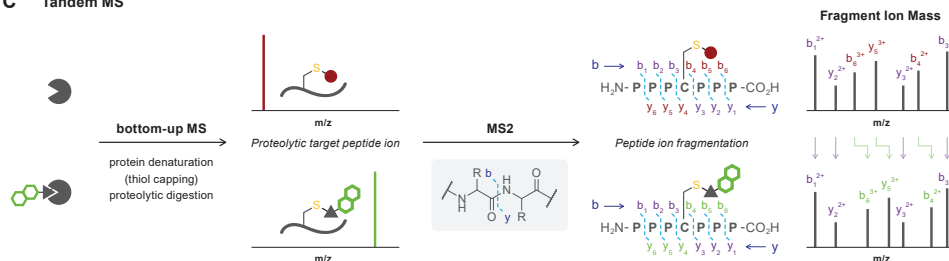


Figure 2 | Schematic overview of MS-based methodologies for detection of covalent protein–drug adducts. **(A)** Intact protein analysis by top-down MS analysis. Samples containing unbound protein (*top*) or covalent adduct (*bottom*) are resolved by liquid chromatography (LC) to promote ligand dissociation in noncovalent complexes and remove free ligand. Intact protein and covalent adduct are subjected to MS analysis, where they are ionized multiple times ($z \geq 1$) generating an ionization envelope originating from the various charge states, from which the deconvoluted total mass is calculated. The covalent protein–drug adduct has a higher deconvoluted mass than the unbound protein. **(B)** Bottom-up MS analysis. Samples containing unbound protein (*top*) or covalent adduct (*bottom*) are subjected to proteolytic digestion, with optional capping of free thiols using thiol-reactive reagent iodoacetamide (IAC) before or after digestion, followed by MS analysis. Proteolytic peptide ions originating from unmodified protein sequences are identical in both samples whereas different mass is observed for peptide ions containing the covalently modified amino acid residue. **(C)** Tandem MS or MS/MS. Following bottom-up MS analysis, proteolytic peptide ions (MS1) are exposed to fragmentation conditions that break the amide bonds, producing one out of two possible fragment ions for each broken peptide bond. The resulting fragment ions are annotated with increasing numbers from the N-terminus (b-fragment ions) or C-terminus (y-fragment ions). A mass difference is observed for fragment ions (MS2) containing the modified amino acid, thereby aiding identification of the modified amino acid residue.

2.1. Top-Down MS

It should not be surprising that intact protein analysis by top-down MS is the most popular technique to validate covalent adduct formation with a wide variety of targets:⁶³ most (academic) drug discovery labs are equipped with an LC-MS system (**Figure 2A**), and sample preparation is relatively straightforward when the protein–ligand adduct is formed using recombinant purified protein. Benchmark protocols are composed by Donnelly and co-workers for intact protein analysis by top-down MS.⁷⁷ Generally, the unbound protein and protein (adduct) are ionized after denaturation and removal of unbound ligand on the LC, generating (positively or negatively) charged ions ($z > 1$) detectable by MS. The total mass of the parent protein or adduct is calculated by deconvolution of the charge states in the ionization envelope. It is important to note that sample preparation is conducted under denaturing conditions that ensure noncovalent interactions are disrupted, but detection of noncovalently bound protein–ligand complexes is theoretically possible with native MS, with dedicated conditions to ensure noncovalent interactions are maintained.^{78–79} The main practical limitations to intact protein analysis by top-down MS are the incompatibility with larger proteins (>50 kDa), proteins that ionize poorly,^{80–81} proteins that require MS-incompatible detergents or surfactants, and complex (cellular) mixtures that have not been enriched for the protein target: bottom-up MS analysis (section 2.2) might be more suitable as ionization of peptides is often better.

Intact protein analysis by top-down MS is one of the less informative methods as it does not reveal the bond layout or identifies which amino acid residue is covalently modified. Mitigation of covalent adduct formation by site-directed mutagenesis provides (indirect) evidence on the modified amino acid residue, and is part of most covalent drug development workflows.⁶³ Intact protein analysis has recently been employed to validate covalent adduct formation with proteases,^{82–85} recombinant kinase domains,^{74, 86–88} and other (potential) clinical targets.^{89–91} Biophysical confirmation of covalent binding is also an important step in the ongoing industrial efforts to develop covalent kinase inhibitors with an improved selectivity/potency profile, as illustrated by intact protein analysis of covalent adducts between the BTK kinase domain and clinical candidates **evobrutinib** (Merck),⁸⁶ **remibrutinib** (Novartis),⁸⁷ and **tirabrutinib** (Ono Pharmaceutical/Gilead Sciences).⁸⁸

Covalent Fragment-Based Drug Development (FBDD). Intact protein analysis has a prominent role in target-directed covalent fragment-based ligand discovery (FBLD). MS analysis is utilized to identify (cysteine-reactive) covalent ligands, that serve as a starting point for medicinal chemistry optimization after validation of inhibitory properties associated with biophysical binding.^{36, 38–39} Kathman and co-workers developed an MS-based assay to screen mixtures of fragments containing a vinyl methyl ester (VME) warhead for covalent adduct formation with cysteine protease papain.^{42, 85} This assay has since successfully been employed to screening cysteine-reactive covalent fragment libraries with mixed electrophile chemotypes,^{41, 92} covalent ligand identification for E3 ligases,^{90–91} and several other recombinant protein targets.^{38, 93}

Reversible binding mode. The reversible covalent adduct of **odanacatib** (ODN, MK-0822) bound to recombinant cysteine protease CatK can be detected by top-down MS analysis (**Figure 3A**),⁸³ provided that inhibitor concentration exceeds its steady-state

equilibrium constant K_i^{app} to ensure sufficient covalent occupancy.⁶¹ The reversible covalent cTnC-levosimendan adduct was detected in endogenous thin and thick filament proteins extracted from porcine cardiomyofibrils.⁹⁴ However, sample preparation and denaturing conditions can induce inhibitor dissociation, and detection of the reversible covalent protein–inhibitor adduct is not possible if the covalent dissociation rate is relatively fast. Incubation of RSK2 with cyanoacrylamide **CN-NHiPr** failed to produce a detectable adduct,^{54, 72} but this is highly context-dependent as covalent adducts with other cyanoacrylamides have since successfully been detected by top-down MS.^{57, 74}

Reversibility assays. The Rauh group developed a top-down MS-based reversibility assay,⁷⁴ illustrated for reversible covalent EGFR inhibitors bearing a cyanoacrylamide warhead (**Figure 3B**). EGFR kinase domain and reversible covalent inhibitor (CRI) are incubated to form the covalent EGFR–CRI adduct, followed by incubation with excess chaser COV2 – an irreversible covalent ligand selectively targeting the same amino acid residue (e.g. **osimertinib**, **ibrutinib**) – that displaces the reversible covalent inhibitor, forming a covalent EGFR–COV2 adduct. Top-down MS analysis reveals a deconvoluted mass corresponding to the EGFR–COV2 adduct. Displacement is indicative of a reversible binding mode, as it is not possible to displace an irreversibly bound covalent inhibitor. Prerequisites to this reversibility assay are the availability of a selective irreversible chaser targeting the same amino acid residue, that forms a covalent protein–chaser adduct with a mass difference to the protein–inhibitor adduct. This method has since been employed in the preclinical development of irreversible covalent BTK inhibitor **tirabrutinib** (ONO-4059),^{88, 95} which was resistant to chasing with **ibrutinib**. Indirect methods with MS detection of released free ligand upon induction of ligand dissociation (e.g. dilution, dialysis, washout, denaturation, competition) will not be further discussed here.⁵⁴

Quantification covalent occupancy. The research groups of House (Crick–GSK Biomedical LinkLabs) and Rittinger (Francis Crick Institute) recently reported a quantitative covalent occupancy assay for kinetic analysis of irreversible ligand binding to the RBR domain of HOIP, an RBR E3 ubiquitin ligase for which quantitative HTS activity assays are not available (**Figure 3C**).⁹⁰ Time-dependent covalent occupancy was calculated from the total ion count (TIC) of the deconvoluted mass of covalent HOIP–fragment adduct relative to the unbound HOIP. LC-MS approaches were also employed to assess the potency of covalent KRAS^{G12C} inhibitors.^{36, 96–97} Differences in ionization efficiency of the unbound protein and adduct are only a minor concern as the protein size is significantly larger than the covalent ligand. Alternatively, indirect methods based on quantification of remaining unbound inhibitor (excess protein)⁹⁶ or unbound protein (excess inhibitor)^{89, 98} are employed to assess the biochemical rate of covalent target modification.

2.2. Bottom-Up MS

Bottom-up MS analysis (**Figure 2B**) is the preferred method to verify covalent adduct formation with large proteins (>50 kDa), proteins that are poorly ionized, and for detection of covalent adducts in complex mixtures (e.g. cell lysates and samples from living organisms). A comprehensive overview of bottom-up MS methodologies is available elsewhere.⁷⁶ Generally, the protein–ligand adduct and the unbound protein are subjected to a thiol-alkylating reagent

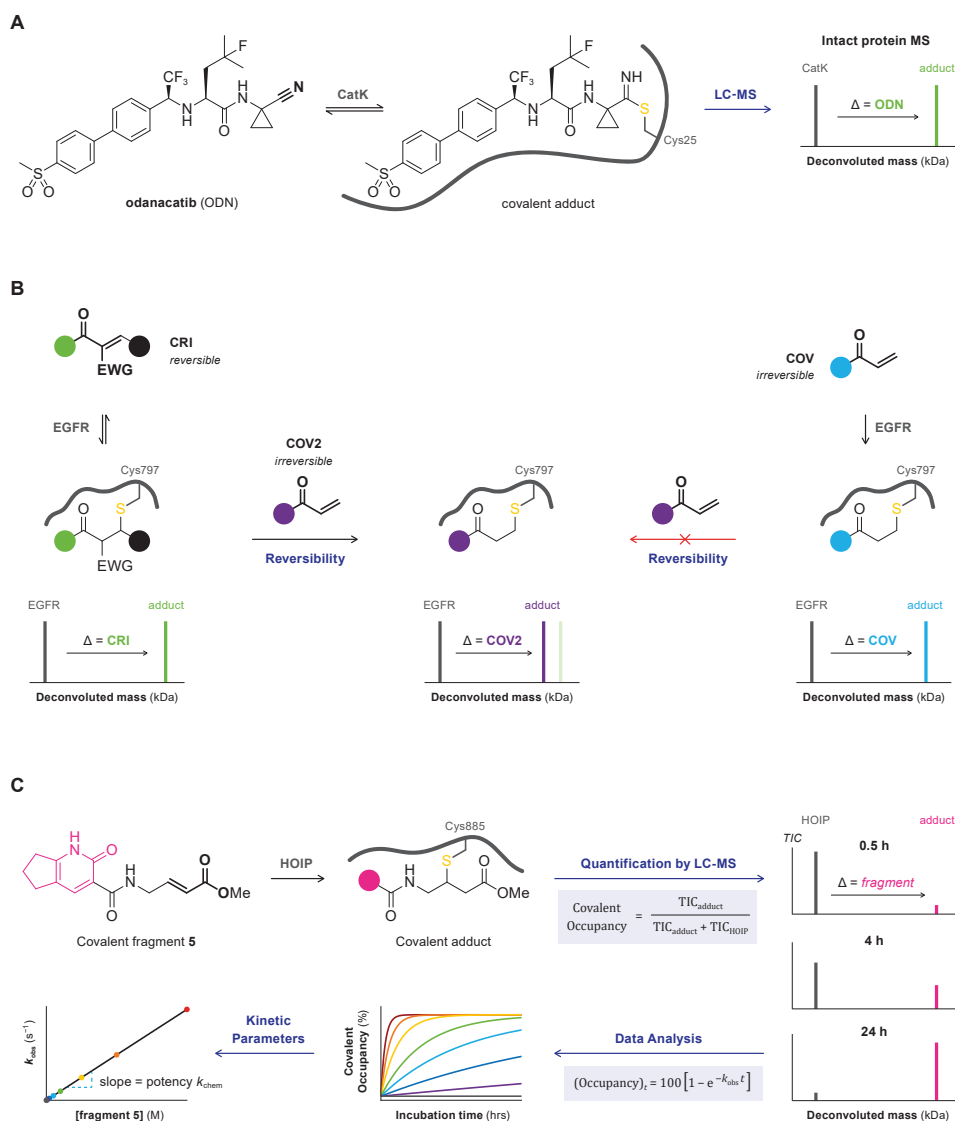


Figure 3 | Biophysical covalent adduct detection with intact protein analysis by top-down MS. **(A)** Covalent CatK–ODN adduct formation between recombinant purified cathepsin K (CatK) and reversible covalent CatK inhibitor odanacatib (ODN) is confirmed by intact protein analysis:⁸³ the higher deconvoluted mass for the adduct compared to unbound CatK is in agreement with covalent ODN binding. **(B)** MS-based reversibility assay illustrated with recombinant purified EGFR^{T790M/L858R} mutant and reversible cyanoacrylamide-based inhibitor CRI.⁷⁴ Detection of covalent EGFR–COV2 adduct rather than reversible covalent EGFR–CRI adduct upon competition with irreversible covalent chaser COV2 is indicative of a reversible binding mode. **(C)** Quantitative biochemical covalent occupancy assay illustrated for incubation of recombinant purified HOIP(RBR)^{WT} protein with excess covalent fragment 5.⁹⁰ Adduct formation is detected by top-down MS and covalent occupancy is quantified from the total ion count (TIC) of covalent adduct and unbound HOIP(RBR). Biochemical rate of covalent target engagement k_{obs} is calculated for each fragment concentration from the time-dependent covalent occupancy, which can be used to calculate the kinetic rate constant reflecting binding efficiency of an irreversible covalent fragment (more details in reference^{60–61}).

such as **iodoacetamide** (IAc) to cap free cysteine thiols (and sometimes lysine amines) with a carbamidomethyl group (+57.021 u), followed by trypsin- or pepsin-mediated proteolytic digestion (other proteases are also possible).⁷⁶ The proteolytic peptides are separated by LC, ionized and the peptide ions are detected by MS (**Figure 4**). Each parent peptide will be charged once or multiple times ($z \geq 1$) to generate ionized peptides, and a database is used to correlate found m/z values with the predicted mass of various amino acid stretches. In the protein–ligand adduct, a peptide with a covalently modified amino acid residue appears along with a decrease or even disappearance of the (capped) unmodified peptide with the same sequence. Consequently, not only the covalent adduct is validated, but the peptide sequence containing the modified amino acid is also identified.⁷⁶ Optionally, sample preparation may involve purification by gel electrophoresis prior to capping and proteolytic digestion, to remove unbound ligand and MS-incompatible buffer components (e.g. surfactants, detergents), and enrich the sample for the desired protein (adduct). Proteolytic digestion ensures that only stable covalent adducts are detected but these harsh conditions also have a drawback: incompatibility with sensitive/labile functional groups. LC-MS detection of unbound inhibitor after digestion-induced inhibitor dissociation is commonly used to assess binding reversibility,^{49, 55} but does not involve direct detection of the covalent adduct.

Bottom-up MS analysis is compatible with complex mixtures and native systems. For example, covalent adduct formation of clinically approved covalent KRAS^{G12C} inhibitor **sotorasib** (AMG 510) in (*in vitro* or *in vivo*) treated tumor cells was detected (**Figure 4A**).^{18, 99} RAS proteins were isolated from lysates by immunocapture on anti-RAS beads, eluted proteins were denatured (8M urea), free thiols were alkylated with IAc, and proteins were digested with trypsin prior to mass spectrometric analysis. Bottom-up MS analysis has been employed to validate covalent adduct formation with oncogenic KRAS^{G12C},^{18, 99–100} proteases,^{83–84} and various other (potential) clinical targets.^{70, 90}

The detected mass of the modified proteolytic peptide ions can correspond to the simple adduct (peptide + full ligand) but can be smaller if the ligand contains bonds sensitive to proteolysis (e.g. amide bonds), as is frequently seen for binding of ubiquitin(-like modifiers).⁸² Furthermore, covalent modification can block the proteolytic cleavage site, resulting in a missed cleavage and larger peptide sequences in the ligand-treated sample compared to the untreated (free protein) sample. This is exemplified by bottom-up MS analysis of the covalent adduct of SUMO-activating enzyme (SUMO E1 or SAE) with inhibitor **COH000** (**Figure 4B**):⁷⁰ pepsin-mediated proteolytic peptide A14–L32 was found in the untreated sample, but covalent modification of Cys30 in the COH000-treated sample blocked access to the pepsin cleavage site after Leu32, resulting in the simple COH000 adduct of peptide A14–N35 (missed cleavage). More importantly, bottom-up MS analysis was instrumental in the initial identification of the unexpected allosteric Cys30 modification: COH000 was expected to modify the catalytic (nucleophilic) Cys173 residue but only the unmodified A131–C185 peptide was found. The importance of careful interpretation of MS data is further illustrated by the work of Pettinger and co-workers.⁷¹ Their covalent acrylamide ligand was designed to covalently target Cys17 in stress-inducible ATPase molecular chaperone heat shock 70 kDa protein 1 (HSP72) but no evidence of Cys17 labeling was found after proteolytic digestion and subsequent MS analysis

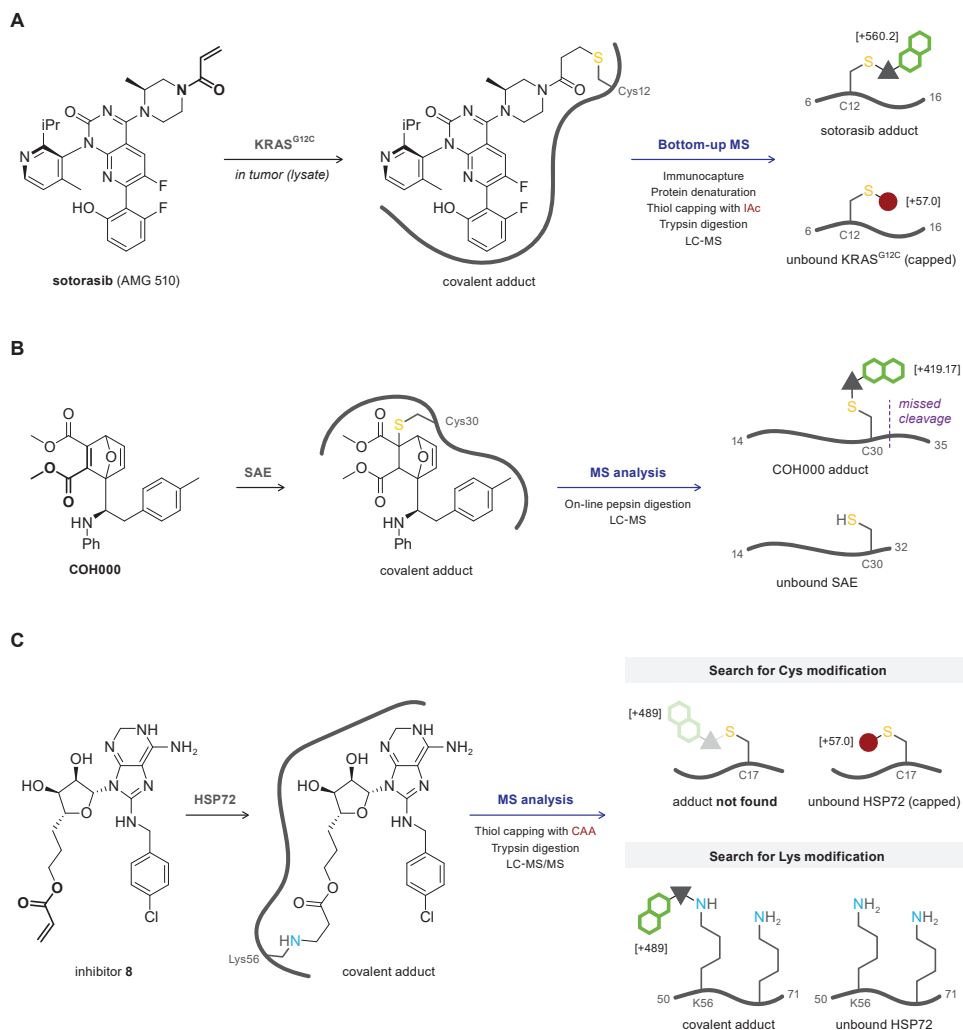


Figure 4 | Bottom-up MS analysis of covalent protein–drug adducts. **(A)** Detection of covalent adduct formation for clinically approved covalent inhibitor sotorasib (AMG 510) with KRAS^{G12C} isolated from lysates originating from (*in vitro* or *in vivo*) treated tumor cells.^{18, 99} **(B)** Identification of covalent adduct formation between recombinant SUMO E1 (SAE) and covalent inhibitor COH000 reveals unexpected modification of allosteric Cys30 rather than catalytic Cys17.⁷⁰ Covalent modification of Cys30 interferes with pepsin-mediated digestion, generating longer adduct peptides than in the untreated control (missed cleavage). **(C)** Bottom-up MS analysis of recombinant HSP72 incubated with covalent acrylamide ligand **8** revealed covalent modification of Lys56 rather than catalytic Cys17.⁷¹ Data analysis focused on lysine modification resulted in detection of the simple adduct of L50–K71 peptide.

(Figure 4C). Instead, they found evidence suggesting allosteric Cys267 was modified, but site-directed mutagenesis revealed that this modification only contributed to a minor covalent adduct and is not responsible for inhibition of protein function. Finally, expanding the search to modification of other nucleophiles (lysine) revealed the modified L50–K71 peptide. A reliable

MS2 spectrum confirming Lys56 as the modified amino acid could not be obtained (details on MS/MS analysis in *section 2.3*) but the unanticipated covalent modification of Lys56 driving the inhibitory activity was confirmed with the HSP72^{K56A} mutant.

Reversible binding mode. Direct detection of the proteolytic peptide modified with a reversible covalent inhibitor (CRI) or ligand can be challenging because denaturation and proteolytic digestion are known to promote CRI dissociation,^{49, 54-55} and treatment with alkylating reagent to cap free thiols can block CRI rebinding. As such, modified peptides are more likely to be detected when thiol capping precedes denaturation and proteolytic digestion. It is possible to detect the modified tryptic peptides with bottom-up MS-based methods if the dissociation rate of the reversible covalent modifier is slow enough, as exemplified by detection of proteolytic UCHL1 peptides modified with reversible covalent cyanimide IMP-1710.⁸⁴

Quantification of covalent occupancy. Covalent target engagement is often quantified indirectly from depletion of the unmodified proteolytic peptide in the treated sample relative to the untreated sample^{96-97, 101} as unbiased quantification of the (modified) proteolytic peptide can be challenging; ionization efficiency differences can occur following modification with a covalent ligand. The (LC-)MS/MS methods to overcome this bias will be discussed in the next section.¹⁰⁰

2.3. MS/MS or Tandem MS

Bottom-up MS analysis is frequently coupled to a subsequent MS analysis (tandem MS or MS/MS) to enable identification of the covalently modified amino acid (**Figure 2C**). Specific precursor peptides are isolated after MS1 and subjected to collision/fragmentation conditions to generate charged fragment ions. The amino acid sequence of the precursor ion can be deduced from the mass of the fragment ions in MS2. For covalent adducts, fragment ions containing the covalently modified amino acid residue have a higher mass than fragment ions derived from the unmodified peptide, thus enabling identification of the modified amino acid. Tandem MS procedures have frequently been used to identify the modified amino acid residue covalent inhibitors targeting different protein classes.^{18, 22, 70-71, 83-84, 90} For the unambiguous assignment of the modified noncatalytic cysteine in the kinase domains affected by clinical irreversible covalent EGFR inhibitor **afatinib** (BIBW 2992) (**Figure 5A**),¹⁴ LC-MS/MS analysis following pepsin-digestion proved very valuable. The modified Q791-L798 peptide ion was found in the MS1 spectrum, and MS2 data identified Cys797 as the modified amino acid. Moreover, detection of a sulfurized afatinib fragment ion resultant from fragmentation of the C-S bond between the thiol and the cysteine β -carbon further confirmed covalent thiol addition to **afatinib**.

Covalent modification can (negatively) affect the ionization of peptide fragments, and it is not uncommon to only detect unmodified fragment ions (**Figure 5B**).^{83-84, 88} LC-MS/MS analysis of the trypsin- and GluC-digested covalent adduct confirmed covalent binding of **tirabrutinib** (ONO-4059)⁸⁸ to the BTK Y476-R487 precursor peptide (MS1) but only unmodified fragment ions and fragmentation of the parent inhibitor were found in MS2. However, the unmodified fragment ions indicate ligand modification occurred at one of the A478-C481 residues, of which Cys481 is the most nucleophilic residue. Interpretation of MS/MS data is usually tailored

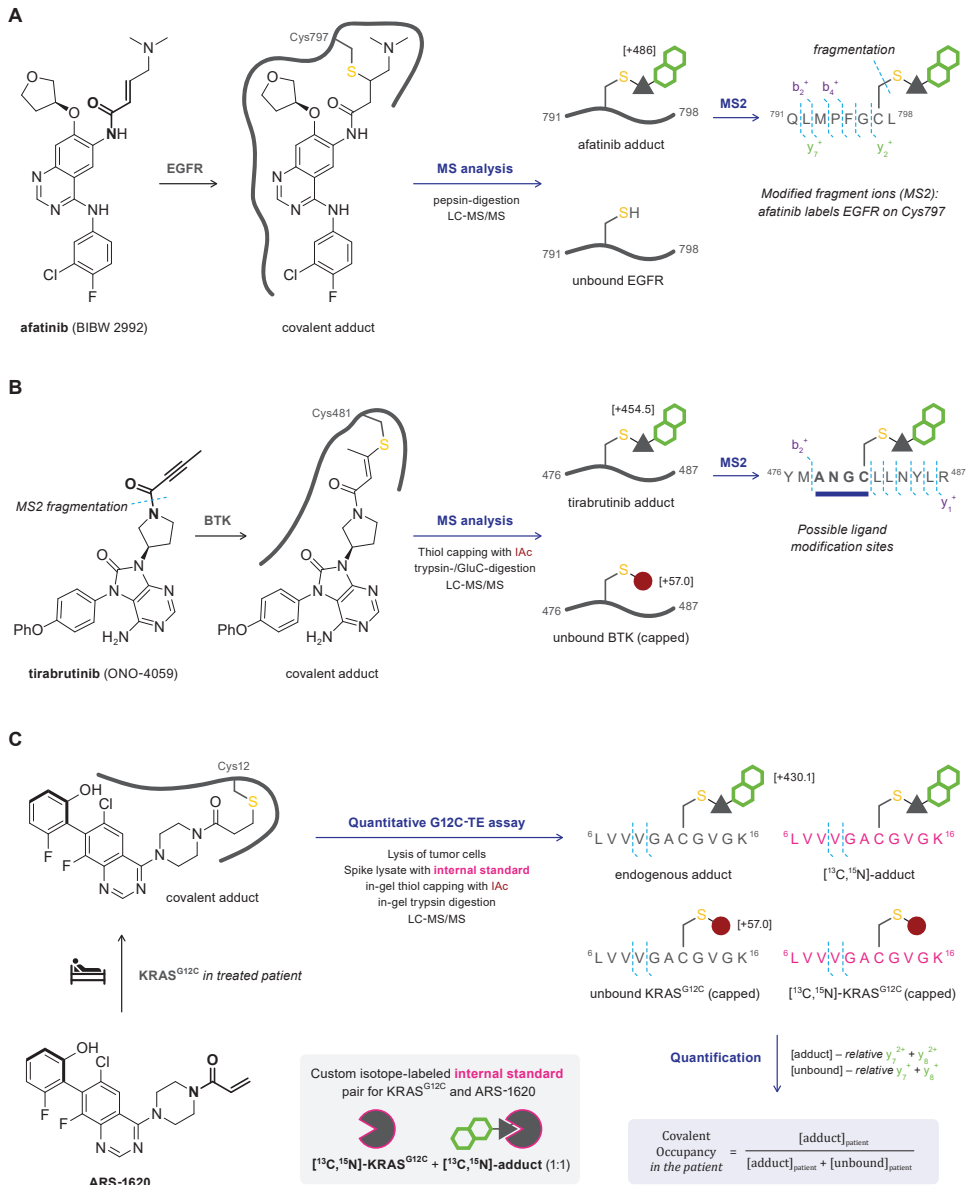


Figure 5 | LC-MS/MS analysis of covalent protein–drug adducts. (A) Identification of the covalently targeted amino acid by afatinib (BIBW 2992) in purified EGFR^{T790M/L858R} kinase domain.¹⁴ **(B)** Identification of covalently modified amino acids in recombinant BTK kinase domain by BTK inhibitor tirabrutinib (ONO-4059).⁸⁵ MS2 detection only showed unmodified fragment ions, and ligand fragmentation of unbound tirabrutinib. **(C)** Internally-controlled quantitative KRAS^{G12C} Target Engagement (G12C-TE) assay illustrated with KRAS^{G12C} inhibitor ARS-1620.¹⁰⁰ Lysates originating from clinical tumor biopsies are spiked with an internal standard: recombinant stable isotope-labeled KRAS^{G12C} (1-169) internal standard – consisting of a 1:1 mixture of free [¹³C,¹⁵N]-KRAS^{G12C} and covalent [¹³C,¹⁵N]-KRAS^{G12C}–ARS-1620 adduct – to calculate the *in vivo* covalent target occupancy from the relative abundance of fragment ions corresponding to endogenous or stable isotope-labeled adducts as well as unbound KRAS^{G12C}.

towards modification of a specific amino acid class or even a single specific residue, searching only for modification of cysteine residues and performing MS2 for peptide ions containing the catalytic cysteine. Although it is generally safe to assume that covalent inhibitors bearing a thiol-reactive electrophile will target the nucleophilic catalytic cysteine residue, covalent modification of less reactive cysteines^{70, 91} or even unexpected amino acids has been reported in exceptional cases.⁷¹

Reversible binding mode. Limitations and challenges for reversible covalent ligands are similar to bottom-up MS analysis. Detection of the simple adduct and fragment ions containing the covalent ligand has been reported for reversible covalent ligands with a slow dissociation rate.⁸⁴

Quantification of covalent occupancy. Covalent *in vivo* target engagement of clinically approved KRAS^{G12C} inhibitor **sotorasib** (AMG 510) has been quantified from the percentage of modified KRAS^{G12C} peptide normalized to the total KRAS^{G12C} peptides in tumor cells recovered from treated mice.¹⁸ However, caution is advised as this method does not take the possible effect of covalent modification onto the ionization of the fragment ions into account. Quantitative covalent KRAS^{G12C} Target Engagement (G12C-TE) assays are typically indirect, using [¹³C,¹⁵N]-KRAS^{G12C} (peptide) as an internal control to determine the absolute level of unoccupied KRAS^{G12C}.^{89, 98, 101-103} Cellular/biochemical occupancy is calculated from comparison of unbound KRAS^{G12C} levels in the treated sample to the untreated control. However, these indirect methods are not compatible with clinical development of solid tumor treatment because pretreatment or patient-matched reference biopsies are typically not available.¹⁰⁴ Scientists at Wellspring Biosciences developed a direct internally controlled quantitative MS/MS-method for the accurate determination of target occupancy in FFPE (formalin fixed paraffin embedded) samples prepared from clinical tumor biopsies without the requirement of pretreatment or untreated controls, illustrated for KRAS^{G12C} inhibitor **ARS-1620** as a proof-of-concept study (**Figure 5C**).¹⁰⁰ Here, tumor-derived lysates are spiked with an internal standard consisting of a 1:1 mixture of unbound [¹³C,¹⁵N]-KRAS^{G12C} and covalent [¹³C,¹⁵N]-KRAS^{G12C}-inhibitor adduct, thereby enabling absolute quantification of endogenous unbound as well as modified KRAS^{G12C} peptide ions. The samples were then exposed to reducing conditions, with in-gel thiol capping with IAc and trypsin digestion of RAS proteins, before being submitted to LC-MS/MS analysis. Using the respective internal standard peptides, the ratio between endogenous unbound and ARS-1620-bound KRAS^{G12C} could be determined, allowing calculation of *in vivo* covalent occupancy. This method is generally applicable for proteins with endogenous expression levels well above the limit of quantification by MS, but its application will be practically limited by the production of recombinant stable isotope-labeled protein and the required generation of an isotope-labeled internal standard for each individual inhibitor.

3. Protein Crystallography

X-ray crystallography is a technique used to elucidate the 3D structure of crystalline compounds, from small molecules to (large) proteins.¹⁰⁵ Protein crystallography is the most informative technique discussed in this work: providing biophysical evidence on the covalent adduct

along with detailed structural information on the modified amino acid residue and the bond layout of the protein-bound ligand. Covalent bonds between individual atoms are not directly observed: the distance between the individual atoms is detected, along with a continuous electron density, from which the likeliness that these atoms are involved in a covalent bond is determined, with performance of orthogonal experiments (e.g. MS, mutagenesis) to validate covalency (**Figure 6**). One of the major practical drawbacks is the consumption of large amounts of highly pure soluble protein, and not all soluble proteins (or protein complexes) form suitably, diffracting crystals (if any at all).¹⁰⁵ In comparison: smaller protein amounts (of lower purity) are sufficient for less informative methods (e.g. MS). For protein crystals with appropriate diffraction, resolved macromolecular (ligand-bound) structures are deposited to the publicly accessible Protein Data Bank (PDB),^{106–107} enabling other researchers to access this wealth of structural information. High resolution structures of covalent adducts are available for various clinically approved TCIs including BTK inhibitors **ibrutinib** (PCI-32765, PDB: 5P9J)¹⁰⁸ and **zanubrutinib** (BGB-3111, PDB: 6J6M),¹² EGFR inhibitor **afatinib** (BIBW 2992, PDB: 4G5J),¹⁴ and proteasome inhibitor **bortezomib** (PS-341, PDB: 2F16).⁵¹ The structural binding information can be used to gain insight on ligand binding driving target selectivity and/or reactivity,^{109–112} and can be combined with (covalent) docking studies^{44, 113–115} to aid structure-based design of covalent ligands with improved potency and/or selectivity.^{22, 45, 116–119} Structure-guided drug design approaches are employed to optimize the proximity of the electrophilic warhead to the nucleophilic amino acid.²² The potency of clinical candidate **ARS-1620** (PDB: 5V9U) was improved by surface groove occupation, resulting in enhanced interactions with the KRAS^{G12C} protein eventually leading to the development of clinically approved KRAS^{G12C} inhibitor **sotorasib** (AMG 510, PDB: 6OIM).⁹⁹

A typical workflow starts with expression and purification of recombinant protein (domain), treatment with ligand, and screening hundreds of crystallization conditions to produce sufficiently large, singular crystals.^{105, 120} Suitable crystals of, hopefully, the covalent protein–ligand adduct are then fished, flash frozen and exposed to an X-ray beam at a synchrotron. The atoms in a crystalline structure (at low temperature) ideally are stationary and the diffraction pattern will be collected over different angles. The resulting intensities are then indexed to determine the space group and integrated into a dataset from which a 3D model can be determined.¹²¹ As the dataset is a reciprocal space representation of the structure, the intensities need phases to actually solve the structure. Using one of the various phasing methods,¹²² an initial structure model and electron density can be determined. After various iterative rounds of model building and structure refinement, hopefully electron density for the (covalent) ligand can be seen. The most reliable structural data is obtained when the model is first refined for the protein (based on the apo structure of the protein) and the remaining electron density is used to fit the ligand as this minimizes the bias for the inhibitor binding site.¹²³ For some proteins, apo structures of active, uninhibited enzymes may not be available for autoproteolytic/cannibalistic reasons (e.g. cysteine cathepsins) due to self-proteolysis or autodigestion.^{124–125} Occasionally, one can obtain mixed crystals, consisting of free protein, noncovalent protein–ligand complex and covalent protein–ligand adduct. This can decrease the quality and may impair detection altogether, thus requiring purification of the covalent adduct prior to crystallization. Alternatively, crystals of the free (apo) protein are allowed to form,

before soaking in the (covalent) ligand, though this can result in mixed crystals or cracking of the crystal.¹⁴ Soaking is popular in structure-based ligand screens as it conveniently sidesteps the optimization of crystallization conditions for each individual ligand.¹²⁶ However, soaking is not recommended for covalent drugs as the rigid crystalline protein can hinder formation of a covalent adduct, especially if the crystalline apo protein is in the incorrect conformation for ligand binding; if noncovalent ligand binding induces a conformational change before covalent adduct formation,¹²⁷ or if the crystalline protein has lost its catalytic activity essential for covalent adduct formation with mechanism-based inhibitors.⁴⁸

Non-crystallographers are advised to consult the works of Wlodawer and co-workers on interpretation and critical evaluation of structural data.^{120, 128} The value for resolution is expressed as the smallest resolved distance (in Å = 10⁻¹⁰ m) in the structure model. Although the resolution applies to the whole map, parts of the structure may suffer from disorder and have high temperature factors as a result.¹²⁸⁻¹²⁹ It is important to realize that covalent adduct formation in these areas may not be reliably detected. Hydrogen atoms are not shown in most structures obtained by X-ray diffraction (XRD) as hydrogen atoms only weakly scatter X-ray beams: they only have one electron, which is always involved in a bond with another atom, and are therefore not precisely localized at the usual resolution.¹³⁰ A low numeric value correlates with a high resolution: individual atoms (including some hydrogens) can be observed at <1.2 Å, most backbones and sidechains are clear at 2.5 Å, while only the general backbone can be solved at a resolution of 5 Å.¹²¹ The average distance between a thiol atom and carbon atom in a covalent single C-S bond is 1.82 Å and cannot reliably be observed when the resolution at the ligand binding site is too low.

Protein crystallography has revealed unexpected modification of noncatalytic (allosteric) cysteine residues rather than the catalytic cysteine residue. Solving the crystal structure provided molecular insight on why a covalent E3 ubiquitin ligase Nedd4-1 inhibitor (PDB: 5C91) inhibits elongation of polyubiquitin chains but does not completely inhibit all catalytic activity:⁹¹ the inhibitor targets allosteric Cys627 positioned at the substrate binding site rather than the more nucleophilic catalytic cysteine residue Cys867 (**Figure 6A**). Covalent modification of allosteric Cys30 in SUMO-activating enzyme (SUMO E1 or SAE) by **COH000** rather than catalytic Cys173 (see **Figure 4B**) was validated by solving the X-ray crystal structure (PDB: 6CWY).¹³¹ Of note, protein crystallography reveals the protein-ligand complex or adduct that forms the best crystals but this does not have to be the most prevalent binding mode in solution: modification of a specific (unexpected) amino acid should be validated with orthogonal techniques to ensure the modification is representative for ligand binding in solution. As such, protein crystallography is not a suitable technique for quantification of covalent adduct formation.

Natural product **salinosporamide A** (SalA, NPI-0052, marizomib) is a clinically approved covalent 20S proteasome inhibitor with an irreversible binding mode, whereas closely related natural product **salinosporamide B** (SalB, NPI-0047) has a reversible binding mode (**Figure 6B**).¹³² Crystal structure analysis reveals that threonine Thr1 addition to the chloroalkyl β-lactone in SalA resulting in β-lactone ring opening is followed by intramolecular nucleophilic substitution to irreversibly form a stable cyclic tetrahydrofuran (THF) ring (PDB: 2FAK)¹¹⁰

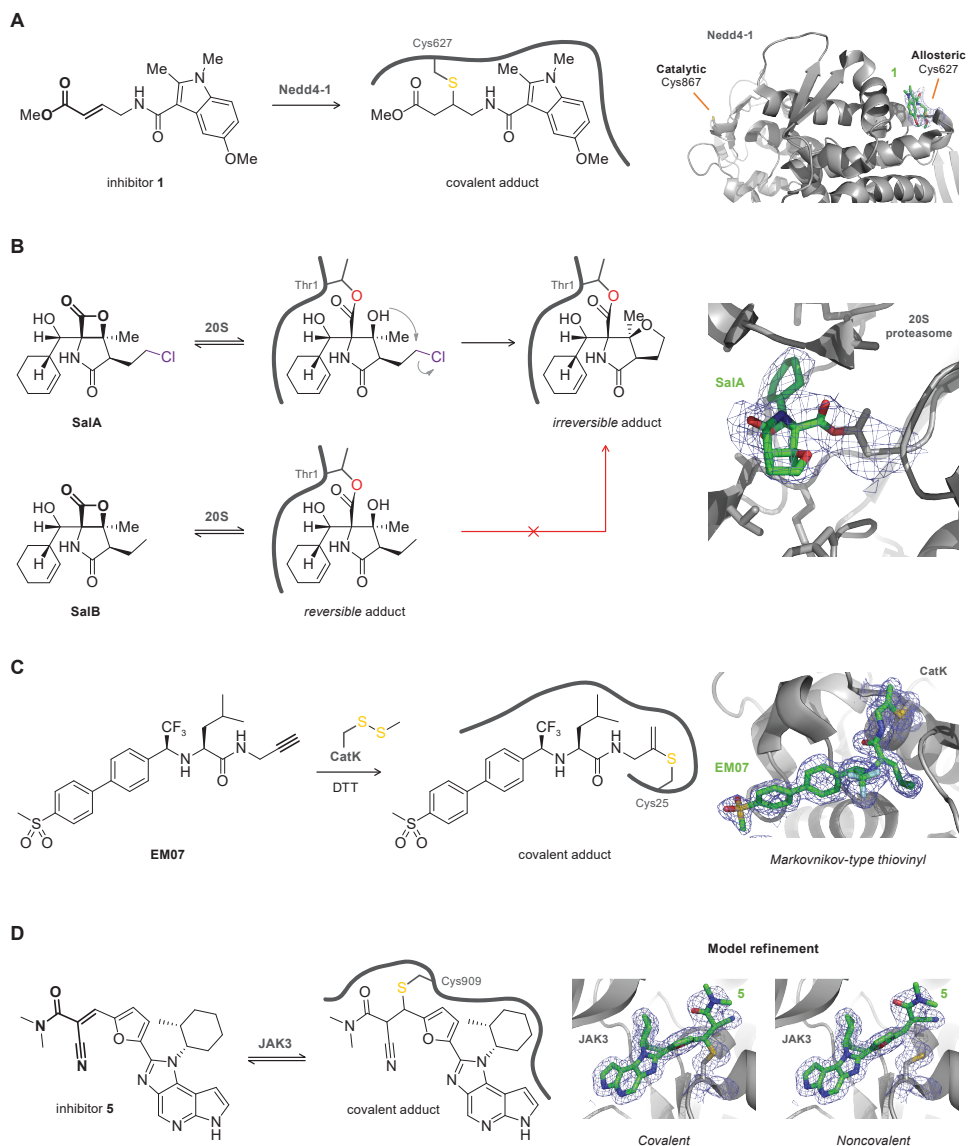


Figure 6 | Structural binding information on covalent protein–drug adducts obtained by protein crystallography. **(A)** Identification of unexpected modified amino acid residue. Inhibitor 1 modifies ubiquitin E3 ligase Nedd4-1 on allosteric cysteine residue Cys627 rather than the more nucleophilic catalytic cysteine residue Cys867 (PDB: 5C91).⁹¹ **(B)** Protein crystallography of closely related 20S proteasome inhibitors salinosporamide A (SalA, NPI-0052, marizomib) and salinosporamide B (SalB) aids mechanistic understanding on their different binding modes:¹³² SalA (PDB: 2FAK)¹¹⁰ forms an irreversible adduct by ring closure with a chloride leaving group following initial formation of the reversible covalent acyl ester with Thr1-OH. **(C)** Structural analysis confirms thiol–alkyne addition of catalytic Cys25 in human cathepsin K (hCatK) to the internal alkyne carbon on odanacatib derivative EM07 (PDB: 6QBS),⁸³ forming a covalent adduct with a Markovnikov thiovinyl bond lay-out similar to the thioimide adduct of hCatK with the odanacatib nitrile (PDB: 5TDI).¹³⁴ **(D)** Refined electron density maps assuming covalent ligand binding (*left*) or noncovalent ligand binding (*right*) indicate a mixture of both states upon co-crystallization of reversible covalent cyanoacrylamide 5 with Janus kinase JAK3 (PDB: 5LWN).¹³⁵

that cannot be formed with the β -lactone of SalB, thus aiding molecular understanding of the irreversible binding mode of SalA. Similarly, protein crystallography provided mechanistic insight on the superior selectivity of clinical multiple myeloma drug **carfilzomib** (PR-171, PDB: 4R67)¹³³ for the 20S proteasome over non-proteasomal proteases, and why such selectivity is not observed for **bortezomib** (PS-341, PDB: 2F16).⁵¹ **Carfilzomib** forms a dual covalent adduct with the 20S proteasome, and the additional engagement of the Thr1 primary amine is specific for proteasomal proteins.

Protein crystallography provides valuable structural information on the bond layout of the covalently bound ligand. In our group, solving the crystal structure of ABP (activity-based probe) Ub-**Prg** with cysteine protease vOTU revealed an unexpected Markovnikov-type thiovinyl bond between the active site cysteine thiol in the protease and internal carbon of the alkyne in Ub-**Prg** (PDB: 3ZNH).¹³⁶ This thiovinyl bond layout has since been observed for propargyl-containing ABPs targeting various cysteine proteases (listed in Table S1 of citation⁸²), and for small molecule CatK inhibitor **EM07** (PDB: 6QBS)⁸³ (**Figure 6C**). Active, mature CatK had to be treated with *S*-methyl methanethiosulfonate (MMTS) to prevent autodigestion in absence of competing substrate, which is more prone to occur at the high concentrations (>10 mg/mL) used for crystallography.^{124-125, 137} The thiomethyl protecting group is removed with reducing agent (e.g. BME, TCEP or DTT), to allow covalent adduct formation with the simultaneously added inhibitor.¹³⁸

Reversible binding mode. Crystallography does not involve stringent washing or (harsh) denaturing conditions that would promote ligand dissociation, thus being particularly suitable for the evaluation of reversible covalent ligands. Co-crystallization of clinical Hepatitis C virus (HCV) drug **telaprevir** (VX-950) with serine protease NS3/4A confirms the bond layout of catalytic Ser139 bound to the C- α carbon of the ketoamide warhead (PDB: 3SV6).⁵³ Ligand interactions with frequently mutated protein sites provides a molecular basis for clinically occurring drug resistance. Importantly, covalent and noncovalently bound ligand may coexist in the crystal structure, as was demonstrated for reversible covalent cyanoacrylamide inhibitors targeting nonconserved cysteine Cys909 in an induced fit binding pocket of Janus kinase JAK3 (PDB: 5LWN)¹³⁵ (**Figure 6D**). Other notable examples of reversible inhibitors are crystal structures of the first clinically approved proteasome inhibitor **bortezomib** (PS-341)¹³⁹ covalently bound to Thr1 of yeast 20S proteasome through the boronic acid moiety (PDB: 2F16),⁵¹ COVID-19 drug **nirmatrelvir** (PF-07321332) forming a reversible covalent thioimide adduct with catalytic cysteine Cys145 of SARS-CoV-2 M^{pro} protease in crystals formed by co-crystallization (PDB: 7RFW) or by soaking the apo crystal (PDB: 7RFS),⁴⁷ and the structure-based design of reversible covalent BTK inhibitors with tunable residence times.⁵⁵

4. Intrinsic Fluorescence/Absorbance

Covalent thiol addition can change the intrinsic spectroscopic properties of certain ligands and can be used to monitor covalent adduct formation in plate-based fluorescence/absorbance assays. Generating a chemical tool or an activity-based probe (ABP) by introducing a fluorophore or fluorescent leaving group to the ligand core will be discussed in more detail in

section 6. Here, we will focus on ligands that do not require late-stage structural modifications because they contain a structural motif that has intrinsic spectroscopic properties,¹⁴⁰ with a detectable change upon covalent adduct formation. This method is not generally applicable as there are strict structural limitations to the ligand core and nature of the electrophilic warhead. The main advantage of intrinsic spectroscopic methods is the compatibility with plate-based HTS assays, and catalytic activity is not required. The latter is directly a major drawback as it is impossible to discriminate between desired adduct formation with the intended cysteine thiol and undesired adduct formation with untargeted thiols present in the reaction buffer. This method always needs orthogonal validation as noncovalent binding events can also induce detectable changes in intrinsic fluorescence/absorbance.¹⁴¹

The Rauh group reported a plate-based assay for direct detection of covalent bond formation of quinazoline- and quinoline-based kinase inhibitors with an attached conjugated electron-deficient group such as an acrylamide warhead (**Figure 7A**).¹⁴² The unbound inhibitor exhibits

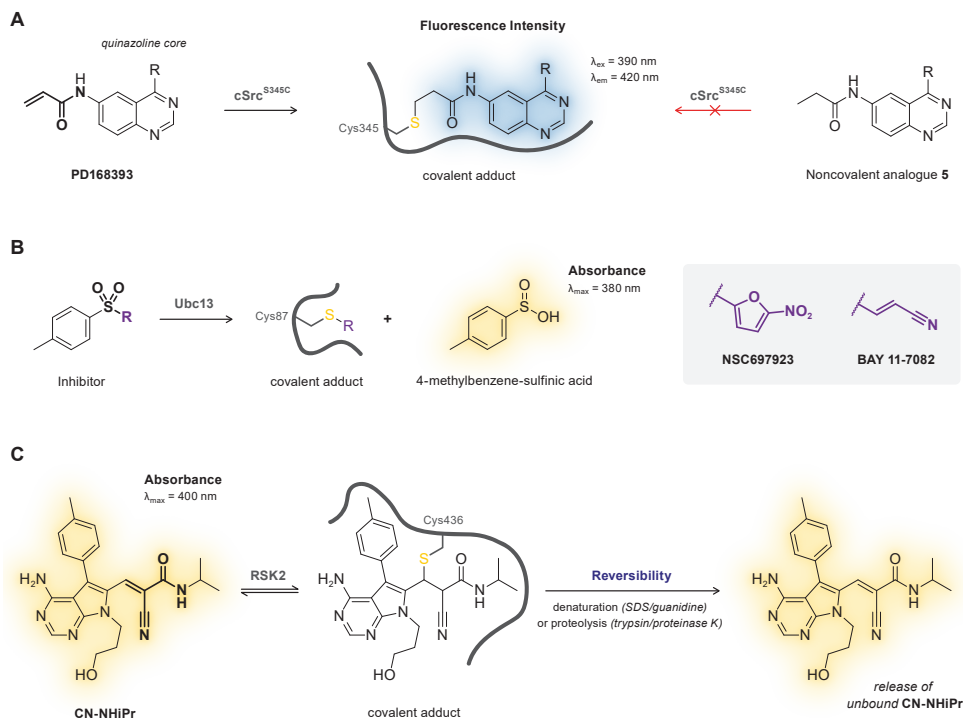


Figure 7 | Direct detection of changes in intrinsic spectroscopic properties upon covalent thiol addition. **(A)** Thiol addition increases intrinsic fluorescence intensity of quinazoline and quinoline cores with an attached conjugated Michael acceptor.¹⁴² Detection of increased fluorescence intensity upon adduct formation for irreversible covalent inhibitor PD168393 with cSrc^{S345C} but not for the noncovalent analogue. **(B)** Release of 4-methylbenzene-sulfinic acid results in a detectable absorption increase upon covalent adduct formation of inhibitors NSC697923 and BAY 11-7082 with Ubc13^{WT}.¹⁴³ **(C)** Intrinsic absorption in the UV-visible spectrum of *N*-isopropyl cyanoacrylamide CN-NHiPr decreases upon nucleophilic thiol addition.⁵⁴ Reappearance of signal upon protein denaturation or proteolysis-induced inhibitor dissociation is indicative of a reversible covalent binding mode.

weak fluorescence emission upon excitation due to photo-induced electron transfer (PET) from the quinazoline/quinoline core to the attached conjugated Michael acceptor. Covalent thiol adduct formation enhances the quantum yield and can thus be detected as an increase in the fluorescence emission as was illustrated for recombinant cSrc^{S345C} mutant (Cys345 mutation on an isostructural position to Cys797 in EGFR) with irreversible covalent quinazoline **PD168393**, but not with a noncovalent analogue.¹⁴²

Analogous to fluorogenic substrates that release a fluorescent group upon proteolytic cleavage,¹⁴⁴ certain irreversible covalent inhibitors release a (detectable) leaving group upon covalent thiol addition (**Figure 7B**). This concept has been utilized to monitor covalent adduct formation of covalent inhibitors **NSC697923** and **BAY 11-7082** with E2 ubiquitin conjugating enzyme Ubc13:¹⁴³ elimination of 4-methylbenzene-sulfinic acid upon covalent thiol addition can be monitored by an increase in absorbance in the UV-visible spectrum directly related to covalent adduct formation.

Reversible covalent binding and reversibility assays. Detection of intrinsic fluorescence is compatible with reversible covalent inhibition, as demonstrated by the Taunton group for thiol addition to reversible covalent kinase inhibitors bearing a cyanoacrylamide warhead.⁵⁴ They report that unbound *N*-isopropyl cyanoacrylamide **CN-NHiPr** has a strong intrinsic absorption in the UV-visible spectrum which disappears upon treatment with excess recombinant RSK2 kinase domain, consistent with nucleophilic thiol addition of Cys436 to the cyanoacrylamide warhead (**Figure 7C**). Reversibility could then be assessed by exposing covalent RSK2–CN-NHiPr adduct to denaturing conditions or proteolytic digestion to induce target dissociation: reappearance of the absorption peak (and LC-MS detection of recovered unbound cyanoacrylamide) is in agreement with a reversible covalent binding mode.

5. Nuclear Magnetic Resonance (NMR)

Ligand-observed NMR analysis was the predominant method to detect covalent bond formation between enzyme and covalent inhibitor prior to rise in popularity of MS analysis or protein crystallography.³⁹ A change in the chemical environment resultant from (non) covalent interactions causes a detectable change in the resonance frequency (typically reported as ‘chemical shift’) and the coupling of nuclei with a nonzero nuclear spin (e.g. ¹H, ¹³C, ¹⁵N) in the magnetic field of the NMR spectrometer. An overview of NMR spectroscopy principles for protein–ligand interactions can be found elsewhere.^{145–147} Nowadays, NMR studies are employed in structure-based drug discovery and NMR screening for covalent (fragment) ligands,^{148–149} and have been used for ligand binding site mapping and structural elucidation of various covalent ligands. In macromolecular structure determination, NMR and protein crystallography can be complementary techniques,^{150–151} and NMR-resolved macromolecular (ligand-bound) structures are also deposited to the publicly accessible Protein Data Bank (PDB). Solution structures of covalent adducts have been deposited for compounds bound to protein targets,^{152–154} but also to minor groove duplex DNA: for example the covalent adduct of chemotherapy drug **mitomycin C** (UGN-101) a DNA 9-mer (PDB: 199D),¹⁵⁵ and alkylating agent **duocarmycin A** covalently bound to a DNA 7-mer (PDB: 107D).¹⁵⁶ Contrary to protein

crystallography, NMR techniques are compatible with characterization of binding mode reversibility by performing dialysis experiments,¹⁵⁴ and ligand binding can be quantified to determine kinetic parameters (e.g. dissociation constant K_D).^{146-147, 157-158}

Macromolecular structure determination typically involves multiple different NMR experiments to interrogate the different facets of the covalent adduct but 2D NMR techniques that detect the scalar (through multiple bonds) correlation of protein atoms to ligand atoms (e.g. pulse programs based on (HSQC)-TOCSY, HMBC, or 2D-INADEQUATE) ultimately provide the most conclusive NMR-based evidence of a covalent protein–drug adduct because these correlations are exclusive to covalent adducts. Unfortunately, protein signals often overlap with ligand signals, making it practically impossible to confidently discern the correlation between a ligand atom and a protein atom in a covalent adduct because correlations of atoms residing in the same ligand/protein (that do not require a covalent adduct) overlap.¹⁵³ Here we will feature the two main detection principles: protein-observed NMR (section 5.1) and ligand-observed NMR (section 5.2).

5.1. Protein-Observed NMR

Protein-observed NMR experiments compare the signals originating from the protein in unbound state to the protein–ligand complex; ligand binding changes the chemical environment of amino acids in proximity of the ligand, inducing chemical shift perturbations that can be used to map the ligand binding site onto the protein structure.^{147, 159} Given the vast number of atoms in a protein and the low natural abundance of the most suitable isotopes (e.g. ^{13}C , ^{15}N), protein-observed NMR spectroscopy typically involves production and purification of a uniform isotope-labeled protein along with recording a reference spectrum of the unbound protein to enable assignment of peaks to specific protein atoms.¹⁶⁰⁻¹⁶² Protein-observed NMR experiments can be used to gain structural insight to ligand binding in solution, which is particularly useful for targets that are not compatible with crystallization or conformations that do not crystallize: for example, solution protein-observed NMR spectra revealed that noncovalent kinase inhibitor **imatinib** binds to c-Abl in an previously unidentified open state.¹⁶³ Furthermore, NMR experiments were employed to identify the binding site of covalent inhibitors of the *S. aureus* Sortase A enzyme (*Sa*-SrtA), and used to solve the structure of the covalent adduct (PDB: 2MLM, 6R1V).¹⁵³⁻¹⁵⁴ However, protein-observed NMR techniques are typically restricted to relatively small proteins (<50 kDa), and most techniques used in macromolecular structure elucidation (e.g. ^{15}N , ^1H]-HSQC) cannot directly discriminate between a covalent or a noncovalent ligand. It is advisable to employ additional ligand-observed experiments or orthogonal techniques (e.g. MS analysis) for covalent adduct validation: protein-observed NMR experiments technically only provide indirect evidence on covalency.^{160, 164}

5.2. Ligand-Observed NMR

In ligand-observed NMR experiments, a change in chemical shift of the ligand signals in the protein–ligand complex is compared relative to the ligand signals in the unbound ligand. The most popular ligand-observed NMR techniques for fragment screening (e.g. saturation transfer difference spectroscopy and its variants) are based on the NOE principle (proximity in

space) and thus cannot discriminate between binding of covalent or noncovalent ligands.^{160, 165} ¹H NMR chemical shift perturbations can be employed to distinguish unbound ligand from a covalent protein–ligand adduct, and support identification of the adduct isoform.¹⁶⁶ In addition to validation of covalent adduct formation, 1D ¹H NMR approaches enable indirect quantification of covalent occupancy by integration of the disappearing unbound ligand signals.^{39, 167} However, ¹H NMR experiments are typically only performed for adduct formation with small molecule thiol reagents (e.g. GSH) as overlapping background signals originating from protein hydrogens limit the practical application.

Direct detection of the covalent protein–ligand adduct by ligand-observed ¹³C NMR experiments is a more feasible approach but requires chemical synthesis of ligand with a ¹³C-labeled warhead to improve the signal over the background, otherwise the naturally occurring ¹³C signals in the ligand will be lost among those originating from the protein. The ¹³C chemical shift perturbations of adjacent carbons in the electrophilic warhead can be significant upon covalent thiol modification, especially compared to the less pronounced shifts induced by noncovalent binding interactions.¹⁶⁸ Detection of chemical shift perturbations of (isotope-labeled) epoxy succinyl peptides upon cysteine protease papain binding was successfully utilized to detect the covalent adduct along with identification of the covalent modification site.¹⁶⁹ Moreover, ¹³C NMR APT (attached proton test) experiments can be indicative of covalent adduct formation with unsaturated electrophiles (e.g. acrylamide): the phasing of the vinyl carbon adjacent to the reactive carbon in the unbound acrylamide is negative (CH) but is positive (CH₂) in the covalent adduct. Future application of ligand-observed NMR may be extended beyond the common ¹³C NMR and ¹H NMR without chemical introduction of an isotope-labeled atom for warheads bearing naturally abundant reactive atoms compatible with NMR (e.g. ³¹P or ¹⁹F in fluorophosphonates). ¹¹B NMR has been employed to detect the tetrahedral adduct of boronic acid covalently bound to Ser195 in serine protease trypsin.¹⁷⁰ This label-free approach has only been employed to study model reagents,¹⁷¹ but there still are seemingly unexplored opportunities for covalent adduct detection of boronic acid-bearing inhibitors (e.g. proteasome inhibitor **bortezomib**) with their pharmaceutical target.

Reversible inhibition and reversibility assays. An important advantage of NMR-based detection of covalent adducts is the compatibility with reversible covalent inhibitors. Especially when the covalent adduct is too short-lived to be isolated or detected due to rapid inhibitor dissociation under MS/sample preparation conditions as NMR enables detection in (aqueous) solution.⁵⁴ Ligand-observed ¹³C NMR analyses were already performed in 1986 to obtain evidence for the formation of a thioimide ester adduct between a nitrile ligand and the active site sulfhydryl of cysteine protease papain (**Figure 8A**).¹⁷² Incubation of active papain with the [¹³C]-labeled nitrile inhibitor resulted in appearance of a resonance signal at 182 ppm in ¹³C NMR in accordance with a covalent thioimide ester adduct. Rapid disappearance of the thioimide signal and increase of unbound inhibitor signal (~117 ppm) was detected upon treatment of the covalent adduct with glacial acid (AcOH) and thiol-trapping reagent 2,2'-dipyridyldisulfide (DPS), indicative of a reversible covalent binding mode. Similar ligand-observed NMR studies have been performed to provide evidence for reversible covalent adduct formation of cathepsin K with a [¹³C, ¹⁵N₄]-diacylhydrazine,¹⁷⁴ and papain with a [¹³C]-cyanamide.¹⁷⁵ A more

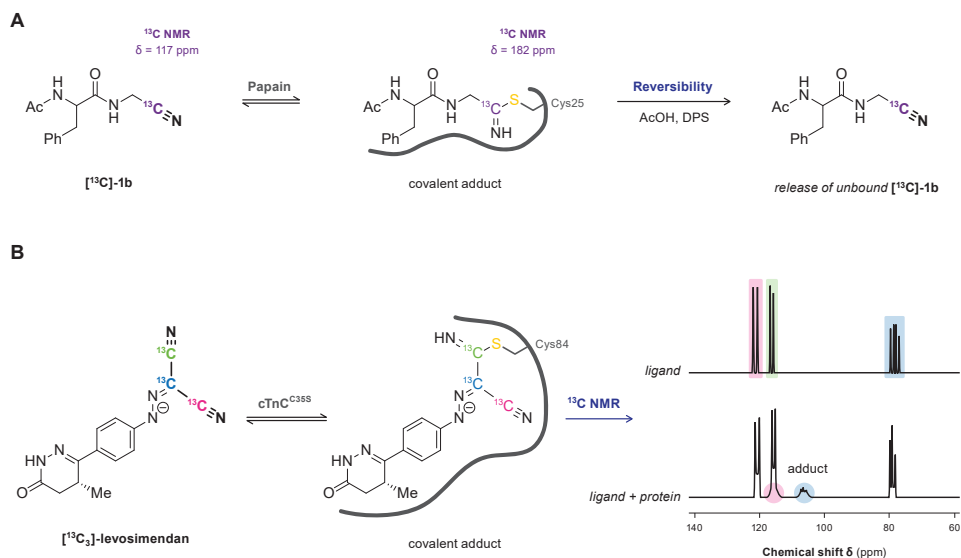


Figure 8 | Ligand-observed ^{13}C NMR detection of covalent protein–drug adducts. **(A)** Chemical shift perturbation of the electrophilic ^{13}C -labeled carbon in unbound nitrile ^{13}C -1b relative to the thioimide ester adduct provides evidence of a covalent papain–nitrile adduct.¹⁷² Detection of unbound nitrile upon treatment with glacial acid (AcOH) and thiol-trapping reagent 2,2'-dipyridyldisulfide (DPS) is indicative of a reversible covalent binding mode. **(B)** Ligand-observed NMR studies with ^{13}C -labeled levosimendan provide evidence for reversible covalent binding to a cysteine thiol in cardiac troponin C (cTnC).¹⁷³

recent example of ligand-observed NMR analysis aided elucidation of the binding mode of Ca^{2+} sensitizer **levosimendan** (**Figure 8B**), a clinical drug for heart failure treatment whose exact mechanism of action remained elusive for over 20 years after its discovery. Formation of a thioimide bond between the electrophilic malonitrile moiety and cardiac troponin C (cTnC) was always assumed to have an important role,¹⁷⁶ but evidence for this reversible covalent binding mode was finally provided in 2016 by employing $^{13}\text{C}_3$ -levosimendan in ligand-observed ^{13}C NMR studies.¹⁷³ Disappearance of unbound $^{13}\text{C}_3$ -levosimendan signals (~ 120 ppm) along with appearance of new signals (~ 160 ppm) is in agreement with predicted chemical shifts for a thioimide adduct between the electrophilic malonitrile moiety on **levosimendan** and a cysteine thiol in cTnC. Lack of adduct in presence of cTnC^{C84S} but not cTnC^{C35S} validates Cys84 as the covalently modified amino acid residue.²⁹

6. Activity-Based Protein Profiling (ABPP)

Activity-based protein profiling (ABPP) is a chemical biology technique that employs covalent activity-based probes (ABPs) to characterize covalent enzyme modification in relevant biological systems (e.g. live cells, *in vivo*).^{104, 177–180} Pioneered in the labs of Cravatt¹⁸¹ and Bogoy,¹⁸² ABPs entailed a reactive group with a detection tag (e.g. fluorophore, radiolabeled isotope) or enrichment handle (e.g. biotin), that covalently modified catalytic serine/cysteine residues in active and uninhibited enzyme.^{178, 183} This general structure design is mostly maintained in

modern ABPs which typically comprise of an electrophilic moiety that forms a covalent bond with a nucleophilic amino acid residue, a reporter group (e.g. fluorophore, enrichment handle, bioorthogonal handle) to detect the covalent adduct, and, optionally, a recognition element for target/class-selectivity (**Figure 9A**). Nowadays, ABP development is not limited to catalytic amino acid residues: a wide range of ABPs is available from general residue-specific agents (e.g. **iodoacetamide** (IAc)-based thiol-alkylating reagents for cysteines)¹⁸⁴ and class-specific ABPs (e.g. fluorophosphonate-based probes for serine hydrolyses,¹⁷⁷ ubiquitin-based probes for DUBs,¹⁸⁵ ATP-based probes for kinases¹⁸⁶) to target-selective ABPs³⁷ (e.g. **ibrutinib**-based ABPs for BTK).¹¹ The field has since expanded beyond truly activity-based probes: ABPs targeting noncatalytic residues also label catalytic inactive mutants thus not requiring catalytic activity.¹¹ ABPP has a prominent role in covalent drug discovery:^{178, 187-188} not only for identification of new covalent hits in covalent (fragment) screening,¹⁸⁹ but also to identify cellular/*in vivo* covalently modified (off-target) proteins thereby derisking covalent inhibitor development.^{2, 190} The latter is emphasized by the recent work of van Esbroeck *et al.*:¹⁹¹ multiple off-target lipases targeted by fatty acid amide hydrolase (FAAH) inhibitor **BIA 10-2474** were identified by competitive ABPP, providing a possible explanation to the clinical neurotoxicity with lethal outcome for one of the human subjects in the phase I clinical trial (2016). This tragic example highlights why identification of potential covalent off-target modifications by (competitive) ABPP is recommended to be an integral part of early-stage covalent drug development.¹⁹⁰

Drug-derived ABPs are designed in two flavors: one-step ABPs (**Figure 9B**) and two-step ABPs (**Figure 9C**). One-step ABPs are generated by introduction of a (fluorescent) detection tag or an enrichment handle onto the parent drug by chemical synthesis. The tag or handle is introduced in a position that does not interfere with target binding, as indicated by structural data (e.g. crystal structure, docking simulations) or SAR analysis. Introduction of a large tag/handle can modify ligand reactivity, target selectivity, as well as cell permeability.¹⁹² Similar to one-step ABPs, two-step ABPs are generated from the parent drug but now a small and nonperturbing bioorthogonal handle is introduced, to which the actual detection group (fluorophore, enrichment handle) is clicked in the second step (**Figure 9C**). This bioorthogonal handle is less likely to have a pronounced effect on ligand selectivity, which is why evaluation of a two-step ABP is recommended in an early stage of covalent drug development to identify potential off-target effects.^{32, 190} Here, the proteome is treated with the two-step ABP bearing a small bioorthogonal handle (*step 1*), followed by coupling of a relatively large detection tag or enrichment handle (*step 2*). Traditionally, the coupling reaction employs the Huisgen Copper-catalyzed Alkyne–Azide Cycloaddition (CuAAC) reaction between alkynes and azides ('Click' chemistry)¹⁹³⁻¹⁹⁴ but alternative bioorthogonal reactions are available^{192, 195} – such as the Strain-Promoted Alkyne–Azide Cycloaddition (SPAAC) between strained alkynes and azides,¹⁹⁶ or the Inverse Electron Demand Diels Alder (IEDDA) reaction between (fluorogenic) tetrazines and strained dienophiles.¹⁹⁷ Two-step ABPs enable incubation in the native environment and are more likely to retain the membrane penetrating properties of the parent inhibitor, and are thus compatible with *in situ* and *in vivo* applications.^{104, 198} The success of this approach has recently been illustrated for inhibitors with various targets³² among which are BTK inhibitor **ibrutinib**,^{11, 190} JAK3 inhibitor **ritlecitinib** (PF-06651600),¹⁹⁹ KRAS^{G12C} inhibitor **adagrasib** (MRTX849),¹⁰³ and anti-obesity drug **orlistat**.²⁰⁰ Two-step clickable ABPs facilitate

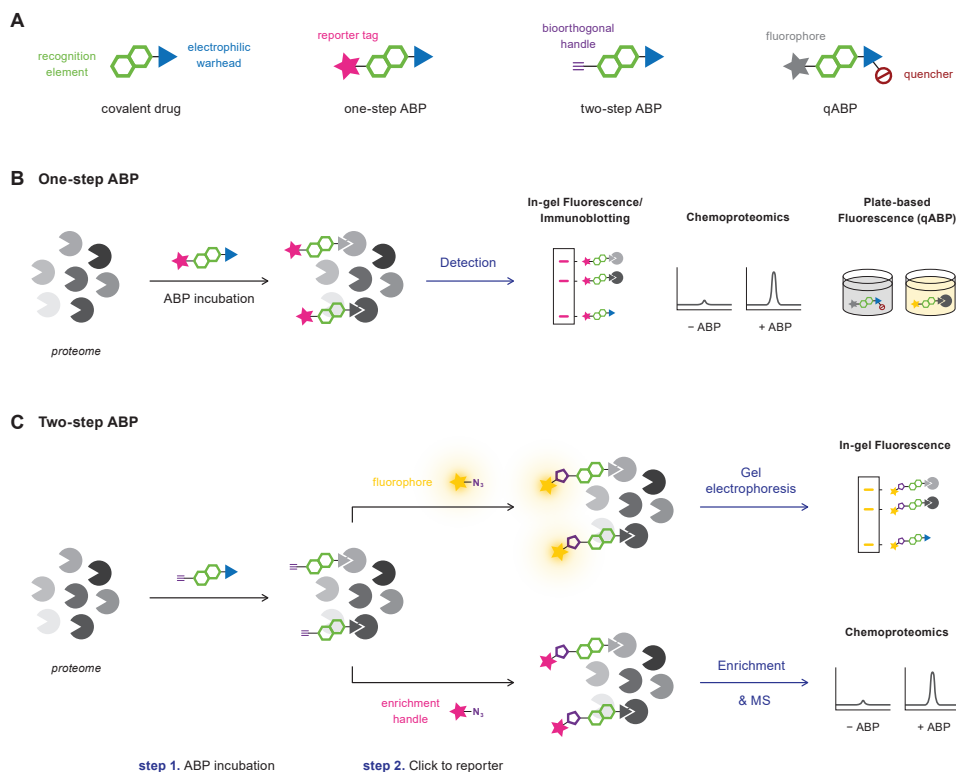


Figure 9 | Strategies for covalent adduct detection with drug-derived activity-based probes (ABPs). **(A)** General design principle for covalent drug-derived ABPs. A fluorophore, detection tag or enrichment handle is introduced onto the parent covalent drug bearing a recognition element and a covalent warhead. **(B)** Detection of covalently modified proteins with one-step ABPs. Proteome is treated with one-step ABP, proteins are resolved by gel electrophoresis or affinity purification, and modified targets are detected by in-gel fluorescence or immunoblotting, or by chemoproteomic evaluation. **(C)** Detection of covalent adducts with two-step ABPs. Proteome is incubated with a two-step ABP bearing a small bioorthogonal handle (*step 1*), followed by bioorthogonal coupling of a fluorophore, detection tag or enrichment handle (*step 2*), with subsequent analysis as shown for the one-step ABPs.

the coupling of a dual biotin/TAMRA-azide, allowing both in-gel fluorescence scanning for the TAMRA fluorophore and immunoblotting for the biotin tag in gel-based evaluation, and the biotin tag can also be utilized as an enrichment handle in chemoproteomic evaluation. The success of this dual approach is illustrated by clickable two-step ABPs equipped with a bioorthogonal alkyne handle: ABP PF-06789402 based on the scaffold of JAK3/TEC family kinase inhibitor **ritlectinib** (PF-06651600),¹⁹⁹ and ABP selinexor-yne derived from clinically approved covalent XPO1 inhibitor **selinexor** (KPT-330).²⁰¹

Altogether, ABPP is a powerful tool to identify (un)desired covalent modification in a relevant biological setting. Here, we will discuss the detection of the covalent adduct in whole proteome with gel electrophoresis platforms (section 6.1), chemoproteomic platforms (section 6.2), and homogeneous (plate-based) platforms (section 6.3).

6.1. Gel Electrophoresis Platforms (In-gel Fluorescence, Immunoblotting)

Gel electrophoresis platforms were among the earliest ABPP methods to interrogate enzyme activity in complex mixtures, and are still a common method for rapid evaluation of inhibitor specificity.¹⁷⁸ A typical workflow (**Figure 9B**) involves incubation of recombinant protein or a whole proteome (e.g. cell lysate) with a one-step ABP followed by sample preparation under denaturing conditions (e.g. heating in presence of a reducing agent such as BME or TCEP) to simultaneously remove unreacted ABP and promote dissociation of noncovalent complexes. Then, the treated proteome is submitted to gel electrophoresis, and covalent adducts are visualized by in-gel fluorescence scanning for the fluorophore (e.g. TAMRA, Cy5, BODIPY)^{11, 199, 201} or immunoblotting for a reporter tag or enrichment handle (e.g. biotin, GST, His),²⁰² with a band appearing at the adduct mass (kDa). Gel-based ABP analysis is fast but is less informative than chemoproteomic approaches (discussed in *section 6.2*). Identifying the exact protein target in a proteome can be challenging as proteins of similar mass may overlap on gel, which may be addressed by comparative ABPP with knock-out cell lines.²⁰³⁻²⁰⁴ Competitive ABPP experiments are typically conducted to validate that the ABP has the same specificity as the inhibitor:^{11, 199, 205-206} treatment with parent BTK inhibitor **ibrutinib** (PCI-32765) precludes labeling with cell permeable fluorescent ABP PCI-33380 (**Figure 10A**).¹¹ The modified amino acid can be identified indirectly by treatment of (recombinant) protein with a single point-mutation,^{11, 82, 206} or by competitive labeling of the parent inhibitor with a validated residue-selective ABP (that is not derived from the parent inhibitor of interest).^{178, 207}

Reversible inhibition. ABPs bearing a reversible covalent warhead are compatible with gel-based analysis, as illustrated with cyanimide-based ABPs **IMP-1710**⁸⁴ and **8RK59**²⁰⁷ targeting deubiquitinating enzyme UCHL1 (**Figure 10B**). Importantly, reaction conditions required careful optimization as the fluorescent covalent adduct could not be detected after sample preparation under denaturing conditions: heating BODIPY-labeled UCHL1–8RK59 adduct to 94 °C in presence of reducing agent β -mercaptoethanol (BME) promoted covalent target disengagement.²⁰⁷ Subsequent chemoproteomic evaluation with **8RK64**, an alkyne-bearing derivative of 8RK59, revealed enrichment of not only UCHL1 but also protein deglycase PARK7/DJ1, an attractive target in Parkinson's disease with a similar molecular mass that overlaps with UCHL1 by gel analysis. The discovery of this off-target modification has since aided the development of selective chemical tools to study PARK7 activity.²⁰⁸

Quantification of covalent occupancy. Direct gel-based strategies are typically used for qualitative (visual) identification of binding partners in cellular proteomes as low throughput gel electrophoresis strategies are associated with large error margins originating from deviations in gel loading volumes and protein distribution on gel. Our group reported a direct quantitative approach to calculate relevant kinetic parameters from time-dependent covalent occupancy of purified recombinant cysteine protease USP16 with irreversible covalent Rho-Ub-ABPs.⁸² Cellular JAK3 occupancy after pretreatment with **ritlecitinib** (PF-06651600) was assessed with two-step ABP **PF-06789402**.¹⁹⁹ Lysate was treated with biotin/TAMRA-azide, enriched for ABP-modified uninhibited proteins by pulldown with streptavidin beads and resolved by SDS-PAGE gel electrophoresis to quantify the remaining uninhibited JAK3 by immunoblotting. A popular though indirect approach in preclinical development of BTK inhibitors is to derive

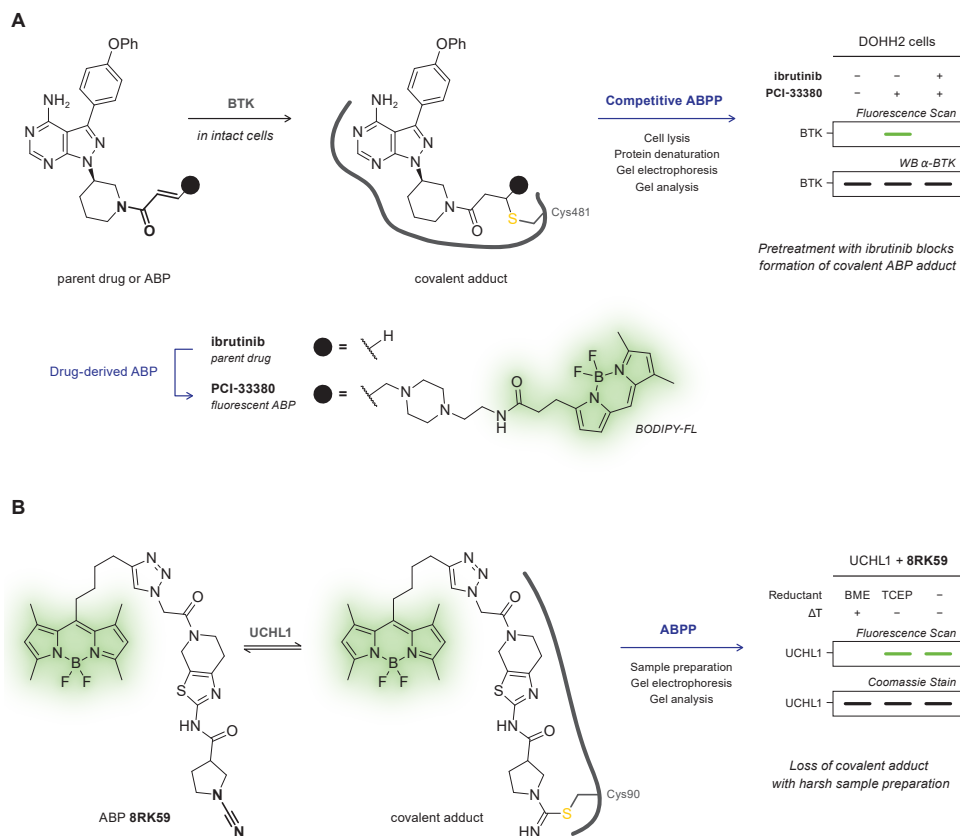


Figure 10 | Gel electrophoresis platforms for covalent adduct detection with drug-derived ABPs. **(A)** Competitive ABPP. Fluorescence scan for the BODIPY-FL fluorophore reveals that BTK labeling in DOHH2 cells by cell permeable one-step fluorescent ABP PCI-33380 is precluded by pretreatment with irreversible covalent parent inhibitor ibrutinib.¹¹ **(B)** Fluorescent signal originating from reversible covalent UCHL1–8RK59 adduct is not observed using harsh sample treatment prior to gel electrophoresis (boiling in presence of reducing agent BME) but can be observed using milder conditions (sample preparation in presence of reducing agent TCEP).²⁰⁷ Covalent adduct formation is validated using intact protein analysis by top-down MS.

inhibitor target engagement from the fluorescent ABP labeling of the remaining unbound protein: inhibitor-treated proteome is incubated with a target-selective fluorescent ABP (not necessarily derived from the parent inhibitor) that only binds to unbound BTK, and ABP-bound BTK is quantified by in-gel fluorescence. This approach has been successfully applied with irreversible BODIPY-labeled BTK-selective ABP **PRN-933** to assess occupancy of reversible covalent BTK inhibitor **rilzabrutinib** (PRN-1008) in human PBMCs,⁴⁹ and in the development of reversible covalent BTK inhibitors with irreversible ABP PP-BODIPY.⁵⁵ Competition with general thiol-reactive ABPs (e.g. IAc-alkyne, TMR-maleimide) is of little use in gel-based ABP analysis as blocking a single cysteine residue will not perturb the ABP from binding to other available cysteine residues in the same protein, thus still resulting in a detectable signal on gel. Importantly, blocking adduct formation with a selective irreversible ABP provides indirect

evidence on the ligand binding site⁷⁴ but is not suitable to identify the modified amino acid by itself as this also could be a(n) (allo)steric effect.

DNA electrophoretic mobility shift. Most covalent drugs are too small to cause a detectable shift in electrophoresis upon covalent adduct formation, thus requiring modification with a detection group. Nevertheless, a DNA electrophoretic mobility shift assay has been employed to validate the covalent binding mode of **lurbinectedin** (PM01183) to naked DNA, despite its relatively small mass (785 Da).²⁰⁹

Scintillation autoradiography (fluorography). Drug-derived ABPs bearing a radioisotope tag (e.g. ¹²⁵I) used to be the primary mode for detection of catalytically active cysteine proteases.²¹⁰⁻²¹¹ Nowadays, radiolabeled inhibitors prepared for *in vivo* ADME (Absorption, Distribution, Metabolism, and Excretion) studies and PK (pharmacokinetic) profiling are occasionally employed as radioactive ABPs where the radioactive atom (typically ¹⁴C or ³H) serves as a small, non-perturbing tag.²¹² Radioactivity originating from the radiolabeled covalent adduct is detected after removal of unbound and noncovalently bound radiolabeled inhibitor by gel electrophoresis on polyacrylamide/SDS-PAGE gels (fluorography),²¹³ or filtration with stringent washing (liquid scintillation counting).²¹³⁻²¹⁴ This technique has recently been employed to validate covalent adduct formation of **neratinib** (HKI-272) with HER2, using neratinib-derived ABP [¹⁴C]HKI-272 ([¹⁴C]-25o) on recombinant HER2 cytoplasmic domain or in intact BT474 cells.^{16, 215} Similarly, *in vivo* covalent alkylation of hemoglobin by **RRx-001** (ABDNAZ) in red blood cells from various species was detected using radiolabeled ABP [¹⁴C]RRx-001.²¹⁶ Finally, scientists at Takeda Pharmaceutical developed a direct competitive ligand binding assay to quantify covalent occupancy and determine the inhibitory potency (k_{inact}/K_i) of irreversible EGFR inhibitor **canertinib** (CI-1033): time-dependent radioactivity originating from the covalent adduct with radiolabeled ABP [³H]CI-1033 (after filtration to remove unbound ABP) was quantified by liquid scintillation counting.²¹⁴

6.2. Chemoproteomic Platforms

Chemoproteomic protein profiling is a sensitive MS-based ABPP technology predominantly employed to evaluate the proteome-wide selectivity of covalent inhibitors, and identify (undesired) cellular targets for covalent modification in complex mixtures (e.g. cell lysate, live cells, tissue).^{38, 104, 190-191, 217-219} A general chemoproteomic procedure involves incubation of the proteome with ABP, coupling to an enrichment handle (e.g. biotin-azide, **Figure 9C**) and pull-down to enrich for ABP-modified proteins on beads. Stringent washing is performed to remove noncovalently bound proteins and eliminate nonspecific binders, followed by bottom-up MS/MS analysis of modified proteins. Quantification of relative protein abundance can be achieved with label-free quantitative methods comparing relative changes in two (or more) individual biological samples.^{198, 219-221} The majority of chemoproteomic formats are indirect, detecting proteins modified by a general thiol-alkylating or cysteine ABP (e.g. IAc-alkyne)^{189, 222-223} in presence and absence of the covalent inhibitor of interest, from which inhibitor-binding is deduced.^{178, 187, 222} For this purpose, classic broad-spectrum reactivity ABPs targeting various amino acid residues and protein classes are (commercially) available.^{37-38, 186, 222} In particular, the isoTOP-ABPP (isotopic Tandem Orthogonal Activity-Based Protein Profiling) platform is

an established indirect competitive method to simultaneously identify (off-)target modified proteins in whole proteomes together with the exact site of protein modification.^{2, 190, 223–224} The role of predominantly indirect competition methods in (fragment-based) covalent drug development has been reviewed before.^{37–38, 225} Here, we will focus on the (less prevalent) direct approaches in which the ABP is derived from the parent inhibitor. Direct (competitive) chemoproteomic approaches with drug-derived ABPs have the potential to identify lower abundance protein targets and can overcome the bias in global cysteine reactivity experiments with general thiol-reactive ABPs: inhibitor binding is only detected if the amino acid residue is targeted by the competing ABP, even though the inhibitor might be interacting with other amino acid residues.^{103, 198} Such targetable, or ‘druggable’, cysteines in human proteins were recently collated in publicly available curated repository CysDB.²²⁶ The success of a direct approach is illustrated by FDA-approved anti-obesity drug **tetrahydrolipstatin** (THL, orlistat) (**Figure 11A**): MS analysis following pull-down of modified proteins in cancer cell lines treated with two-step ABP **THL-R** did not only confirm binding to fatty acid synthase (FAS) but also identified other (off-)target proteins that aid its early development as an anticancer agent.²⁰⁰

High-throughput proteomic methods may involve metabolic or chemical labeling with stable heavy isotopes prior to MS analysis to enable absolute protein quantification and multiplexed measurements (mix of multiple samples/reaction conditions) minimizing run-to-run deviations.^{221, 227} The popular SILAC-ABPP platform (**Figure 11B**) is a metabolic isotope-labeling methodology, thus being restricted to stable cell lines as generating isotope-labeled controls is challenging for tissue or primary cell line samples.²²⁸ SILAC-ABPP combines ABPP with SILAC (stable-isotope labeling by amino acids in cell culture)^{229–230} to assess identity of covalent modified proteins. Cells are cultured in normal (*light*) or isotope-labeled (*heavy*) medium, treated with DMSO or ABP, and mixed after lysis. Modified proteins are detected by bottom-up LC/LC-MS/MS analysis after enrichment for covalent protein–ABP adducts. The isotope-labeling of the proteome is crucial to calculate the SILAC ratio compared to the untreated sample – proteins with SILAC ratios ≥ 3 –5 are designated as targeted. SILAC-ABPP analysis with an ibrutinib-derived ABP identified established off-target kinases as well as specific non-kinase targets from structurally and functionally diverse protein families in Ramos cells, including the uncharacterized protein FAM213A.¹⁹⁰ Typically, complementary competitive SILAC-ABPP experiments are performed to ensure that the drug-derived ABP has the same selectivity as the parent inhibitor: over 400 proteins were identified in a SILAC experiment with an adagrasib-derived ABP (**Figure 11B**) but only KRAS^{G12C} significantly decreased upon pretreatment with clinically approved KRAS^{G12C} inhibitor **adagrasib** (MRTX849).¹⁰³

Chemical isotope-labeling methods are compatible with samples that are not amendable for SILAC, such as endogenous (human) tissue samples,¹⁰⁴ as chemical isotope-labeling can be performed during sample preparation.^{221, 232} These methods are typically indirect, employing classic broad-spectrum reactivity ABPs. Among the most popular methodologies are isoTOP-ABPP^{223–224, 233} using isotope-labeled TEV protease-cleavable Click reagents (TEV tags), isoDTB-ABPP employing isotopically-labeled desthiobiotin (isoDTB) tags,²³⁴ rdTOP-ABPP²³⁵ employing stable-isotope diMe labeling of primary amines (peptide N-terminus and lysine ϵ -amino group),^{227, 236–239} and TMT-ABPP²⁴⁰ employing tandem mass tags (TMT) such as

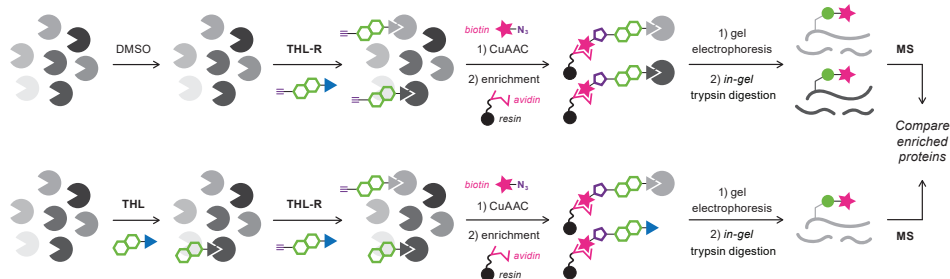
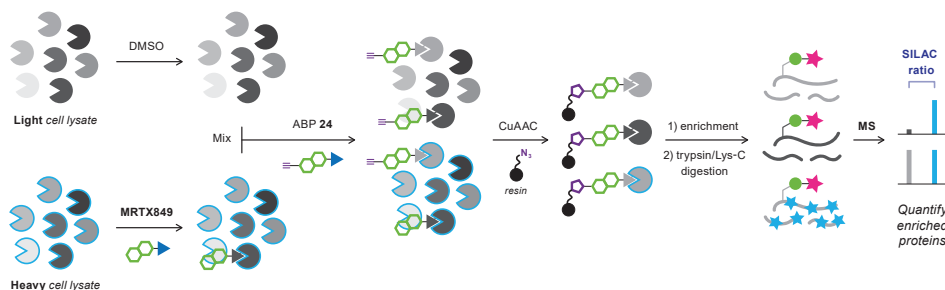
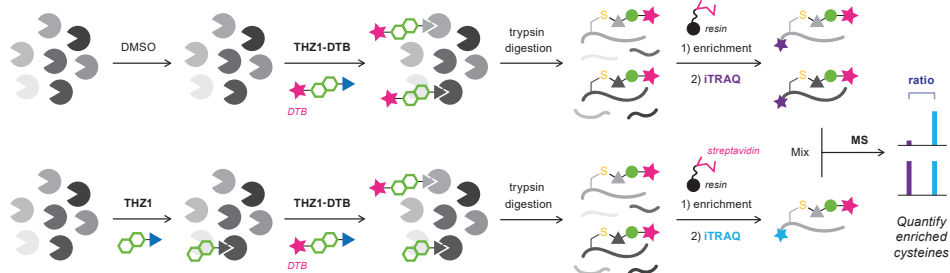
A Label-free Competitive ABPP (pull-down)**B Competitive SILAC-ABPP****C Covalent Inhibitor Target-site Identification (CiTe-ID)**

Figure 11 | Chemoproteomic approaches to identify covalently modified (off)-target proteins in whole proteomes. Pretreatment with inhibitor blocks drug-derived ABP binding and protein enrichment, resulting in a lower abundance of target protein compared to the DMSO-treated (control) sample. **(A)** Label-free protein target detection in pull-down experiment with drug-derived ABP THL-R to identify *in situ* protein targets of orlistat (THL) in HepG2 cells.²⁰⁰ ABP-bound proteins are enriched on avidin-agarose beads and submitted to bottom-up MS/MS evaluation. **(B)** Multiplexed detection of cellular protein targets of KRAS^{G12C} inhibitor adagrasib (MRTX849) in a competitive SILAC-ABPP experiment.¹⁰³ NCI-H358 cells cultured in normal (*light*) or isotope-labeled (*heavy*) medium (metabolic stable isotope-labeling) are incubated with DMSO or adagrasib, and the mixture of heavy and light lysate is then treated with drug-derived two-step ABP 24. ABP-bound proteins are enriched on azide-labeled agarose beads and submitted to bottom-up MS/MS for identification and relative quantitation of enriched protein abundance (SILAC ratio). **(C)** Identification and quantitation of novel protein targets for inhibitor THZ1 in a CiTe-ID experiment.²³¹ Cell lysates preincubated with DMSO (*control*) or THZ1 (*inhibitor*) are treated with drug-derived desthiobiotinylated ABP THZ1-DTB and enriched for DTB-modified proteolytic peptides. Primary amines are labeled with a unique isobaric iTRAQ reagent in each sample, and samples are combined for multiplexed RP-SAXRP MS/MS analysis.

isobaric amine-reactive iTRAQ or TMT™ multiplex tags.^{221, 241} Recently, the CITE-Id (Covalent Inhibitor Target-site Identification) platform was reported,²³¹ enabling unbiased identification and detection of modified proteins and inhibitor target site in the whole proteome by competing drug-derived desthiobiotinylated ABP with its parent inhibitor (**Figure 11C**). The success of this approach was illustrated with one-step ABP THZ1-DTB, a desthiobiotinylated analog of cyclin-dependent kinase (CDK7) inhibitor **THZ1**, focusing on the eight cysteine residues competitively modified by THZ1 in a dose-dependent manner.²³¹ Among the newly identified THZ1 targets was Cys840 of PKN3, and CITE-Id streamlined the development of first-in-class PKN3 inhibitor **JZ128**. Proteomic platforms to assess the global electrophile selectivity are under development,²⁴² as are improved competitive platforms to assess covalent inhibitor reactivity.²²⁵

Reversible inhibition and reversibility assays. Direct chemoproteomic assessment of cellular protein targets can be challenging for reversible covalent inhibitors, as proteolytic digestion can induce dissociation of a reversible covalent ABP. Successful identification of cellular targets for (slow) reversible covalent cyanimides targeting UCHL1 has been reported with biotinylated one-step ABP **11RK73**, and clickable two-step ABP **8RK64**.²⁰⁷ The proteome-wide reactivity of reversible inhibitors is typically assessed in competitive proteomic experiments, which is not necessarily restricted to covalent binding modes.^{239, 243} The Cravatt group demonstrated that it is not only possible to evaluate the proteome-wide reactivity of reversible covalent cysteine-directing compounds with competitive isoTOP-ABPP,²⁴⁴ but that adaptation of this method by introduction of a gel filtration (GF) step before treatment with the thiol-alkylating ABP can be employed to evaluate reversibility of the covalent adduct.

Quantification of covalent occupancy. Application of broad-spectrum ABPs to monitor ligand target engagement in native systems has been performed in model organisms (*in vivo* or *ex vivo*) and human tissue (*ex vivo*), with quantitation of relative protein abundance in presence of inhibitor compared to an untreated sample.^{104, 198} Occasionally, competitive ABPP experiments with drug-derived ABPs are performed. Time-dependent JAK3 occupancy in mouse spleen upon oral administration of **ritlecitinib** (PF-06651600) was monitored with drug-derived ABP PF-06789402: homogenized spleens were treated (*ex vivo*) with drug-derived ABP PF-06789402, enriched for ABP-modified proteins, and each sample was treated with a unique isobaric TMT-10plex tag to label (reactive) amines for multiplexed MS/MS analysis.¹⁹⁹

6.3. Homogeneous (Plate-based) Platforms

Gel electrophoresis (*section 6.1*) and chemoproteomic platforms (*section 6.2*) require removal of the unbound ABP or enrichment for modified proteins prior to detection of a covalent protein–ABP adduct. In this section we will discuss a few approaches that enable covalent adduct detection directly in a complex mixture (compatible with *in situ*/live cell imaging).

Traditional fluorescent ABPs suffer from a high fluorescence background as they are also fluorescent in their unbound state, and are thus less suitable for homogeneous applications that do not involve removal of the unbound ABP (e.g. live cell imaging, *in vitro* microplate-based activity assays). Turn-on fluorogenic and quenched ABPs are a subtype of fluorescent ABPs that only become fluorescent after covalent adduct formation. Quenched ABPs (qABPs) were

originally developed in the Bogoy lab to enable dynamic imaging of cysteine protease activity in living cells.²⁴⁵ Here, the fluorophore is 'dark' until the quencher is released or removed upon covalent thiol addition, generating a fluorescent covalent enzyme-inhibitor adduct. Adduct formation can be monitored by traditional in-gel fluorescence, but the low intrinsic fluorescent background also enables monitoring fluorescence intensity in homogeneous plate-based read-outs and even non-invasive real time optical imaging of cysteine protease activity in intact (live) cells, and even *in vivo*.²⁴⁶⁻²⁴⁸ Most qABPs targeting serines²⁴⁹⁻²⁵⁰ and cysteines^{182, 246-248, 251} were developed as chemical tools to study enzyme activity. Target-selective qABP typically have a peptidic recognition element with exception of a BTK kinase qABP derived from the noncovalent core of **ibrutinib** (Figure 12A).²⁵² The major drawback to turning a covalent ligand into a qABP is the mandatory replacement of the warhead with a suitable electrophile (e.g. acyloxymethyl ketone (AOMK),²⁴⁵ phenoxymethyl ketone (PMK),²⁴⁶ urea triazole^{249, 253}) to enable nucleophilic substitution of the quencher/fluorophore upon covalent adduct formation: the thiol-reactive electrophiles consisting of a carbonyl with a leaving group on the α -carbon may have a different intrinsic chemical reactivity than the original warhead,^{26-27, 254} and the qABP (thiol) reactivity may no longer be representative of the parent ligand.

A more generally applicable approach for thiol-reactive covalent small molecule inhibitors was recently reported by the London group:²⁵⁵ CoLDR (Covalent Ligand Directed Release) turn-on ABPs were generated by late-stage functionalization of covalent inhibitors containing the popular acrylamide warhead (Figure 12B). Modification of the acrylamide warhead on the α -carbon generates substituted α -methacrylamide warheads that release a (detectable) leaving group upon thiol addition turning the covalent inhibitor into a turn-on fluorogenic, chemiluminescent or otherwise functionalized ABP while maintaining the acrylamide geometry. This elegant approach was illustrated with a turn-on fluorogenic ABP based on covalent BTK inhibitor **ibrutinib**, that releases a fluorescent 7-hydroxycoumarin group upon thiol addition allowing homogeneous plate-based kinetic detection of irreversible covalent adduct formation.²⁵⁵ The versatility of this approach was illustrated with turn-on fluorogenic ABPs based on covalent EGFR inhibitor **afatinib**, covalent KRAS^{G12C} inhibitor **AMG-510** functionalized with coumarin, and a chemiluminescent **ibrutinib**-based ABP.²⁵⁵ Note: turn-on fluorescence probes do not have to be covalent as there are examples of increased fluorescence induced by a noncovalent binding event,²⁵⁶ and covalent adduct formation with the desired target should be validated with orthogonal techniques such as intact protein analysis. Alternatively, two-step ABPs can be employed as bioorthogonal fluorogenic probes, reacting the protein-ABP adduct with photophysically quenched fluorogenic dyes (e.g. azido-BODIPY, dibenzocyclooctyne) that are activated by Click chemistry.^{197, 257-258}

To date, qABPs and turn-on ABPs have limited clinical applications as the optical signal of most fluorophores is plagued by insufficient tissue penetration, thus obstructing their application as non-invasive diagnostic tools in living patients.²⁵⁹ Research endeavors in the qABP field have since progressed to advanced theranostic probes that combine detection and simultaneous inhibition of cathepsins with induction of sensitivity to Photodynamic Therapy (PDT) at the sites with high (aberrant) protease activity,²⁶⁰⁻²⁶¹ which may one day find application in non-invasive (clinical) diagnosis and treatment.

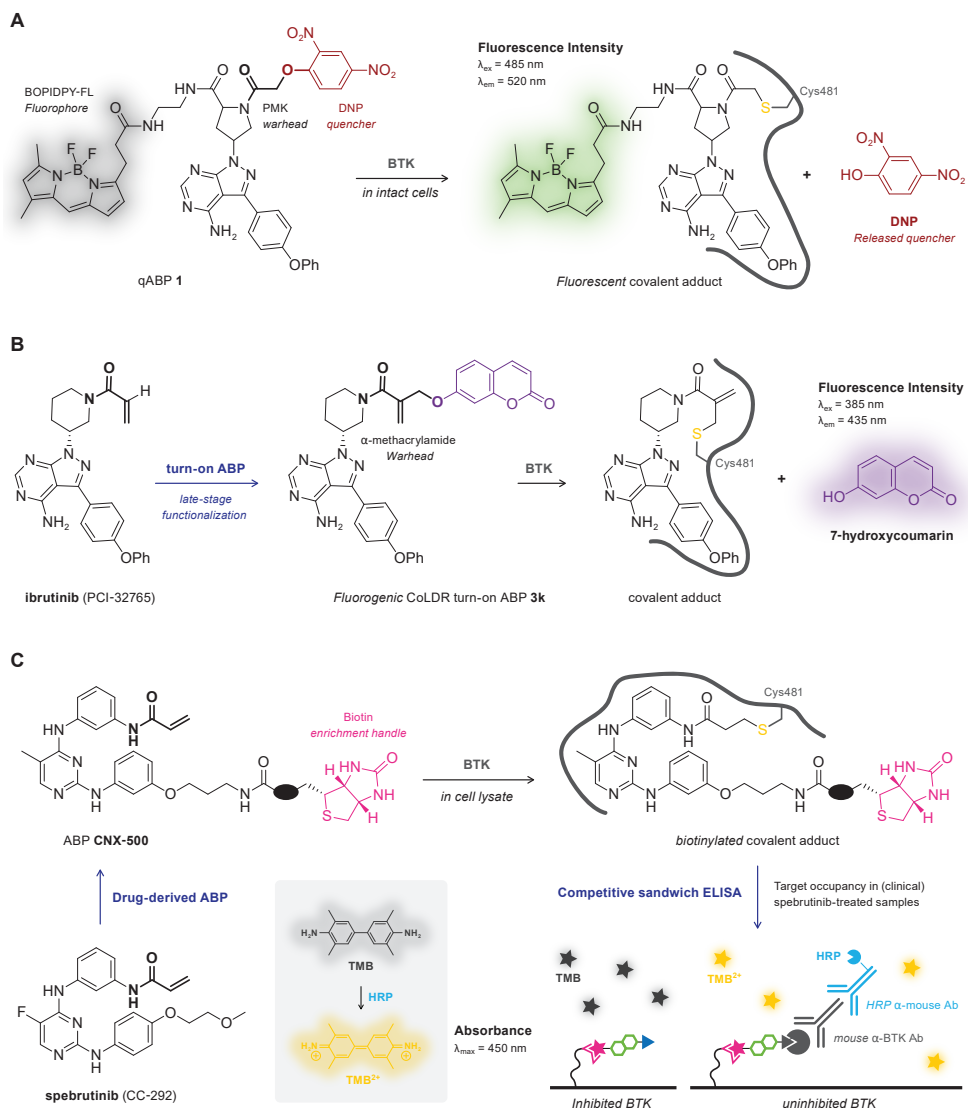


Figure 12 | Homogeneous (plate-based) approaches to detect covalent adduct formation with drug-derived ABPs. **(A)** Quenched fluorescent ABP (qABP) with a recognition element based on the ibrutinib scaffold to selectively target BTK.²⁵² The DNP (2,4-dinitrophenyl) quencher is expelled upon covalent adduct formation, enabling fluorescence detection of the BODIPY-FL fluorophore in the covalent adduct but not in unbound or noncovalently-bound qABP. **(B)** Covalent ligand directed release (CoLDR) chemistry to generate turn-on fluorogenic ABPs.²⁵⁵ Adduct formation of ibrutinib-derived turn-on fluorogenic ABP 3k with BTK is monitored by fluorescence intensity as thiol addition to the substituted α -methacrylamide warhead results in release of fluorescent 7-hydroxycoumarin. **(C)** Quantification of cellular BTK occupancy in a competitive ELISA (enzyme-linked immunosorbent assay) experiment.²⁶² Spebrutinib-treated lysates originating from tissue (culture) or clinical samples are incubated with biotinylated ABP CNX-500 to detect free, uninhibited BTK. Biotinylated BTK–ABP adducts are captured on streptavidin-coated ELISA plate, treated with primary mouse α -BTK Ab and secondary HRP anti-mouse Ab, and developed by addition of HRP substrate tetramethyl benzidine (TMB). Uninhibited BTK is quantified from the concentration of the yellow HRP product TMB^{2+} , with calculation of inhibitor occupancy from the OD_{450} in the treated samples relative to the untreated sample.

Quantification of covalent occupancy. Relative fluorescence intensity can be employed to quantify covalent adduct formation with quenched or turn-on fluorogenic ABPs.²⁵⁵ Recently, competitive approaches employing biotin-labeled ABPs to assess the cellular occupancy of covalent BTK inhibitors have been reported, where detection of the covalent BTK–ABP adduct is facilitated by an enzyme-linked immunosorbent assay (ELISA)^{12–13, 262} or an Amplified Luminescent Proximity Homogeneous Assay (ALPHA).⁵⁵ BTK occupancy is calculated from normalization of the signal in inhibitor-treated samples to the untreated control as covalent BTK–ABP adduct is formed with uninhibited BTK but not with BTK–inhibitor adducts. Cellular BTK occupancy of irreversible covalent BTK inhibitor **spebrutinib** (CC-292) in human B cell lysate was assessed using spebrutinib-derived biotinylated ABP **CNX-500** (**Figure 12C**),²⁶² with capture of the biotinylated BTK–ABP adducts on streptavidin-coated ELISA plates. Uninhibited BTK was quantified from the optical density (OD₄₅₀) originating from BTK–ABP adduct after subsequent addition of a primary BTK antibody, a secondary antibody modified with horseradish peroxidase (HRP), and development with the HRP substrate. The stepwise ELISA technology has been employed to assess BTK occupancy in the development of various covalent BTK inhibitors: **zanubrutinib** (BGB-3111) with zanubrutinib-derived biotinylated ABP **P-1** on neutravidin-coated ELISA plates,¹² and **acalabrutinib** (ACP-196) with acalabrutinib-derived biotinylated ABP ACP-4016 on BTK antibody-coated ELISA plates with HRP-linked streptavidin.¹³ The general BTK-selective biotinylated ABP S49 was employed rather than a drug-derived ABP to quantify cellular BTK occupancy of **remibrutinib** (LOU064).⁸⁷ Finally, the Taunton lab reports a high-throughput method using AlphaScreen technology based on competition with ibrutinib-derived ABP PP-biotin,⁵⁵ that does not only enable quantification of cellular BTK occupancy in Ramos cells but also elucidates inhibitor reversibility in washout experiments. An indirect competitive AlphaScreen methodology has since been used in the preclinical development of reversible BTK inhibitor **rilzabrutinib** (PRN-1008): BTK target occupancy in Ramos B cells was assessed using a BTK-selective biotinylated ABP.⁴⁹ Importantly, these methods do not provide direct evidence on inhibitor or ABP covalency: a stringent washing step promoting dissociation of noncovalent complexes has to be introduced to discriminate between noncovalent or covalent ligands. Nevertheless, these methodologies are attractive as they are complementary proteomic approaches by using the same biotin-labeled ABPs.

7. Conclusions, Current Challenges, and Future Directions

Biophysical detection of covalent adduct is an important step in covalent drug development as a(n) (ir)reversible covalent binding mode affects the SAR analysis and kinetic behavior.⁶¹ In this work, we reviewed the available methods for direct detection of the covalent protein–drug adduct, as opposed to deduction from the decrease of unbound drug. To ensure the detected signal originates from a covalent protein–drug adduct, covalent adduct formation is validated with at least two orthogonal methods. With a wide variety of techniques to choose from (**Table 1**), method selection is dictated by compatibility with (fluorescent) read-out, the available amount and purity of protein, the complexity of the reaction mixture (from purified recombinant protein to *in vivo*), and the desired level of information. Beyond the

simple detection of a covalent adduct, some techniques can aid identification of the targeted amino acid residue, but often protein mutagenesis is key to validate the modified amino acid residue.⁷⁰⁻⁷¹

This work includes most commonly used methods but is by no means complete. Although protein crystallography thus far remains the most informative method for structural evaluation of covalent adducts, there is a shift towards cryo-EM for the structural elucidation of large protein complexes.²⁶³ The phenomenal progress in the past years has resulted in highly detailed structures, wherein features of <2 Å can be resolved, but structure determination of small (<50 kDa) proteins is often hindered by intrinsically noisy micrographs and low image contrast.²⁶⁴⁻²⁶⁵ Currently, single-particle cryo-EM can successfully map small molecule ligands onto (large) proteins,²⁶⁶⁻²⁷⁰ making it a promising technique to also resolve covalent adducts or as an alternative for protein–drug complexes that are difficult to crystallize. Depending on the system studied, scientists may employ a plethora of different biophysical techniques. Importantly, covalent adduct formation may require identification of the reactive metabolite to assess covalent adduct formation with prodrugs (e.g. **omeprazole**).²⁷¹ Other challenging systems involve (membrane-bound) protein targets that are inactive in isolation, in absence of the other components of a protein complex. The majority of the described techniques are compatible with detection of reversible covalent adducts. The main challenge is to maintain intact protein–drug adduct during sample preparation as the sample preparation can induce target dissociation for reversible covalent ligands. The intrinsic property of a reversible covalent ligand to dissociate from its protein target upon protein denaturation^{54, 72} or chasing with an irreversible competitive tracer⁷⁴ has been exploited to assess binding reversibility of various clinically approved TCIs.^{88, 95}

The unavailability of (quantitative) high-throughput screening techniques can hamper widespread discovery of covalent ligands. Although intact protein MS can overcome this hurdle,⁹⁰ potential hits should always be validated in a functional assay,³⁹ as covalent modification does not necessarily mean altered protein function.⁷¹ Another factor is monitoring and quantifying covalent target occupancy in living patients, since drug levels in serum are not representative for irreversible binders. MS-based assays to quantify target engagement have been developed,¹⁰⁰ though practical application in clinical drug development is still limited by the optimization for each individual protein–drug pair.

Beyond validation of covalent adduct formation with the desired target, ABPP has the advantage that it can be used to evaluate target selectivity in a cellular, biological setting. Although detection of low-abundance proteins remains a challenge, the approach is in general very versatile, as evidenced by the various subtypes of ABPP. The prevalent indirect competitive ABPP with a general reactive ABP has successfully been employed to identify off-target protein targets in patient-derived samples,¹⁹¹ but this indirect approach is biased for the amino acid residues labeled by the general ABP and will miss labeling of other residues. Here, opportunities arise for the lower throughput use of drug-derived ABPs that have the same selectivity as the parent drug. Drug-derived ABPs have since been used to evaluate target selectivity of approved

covalent inhibitors,^{103, 200} quantify target occupancy in inhibitor-treated patients,^{199, 262} and identify novel targetable proteins.²⁰⁷ Further developments in ABP techniques are always limited by the deviation from the parent drug when a tag/handle is introduced. At the same time, general indirect (non-drug derived) methods to evaluate proteome-wide electrophile reactivity towards other nucleophilic residues (e.g. histidine, arginine, lysine)^{23, 242, 272-276} are becoming more prevalent as these nucleophilic residues are attractive targets (e.g. Arg12 in oncogenic mutant KRAS^{G12R},²⁷⁷ catalytic Lys271 in BCR-ABL²⁷⁸). All taken together, such chemoproteomic approaches will likely become an integral part of standard covalent drug development to identify covalently modified off-target proteins at an early stage,²⁷⁹ thereby further derisking covalent drug development.²

To conclude, there is a broad toolbox available for the evaluation and detection of covalent protein–drug adducts, ranging from recombinant protein to live patients. The techniques are instrumental in the evaluation of covalent drug reactivity and selectivity, and have guided covalent drug development programs and SAR optimization studies. Beyond the methods covered in this review, novel techniques will continue to be developed and improved to cater to the exciting and fast-paced field of covalent drug development.

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Author Contributions

E.M.: Conceptualization, Investigation, Visualization, Writing – Original Draft, Writing – Review & Editing. R.Q.K.: Writing – Review & Editing. M.P.C.M.: Supervision, Writing – Review & Editing.

8. References

1. Singh, J.; Petter, R.C.; Baillie, T.A.; Whitty, A. The Resurgence of Covalent Drugs. *Nat. Rev. Drug Discov.* **2011**, *10*, 307-317. doi: 10.1038/nrd3410.
2. Johnson, D.S.; Weerapana, E.; Cravatt, B.F. Strategies for Discovering and Derisking Covalent, Irreversible Enzyme Inhibitors. *Future Med. Chem.* **2010**, *2*, 949-964. doi: 10.4155/fmc.10.21.
3. Vane, J.R.; Botting, R.M. The Mechanism of Action of Aspirin. *Thromb. Res.* **2003**, *110*, 255-258. doi: 10.1016/S0049-3848(03)00379-7.
4. Yocum, R.R.; Rasmussen, J.R.; Strominger, J.L. The Mechanism of Action of Penicillin. Penicillin Acylates the Active Site of Bacillus Stearothermophilus D-Alanine Carboxypeptidase. *J. Biol. Chem.* **1980**, *255*, 3977-3986. doi: 10.1016/S0021-9258(19)85621-1.
5. Ding, Z.; Kim, S.; Dorsam, R.T.; Jin, J.; Kunapuli, S.P. Inactivation of the Human P2Y12 Receptor by Thiol Reagents Requires Interaction with Both Extracellular Cysteine Residues, Cys17 and Cys270. *Blood* **2003**, *101*, 3908-3914. doi: 10.1182/blood-2002-10-3027.
6. Shin, J.M.; Cho, Y.M.; Sachs, G. Chemistry of Covalent Inhibition of the Gastric (H⁺, K⁺)-ATPase by Proton Pump Inhibitors. *J. Am. Chem. Soc.* **2004**, *126*, 7800-7811. doi: 10.1021/ja049607w.
7. Bauer, R.A. Covalent Inhibitors in Drug Discovery: From Accidental Discoveries To Avoided Liabilities and Designed Therapies. *Drug Discov. Today* **2015**, *20*, 1061-1073. doi: 10.1016/j.drudis.2015.05.005.
8. Gehringer, M. Covalent Inhibitors: Back On Track? *Future Med. Chem.* **2020**, *12*, 1363-1368. doi: 10.4155/fmc-2020-0118.
9. Barf, T.; Kaptein, A. Irreversible Protein Kinase Inhibitors: Balancing the Benefits and Risks. *J. Med. Chem.* **2012**, *55*, 6243-6262. doi: 10.1021/jm3003203.
10. Chaikuad, A.; Koch, P.; Laufer, S.A.; Knapp, S. The Cysteinome of Protein Kinases as a Target in Drug Development. *Angew. Chem. Int. Ed.* **2018**, *57*, 4372-4385. doi: 10.1002/anie.201707875.

11. Honigberg, L.A.; Smith, A.M.; Sirisawad, M.; Verner, E.; Louny, D.; Chang, B.; Li, S.; Pan, Z.; Thamm, D.H.; Miller, R.A.; Buggy, J.J. The Bruton Tyrosine Kinase Inhibitor PCI-32765 Blocks B-cell Activation and is Efficacious in Models of Autoimmune Disease and B-cell Malignancy. *Proc. Natl. Acad. Sci.* **2010**, *107*, 13075–13080. doi: 10.1073/pnas.1004594107.
12. Guo, Y.; Liu, Y.; Hu, N.; Yu, D.; Zhou, C.; Shi, G.; Zhang, B.; Wei, M.; Liu, J.; Luo, L., et al. Discovery of Zanubrutinib (BGB-3111), a Novel, Potent, and Selective Covalent Inhibitor of Bruton's Tyrosine Kinase. *J. Med. Chem.* **2019**, *62*, 7923–7940. doi: 10.1021/acs.jmedchem.9b00687.
13. Barf, T.; Covey, T.; Izumi, R.; van de Kar, B.; Gulrajani, M.; van Lith, B.; van Hoek, M.; de Zwart, E.; Mittag, D.; Demont, D.; Verkaik, S.; Krantz, F.; Pearson, P.G.; Ulrich, R.; Kaptein, A. Acalabrutinib (ACP-196): A Covalent Bruton Tyrosine Kinase Inhibitor with a Differentiated Selectivity and In Vivo Potency Profile. *J. Pharmacol. Exp. Ther.* **2017**, *363*, 240. doi: 10.1124/jpet.117.242909.
14. Solca, F.; Dahl, G.; Zoephel, A.; Bader, G.; Sanderson, M.; Klein, C.; Kraemer, O.; Himmelsbach, F.; Haakma, E.; Adolf, G.R. Target Binding Properties and Cellular Activity of Afatinib (BIBW 2992), an Irreversible ErbB Family Blocker. *J. Pharmacol. Exp. Ther.* **2012**, *343*, 342–350. doi: 10.1124/jpet.112.197756.
15. Cross, D.A.E.; Ashton, S.E.; Ghioreghiu, S.; Eberlein, C.; Nebhan, C.A.; Spitzler, P.J.; Orme, J.P.; Finlay, M.R.V.; Ward, R.A.; Mellor, M.J., et al. AZD9291, an Irreversible EGFR TKI, Overcomes T790M-Mediated Resistance to EGFR Inhibitors in Lung Cancer. *Cancer Discov.* **2014**, *4*, 1046–1061. doi: 10.1158/2159-8290.Cd-14-0337.
16. Rabindran, S.K.; Discifani, C.M.; Rosfjord, E.C.; Baxter, M.; Floyd, M.B.; Golas, J.; Hallett, W.A.; Johnson, B.D.; Nilakantan, R.; Overbeek, E.; Reich, M.F.; Shen, R.; Shi, X.; Tsou, H.-R.; Wang, Y.-F.; Wissner, A. Antitumor Activity of HKI-272, an Orally Active, Irreversible Inhibitor of the HER-2 Tyrosine Kinase. *Cancer Res.* **2004**, *64*, 3958–3965. doi: 10.1158/0008-5472.CAN-03-2868.
17. Gonzales, A.J.; Hook, K.E.; Althaus, I.W.; Ellis, P.A.; Trachet, E.; Delaney, A.M.; Harvey, P.J.; Ellis, T.A.; Amato, D.M.; Nelson, J.M., et al. Antitumor Activity and Pharmacokinetic Properties of PF-00299804, a Second-Generation Irreversible Pan-erbB Receptor Tyrosine Kinase Inhibitor. *Mol. Cancer Ther.* **2008**, *7*, 1880–1889. doi: 10.1158/1535-7163.Mct-07-2232.
18. Lanman, B.A.; Allen, J.R.; Allen, J.G.; Amegadzie, A.K.; Ashton, K.S.; Booker, S.K.; Chen, J.J.; Chen, N.; Frohn, M.J.; Goodman, G., et al. Discovery of a Covalent Inhibitor of KRAS^{G12C} (AMG 510) for the Treatment of Solid Tumors. *J. Med. Chem.* **2020**, *63*, 52–65. doi: 10.1021/acs.jmedchem.9b01180.
19. Gonzalez, F.; Vincent, S.; Baker, T.E.; Gould, A.E.; Li, S.; Wardwell, S.D.; Nadworny, S.; Ning, Y.; Zhang, S.; Huang, W.-S., et al. Mobocertinib (TAK-788): A Targeted Inhibitor of EGFR Exon 20 Insertion Mutants in Non-Small Cell Lung Cancer. *Cancer Discov.* **2021**, *11*, 1672–1687. doi: 10.1158/2159-8290.Cd-20-1683.
20. De Vita, E. 10 Years into the Resurgence of Covalent Drugs. *Future Med. Chem.* **2021**, *13*, 193–210. doi: 10.4155/fmc-2020-0236.
21. Ward, R.A.; Grimster, N.P. The Design of Covalent-Based Inhibitors. *Annu. Rep. Med. Chem.* **2021**, *56*, 2–284. <https://www.sciencedirect.com/bookseries/annual-reports-in-medicinal-chemistry/vol/56/suppl/C>
22. Borsari, C.; Keles, E.; McPhail, J.A.; Schaefer, A.; Sriramaratnam, R.; Goch, W.; Schaefer, T.; De Pascale, M.; Bal, W.; Gstaiger, M.; Burke, J.E.; Wymann, M.P. Covalent Proximity Scanning of a Distal Cysteine to Target PI3Kα. *J. Am. Chem. Soc.* **2022**, *144*, 6326–6342. doi: 10.1021/jacs.1c13568.
23. Gehring, M.; Laufer, S.A. Emerging and Re-Emerging Warheads for Targeted Covalent Inhibitors: Applications in Medicinal Chemistry and Chemical Biology. *J. Med. Chem.* **2019**, *62*, 5673–5724. doi: 10.1021/acs.jmedchem.8b01153.
24. Shindo, N.; Ojida, A. Recent Progress in Covalent Warheads for In Vivo Targeting of Endogenous Proteins. *Bioorg. Med. Chem.* **2021**, *47*, 116386. doi: 10.1016/j.bmc.2021.116386.
25. Ray, S.; Murkin, A.S. New Electrophiles and Strategies for Mechanism-Based and Targeted Covalent Inhibitor Design. *Biochemistry* **2019**, *58*, 5234–5244. doi: 10.1021/acs.biochem.9b00293.
26. Martin, J.S.; MacKenzie, C.J.; Fletcher, D.; Gilbert, I.H. Characterising Covalent Warhead Reactivity. *Bioorg. Med. Chem.* **2019**, *27*, 2066–2074. doi: 10.1016/j.bmc.2019.04.002.
27. Lonsdale, R.; Burgess, J.; Colclough, N.; Davies, N.L.; Lenz, E.M.; Orton, A.L.; Ward, R.A. Expanding the Armory: Predicting and Tuning Covalent Warhead Reactivity. *J. Chem. Inf. Model.* **2017**, *57*, 3124–3137. doi: 10.1021/acs.jcim.7b00553.
28. Guan, I.; Williams, K.; Pan, J.; Liu, X. New Cysteine Covalent Modification Strategies Enable Advancement of Proteome-wide Selectivity of Kinase Modulators. *Asian J. Org. Chem.* **2021**, *10*, 949–963. doi: 10.1002/ajoc.202100036.
29. Oballa, R.M.; Truchon, J.-F.; Bayly, C.I.; Chaurat, N.; Day, S.; Crane, S.; Berthelette, C. A Generally Applicable Method for Assessing the Electrophilicity and Reactivity of Diverse Nitrile-Containing Compounds. *Bioorg. Med. Chem. Lett.* **2007**, *17*, 998–1002. doi: 10.1016/j.bmcl.2006.11.044.
30. Péczka, N.; Orgován, Z.; Ábrányi-Balogh, P.; Keserű, G.M. Electrophilic Warheads in Covalent Drug Discovery: An Overview. *Expert Opin. Drug Discov.* **2022**, *17*, 413–422. doi: 10.1080/17460441.2022.2034783.
31. McAulay, K.; Bilsland, A.; Bon, M. Reactivity of Covalent Fragments and Their Role in Fragment Based Drug Discovery. *Pharmaceuticals* **2022**, *15*, 1366. doi: 10.3390/ph15111366.
32. De Cesco, S.; Kurian, J.; Dufresne, C.; Mittermaier, A.K.; Moitessier, N. Covalent Inhibitors Design and Discovery. *Eur. J. Med. Chem.* **2017**, *138*, 96–114. doi: 10.1016/j.ejmech.2017.06.019.
33. Engel, J.; Richters, A.; Getlik, M.; Tomassi, S.; Keul, M.; Termathe, M.; Lategahn, J.; Becker, C.; Mayer-Wrangowski, S.; Grütter, C., et al. Targeting Drug Resistance in EGFR with Covalent Inhibitors: A Structure-Based Design Approach. *J. Med. Chem.* **2015**, *58*, 6844–6863. doi: 10.1021/acs.jmedchem.5b01082.
34. Zhang, T.; Hatcher, J.M.; Teng, M.; Gray, N.S.; Kostic, M. Recent Advances in Selective and Irreversible Covalent Ligand Development and Validation. *Cell Chem. Biol.* **2019**, *26*, 1486–1500. doi: 10.1016/j.chembiol.2019.09.012.
35. Chen, S.; Lovell, S.; Lee, S.; Fellner, M.; Mace, P.D.; Bogoy, M. Identification of Highly Selective Covalent Inhibitors by Phage Display. *Nat. Biotechnol.* **2021**, *39*, 490–498. doi: 10.1038/s41587-020-0733-7.
36. Campuzano, I.D.G.; San Miguel, T.; Rowe, T.; Onea, D.; Cee, V.J.; Arvedson, T.; McCarter, J.D. High-Throughput Mass Spectrometric Analysis of Covalent Protein-Inhibitor Adducts for the Discovery of Irreversible Inhibitors: A Complete Workflow. *J. Biomol. Screen.* **2015**, *21*, 136–144. doi: 10.1177/1087057115621288.
37. Dalton, S.E.; Campos, S. Covalent Small Molecules as Enabling Platforms for Drug Discovery. *ChemBioChem* **2020**, *21*, 1080–1100. doi: 10.1002/cbic.201900674.
38. Lu, W.; Kostic, M.; Zhang, T.; Che, J.; Patricelli, M.P.; Jones, L.H.; Chouchani, E.T.; Gray, N.S. Fragment-Based Covalent Ligand Discovery. *RSC Chem. Biol.* **2021**, *2*, 354–367. doi: 10.1039/D0CB00222D.
39. Keeley, A.; Petri, L.; Ábrányi-Balogh, P.; Keserű, G.M. Covalent Fragment Libraries in Drug Discovery. *Drug Discov. Today* **2020**, *25*, 983–996. doi: 10.1016/j.drudis.2020.03.016.
40. Miller, R.M.; Paavilainen, V.O.; Krishnan, S.; Serafimova, I.M.; Taunton, J. Electrophilic Fragment-Based Design of Reversible Covalent Kinase Inhibitors. *J. Am. Chem. Soc.* **2013**, *135*, 5298–5301. doi: 10.1021/ja401221b.

41. Resnick, E.; Bradley, A.; Gan, J.; Douangamath, A.; Krojer, T.; Sethi, R.; Geurink, P.P.; Aimon, A.; Amitai, G.; Bellini, D., et al. Rapid Covalent-Probe Discovery by Electrophile-Fragment Screening. *J. Am. Chem. Soc.* **2019**, *141*, 8951-8968. doi: 10.1021/jacs.9b02822.
42. Kathman, S.G.; Statsyuk, A.V. Methodology for Identification of Cysteine-Reactive Covalent Inhibitors. *Methods Mol. Biol.* **2019**, *1967*, 245-262. doi: 10.1007/978-1-4939-9187-7_15.
43. Kathman, S.G.; Statsyuk, A.V. Covalent Tethering of Fragments for Covalent Probe Discovery. *MedChemComm* **2016**, *7*, 576-585. doi: 10.1039/C5MD00518C.
44. Kumalo, H.M.; Bhakat, S.; Soliman, M.E.S. Theory and Applications of Covalent Docking in Drug Discovery: Merits and Pitfalls. *Molecules* **2015**, *20*, 1984-2000. doi: 10.3390/molecules20021984.
45. Lonsdale, R.; Ward, R.A. Structure-Based Design of Targeted Covalent Inhibitors. *Chem. Soc. Rev.* **2018**, *47*, 3816-3830. doi: 10.1039/C7CS00220C.
46. Zaidman, D.; Gehrtz, P.; Filep, M.; Fearon, D.; Gabizon, R.; Douangamath, A.; Prilusky, J.; Duberstein, S.; Cohen, G.; Owen, C.D.; Resnick, E.; Strain-Damerell, C.; Lukacik, P.; Barr, H.; Walsh, M.A.; von Delft, F.; London, N. An Automatic Pipeline for the Design of Irreversible Derivatives Identifies a Potent SARS-CoV-2 M^{pro} Inhibitor. *Cell Chem. Biol.* **2021**, *28*, 1795-1806.e5. doi: 10.1016/j.chembiol.2021.05.018.
47. Owen, D.R.; Allerton, C.M.N.; Anderson, A.S.; Aschenbrenner, L.; Avery, M.; Berritt, S.; Boras, B.; Cardin, R.D.; Carlo, A.; Coffman, K.J., et al. An Oral SARS-CoV-2 M^{pro} Inhibitor Clinical Candidate for the Treatment of COVID-19. *Science* **2021**, *374*, 1586-1593. doi: 10.1126/science.abc4784.
48. Tuley, A.; Fast, W. The Taxonomy of Covalent Inhibitors. *Biochemistry* **2018**, *57*, 3326-3337. doi: 10.1021/acs.biochem.8b00315.
49. Langrish, C.L.; Bradshaw, J.M.; Francesco, M.R.; Owens, T.D.; Xing, Y.; Shu, J.; LaStant, J.; Bisconte, A.; Outerbridge, C.; White, S.D.; Hill, R.J.; Brameld, K.A.; Goldstein, D.M.; Nunn, P.A. Preclinical Efficacy and Anti-Inflammatory Mechanisms of Action of the Bruton Tyrosine Kinase Inhibitor Rilzabrutinib for Immune-Mediated Disease. *J. Immunol.* **2021**, *206*, 1454-1468. doi: 10.4049/jimmunol.2001130.
50. Gauthier, J.Y.; Charet, N.; Cromlish, W.; Desmarais, S.; Duong, L.T.; Falguyet, J.-P.; Kimmel, D.B.; Lamontagne, S.; Léger, S.; LeRiche, T.; Li, C.S., et al. The Discovery of Odanacatib (MK-0822), a Selective Inhibitor of Cathepsin K. *Bioorg. Med. Chem. Lett.* **2008**, *18*, 923-928. doi: 10.1016/j.bmcl.2007.12.047.
51. Groll, M.; Berkers, C.R.; Ploegh, H.L.; Ova, H. Crystal Structure of the Boronic Acid-Based Proteasome Inhibitor Bortezomib in Complex with the Yeast 20S Proteasome. *Structure* **2006**, *14*, 451-456. doi: 10.1016/j.str.2005.11.019.
52. Perni, R.B.; Almqvist, S.J.; Byrn, R.A.; Chandorkar, G.; Chaturvedi, P.R.; Courtney, L.F.; Decker, C.J.; Dinehart, K.; Gates, C.A.; Harbeson, S.L., et al. Preclinical Profile of VX-950, a Potent, Selective, and Orally Bioavailable Inhibitor of Hepatitis C Virus NS3-4A Serine Protease. *Antimicrob. Agents Chemother.* **2006**, *50*, 899-909. doi: 10.1128/AAC.50.3.899-909.2006.
53. Romano, K.P.; Ali, A.; Aydin, C.; Soumana, D.; Özen, A.; Deveau, L.M.; Silver, C.; Cao, H.; Newton, A.; Petropoulos, C.J.; Huang, W.; Schiffer, C.A. The Molecular Basis of Drug Resistance against Hepatitis C Virus NS3/4A Protease Inhibitors. *PLoS Pathog.* **2012**, *8*, e1002832. doi: 10.1371/journal.ppat.1002832.
54. Serafimova, I.M.; Pufall, M.A.; Krishnan, S.; Duda, K.; Cohen, M.S.; Maglathlin, R.L.; McFarland, J.M.; Miller, R.M.; Frödin, M.; Taunton, J. Reversible Targeting of Noncatalytic Cysteines with Chemically Tuned Electrophiles. *Nat. Chem. Biol.* **2012**, *8*, 471-476. doi: 10.1038/nchembio.925.
55. Bradshaw, J.M.; McFarland, J.M.; Paavilainen, V.O.; Bisconte, A.; Tam, D.; Phan, V.T.; Romanov, S.; Finkle, D.; Shu, J.; Patel, V., et al. Prolonged and Tunable Residence Time using Reversible Covalent Kinase Inhibitors. *Nat. Chem. Biol.* **2015**, *11*, 525-531. doi: 10.1038/nchembio.1817.
56. Krenke, E.H.; Petter, R.C.; Houk, K.N. Kinetics and Thermodynamics of Reversible Thiol Additions to Mono- and Diactivated Michael Acceptors: Implications for the Design of Drugs That Bind Covalently to Cysteines. *J. Org. Chem.* **2016**, *81*, 11726-11733. doi: 10.1021/acs.joc.6b02188.
57. Basu, D.; Richters, A.; Rauh, D. Structure-Based Design and Synthesis of Covalent-Reversible Inhibitors to Overcome Drug Resistance in EGFR. *Bioorg. Med. Chem.* **2015**, *23*, 2767-2780. doi: 10.1016/j.bmc.2015.04.038.
58. Forster, M.; Gehringer, M.; Laufer, S.A. Recent Advances in JAK3 Inhibition: Isoform Selectivity by Covalent Cysteine Targeting. *Bioorg. Med. Chem. Lett.* **2017**, *27*, 4229-4237. doi: 10.1016/j.bmcl.2017.07.079.
59. U.S. Food & Drug Administration. Coronavirus (COVID-19) Update: FDA Authorizes First Oral Antiviral for Treatment of COVID-19. *FDA News Release* [Online], 22 December 2021. <https://www.fda.gov/news-events/press-announcements/coronavirus-covid-19-update-fda-authorizes-first-oral-antiviral-treatment-covid-19>.
60. Strelow, J.M. A Perspective on the Kinetics of Covalent and Irreversible Inhibition. *SLAS Discov.: Adv. Life Sci. R&D* **2017**, *22*, 3-20. doi: 10.1177/1087057116671509.
61. Mons, E.; Roet, S.; Kim, R.Q.; Mulder, M.P.C. A Comprehensive Guide for Assessing Covalent Inhibition in Enzymatic Assays Illustrated with Kinetic Simulations. *Curr. Protoc.* **2022**, *2*, e419. doi: 10.1002/cpz1.419.
62. McWhirter, C. Chapter One - Kinetic Mechanisms of Covalent Inhibition. *Annu. Rep. Med. Chem.* **2021**, *56*, 1-31. doi: 10.1016/bs.armc.2020.11.001.
63. Harris, C.M.; Foley, S.E.; Goedken, E.R.; Michalak, M.; Murdock, S.; Wilson, N.S. Merits and Pitfalls in the Characterization of Covalent Inhibitors of Bruton's Tyrosine Kinase. *SLAS Discov.: Adv. Sci. Drug Discov.* **2018**, *23*, 1040-1050. doi: 10.1177/2472555218787445.
64. Al-Khafaji, K.; Al-Duaidahawi, D.; Taskin Tok, T. Using Integrated Computational Approaches to Identify Safe and Rapid Treatment for SARS-CoV-2. *J. Biomol. Struct. Dyn.* **2021**, *39*, 3387-3395. doi: 10.1080/07391102.2020.1764392.
65. Yang, X.; Dilweg, M.A.; Osemwengie, D.; Burggraaf, L.; van der Es, D.; Heitman, L.H.; Ijzerman, A.P. Design and Pharmacological Profile of a Novel Covalent Partial Agonist for the Adenosine A1 Receptor. *Biochem. Pharmacol.* **2020**, *180*, 114144. doi: 10.1016/j.bcp.2020.114144.
66. Yang, X.; van Veldhoven, J.P.D.; Offringa, J.; Kuiper, B.J.; Lenselink, E.B.; Heitman, L.H.; van der Es, D.; Ijzerman, A.P. Development of Covalent Ligands for G Protein-Coupled Receptors: A Case for the Human Adenosine A3 Receptor. *J. Med. Chem.* **2019**, *62*, 3539-3552. doi: 10.1021/acs.jmedchem.8b02026.
67. Weichert, D.; Kruse, A.C.; Manglik, A.; Hiller, C.; Zhang, C.; Hübner, H.; Kobilka, B.K.; Grmeiner, P. Covalent Agonists for Studying G Protein-Coupled Receptor Activation. *Proc. Natl. Acad. Sci.* **2014**, *111*, 10744-10748. doi: 10.1073/pnas.1410415111.
68. Grimster, N.P. Covalent PROTACs: The Best of Both Worlds? *RSC Med. Chem.* **2021**, *12*, 1452-1458. doi: 10.1039/D1MD00191D.
69. Gabizon, R.; Shraga, A.; Gehrtz, P.; Livnah, E.; Shorer, Y.; Gurwicz, N.; Avram, L.; Unger, T.; Aharoni, H.; Albeck, S.; Brandis, A.; Shulman, Z.; Katz, B.-Z.; Herishanu, Y.; London, N. Efficient Targeted Degradation via Reversible and Irreversible Covalent PROTACs. *J. Am. Chem. Soc.* **2020**, *142*, 11734-11742. doi: 10.1021/jacs.9b13907.
70. Li, Y.-J.; Du, L.; Wang, J.; Vega, R.; Lee, T.D.; Miao, Y.; Aldana-Masangkay, G.; Samuels, E.R.; Li, B.; Ouyang, S.X.; Colayco, S.A.; Bobkova, E.V.; Divlianska, D.B.; Sergienko, E.; Chung, T.D.X.; Fakhri, M.; Chen, Y. Allosteric Inhibition of Ubiquitin-like Modifications by a Class of Inhibitor of SUMO-Activating Enzyme. *Cell Chem. Biol.* **2019**, *26*, 278-288.e6. doi: 10.1016/j.chembiol.2018.10.026.
71. Pettinger, J.; Le Bihan, Y.-V.; Widya, M.; van Montfort, R.L.M.; Jones, K.; Cheeseman, M.D. An Irreversible Inhibitor of HSP72 that Unexpectedly Targets Lysine-56. *Angew. Chem. Int. Ed.* **2017**, *56*, 3536-3540. doi: 10.1002/anie.201611907.
72. Abdeldayem, A.; Raouf, Y.S.; Constantinescu, S.N.; Moriggi, R.; Gunning, P.T. Advances in Covalent Kinase Inhibitors. *Chem. Soc. Rev.* **2020**, *49*, 2617-2687. doi: 10.1039/C9CS00720B.

73. Copeland, R.A.; Basavapathruni, A.; Moyer, M.; Scott, M.P. Impact of Enzyme Concentration and Residence Time on Apparent Activity Recovery in Jump Dilution Analysis. *Anal. Biochem.* **2011**, *416*, 206–210. doi: 10.1016/j.ab.2011.05.029.
74. Smith, S.; Keul, M.; Engel, J.; Basu, D.; Eppmann, S.; Rauh, D. Characterization of Covalent-Reversible EGFR Inhibitors. *ACS Omega* **2017**, *2*, 1563–1575. doi: 10.1021/acsomega.7b00157.
75. Tailor, A.; Waddington, J.C.; Meng, X.; Park, B.K. Mass Spectrometric and Functional Aspects of Drug–Protein Conjugation. *Chem. Res. Toxicol.* **2016**, *29*, 1912–1935. doi: 10.1021/acs.chemrestox.6b00147.
76. Zhang, Y.; Fonslow, B.R.; Shan, B.; Baek, M.-C.; Yates, J.R. Protein Analysis by Shotgun/Bottom-Up Proteomics. *Chem. Rev.* **2013**, *113*, 2343–2394. doi: 10.1021/cr3003533.
77. Donnelly, D.P.; Rawlins, C.M.; DeHart, C.J.; Fornelli, L.; Schachner, L.F.; Lin, Z.; Lippens, J.L.; Aluri, K.C.; Sarin, R.; Chen, B., et al. Best Practices and Benchmarks for Intact Protein Analysis for Top-Down Mass Spectrometry. *Nat. Meth.* **2019**, *16*, 587–594. doi: 10.1038/s41592-019-0457-0.
78. Tokmina-Lukaszewska, M.; Patterson, A.; Berry, L.; Scott, L.; Balasubramanian, N.; Bothner, B. The Role of Mass Spectrometry in Structural Studies of Flavin-Based Electron Bifurcating Enzymes. *Front. Microbiol.* **2018**, *9*, 1397. doi: 10.3389/fmicb.2018.01397.
79. Hilton, G.R.; Benesch, J.L.P. Two Decades of Studying Non-covalent Biomolecular Assemblies by Means of Electrospray Ionization Mass Spectrometry. *J. R. Soc. Interface* **2012**, *9*, 801–816. doi: 10.1098/rsif.2011.0823.
80. Han, X.; Jin, M.; Breuker, K.; McLafferty, F.W. Extending Top-Down Mass Spectrometry to Proteins with Masses Greater Than 200 Kilodaltons. *Science* **2006**, *314*, 109–112. doi: 10.1126/science.1128868.
81. Melby, J.A.; Roberts, D.S.; Larson, E.J.; Brown, K.A.; Bayne, E.F.; Jin, S.; Ge, Y. Novel Strategies to Address the Challenges in Top-Down Proteomics. *J. Am. Soc. Mass Spectrom.* **2021**, *32*, 1278–1294. doi: 10.1021/jasms.1c00099.
82. Mons, E.; Kim, R.Q.; van Doodewaerd, B.R.; van Veelen, P.A.; Mulder, M.P.C.; Ovaa, H. Exploring the Versatility of the Covalent Thiol–Alkyne Reaction with Substituted Propargyl Warheads: A Deciding Role for the Cysteine Protease. *J. Am. Chem. Soc.* **2021**, *143*, 6423–6433. doi: 10.1021/jacs.0c10513.
83. Mons, E.; Jansen, I.D.C.; Loboda, J.; van Doodewaerd, B.R.; Hermans, J.; Verdoes, M.; van Boeckel, C.A.A.; van Veelen, P.A.; Turk, B.; Turk, D.; Ovaa, H. The Alkyne Moiety as a Latent Electrophile in Irreversible Covalent Small Molecule Inhibitors of Cathepsin K. *J. Am. Chem. Soc.* **2019**, *141*, 3507–3514. doi: 10.1021/jacs.8b11027.
84. Panyain, N.; Godinat, A.; Lanyon-Hogg, T.; Lachiondo-Ortega, S.; Will, E.J.; Soudy, C.; Mondal, M.; Mason, K.; Elkhalfa, S.; Smith, L.M.; Harrigan, J.A.; Tate, E.W. Discovery of a Potent and Selective Covalent Inhibitor and Activity-Based Probe for the Deubiquitylating Enzyme UCHL1, with Antifibrotic Activity. *J. Am. Chem. Soc.* **2020**, *142*, 12020–12026. doi: 10.1021/jacs.0c04527.
85. Kathman, S.G.; Xu, Z.; Statsyuk, A.V. A Fragment-Based Method to Discover Irreversible Covalent Inhibitors of Cysteine Proteases. *J. Med. Chem.* **2014**, *57*, 4969–4974. doi: 10.1021/jm500345q.
86. Caldwell, R.D.; Qiu, H.; Askew, B.C.; Bender, A.T.; Brugger, N.; Camps, M.; Dhanabal, M.; Dutt, V.; Eichhorn, T.; Gardberg, A.S., et al. Discovery of Evobrutinib: An Oral, Potent, and Highly Selective, Covalent Bruton's Tyrosine Kinase (BTK) Inhibitor for the Treatment of Immunological Diseases. *J. Med. Chem.* **2019**, *62*, 7643–7655. doi: 10.1021/acs.jmedchem.9b00794.
87. Angst, D.; Gessier, F.; Janser, P.; Vulpetti, A.; Wälchli, R.; Beerli, C.; Littlewood-Evans, A.; Dawson, J.; Nuesslein-Hildesheim, B.; Wieczorek, G.; Gutmann, S.; Scheufler, C.; Hinniger, A.; Zimmerlin, A.; Funhoff, E.G.; Pulz, R.; Cenni, B. Discovery of LOU064 (Remibrutinib), a Potent and Highly Selective Covalent Inhibitor of Bruton's Tyrosine Kinase. *J. Med. Chem.* **2020**, *63*, 5102–5118. doi: 10.1021/acs.jmedchem.9b01916.
88. Liclican, A.; Serafini, L.; Xing, W.; Czerwieniec, G.; Steiner, B.; Wang, T.; Brendza, K.M.; Lutz, J.D.; Keegan, K.S.; Ray, A.S.; Schultz, B.E.; Sakowicz, R.; Feng, J.Y. Biochemical Characterization of Tirabrutinib and Other Irreversible Inhibitors of Bruton's Tyrosine Kinase Reveals Differences in On- and Off - Target Inhibition. *Biochim. Biophys. Acta, Gen. Subj.* **2020**, *1864*, 129531. doi: 10.1016/j.bbagen.2020.129531.
89. Patricelli, M.P.; Janes, M.R.; Li, L.-S.; Hansen, R.; Peters, U.; Kessler, L.V.; Chen, Y.; Kucharski, J.M.; Feng, J.; Ely, T.; Chen, J.H.; Firdaus, S.J.; Babbar, A.; Ren, P.; Liu, Y. Selective Inhibition of Oncogenic KRAS Output with Small Molecules Targeting the Inactive State. *Cancer Discov.* **2016**, *6*, 316–329. doi: 10.1158/2159-8290.Cd-15-1105.
90. Johansson, H.; Tsai, Y.-C.I.; Fantom, K.; Chung, C.-W.; Kümpfer, S.; Martino, L.; Thomas, D.A.; Eberl, H.C.; Muelbauer, M.; House, D.; Rittinger, K. Fragment-Based Covalent Ligand Screening Enables Rapid Discovery of Inhibitors for the RBR E3 Ubiquitin Ligase HOIP. *J. Am. Chem. Soc.* **2019**, *141*, 2703–2712. doi: 10.1021/jacs.8b13193.
91. Kathman, S.G.; Span, I.; Smith, A.T.; Xu, Z.; Zhan, J.; Rosenzweig, A.C.; Statsyuk, A.V. A Small Molecule That Switches a Ubiquitin Ligase From a Processive to a Distributive Enzymatic Mechanism. *J. Am. Chem. Soc.* **2015**, *137*, 12442–12445. doi: 10.1021/jacs.5b06839.
92. Dubiella, C.; Pinch, B.J.; Koikawa, K.; Zaidman, D.; Poon, E.; Manz, T.D.; Nabet, B.; He, S.; Resnick, E.; Rogel, A., et al. Sulfipin is a Covalent Inhibitor of Pin1 that Blocks Myc-Driven Tumors In Vivo. *Nat. Chem. Biol.* **2021**, *17*, 954–963. doi: 10.1038/s41589-021-00786-7.
93. Douangamath, A.; Fearon, D.; Gehrtz, P.; Krojer, T.; Lukacik, P.; Owen, C.D.; Resnick, E.; Strain-Damerell, C.; Aimon, A.; Ábrányi-Balogh, P., et al. Crystallographic and Electrophilic Fragment Screening of the SARS-CoV-2 Main Protease. *Nat. Commun.* **2020**, *11*, 5047. doi: 10.1038/s41467-020-18709-w.
94. Klein, B.A.; Reiz, B.; Robertson, I.M.; Irving, M.; Li, L.; Sun, Y.-B.; Sykes, B.D. Reversible Covalent Reaction of Levosimendan with Cardiac Troponin C *in Vitro* and *in Situ*. *Biochemistry* **2018**, *57*, 2256–2265. doi: 10.1021/acs.biochem.8b00109.
95. Dhillon, S. Tirabrutinib: First Approval. *Drugs* **2020**, *80*, 835–840. doi: 10.1007/s40265-020-01318-8.
96. Hansen, R.; Peters, U.; Babbar, A.; Chen, Y.; Feng, J.; Janes, M.R.; Li, L.-S.; Ren, P.; Liu, Y.; Zarrinkar, P.P. The Reactivity-Driven Biochemical Mechanism of Covalent KRAS^{G12C} Inhibitors. *Nat. Struct. Mol. Biol.* **2018**, *25*, 454–462. doi: 10.1038/s41594-018-0061-5.
97. Li, K.S.; Quinn, J.G.; Saabye, M.J.; Guerrero, J.F.S.; Nonomiya, J.; Lian, Q.; Phung, W.; Izrayelty, Y.; Walters, B.T.; Gustafson, A.; Endres, N.F.; Beresini, M.H.; Mulvihill, M.M. High-Throughput Kinetic Characterization of Irreversible Covalent Inhibitors of KRAS^{G12C} by Intact Protein MS and Targeted MRM. *Anal. Chem.* **2022**, *94*, 1230–1239. doi: 10.1021/acs.analchem.1c04463.
98. Janes, M.R.; Zhang, J.; Li, L.-S.; Hansen, R.; Peters, U.; Guo, X.; Chen, Y.; Babbar, A.; Firdaus, S.J.; Darjanina, L., et al. Targeting KRAS Mutant Cancers with a Covalent G12C-Specific Inhibitor. *Cell* **2018**, *172*, 578–589.e17. doi: 10.1016/j.cell.2018.01.006.
99. Canon, J.; Rex, K.; Saiki, A.Y.; Mohr, C.; Cooke, K.; Bagal, D.; Gaida, K.; Holt, T.; Knutson, C.G.; Koppada, N., et al. The Clinical KRAS(G12C) Inhibitor AMG 510 Drives Anti-Tumour Immunity. *Nature* **2019**, *575*, 217–223. doi: 10.1038/s41586-019-1694-1.
100. Hansen, R.; Firdaus, S.J.; Li, S.; Janes, M.R.; Zhang, J.; Liu, Y.; Zarrinkar, P.P. An Internally Controlled Quantitative Target Occupancy Assay for Covalent Inhibitors. *Sci. Rep.* **2018**, *8*, 14312. doi: 10.1038/s41598-018-32683-w.
101. Kantae, V.; Polanski, R.; Lewis, H.J.; Halder, A.; Barratt, D.; Srinivasan, B. Accelerating the Validation of Endogenous On-Target Engagement and In Cellulo Kinetic Assessment for Covalent Inhibitors of KRAS^{G12C} in Early Drug Discovery. *ACS Chem. Biol.* **2022**, *17*, 2366–2376. doi: 10.1021/acscchembio.2c00483.
102. Lito, P.; Solomon, M.; Li, L.-S.; Hansen, R.; Rosen, N. Allele-Specific Inhibitors Inactivate Mutant KRAS G12C By a Trapping Mechanism. *Science* **2016**, *351*, 604–608. doi: 10.1126/science.1286204.

103. Fell, J.B.; Fischer, J.P.; Baer, B.R.; Blake, J.F.; Bouhana, K.; Briere, D.M.; Brown, K.D.; Burgess, L.E.; Burns, A.C.; Burkard, M.R., et al. Identification of the Clinical Development Candidate MRTX849, a Covalent KRASG12C Inhibitor for the Treatment of Cancer. *J. Med. Chem.* **2020**, *63*, 6679–6693. doi: 10.1021/acs.jmedchem.9b02052.
104. Simon, G.M.; Niphakis, M.J.; Cravatt, B.F. Determining Target Engagement in Living Systems. *Nat. Chem. Biol.* **2013**, *9*, 200–205. doi: 10.1038/nchembio.1211.
105. McPherson, A.; Gavia, J.A. Introduction to Protein Crystallization. *Acta Crystallogr., Sect. F* **2014**, *70*, 2–20. doi: 10.1107/S2053230X13033141.
106. Berman, H.; Henrick, K.; Nakamura, H. Announcing the Worldwide Protein Data Bank. *Nat. Struct. Mol. Biol.* **2003**, *10*, 980–980. doi: 10.1038/nsb1203-980.
107. Berman, H.M.; Westbrook, J.; Feng, Z.; Gilliland, G.; Bhat, T.N.; Weissig, H.; Shindyalov, I.N.; Bourne, P.E. The Protein Data Bank. *Nucleic Acids Res.* **2000**, *28*, 235–242. doi: 10.1093/nar/28.1.235.
108. Bender, A.T.; Gardberg, A.; Pereira, A.; Johnson, T.; Wu, Y.; Grenningloh, R.; Head, J.; Morandi, F.; Haselmayer, P.; Liu-Bujalski, L. Ability of Bruton's Tyrosine Kinase Inhibitors to Sequester Y551 and Prevent Phosphorylation Determines Potency for Inhibition of Fc Receptor but not B-Cell Receptor Signaling. *Mol. Pharmacol.* **2017**, *91*, 208–219. doi: 10.1124/mol.116.107037.
109. Engel, J.; Smith, S.; Lategahn, J.; Tumbrik, H.L.; Goebel, L.; Becker, C.; Hennes, E.; Keul, M.; Unger, A.; Müller, H.; Baumann, M.; Schultz-Fademrecht, C.; Günther, G.; Hengstler, J.G.; Rauh, D. Structure-Guided Development of Covalent and Mutant-Selective Pyrazolopyrimidines to Target T790M Drug Resistance in Epidermal Growth Factor Receptor. *J. Med. Chem.* **2017**, *60*, 7725–7744. doi: 10.1021/acs.jmedchem.7b00515.
110. Groll, M.; Huber, R.; Potts, B.C.M. Crystal Structures of Salinosporamide A (NPI-0052) and B (NPI-0047) in Complex with the 20S Proteasome Reveal Important Consequences of β -Lactone Ring Opening and a Mechanism for Irreversible Binding. *J. Am. Chem. Soc.* **2006**, *128*, 5136–5141. doi: 10.1021/ja058320b.
111. Uhlenbrock, N.; Smith, S.; Weisner, J.; Landel, I.; Lindemann, M.; Le, T. A.; Hardick, J.; Gontla, R.; Scheinpluf, R.; Czodrowski, P.; Janning, P.; Depta, L.; Quambusch, L.; Müller, M.P.; Engels, B.; Rauh, D. Structural and Chemical Insights into the Covalent-Allosteric Inhibition of the Protein Kinase Akt. *Chem. Sci.* **2019**, *10*, 3573–3585. doi: 10.1039/C8SC05212C.
112. Lockbaum, G.J.; Henes, M.; Lee, J.M.; Timm, J.; Nalivaika, E.A.; Thompson, P.R.; Kurt Yilmaz, N.; Schiffer, C.A. Pan-3C Protease Inhibitor Rupintrivir Binds SARS-CoV-2 Main Protease in a Unique Binding Mode. *Biochemistry* **2021**, *60*, 2925–2931. doi: 10.1021/acs.biochem.1c00414.
113. London, N.; Miller, R.M.; Krishnan, S.; Uchida, K.; Irwin, J.J.; Eidam, O.; Gibold, L.; Cimermančić, P.; Bonnet, R.; Shoichet, B.K.; Taunton, J. Covalent Docking of Large Libraries for the Discovery of Chemical Probes. *Nat. Chem. Biol.* **2014**, *10*, 1066–1072. doi: 10.1038/nchembio.1666.
114. Scarpino, A.; Ferenczy, G.G.; Keserü, G.M. Comparative Evaluation of Covalent Docking Tools. *J. Chem. Inf. Model.* **2018**, *58*, 1441–1458. doi: 10.1021/acs.jcim.8b00228.
115. Gao, M.; Moumbock, A.F.A.; Qaseem, A.; Xu, Q.; Günther, S. CovPDB: A High-Resolution Coverage of the Covalent Protein–Ligand Interactome. *Nucleic Acids Res.* **2022**, *50*, D445–D450. doi: 10.1093/nar/gkab868.
116. Shraga, A.; Olshvang, E.; Davidzohn, N.; Khoshkenar, P.; Germain, N.; Shurrush, K.; Carvalho, S.; Avram, L.; Albeck, S.; Unger, T.; Lefker, B.; Subramanyam, C.; Hudkins, R.L.; Mitchell, A.; Shulman, Z.; Kinoshita, T.; London, N. Covalent Docking Identifies a Potent and Selective MKK7 Inhibitor. *Cell Chem. Biol.* **2019**, *26*, 98–108.e5. doi: 10.1016/j.chembiol.2018.10.011.
117. Wan, X.; Yang, T.; Cuesta, A.; Pang, X.; Ballius, T.E.; Irwin, J.J.; Shoichet, B.K.; Taunton, J. Discovery of Lysine-Targeted eIF4E Inhibitors through Covalent Docking. *J. Am. Chem. Soc.* **2020**, *142*, 4960–4964. doi: 10.1021/jacs.9b10377.
118. Becker, D.; Kaczmarek, C.; Arkona, C.; Schulz, R.; Tauber, C.; Wolber, G.; Hilgenfeld, R.; Coll, M.; Rademann, J. Irreversible Inhibitors of the 3C Protease of Coxsackie Virus Through Templated Assembly of Protein-Binding Fragments. *Nat. Commun.* **2016**, *7*, 12761. doi: 10.1038/ncomms12761.
119. Serafini, R.A.M.; da Silva Santiago, A.; Schwalm, M.P.; Hu, Z.; dos Reis, C.V.; Takarada, J.E.; Mezzomo, P.; Massier, K.B.; Kudolo, M.; Gerstenecker, S.; Chaikwad, A.; Zender, L.; Knapp, S.; Laufer, S.; Couñago, R.M.; Gehring, M. Development of the First Covalent Monopolar Spindle Kinase 1 (MPK1/TTK) Inhibitor. *J. Med. Chem.* **2022**, *65*, 3173–3192. doi: 10.1021/acs.jmedchem.1c01165.
120. Wlodawer, A.; Minor, W.; Dauter, Z.; Jaskolski, M. Protein Crystallography for Aspiring Crystallographers or How to Avoid Pitfalls and Traps in Macromolecular Structure Determination. *FEBS J.* **2013**, *280*, 5705–5736. doi: 10.1111/febs.12495.
121. Rupp, B. *Biomolecular Crystallography: Principles, Practice, and Application to Structural Biology*. Garland Science: New York, 2009. doi: 10.1201/9780429258756.
122. Taylor, G. Introduction to Phasing. *Acta Crystallogr., Sect. D* **2010**, *66*, 325–338. doi: 10.1107/S0907444910006694.
123. Hodel, A.; Kim, S.-H.; Brunger, A.T. Model Bias in Macromolecular Crystal Structures. *Acta Crystallogr., Sect. A* **1992**, *48*, 851–858. doi: 10.1107/S0108767392006044.
124. Ferrall-Fairbanks, M.C.; Kieslich, C.A.; Platt, M.O. Reassessing Enzyme Kinetics: Considering Protease-As-Substrate Interactions in Proteolytic Networks. *Proc. Natl. Acad. Sci.* **2020**, *117*, 3307. doi: 10.1073/pnas.1912207117.
125. Lee, G.M.; Balouch, E.; Goetz, D.H.; Lasic, A.; McKerrow, J.H.; Craik, C.S. Mapping Inhibitor Binding Modes on an Active Cysteine Protease via Nuclear Magnetic Resonance Spectroscopy. *Biochemistry* **2012**, *51*, 10087–10098. doi: 10.1021/bi301305k.
126. Hassell, A.M.; An, G.; Bledsoe, R.K.; Bynum, J.M.; Carter, H.L., II; Deng, S.-J.J.; Gampe, R.T.; Grisard, T.E.; Madauss, K.P.; Nolte, R.T.; Rocque, W.J.; Wang, L.; Weaver, K.L.; Williams, S.P.; Wisely, G.B.; Xu, R.; Shewchuk, L.M. Crystallization of Protein-Ligand Complexes. *Acta Crystallogr., Sect. D* **2007**, *63*, 72–79. doi: 10.1107/S0907444906047020.
127. Wienen-Schmidt, B.; Oebbecke, M.; Ngo, K.; Heine, A.; Klebe, G. Two Methods, One Goal: Structural Differences between CocrySTALLization and Crystal Soaking to Discover Ligand Binding Poses. *ChemMedChem* **2021**, *16*, 292–300. doi: 10.1002/cmdc.202000565.
128. Wlodawer, A.; Minor, W.; Dauter, Z.; Jaskolski, M. Protein Crystallography for Non-crystallographers, or How to Get the Best (But Not More) from Published Macromolecular Structures. *FEBS J.* **2008**, *275*, 1–21. doi: 10.1111/j.1742-4658.2007.06178.x.
129. Martz, E.; Sussman, J.L.; Decatur, W.; Hodis, E.; Jiang, Y.; Prilusky, J. Resolution. In *Proteopedia* [Online]; 2022. Updated 2022-05-16. <https://proteopedia.org/wiki/index.php/Resolution> (accessed 2022-08-21).
130. Woźniak, M.; Grabowski, S.; Dominiak, P.M.; Woźniak, K.; Jayatilaka, D. Hydrogen Atoms Can Be Located Accurately and Precisely by X-ray Crystallography. *Sci. Adv.* **2016**, *2*, e1600192. doi: 10.1126/sciadv.1600192.
131. Lv, Z.; Yuan, L.; Atkinson, J.H.; Williams, K.M.; Vega, R.; Sessions, E.H.; Divlianska, D.B.; Davies, C.; Chen, Y.; Olsen, S.K. Molecular Mechanism of a Covalent Allosteric Inhibitor of SUMO E1 Activating Enzyme. *Nat. Commun.* **2018**, *9*, 5145. doi: 10.1038/s41467-018-07015-1.
132. Obaidat, A.; Weiss, J.; Wahlgren, B.; Manam, R.R.; Macherla, V.R.; McArthur, K.; Chao, T.-H.; Palladino, M.A.; Lloyd, G.K.; Potts, B.C.; Enna, S.J.; Neuteboom, S.T.C.; Hagenbuch, B. Proteasome Regulator Marizomib (NPI-0052) Exhibits Prolonged Inhibition, Attenuated Efflux, and Greater Cytotoxicity than Its Reversible Analogs. *J. Pharmacol. Exp. Ther.* **2011**, *337*, 479–486. doi: 10.1124/jpet.110.177824.

133. Harshbarger, W.; Miller, C.; Diedrich, C.; Sacchettini, J. Crystal Structure of the Human 20S Proteasome in Complex with Carfilzomib. *Structure* **2015**, *23*, 418–424. doi: 10.1016/j.str.2014.11.017.
134. Law, S.; Andrault, P.-M.; Aguda, A.H.; Nguyen, N.T.; Brayer, G.D.; Brömme, D. Identification of Mouse Cathepsin K Structural Elements that Regulate the Potency of Odanacatib. *Biochem. J.* **2017**, *474*, 851–864. doi: 10.1042/BCJ20160985.
135. Forster, M.; Chaikuad, A.; Bauer, S.M.; Holstein, J.; Robers, M.B.; Corona, C.R.; Gehringer, M.; Pfaffenrot, E.; Ghoreschi, K.; Knapp, S.; Laufer, S.A. Selective JAK3 Inhibitors with a Covalent Reversible Binding Mode Targeting a New Induced Fit Binding Pocket. *Cell Chem. Biol.* **2016**, *23*, 1335–1340. doi: 10.1016/j.chembiol.2016.10.008.
136. Ekkebus, R.; van Kasteren, S.I.; Kulathu, Y.; Scholten, A.; Berlin, I.; Geurink, P.P.; de Jong, A.; Goerdal, S.; Neefjes, J.; Heck, A.J.R.; Komander, D.; Ovaa, H. On Terminal Alkynes That Can React with Active-Site Cysteine Nucleophiles in Proteases. *J. Am. Chem. Soc.* **2013**, *135*, 2867–2870. doi: 10.1021/ja309802n.
137. Barbosa da Silva, E.; Dall, E.; Briza, P.; Brandstetter, H.; Ferreira, R.S. Cruzain Structures: Apocruzain and Cruzain Bound to S-Methyl Thiomethanesulfonate and Implications for Drug Design. *Acta Crystallogr., Sect. F* **2019**, *75*, 419–427. doi: 10.1107/S2053230X19006320.
138. Karala, A.R.; Ruddock, L.W. Does S-Methyl Methanethiosulfonate Trap the Thiol–Disulfide State of Proteins? *Antioxid. Redox Signal.* **2007**, *9*, 527–531. doi: 10.1089/ars.2006.1473.
139. Adams, J.; Kauffman, M. Development of the Proteasome Inhibitor Velcade™ (Bortezomib). *Cancer Invest.* **2004**, *22*, 304–311. doi: 10.1081/CNV-120030218.
140. Kabir, M.L.; Wang, F.; Clayton, A.H.A. Intrinsically Fluorescent Anti-Cancer Drugs. *Biology* **2022**, *11*, 1135. doi: 10.3390/biology11081135.
141. Wilson, J.N.; Liu, W.; Brown, A.S.; Landgraf, R. Binding-Induced, Turn-On Fluorescence of the EGFR/ERBB Kinase Inhibitor, Lapatinib. *Org. Biomol. Chem.* **2015**, *13*, 5006–5011. doi: 10.1039/C5OB00239G.
142. Klüter, S.; Simard, J.R.; Rode, H.B.; Grütter, C.; Pawar, V.; Raaijmakers, H.C.A.; Barf, T.A.; Rabiller, M.; van Otterlo, W.A.L.; Rauh, D. Characterization of Irreversible Kinase Inhibitors by Directly Detecting Covalent Bond Formation: A Tool for Dissecting Kinase Drug Resistance. *ChemBioChem* **2010**, *11*, 2557–2566. doi: 10.1002/cbic.201000352.
143. Hodge, C.D.; Edwards, R.A.; Markin, C.J.; McDonald, D.; Pulvino, M.; Huen, M.S.Y.; Zhao, J.; Spyrapoulos, L.; Hendzel, M.J.; Glover, J.N.M. Covalent Inhibition of Ubc13 Affects Ubiquitin Signaling and Reveals Active Site Elements Important for Targeting. *ACS Chem. Biol.* **2015**, *10*, 1718–1728. doi: 10.1021/acschembio.5b00222.
144. Niu, L.-Y.; Chen, Y.-Z.; Zheng, H.-R.; Wu, L.-Z.; Tung, C.-H.; Yang, Q.-Z. Design Strategies of Fluorescent Probes for Selective Detection Among Biothiols. *Chem. Soc. Rev.* **2015**, *44*, 6143–6160. doi: 10.1039/C5CS00152H.
145. Kwan, A.H.; Mobli, M.; Gooley, P.R.; King, G.F.; Mackay, J.P. Macromolecular NMR Spectroscopy for the Non-Spectroscopist. *FEBS J.* **2011**, *278*, 687–703. doi: 10.1111/j.1742-4658.2011.08004.x.
146. Maitly, S.; Gundampati, R.K.; Suresh Kumar, T.K. NMR Methods to Characterize Protein–Ligand Interactions. *Nat. Prod. Commun.* **2019**, *14*, 1–17. doi: 10.1177/1934578X19849296.
147. Keiffer, S.; Carneiro, M.G.; Hollander, J.; Kobayashi, M.; Pogorelyev, D.; Ab, E.; Theissen, S.; Müller, G.; Siegal, G. NMR in Target Driven Drug Discovery: Why Not? *J. Biomol. NMR* **2020**, *74*, 521–529. doi: 10.1007/s10858-020-00343-9.
148. Ishima, R. Protein–Inhibitor Interaction Studies Using NMR. *Appl. NMR Spectrosc.* **2015**, *1*, 143–181. doi: 10.2174/9781608059621115010007.
149. Sugiki, T.; Furuuta, K.; Fujiwara, T.; Kojima, C. Current NMR Techniques for Structure-Based Drug Discovery. *Molecules* **2018**, *23*, 148. doi: 10.3390/molecules23010148.
150. Krishnan, V.V.; Rupp, B. Macromolecular Structure Determination: Comparison of X-ray Crystallography and NMR Spectroscopy. In *eLS* [Online]; John Wiley & Sons, 2012. Updated 2012-06-15. doi: 10.1002/9780470015902.a0002716.pub2. (accessed 2023-03-26).
151. Schirò, A.; Carlon, A.; Parigi, G.; Murshudov, G.; Calderone, V.; Ravera, E.; Luchinat, C. On the Complementarity of X-ray and NMR Data. *J. Struct. Biol.: X* **2020**, *4*, 100019. doi: 10.1016/j.jsbx.2020.100019.
152. Bjorn Dahl, T.C.; Andrew, L.C.; Semenchenko, V.; Wishart, D.S. NMR Solution Structures of the Apo and Peptide-Inhibited Human Rhinovirus 3C Protease (Serotype 14): Determination and Dynamic Comparison. *Biochemistry* **2007**, *46*, 12945–12958. doi: 10.1021/bi7010866.
153. Zhulenkova, D.; Rudevica, Z.; Jaudzems, K.; Turks, M.; Leonchiks, A. Discovery and Structure–Activity Relationship Studies of Irreversible Benzisothiazolinone-Based Inhibitors Against *Staphylococcus aureus* Sortase A Transpeptidase. *Bioorg. Med. Chem.* **2014**, *22*, 5988–6003. doi: 10.1016/j.bmc.2014.09.011.
154. Jaudzems, K.; Kurbatska, V.; Jēkabsons, A.; Bobrovs, R.; Rudevica, Z.; Leonchiks, A. Targeting Bacterial Sortase A with Covalent Inhibitors: 27 New Starting Points for Structure-Based Hit-to-Lead Optimization. *ACS Infect. Dis.* **2020**, *6*, 186–194. doi: 10.1021/acsinfecdis.9b00265.
155. Sastry, M.; Fiala, R.; Lipman, R.; Tomasz, M.; Patel, D.J. Solution Structure of the Monoalkylated Mitomycin C–DNA Complex. *J. Mol. Biol.* **1995**, *247*, 338–359. doi: 10.1006/jmbi.1994.0143.
156. Lin, C.H.; Patel, D.J. Solution Structure of the Covalent Duocarmycin A–DNA Duplex Complex. *J. Mol. Biol.* **1995**, *248*, 162–179. doi: 10.1006/jmbi.1995.0209.
157. Meyer, B.; Peters, T. NMR Spectroscopy Techniques for Screening and Identifying Ligand Binding to Protein Receptors. *Angew. Chem. Int. Ed.* **2003**, *42*, 864–890. doi: 10.1002/anie.200390233.
158. Furukawa, A.; Konuma, T.; Yanaka, S.; Sugase, K. Quantitative Analysis of Protein–Ligand Interactions by NMR. *Prog. Nucl. Magn. Reson. Spectrosc.* **2016**, *96*, 47–57. doi: 10.1016/j.pnmrs.2016.02.002.
159. Olp, M.D.; Sprague, D.J.; Goetz, C.J.; Kathman, S.G.; Wynia-Smith, S.L.; Shishodia, S.; Summers, S.B.; Xu, Z.; Stasyuk, A.V.; Smith, B.C. Covalent-Fragment Screening of BRD4 Identifies a Ligandable Site Orthogonal to the Acetyl–Lysine Binding Sites. *ACS Chem. Biol.* **2020**, *15*, 1036–1049. doi: 10.1021/acschembio.0c00058.
160. Becker, W.; Bhattiprolu, K.C.; Gubensäk, N.; Zangger, K. Investigating Protein–Ligand Interactions by Solution Nuclear Magnetic Resonance Spectroscopy. *ChemPhysChem* **2018**, *19*, 895–906. doi: 10.1002/cphc.201701253.
161. Felli, I.C.; Pierattelli, R. ¹³C Direct Detected NMR for Challenging Systems. *Chem. Rev.* **2022**, *122*, 9468–9496. doi: 10.1021/acs.chemrev.1c00871.
162. Cook, E.C.; Usher, G.A.; Shwalter, S.A. Chapter Five - The Use of ¹³C Direct-Detect NMR to Characterize Flexible and Disordered Proteins. *Meth. Enzymol.* **2018**, *611*, 81–100. doi: 10.1016/bs.mie.2018.08.025.
163. Skora, L.; Mestán, J.; Fabbro, D.; Jahnke, W.; Grzesiek, S. NMR Reveals the Allosteric Opening and Closing of Abelson Tyrosine Kinase by ATP-Site and Myristoyl Pocket Inhibitors. *Proc. Natl. Acad. Sci.* **2013**, *110*, E4437–E4445. doi: 10.1073/pnas.1314712110.
164. Metzler, W.J.; Yanchunas, J.; Weigelt, C.; Kish, K.; Klei, H.E.; Xie, D.; Zhang, Y.; Corbett, M.; Tamura, J.K.; He, B.; Hamann, L.G.; Kirby, M.S.; Marcinkiewicz, J. Involvement of DPP-IV Catalytic Residues in Enzyme–Saxagliptin Complex Formation. *Protein Sci.* **2008**, *17*, 240–250. doi: 10.1110/ps.073253208.

165. Harner, M.J.; Frank, A.O.; Fesik, S.W. Fragment-Based Drug Discovery Using NMR Spectroscopy. *J. Biomol. NMR* **2013**, *56*, 65–75. doi: 10.1007/s10858-013-9740-z.
166. Gowda, G.A.N.; Pascua, V.; Neto, F.C.; Raftery, D. Hydrogen–Deuterium Addition and Exchange in N-Ethylmaleimide Reaction with Glutathione Detected by NMR Spectroscopy. *ACS Omega* **2022**, *7*, 26928–26935. doi: 10.1021/acsomega.2c03482.
167. Krishnan, S.; Miller, R.M.; Tian, B.; Mullins, R.D.; Jacobson, M.P.; Taunton, J. Design of Reversible, Cysteine-Targeted Michael Acceptors Guided by Kinetic and Computational Analysis. *J. Am. Chem. Soc.* **2014**, *136*, 12624–12630. doi: 10.1021/ja505194w.
168. Boudreau, E.A.; Pelczar, I.; Borer, P.N.; Heffron, G.J.; LaPlante, S.R. Changes in Drug ¹³C NMR Chemical Shifts as a Tool for Monitoring Interactions with DNA. *Biophys. Chem.* **2004**, *109*, 333–344. doi: 10.1016/j.bpc.2003.12.005.
169. Yabe, Y.; Guillaume, D.; Rich, D.H. Irreversible Inhibition of Papain by Epoxysuccinyl Peptides. Carbon-13 NMR Characterization of the Site of Alkylation. *J. Am. Chem. Soc.* **1988**, *110*, 4043–4044. doi: 10.1021/ja00220a056.
170. Transue, T.R.; Krahn, J.M.; Gabel, S.A.; DeRose, E.F.; London, R.E. X-ray and NMR Characterization of Covalent Complexes of Trypsin, Borate, and Alcohols. *Biochemistry* **2004**, *43*, 2829–2839. doi: 10.1021/bi035782y.
171. Glynn, S.J.; Gaffney, K.J.; Sainz, M.A.; Louie, S.G.; Petasis, N.A. Molecular Characterization of the Boron Adducts of the Proteasome Inhibitor Bortezomib with Epigallocatechin-3-gallate and Related Polyphenols. *Org. Biomol. Chem.* **2015**, *13*, 3887–3899. doi: 10.1039/C4OB02512A.
172. Moon, J.B.; Coleman, R.S.; Hanzlik, R.P. Reversible Covalent Inhibition of Papain by a Peptide Nitrile. ¹³C NMR Evidence for a Thioimide Ester Adduct. *J. Am. Chem. Soc.* **1986**, *108*, 1350–1351. doi: 10.1021/ja00266a066.
173. Robertson, I.M.; Pineda-Sanabria, S.E.; Yan, Z.; Kampourakis, T.; Sun, Y.-B.; Sykes, B.D.; Irving, M. Reversible Covalent Binding to Cardiac Troponin C by the Ca²⁺-Sensitizer Levosimendan. *Biochemistry* **2016**, *55*, 6032–6045. doi: 10.1021/acs.biochem.6b00758.
174. Thompson, S.K.; Halbert, S.M.; Bossard, M.J.; Tomaszek, T.A.; Levy, M.A.; Zhao, B.; Smith, W.W.; Abdel-Meguid, S.S.; Janson, C.A.; D'Alessio, K.J., et al. Design of Potent and Selective Human Cathepsin K Inhibitors That Span the Active Site. *Proc. Natl. Acad. Sci.* **1997**, *94*, 14249. doi: 10.1073/pnas.94.26.14249.
175. Falgout, J.-P.; Oballa, R.M.; Okamoto, O.; Wesolowski, G.; Aubin, Y.; Ryzewski, R.M.; Prasit, P.; Riendeau, D.; Rodan, S.B.; Percival, M.D. Novel, Nonpeptidic Cyanamides as Potent and Reversible Inhibitors of Human Cathepsins K and L. *J. Med. Chem.* **2001**, *44*, 94–104. doi: 10.1021/jm0003440.
176. Sorsa, T.; Heikkinen, S.; Abbott, M.B.; Abusamhadneh, E.; Laakso, T.; Tilgmann, C.; Serimaa, R.; Annala, A.; Rosevear, P.R.; Drakenberg, T.; Pollesello, P.; Kilpeläinen, I. Binding of Levosimendan, a Calcium Sensitizer, to Cardiac Troponin C. *J. Biol. Chem.* **2001**, *276*, 9337–9343. doi: 10.1074/jbc.M007484200.
177. Liu, Y.; Patricelli, M.P.; Cravatt, B.F. Activity-Based Protein Profiling: The Serine Hydrolases. *Proc. Natl. Acad. Sci.* **1999**, *96*, 14694–14699. doi: 10.1073/pnas.96.26.14694.
178. Cravatt, B.F.; Wright, A.T.; Kozarich, J.W. Activity-Based Protein Profiling: From Enzyme Chemistry to Proteomic Chemistry. *Annu. Rev. Biochem.* **2008**, *77*, 383–414. doi: 10.1146/annurev.biochem.75.101304.124125.
179. Nomura, D.K.; Dix, M.M.; Cravatt, B.F. Activity-Based Protein Profiling for Biochemical Pathway Discovery in Cancer. *Nat. Rev. Cancer* **2010**, *10*, 630–638. doi: 10.1038/nrc2901.
180. Cravatt, B.F.; Hsu, K.-L.; Weerapana, E. Activity-Based Protein Profiling. *Curr. Top. Microbiol. Immunol.* **2019**, *420*, 1–417. doi: 10.1007/978-3-030-11143-4.
181. Bachovchin, D.A.; Cravatt, B.F. The Pharmacological Landscape and Therapeutic Potential of Serine Hydrolases. *Nat. Rev. Drug Discov.* **2012**, *11*, 52–68. doi: 10.1038/nrd3620.
182. Kato, D.; Boatright, K.M.; Berger, A.B.; Nazif, T.; Blum, G.; Ryan, C.; Chehade, K.A.H.; Salvesen, G.S.; Bogoy, M. Activity-Based Probes that Target Diverse Cysteine Protease Families. *Nat. Chem. Biol.* **2005**, *1*, 33–38. doi: 10.1038/nchembio707.
183. Chakrabarty, S.; Kahler, J.P.; van de Plassche, M.A.T.; Vanhoutte, R.; Verhelst, S.H.L. Recent Advances in Activity-Based Protein Profiling of Proteases. *Curr. Top. Microbiol. Immunol.* **2019**, *420*, 253–281. doi: 10.1007/82_2018_138.
184. Hoch, D.G.; Abegg, D.; Adibekian, A. Cysteine-reactive Probes and their Use in Chemical Proteomics. *Chem. Commun.* **2018**, *54*, 4501–4512. doi: 10.1039/C8CC01485J.
185. Hameed, D.S.; Sapmaz, A.; Ova, H. How Chemical Synthesis of Ubiquitin Conjugates Helps To Understand Ubiquitin Signal Transduction. *Bioconjug. Chem.* **2017**, *28*, 805–815. doi: 10.1021/acs.bioconjugchem.6b00140.
186. Patricelli, M.P.; Szardenings, A.K.; Liyanage, M.; Nomanbhoy, T.K.; Wu, M.; Weissig, H.; Aban, A.; Chun, D.; Tanner, S.; Kozarich, J.W. Functional Interrogation of the Kinome Using Nucleotide Acyl Phosphates. *Biochemistry* **2007**, *46*, 350–358. doi: 10.1021/bi062142x.
187. Deng, H.; Lei, Q.; Wu, Y.; He, Y.; Li, W. Activity-Based Protein Profiling: Recent Advances in Medicinal Chemistry. *Eur. J. Med. Chem.* **2020**, *191*, 112151. doi: 10.1016/j.ejmech.2020.112151.
188. Fang, H.; Peng, B.; Ong, S.Y.; Wu, Q.; Li, L.; Yao, S.Q. Recent Advances in Activity-Based Probes (ABPs) and Affinity-Based Probes (AfBPs) for Profiling of Enzymes. *Chem. Sci.* **2021**, *12*, 8288–8310. doi: 10.1039/D1SC01359A.
189. Backus, K.M.; Correia, B.E.; Lum, K.M.; Forli, S.; Horning, B.D.; González-Pérez, G.E.; Chatterjee, S.; Lanning, B.R.; Teijaro, J.R.; Olson, A.J.; Wolan, D.W.; Cravatt, B.F. Proteome-Wide Covalent Ligand Discovery in Native Biological Systems. *Nature* **2016**, *534*, 570–574. doi: 10.1038/nature18002.
190. Lanning, B.R.; Whitby, L.R.; Dix, M.M.; Douhan, J.; Gilbert, A.M.; Hett, E.C.; Johnson, T.O.; Joslyn, C.; Kath, J.C.; Niessen, S.; Roberts, L.R.; Schnute, M.E.; Wang, C.; Hulce, J.J.; Wei, B.; Whiteley, L.O.; Hayward, M.M.; Cravatt, B.F. A Road Map to Evaluate the Proteome-Wide Selectivity of Covalent Kinase Inhibitors. *Nat. Chem. Biol.* **2014**, *10*, 760–767. doi: 10.1038/nchembio.1582.
191. van Esbroeck, A.C.M.; Janssen, A.P.A.; Cognetta, A.B.; Ogasawara, D.; Shpak, G.; van der Kroeg, M.; Kante, V.; Baggelaar, M.P.; de Vrij, F.M.S.; Deng, H., et al. Activity-Based Protein Profiling Reveals Off-Target Proteins of the FAAH Inhibitor BIA 10-2474. *Science* **2017**, *356*, 1084–1087. doi: 10.1126/science.aaf7497.
192. Bird, R.E.; Lemmel, S.A.; Yu, X.; Zhou, Q.A. Bioorthogonal Chemistry and Its Applications. *Bioconjug. Chem.* **2021**, *32*, 2457–2479. doi: 10.1021/acs.bioconjugchem.1c00461.
193. Rostovtsev, V.V.; Green, L.G.; Fokin, V.V.; Sharpless, K.B. A Stepwise Huisgen Cycloaddition Process: Copper(I)-Catalyzed Regioselective “Ligation” of Azides and Terminal Alkynes. *Angew. Chem. Int. Ed.* **2002**, *41*, 2596–2599. doi: 10.1002/1521-3773(20020715)41:14<2596::AID-ANIE2596>3.0.CO;2-4.
194. Parker, C.G.; Pratt, M.R. Click Chemistry in Proteomic Investigations. *Cell* **2020**, *180*, 605–632. doi: 10.1016/j.cell.2020.01.025.
195. Smeenk, M.L.W.J.; Agramunt, J.; Bongers, K.M. Recent Developments in Bioorthogonal Chemistry and the Orthogonality Within. *Curr. Opin. Chem. Biol.* **2021**, *60*, 79–88. doi: 10.1016/j.cbpa.2020.09.002.

196. Agard, N.J.; Prescher, J.A.; Bertozzi, C.R. A Strain-Promoted [3 + 2] Azide–Alkyne Cycloaddition for Covalent Modification of Biomolecules in Living Systems. *J. Am. Chem. Soc.* **2004**, *126*, 15046–15047. doi: 10.1021/ja044996f.
197. Oliveira, B.L.; Guo, Z.; Bernardes, G.J.L. Inverse Electron Demand Diels–Alder Reactions in Chemical Biology. *Chem. Soc. Rev.* **2017**, *46*, 4895–4950. doi: 10.1039/C7CS00184C.
198. van Rooden, E.J.; Florea, B.I.; Deng, H.; Baggelaar, M.P.; van Esbroeck, A.C.M.; Zhou, J.; Overkleeft, H.S.; van der Stelt, M. Mapping *In Vivo* Target Interaction Profiles of Covalent Inhibitors using Chemical Proteomics with Label-Free Quantification. *Nat. Protoc.* **2018**, *13*, 752–767. doi: 10.1038/nprot.2017.159.
199. Xu, H.; Jesson, M.I.; Seneviratne, U.I.; Lin, T. H.; Sharif, M. N.; Xue, L.; Nguyen, C.; Everley, R.A.; Trujillo, J.I.; Johnson, D.S.; Point, G.R.; Thorarensen, A.; Kilty, I.; Telliez, J.-B. PF-06651600, a Dual JAK3/TEC Family Kinase Inhibitor. *ACS Chem. Biol.* **2019**, *14*, 1235–1242. doi: 10.1021/acscchembio.9b00188.
200. Yang, P.-Y.; Liu, K.; Ngai, M.H.; Lear, M.J.; Wenk, M.R.; Yao, S.Q. Activity-Based Proteome Profiling of Potential Cellular Targets of Orlistat – An FDA-Approved Drug with Anti-Tumor Activities. *J. Am. Chem. Soc.* **2010**, *132*, 656–666. doi: 10.1021/ja907716f.
201. Martin, J.G.; Ward, J.A.; Feyertag, F.; Zhang, L.; Couvartier, S.; Guckian, K.; Huber, K.V.M.; Johnson, D.S. Chemoproteomic Profiling of Covalent XP01 Inhibitors to Assess Target Engagement and Selectivity. *ChemBioChem* **2021**, *22*, 2116–2123. doi: 10.1002/cbic.202100038.
202. Lee, J. H.; Hou, X.; Kumari, E.; Borazjani, A.; Edelmann, M.J.; Ross, M.K. Endocannabinoid Hydrolases in Avian HD11 Macrophages Identified by Chemoproteomics: Inactivation by Small-Molecule Inhibitors and Pathogen-Induced Downregulation of Their Activity. *Mol. Cell. Biochem.* **2018**, *444*, 125–141. doi: 10.1007/s11010-017-3237-0.
203. Gjonaj, L.; Sapmaz, A.; Flierman, D.; Janssen, G.M.C.; van Veelen, P.A.; Ova, H. Development of a DUB-Selective Fluorogenic Substrate. *Chem. Sci.* **2019**, *10*, 10290–10296. doi: 10.1039/C9SC02262K.
204. van Rooden, E.J.; van Esbroeck, A.C.M.; Baggelaar, M.P.; Deng, H.; Florea, B.I.; Marques, A.R.A.; Ottenhoff, R.; Boot, R.G.; Overkleeft, H.S.; Aerts, J.M.F.G.; van der Stelt, M. Chemical Proteomic Analysis of Serine Hydrolase Activity in Niemann–Pick Type C Mouse Brain. *Front. Neurosci.* **2018**, *12*, 440. doi: 10.3389/fnins.2018.00440.205. Li, W.; Blankman, J.L.; Cravatt, B.F. A Functional Proteomic Strategy to Discover Inhibitors for Uncharacterized Hydrolases. *J. Am. Chem. Soc.* **2007**, *129*, 9594–9595. doi: 10.1021/ja073650c.
206. van der Wel, T.; Hilhorst, R.; den Dulk, H.; van den Hooven, T.; Prins, N.M.; Wijnakker, J.A.P.M.; Florea, B.I.; Lenselink, E.B.; van Westen, G.J.P.; Ruijtenbeek, R.; Overkleeft, H.S.; Kaptein, A.; Barf, T.; van der Stelt, M. Chemical Genetics Strategy to Profile Kinase Target Engagement Reveals Role of FES in Neutrophil Phagocytosis. *Nat. Commun.* **2020**, *11*, 3216. doi: 10.1038/s41467-020-17027-5.
207. Kooij, R.; Liu, S.; Sapmaz, A.; Xin, B.-T.; Janssen, G.M.C.; van Veelen, P.A.; Ova, H.; ten Dijke, P.; Geurink, P.P. Small-Molecule Activity-Based Probe for Monitoring Ubiquitin C-Terminal Hydrolase L1 (UCHL1) Activity in Live Cells and Zebrafish Embryos. *J. Am. Chem. Soc.* **2020**, *142*, 16825–16841. doi: 10.1021/jacs.0c07726.
208. Jia, Y.; Kim, R.Q.; Kooij, R.; Ova, H.; Sapmaz, A.; Geurink, P.P. Chemical Toolkit for PARK7: Potent, Selective, and High-Throughput. *J. Med. Chem.* **2022**, *65*, 13288–13304. doi: 10.1021/acs.jmedchem.2c01113.
209. Leal, J.; Martínez-Díez, M.; García-Hernández, V.; Moneo, V.; Domingo, A.; Bueren-Calabuig, J.; Negri, A.; Gago, F.; Guillén-Navarro, M.; Avilés, P.; Cuevas, C.; García-Fernández, L.; Galmarini, C. PM01183, a New DNA Minor Groove Covalent Binder with Potent *In Vitro* and *In Vivo* Anti-Tumour Activity. *Br. J. Pharmacol.* **2010**, *161*, 1099–1110. doi: 10.1111/j.1476-5381.2010.00945.x.
210. Dana, D.; Pathak, S.K. A Review of Small Molecule Inhibitors and Functional Probes of Human Cathepsin L. *Molecules* **2020**, *25*, 698. doi: 10.3390/molecules25030698.
211. Bogoy, M.; Verhelst, S.; Bellingard-Dubouchaud, V.; Toba, S.; Greenbaum, D. Selective Targeting of Lysosomal Cysteine Proteases with Radiolabeled Electrophilic Substrate Analogs. *Chem. Biol.* **2000**, *7*, 27–38. doi: 10.1016/S1074-5521(00)00061-2.
212. Pellegatti, M. Preclinical *In Vivo* ADME Studies in Drug Development: A Critical Review. *Expert Opin. Drug Metab. Toxicol.* **2012**, *8*, 161–172. doi: 10.1517/17425255.2012.652084.
213. Harding, C.R.; Scott, I.R. Fluorography—Limitations on Its Use for the Quantitative Detection of ^3H - and ^{14}C -Labeled Proteins in Polyacrylamide Gels. *Anal. Biochem.* **1983**, *129*, 371–376. doi: 10.1016/0003-2697(83)90564-X.
214. Miyahisa, I.; Sameshima, T.; Hixon, M.S. Rapid Determination of the Specificity Constant of Irreversible Inhibitors (k_{inact}/K_i) by Means of an Endpoint Competition Assay. *Angew. Chem. Int. Ed.* **2015**, *54*, 14099–14102. doi: 10.1002/anie.201505800.
215. Tsou, H.-R.; Overbeek-Klumpers, E.G.; Hallett, W.A.; Reich, M.F.; Floyd, M.B.; Johnson, B.D.; Michalak, R.S.; Nilakantan, R.; Discifani, C.; Golas, J.; Rabindran, S.K.; Shen, R.; Shi, X.; Wang, Y.-F.; Upešlacis, J.; Wissner, A. Optimization of 6,7-Disubstituted-4-[arylamino]quinoline-3-carbonitriles as Orally Active, Irreversible Inhibitors of Human Epidermal Growth Factor Receptor-2 Kinase Activity. *J. Med. Chem.* **2005**, *48*, 1107–1131. doi: 10.1021/jm040159c.
216. Scicinski, J.; Oronsky, B.; Taylor, M.; Luo, G.; Musick, T.; Marini, J.; Adams, C.M.; Fitch, W.L. Preclinical Evaluation of the Metabolism and Disposition of RRx-001, a Novel Investigative Anticancer Agent. *Drug Metab. Dispos.* **2012**, *40*, 1810–1816. doi: 10.1124/dmd.112.046755.
217. McClure, R.A.; Williams, J.D. Impact of Mass Spectrometry-Based Technologies and Strategies on Chemoproteomics as a Tool for Drug Discovery. *ACS Med. Chem. Lett.* **2018**, *9*, 785–791. doi: 10.1021/acsmchemlett.8b00181.
218. Counihan, J.L.; Ford, B.; Nomura, D.K. Mapping Proteome-Wide Interactions of Reactive Chemicals Using Chemoproteomic Platforms. *Curr. Opin. Chem. Biol.* **2016**, *30*, 68–76. doi: 10.1016/j.cbpa.2015.11.007.
219. Maurais, A.J.; Weerapana, E. Reactive-Cysteine Profiling for Drug Discovery. *Curr. Opin. Chem. Biol.* **2019**, *50*, 29–36. doi: 10.1016/j.cbpa.2019.02.010.
220. Neilson, K.A.; Ali, N.A.; Muralidharan, S.; Mirzaei, M.; Mariani, M.; Assadourian, G.; Lee, A.; van Sluyter, S.C.; Haynes, P.A. Less Label, More Free: Approaches in Label-free Quantitative Mass Spectrometry. *Proteomics* **2011**, *11*, 535–553. doi: 10.1002/pmic.201000553.
221. Nikolov, M.; Schmidt, C.; Urlaub, H. Quantitative Mass Spectrometry-Based Proteomics: An Overview. *Methods Mol. Biol.* **2012**, *893*, 85–100. doi: 10.1007/978-1-61779-885-6_7.
222. Benms, H.J.; Wincott, C.J.; Tate, E.W.; Child, M.A. Activity- and Reactivity-Based Proteomics: Recent Technological Advances and Applications in Drug Discovery. *Curr. Opin. Chem. Biol.* **2021**, *60*, 20–29. doi: 10.1016/j.cbpa.2020.06.011.
223. Wang, C.; Weerapana, E.; Blewett, M.M.; Cravatt, B.F. A Chemoproteomic Platform to Quantitatively Map Targets of Lipid-Derived Electrophiles. *Nat. Meth.* **2014**, *11*, 79–85. doi: 10.1038/nmeth.2759.
224. Weerapana, E.; Wang, C.; Simon, G.M.; Richter, F.; Khare, S.; Dillon, M.B.D.; Bachovchin, D.A.; Mowen, K.; Baker, D.; Cravatt, B.F. Quantitative Reactivity Profiling Predicts Functional Cysteines in Proteomes. *Nature* **2010**, *468*, 790–795. doi: 10.1038/nature09472.
225. Kuljanin, M.; Mitchell, D.C.; Schweppe, D.K.; Gikandi, A.S.; Nusinow, D.P.; Bulloch, N.J.; Vinogradova, E.V.; Wilson, D.L.; Kool, E.T.; Mancias, J.D.; Cravatt, B.F.; Gygi, S.P. Reimagining High-Throughput Profiling of Reactive Cysteines for Cell-Based Screening of Large Electrophile Libraries. *Nat. Biotechnol.* **2021**, *39*, 630–641. doi: 10.1038/s41587-020-00778-3.

226. Boatner, L.M.; Palafox, M.F.; Schweppe, D.K.; Backus, K.M. CysDB: A Human Cysteine Database based on Experimental Quantitative Chemoproteomics. *Cell Chem. Biol.* **2023**, *30*, 683-698.e3. doi: 10.1016/j.chembiol.2023.04.004.
227. Hsu, J.-L.; Chen, S.-H. Stable Isotope Dimethyl Labelling for Quantitative Proteomics and Beyond. *Philos. Trans. R. Soc., A* **2016**, *374*, 20150364. doi: 10.1098/rsta.2015.0364.
228. Adibekian, A.; Martin, B.R.; Wang, C.; Hsu, K.-L.; Bachovchin, D.A.; Niessen, S.; Hoover, H.; Cravatt, B.F. Click-Generated Triazole Ureas as Ultrapotent *In Vivo*-Active Serine Hydrolase Inhibitors. *Nat. Chem. Biol.* **2011**, *7*, 469-478. doi: 10.1038/nchembio.579.
229. Mann, M. Functional and Quantitative Proteomics Using SILAC. *Nat. Rev. Mol. Cell Biol.* **2006**, *7*, 952-958. doi: 10.1038/nrm2067.
230. Ong, S.-E.; Blagoev, B.; Kratchmarova, I.; Kristensen, D.B.; Steen, H.; Pandey, A.; Mann, M. Stable Isotope Labeling by Amino Acids in Cell Culture, SILAC, as a Simple and Accurate Approach to Expression Proteomics. *Mol. Cell. Proteom.* **2002**, *1*, 376-386. doi: 10.1074/mcp.M200025-MCP200.
231. Browne, C.M.; Jiang, B.; Ficarro, S.B.; Doctor, Z.M.; Johnson, J.L.; Card, J.D.; Sivakumaren, S.C.; Alexander, W.M.; Yaron, T.M.; Murphy, C.J.; Kwiatkowski, N.P.; Zhang, T.; Cantley, L.C.; Gray, N.S.; Marto, J.A. A Chemoproteomic Strategy for Direct and Proteome-Wide Covalent Inhibitor Target-Site Identification. *J. Am. Chem. Soc.* **2019**, *141*, 191-203. doi: 10.1021/jacs.8b07911.
232. Chen, X.; Sun, Y.; Zhang, T.; Shu, L.; Roepstorff, P.; Yang, F. Quantitative Proteomics Using Isobaric Labeling: A Practical Guide. *Genom. Proteom. Bioinf.* **2021**, *19*, 689-706. doi: 10.1016/j.gpb.2021.08.012.
233. Weerapana, E.; Speers, A.E.; Cravatt, B.F. Tandem Orthogonal Proteolysis-Activity-Based Protein Profiling (TOP-ABPP)—a General Method for Mapping Sites of Probe Modification in Proteomes. *Nat. Protoc.* **2007**, *2*, 1414-1425. doi: 10.1038/nprot.2007.194.
234. Zanon, P.R.A.; Lewald, L.; Hacker, S.M. Isotopically Labeled Dethiobiotin Azide (isoDTB) Tags Enable Global Profiling of the Bacterial Cysteineome. *Angew. Chem. Int. Ed.* **2020**, *59*, 2829-2836. doi: 10.1002/anie.201912075.
235. Yang, F.; Gao, J.; Che, J.; Jia, G.; Wang, C. A Dimethyl-Labeling-Based Strategy for Site-Specifically Quantitative Chemical Proteomics. *Anal. Chem.* **2018**, *90*, 9576-9582. doi: 10.1021/acs.analchem.8b02426.
236. Hsu, J.-L.; Huang, S.-Y.; Chow, N.-H.; Chen, S.-H. Stable-Isotope Dimethyl Labeling for Quantitative Proteomics. *Anal. Chem.* **2003**, *75*, 6843-6852. doi: 10.1021/ac0348625.
237. Boersema, P.J.; Raijmakers, R.; Lemeer, S.; Mohammed, S.; Heck, A.J.R. Multiplex Peptide Stable Isotope Dimethyl Labeling for Quantitative Proteomics. *Nat. Protoc.* **2009**, *4*, 484-494. doi: 10.1038/nprot.2009.21.
238. Li, N.; Kuo, C.-L.; Paniagua, G.; van den Elst, H.; Verdoes, M.; Willems, L.I.; van der Linden, W.A.; Ruben, M.; van Genderen, E.; Gubbens, J.; van Wezel, G.P.; Overkleeft, H.S.; Florea, B.I. Relative Quantification of Proteasome Activity by Activity-Based Protein Profiling and LC-MS/MS. *Nat. Protoc.* **2013**, *8*, 1155-1168. doi: 10.1038/nprot.2013.065.
239. Baggelaar, M.P.; Chameau, P.J.P.; Kantae, V.; Hummel, J.; Hsu, K.-L.; Janssen, F.; van der Wel, T.; Soethoudt, M.; Deng, H.; den Dulk, H.; Allarà, M.; Florea, B.I.; Di Marzo, V.; Wadman, W.J.; Kruse, C.G.; Overkleeft, H.S.; Hankemeier, T.; Werkman, T.R.; Cravatt, B.F.; van der Stelt, M. Highly Selective, Reversible Inhibitor Identified by Comparative Chemoproteomics Modulates Diacylglycerol Lipase Activity in Neurons. *J. Am. Chem. Soc.* **2015**, *137*, 8851-8857. doi: 10.1021/jacs.5b04883.
240. Vinogradova, E.V.; Zhang, X.; Remillard, G.; Lazar, D.C.; Suci, R.M.; Wang, Y.; Bianco, G.; Yamashita, Y.; Crowley, V.M.; Schafroth, M.A., et al. An Activity-Guided Map of Electrophile-Cysteine Interactions in Primary Human T Cells. *Cell* **2020**, *182*, 1009-1026.e29. doi: 10.1016/j.cell.2020.07.001.
241. Werner, T.; Becher, I.; Sweetman, G.; Doce, C.; Savitski, M.M.; Bantscheff, M. High-Resolution Enabled TMT 8-plexing. *Anal. Chem.* **2012**, *84*, 7188-7194. doi: 10.1021/ac301553x.
242. Zanon, P.R.A.; Yu, F.; Musacchio, P.; Lewald, L.; Zollo, M.; Krauskopf, K.; Mrdović, D.; Raunft, P.; Maher, T.E.; Cigler, M.; Chang, C.; Lang, K.; Toste, F.D.; Nesvizhskii, A.I.; Hacker, S.M. Profiling the Proteome-Wide Selectivity of Diverse Electrophiles. *ChemRxiv* **2021**. doi: 10.26434/chemrxiv-2021-w7rss-v2.
243. Adibekian, A.; Martin, B.R.; Chang, J.W.; Hsu, K.-L.; Tsuboi, K.; Bachovchin, D.A.; Speers, A.E.; Brown, S.J.; Spicer, T.; Fernandez-Vega, V.; Ferguson, J.; Hodder, P.S.; Rosen, H.; Cravatt, B.F. Confirming Target Engagement for Reversible Inhibitors *In Vivo* by Kinetically Tuned Activity-Based Probes. *J. Am. Chem. Soc.* **2012**, *134*, 10345-10348. doi: 10.1021/ja303400u.
244. Senkane, K.; Vinogradova, E.V.; Suci, R.M.; Crowley, V.M.; Zaro, B.W.; Bradshaw, J.M.; Brameld, K.A.; Cravatt, B.F. The Proteome-Wide Potential for Reversible Covalency at Cysteine. *Angew. Chem. Int. Ed.* **2019**, *58*, 11385-11389. doi: 10.1002/anie.201905829.
245. Blum, G.; Mullins, S.R.; Keren, K.; Fonović, M.; Jedeszko, C.; Rice, M.J.; Sloane, B.F.; Bogoy, M. Dynamic Imaging of Protease Activity with Fluorescently Quenched Activity-Based Probes. *Nat. Chem. Biol.* **2005**, *1*, 203-209. doi: 10.1038/nchembio728.
246. Edgington-Mitchell, L.E.; Bogoy, M.; Verdoes, M. Live Cell Imaging and Profiling of Cysteine Cathepsin Activity Using a Quenched Activity-Based Probe. *Methods Mol. Biol.* **2017**, *1491*, 145-159. doi: 10.1007/978-1-4939-6439-0_11.
247. Blum, G.; Weimer, R.M.; Edgington, L.E.; Adams, W.; Bogoy, M. Comparative Assessment of Substrates and Activity Based Probes as Tools for Non-invasive Optical Imaging of Cysteine Protease Activity. *PLoS ONE* **2009**, *4*, e6374. doi: 10.1371/journal.pone.0006374.
248. Blum, G.; von Degenfeld, G.; Merchant, M.J.; Blau, H.M.; Bogoy, M. Noninvasive Optical Imaging of Cysteine Protease Activity using Fluorescently Quenched Activity-Based Probes. *Nat. Chem. Biol.* **2007**, *3*, 668-677. doi: 10.1038/nchembio.2007.26.
249. van Rooden, E.J.; Kohsiek, M.; Kreekel, R.; van Esbroeck, A.C.M.; van den Nieuwendijk, A.M.C.H.; Janssen, A.P.A.; van den Berg, R.J.B.H.N.; Overkleeft, H.S.; van der Stelt, M. Design and Synthesis of Quenched Activity-based Probes for Diacylglycerol Lipase and α , β -Hydrolase Domain Containing Protein 6. *Chem. Asian J.* **2018**, *13*, 3491-3500. doi: 10.1002/asia.201800452.
250. Serim, S.; Baer, P.; Verhelst, S.H.L. Mixed Alkyl Aryl Phosphonate Esters as Quenched Fluorescent Activity-Based Probes for Serine Proteases. *Org. Biomol. Chem.* **2015**, *13*, 2293-2299. doi: 10.1039/C4OB02444C.
251. Verdoes, M.; Oresic Bender, K.; Segal, E.; van der Linden, W.A.; Syed, S.; Withana, N.P.; Sanman, L.E.; Bogoy, M. Improved Quenched Fluorescent Probe for Imaging of Cysteine Cathepsin Activity. *J. Am. Chem. Soc.* **2013**, *135*, 14726-14730. doi: 10.1021/ja4056068.
252. Zhang, Q.; Liu, H.; Pan, Z. A General Approach for the Development of Fluorogenic Probes Suitable for No-Wash Imaging of Kinases in Live Cells. *Chem. Commun.* **2014**, *50*, 15319-15322. doi: 10.1039/C4CC07429G.
253. Janssen, A.P.A.; van Hengst, J.M.A.; Béquignon, O.J.M.; Deng, H.; van Westen, G.J.P.; van der Stelt, M. Structure Kinetics Relationships and Molecular Dynamics Show Crucial Role for Heterocycle Leaving Group in Irreversible Diacylglycerol Lipase Inhibitors. *J. Med. Chem.* **2019**, *62*, 7910-7922. doi: 10.1021/acs.jmedchem.9b00686.
254. Petri, L.; Ábrányi-Balogh, P.; Varga, P.R.; Imre, T.; Keserü, G.M. Comparative Reactivity Analysis of Small-Molecule Thiol Surrogates. *Bioorg. Med. Chem.* **2020**, *28*, 115357. doi: 10.1016/j.bmc.2020.115357.
255. Reddi, R.N.; Resnick, E.; Rogel, A.; Rao, B.V.; Gabizon, R.; Goldenberg, K.; Gurwicz, N.; Zaidman, D.; Plotnikov, A.; Barr, H.; Shulman, Z.; London, N. Tunable Methacrylamides for Covalent Ligand Directed Release Chemistry. *J. Am. Chem. Soc.* **2021**, *143*, 4979-4992. doi: 10.1021/jacs.0c10644.

256. Kawahata, W.; Asami, T.; Fujii, I.; Sawa, M. 'Turn On/Off' Fluorescence Probe for the Screening of Unactivated Bruton's Tyrosine Kinase. *Bioorg. Med. Chem. Lett.* **2015**, *25*, 2141–2145. doi: 10.1016/j.bmcl.2015.03.063.
257. Li, X.; Gao, X.; Shi, W.; Ma, H. Design Strategies for Water-Soluble Small Molecular Chromogenic and Fluorogenic Probes. *Chem. Rev.* **2014**, *114*, 590–659. doi: 10.1021/cr300508p.
258. Shie, J.-J.; Liu, Y.-C.; Lee, Y.-M.; Lim, C.; Fang, J.-M.; Wong, C.-H. An Azido-BODIPY Probe for Glycosylation: Initiation of Strong Fluorescence upon Triazole Formation. *J. Am. Chem. Soc.* **2014**, *136*, 9953–9961. doi: 10.1021/ja5010174.
259. Blau, R.; Epshtein, Y.; Pisarevsky, E.; Tiram, G.; Israeli Dangoor, S.; Yeini, E.; Krivitsky, A.; Eldar-Boock, A.; Ben-Shushan, D.; Giori, H., et al. Image-Guided Surgery using Near-Infrared Turn-ON Fluorescent Nanoprobes for Precise Detection of Tumor Margins. *Theranostics* **2018**, *8*, 3437–3460. doi: 10.7150/thno.23853.
260. Ben-Nun, Y.; Merquiol, E.; Brandis, A.; Turk, B.; Scherz, A.; Blum, G. Photodynamic Quenched Cathepsin Activity Based Probes for Cancer Detection and Macrophage Targeted Therapy. *Theranostics* **2015**, *5*, 847–862. doi: 10.7150/thno.10854.
261. Weiss-Sadan, T.; Ben-Nun, Y.; Maimoun, D.; Merquiol, E.; Abd-Elrahman, I.; Gotsman, I.; Blum, G. A Theranostic Cathepsin Activity-Based Probe for Noninvasive Intervention in Cardiovascular Diseases. *Theranostics* **2019**, *9*, 5731–5738. doi: 10.7150/thno.34402.
262. Evans, E.K.; Tester, R.; Aslanian, S.; Karp, R.; Sheets, M.; Labenski, M.T.; Witowski, S.R.; Lounsbury, H.; Chaturvedi, P.; Mazdiasni, H.; Zhu, Z.; Nacht, M.; Freed, M.I.; Petter, R.C.; Dubrovskiy, A.; Singh, J.; Westlin, W.F. Inhibition of Btk with CC-292 Provides Early Pharmacodynamic Assessment of Activity in Mice and Humans. *J. Pharmacol. Exp. Ther.* **2013**, *346*, 219. doi: 10.1124/jpet.113.203489.
263. Callaway, E. Revolutionary Cryo-EM is Taking Over Structural Biology. *Nature* **2020**, *578*, 201. doi: 10.1038/d41586-020-00341-9.
264. Chiu, Y.-H.; Ko, K.-T.; Yang, T.-J.; Wu, K.-P.; Ho, M.-R.; Draczowski, P.; Hsu, S.-T.D. Direct Visualization of a 26 kDa Protein by Cryo-Electron Microscopy Aided by a Small Scaffold Protein. *Biochemistry* **2021**, *60*, 1075–1079. doi: 10.1021/acs.biochem.0c00961.
265. Liu, Y.; Huynh, D.T.; Yeates, T.O. A 3.8 Å Resolution Cryo-EM Structure of a Small Protein Bound To an Imaging Scaffold. *Nat. Commun.* **2019**, *10*, 1864. doi: 10.1038/s41467-019-09836-0.
266. Borgnia, M.J.; Banerjee, S.; Merk, A.; Matthies, D.; Bartesaghi, A.; Rao, P.; Pierson, J.; Earl, L. A.; Falconieri, V.; Subramaniam, S.; Milne, J.L.S. Using Cryo-EM to Map Small Ligands on Dynamic Metabolic Enzymes: Studies with Glutamate Dehydrogenase. *Mol. Pharmacol.* **2016**, *89*, 645. doi: 10.1124/mol.116.103382.
267. Toelzer, C.; Gupta, K.; Yadav, S.K.N.; Borucu, U.; Davidson, A.D.; Kavanagh Williamson, M.; Shoemark, D.K.; Garzoni, F.; Stauffer, O.; Milligan, R.; Capin, J.; Mulholland, A.J.; Spatz, J.; Fitzgerald, D.; Berger, I.; Schaffitzel, C. Free Fatty Acid Binding Pocket in the Locked Structure of SARS-CoV-2 Spike Protein. *Science* **2020**, *370*, 725–730. doi: 10.1126/science.abd3255.
268. Greber, B.J.; Perez-Bertoldi, J.M.; Lim, K.; Iavarone, A.T.; Toso, D.B.; Nogales, E. The Cryoelectron Microscopy Structure of the Human CDK-Activating Kinase. *Proc. Natl. Acad. Sci.* **2020**, *117*, 22849–22857. doi: 10.1073/pnas.2009627117.
269. Robertson, M.J.; van Zundert, G.C.P.; Borrelli, K.; Skiniotis, G. GemSpot: A Pipeline for Robust Modeling of Ligands into Cryo-EM Maps. *Structure* **2020**, *28*, 707–716.e3. doi: 10.1016/j.str.2020.04.018.
270. Chengalroyen, M.D.; Mason, M.K.; Borsellini, A.; Tassoni, R.; Abrahams, G.L.; Lynch, S.; Ahn, Y.-M.; Ambler, J.; Young, K.; Crowley, B.M.; Olsen, D.B.; Warner, D.F.; Barry, C.E.III; Boshoff, H.I.M.; Lamers, M.H.; Mizrahi, V. DNA-Dependent Binding of Nargenacin to DnaE1 Inhibits Replication in *Mycobacterium tuberculosis*. *ACS Infect. Dis.* **2022**, *8*, 612–625. doi: 10.1021/acinfeddis.1c00643.
271. Lindberg, P.; Nordberg, P.; Alminger, T.; Braendstroem, A.; Wallmark, B. The Mechanism of Action of the Antisecretory Agent Omeprazole. *J. Med. Chem.* **1986**, *29*, 1327–1329. doi: 10.1021/jm00158a001.
272. Mukherjee, H.; Grimster, N.P. Beyond Cysteine: Recent Developments in the Area of Targeted Covalent Inhibition. *Curr. Opin. Chem. Biol.* **2018**, *44*, 30–38. doi: 10.1016/j.cbpa.2018.05.011.
273. Jones, L.H. Chapter Four - Design of Next-Generation Covalent Inhibitors: Targeting Residues Beyond Cysteine. *Annu. Rep. Med. Chem.* **2021**, *56*, 95–134. doi: 10.1016/bs.armc.2020.10.001.
274. Gambini, L.; Baggio, C.; Udompholkul, P.; Jossart, J.; Salem, A.F.; Perry, J.J.P.; Pellicchia, M. Covalent Inhibitors of Protein–Protein Interactions Targeting Lysine, Tyrosine, or Histidine Residues. *J. Med. Chem.* **2019**, *62*, 5616–5627. doi: 10.1021/acs.jmedchem.9b00561.
275. Pettinger, J.; Jones, K.; Cheeseman, M.D. Lysine-Targeting Covalent Inhibitors. *Angew. Chem. Int. Ed.* **2017**, *56*, 15200–15209. doi: 10.1002/anie.201707630.
276. Zhang, Z.; Guiley, K.Z.; Shokat, K.M. Chemical Acylation of an Acquired Serine Suppresses Oncogenic Signaling of K-Ras(G12S). *Nat. Chem. Biol.* **2022**, *18*, 1177–1183. doi: 10.1038/s41589-022-01065-9.
277. Zhang, Z.; Morstein, J.; Ecker, A.K.; Guiley, K.Z.; Shokat, K.M. Chemoselective Covalent Modification of K-Ras(G12R) with a Small Molecule Electrophile. *J. Am. Chem. Soc.* **2022**, *144*, 15916–15921. doi: 10.1021/jacs.2c05377.
278. Chen, P.; Sun, J.; Zhu, C.; Tang, G.; Wang, W.; Xu, M.; Xiang, M.; Zhang, C.-J.; Zhang, Z.-M.; Gao, L.; Yao, S.Q. Cell-Active, Reversible, and Irreversible Covalent Inhibitors That Selectively Target the Catalytic Lysine of BCR-ABL Kinase. *Angew. Chem. Int. Ed.* **2022**, *61*, e202203878. doi: 10.1002/anie.202203878.
279. Lill, J.R.; Mathews, W.R.; Rose, C.M.; Schirle, M. Proteomics in the Pharmaceutical and Biotechnology Industry: A Look to the Next Decade. *Expert Rev. Proteom.* **2021**, *18*, 503–526. doi: 10.1080/14789450.2021.1962300.
280. Aranda, J.; Orozco, M. RNA-Dependent RNA Polymerase From SARS-CoV-2. Mechanism Of Reaction And Inhibition By Remdesivir. *bioRxiv* **2020**. doi: 10.1101/2020.06.21.163592.
281. Kalyukina, M.; Yosaatmadja, Y.; Middleditch, M.J.; Patterson, A.V.; Smail, J.B.; Squire, C.J. TAS-120 Cancer Target Binding: Defining Reactivity and Revealing the First Fibroblast Growth Factor Receptor 1 (FGFR1) Irreversible Structure. *ChemMedChem* **2019**, *14*, 494–500. doi: 10.1002/cmdc.201800719.
282. Zhao, Y.; Fang, C.; Zhang, Q.; Zhang, R.; Zhao, X.; Duan, Y.; Wang, H.; Zhu, Y.; Feng, L.; Zhao, J.; Shao, M.; Yang, X.; Zhang, L.; Peng, C.; Yang, K.; Ma, D.; Rao, Z.; Yang, H. Crystal Structure of SARS-CoV-2 Main Protease in Complex with Protease Inhibitor PF-07321332. *Protein Cell* **2022**, *13*, 689–693. doi: 10.1007/s13238-021-00883-2.
283. Shekh-Ahmad, T.; Eckel, R.; Dayalan Naidu, S.; Higgins, M.; Yamamoto, M.; Dinkova-Kostova, A.T.; Kovac, S.; Abramov, A.Y.; Walker, M.C. KEAP1 Inhibition Is Neuroprotective and Suppresses the Development of Epilepsy. *Brain* **2018**, *141*, 1390–1403. doi: 10.1093/brain/awy071.
284. Totir, M.A.; Helfand, M.S.; Carey, M.P.; Sheri, A.; Buynak, J.D.; Bonomo, R.A.; Carey, P.R. Sulbactam Forms Only Minimal Amounts of Irreversible Acylate-Enzyme with SHV-1 β -Lactamase. *Biochemistry* **2007**, *46*, 8980–8987. doi: 10.1021/bi7006146.
285. Shapiro, A.B.; Moussa, S.H.; McLeod, S.M.; Durand-Réville, T.; Miller, A.A. Durlabactam, a New Diazabicyclooctane β -Lactamase Inhibitor for the Treatment of Acinetobacter Infections in Combination With Sulbactam. *Front. Microbiol.* **2021**, *12*, 709974. doi: 10.3389/fmicb.2021.709974.

9. Supporting Information

Table S1 | FDA approved drugs (2020-2023) with a known covalent mechanism of action.^a

Active Ingredient ^b	Drug Brand Name	Sponsor ^c	Approval Year	Target	Indication	Treatment Area	Warhead	Reversibility	Target Residue	Ref
Lurbinectedin (PM01183)	Zepzelca	Pharma Mar	2020	DNA minor groove (alkylation)	Metastatic SCLC	Cancer	Highly reactive carbonylamine (Imine intermediate)	Reversible	Guanine N2	209
Remdesivir (GS-5734) ^d	Veklury	Gilead Sciences	2020	SARS-CoV-2 RdRp SARS-CoV-2 M ^{pro}	COVID-19	Anti-microbial	Nitrile	Reversible	Ser861 (RdRp) Cys145 (M ^{pro})	64, 280
Sotorasib (AMG 510)	Lumakras	Amgen	2021	KRAS ^{G12C}	KRAS ^{G12C} -mutated NSCLC	Cancer	Acrylamide	Irreversible	Cys12	18
Mobocertinib (TAK788)	Exkivity	Takeda	2021	EGFR ^{ex20ins}	Metastatic EGFR ^{ex20ins} -mutated NSCLC	Cancer	Acrylamide	Irreversible	Cys797	19
Futibatinib (TAS-120)	Lytgobi	Taiho Oncology	2022	FGFR1-4	FGFR2 fusion-positive ICC	Cancer	Acrylamide	Irreversible	Cys488 (FGFR1) Cys492 (FGFR2-IIIb)	281
Adagrasib (MRTX849)	Krazati	Mirati Therapeutics	2022	KRAS ^{G12C}	KRAS ^{G12C} -mutated NSCLC	Cancer	Acrylamide	Irreversible	Cys12	103
Nirmatrelvir (PF-07321332)	Paxlovid	Pfizer	2023	SARS-CoV-2 M ^{pro}	COVID-19	Anti-microbial	Nitrile	Reversible	Cys145	47, 282
Omaveloxolone (RTA 408)	Skydarys	Reata Pharmaceuticals	2023	KEAP1	Friedreich's ataxia	Neurodegenerative disease	Cyanoenone	Reversible	Cys151	283
Sulbactam/Durlobactam	Xacduro	Entasis Therapeutics	2023	PBP3 β-lactamase	Bacterial pneumonia	Anti-microbial	β-lactam Diazabicyclooctane	Reversible Reversible	Ser130 (SHV-1) Ser81 (OXA-24/40)	284-285
Ritlecitinib (PF-06651600)	Liftulo	Pfizer	2023	JAK3	Alopecia Areata	Autoimmune	Acrylamide	Irreversible	Cys909	199

Last updated: August 2023. FDA approved covalent drugs (1900-2019) are listed in Table S1 of citation²⁰. ^a Data source: FDA database. ^b Only the New Chemical Entity (NCE) approved in the specified year is reported for combination drugs. ^c Refers to the original sponsor for FDA first approval. ^d Covalent binding mode suspected based on covalent docking studies.