

Alkynes in covalent enzyme inhibitors: down the kinetic rabbit hole

Mons, E.

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Chapter 3



Kinetic Evaluation of Covalent Inhibition in Enzymatic Assays

Adapted from:

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Abstract. Covalent inhibition has become more accepted in the past two decades, as illustrated by the clinical approval of several irreversible inhibitors designed to covalently modify their target. Elucidation of the structure-activity relationship and potency of such inhibitors requires a detailed kinetic evaluation. Here, we elucidate the relationship between the experimental read-out and the underlying inhibitor binding kinetics. Interactive kinetic simulation scripts are employed to highlight the effects of *in vitro* enzyme activity assay conditions and inhibitor binding mode, thereby showcasing which assumptions and corrections are crucial. Four stepwise protocols to assess the biochemical potency of (ir)reversible covalent enzyme inhibitors targeting a nucleophilic active site residue are included, with accompanying data analysis tailored to the covalent binding mode. Together, this will serve as a guide to make an educated decision regarding the most suitable method to assess covalent inhibition potency.



A Comprehensive Guide for Assessing Covalent Inhibition in Enzymatic Assays Illustrated with Kinetic Simulations

1. Introduction

Traditionally, drug design efforts were focused on small molecules that interact with their biological target through noncovalent interactions in a reversible manner. In contrast, covalent inhibitors have the ability to form a much stronger covalent bond with a nucleophilic amino acid residue at the target protein, which is positioned in close proximity to a reactive (electrophilic) moiety in the inhibitor.¹ Risks associated with covalent reactions that can take place not only with the desired target but also with off-target proteins, often undiscovered until late-stage clinical development, resulted in drug discovery programs moving away from candidates bearing intrinsically reactive electrophilic moieties.²⁻³ Nonetheless, the clinical success of covalent drugs that were being used in the clinic long before their mechanism of action was elucidated, which include aspirin and penicillin, along with the more recent clinical approval and success of targeted covalent inhibitors (TCIs) bearing moderately reactive electrophilic warheads, ultimately triggered the current resurgence of covalent drugs.³⁻⁵

The covalent inhibitor development process typically involves identification of noncovalent inhibitors by high-throughput screening (HTS), followed by modification with a moderately reactive electrophilic warhead to improve inhibition potency and selectivity.⁶⁻⁷ Alternatively, an electrophilic fragment that forms a covalent bond with the desired enzyme target is first identified in covalent fragment-based drug discovery,⁸⁻¹⁰ followed by optimization of the noncovalent affinity and positioning of the electrophile. A prerequisite here is that the molecular target must contain a nucleophilic residue (e.g. cysteine, serine, lysine) to form a covalent bond with the electrophilic warhead of the inhibitor.¹¹⁻¹² Whether covalent adduct formation is reversible or irreversible depends on the selected electrophilic warhead.¹³⁻¹⁶ The PK-PD decoupling is one of the major advantages of irreversible inhibition: an infinite target residence time, resulting in a prolonged therapeutic effect after the inhibitor has been cleared from circulation.^{5, 17-19} Here, restoration of enzyme activity can only be achieved by *de novo* protein synthesis. At the same time, if the consequences of continued on-target inhibition are poorly understood, this same property can provide a safety concern. Consequently, inhibitors with a reversible covalent binding mode have become increasingly popular, with (tunable) target residence times ranging from several hours to multiple days.^{13, 20-21}

Although traditional methods to evaluate inhibitor potency, such as determining half-maximal inhibitory concentration (IC₅₀ values), are sufficient to identify hits in high-throughput screens, a more detailed kinetic evaluation is required to elucidate the structure-activity relationship (SAR) of irreversible covalent inhibitors.^{4, 22-23} There are many extensive reviews on the history, development, and success of covalent inhibitors,^{4-5, 12, 24} and experimental methods to assess undesired time-dependent inactivation (TDI) of CYP enzymes have been excellently reviewed,²⁵ but a comprehensive overview of experimental methods compatible with the desired covalent binding mode of TCIs targeting nucleophilic active-site residues has been missing. In *section 2*, we will introduce our customized set of interactive kinetic simulation scripts to study the kinetic concepts of different experimental methods, followed by a general background on (covalent) inhibitor binding modes, the assumptions on experimental enzyme activity assay conditions, and an introduction on time-dependent inhibitor kinetics. In *section 3*, we use kinetic simulations to evaluate four kinetic methods and discuss how assay

conditions affect the outcome in the subsequent data analysis. Stepwise experimental protocols with data analysis protocols tailored to the different covalent binding modes are provided in *section 4*, allowing readers to evaluate their covalent (ir)reversible inhibitor, together with the troubleshooting guidelines in *section 5*.

2. Kinetic Background

This guide has been composed to aid readers that have identified a(n) (ir)reversible covalent inhibitor and are contemplating which experimental method to select for the follow-up SAR analysis. Here, the performance of the enzymatic assay is not expected to be troublesome, but the challenge lies in the design of an assay method that complies with (often implied but not explicitly mentioned) assumptions on experimental conditions, and recognition of artifacts/ errors in the interpretation of experimental outcome. As such, we assume that a functioning enzymatic assay with a robust read-out is already in place, and we will focus on the connection between (algebraic) data analysis methods and the respective assumptions on experimental conditions. It is important to note that this work is tailored to enzyme <u>activity</u> assays with a (fluorescence) read-out upon substrate processing to form a detectable product, and as such may not be compatible with other assay formats such as ligand <u>binding</u> competition assays or direct detection of the covalent enzyme–inhibitor adduct.

In section 2.1, we introduce the interactive kinetic simulation scripts used to illustrate the methods and kinetic concepts in this work. All figures are composed with in silico data generated in kinetic simulations and can be recreated with the information in this section. Section 2.2 provides an overview of the (covalent) inhibition binding modes compatible with the methods in this work. It is paramount to select the appropriate algebraic model for data analysis, as the inhibitor binding mode changes the obtainable parameters as well as the compatibility with experimental methods. Covalent EI* adduct formation should be validated by direct detection with MS, protein crystallography or NMR.^{23, 26-28} Reversibility of covalent adduct formation is commonly assessed in rapid/jump dilution or washout assays with detection of regained enzymatic activity after dilution/washout,²⁹ MS detection of unbound inhibitor upon denaturation or digestion-mediated dissociation,¹³ or competitive binding of a (selective) irreversible (activity-based) probe.^{27,30} It is important to note that noncovalent binding can also irreversibly inhibit enzyme activity by aggregation or precipitation.³¹ Next, we investigated which assumptions on experimental enzyme activity assay conditions are embedded in the algebraic models used for kinetic analysis. Our findings are outlined in section 2.3, highlighting which assumptions are crucial and what the consequences are when these assumptions are violated. Finally, we provide a kinetic background on time-dependent (covalent) inhibition in section 2.4. Readers new to the field of enzyme inhibition kinetics are strongly encouraged to familiarize themselves with the work of Copeland ³²⁻³³ for a general introduction into enzyme kinetics before studying advanced kinetic concepts associated with (ir)reversible covalent enzyme inhibition and their relation to experimental enzyme activity read-out.

2.1. Kinetic Simulations

Keeping assay requirements in mind, it may seem a daunting task to design, perform, and analyze proper inhibition experiments. In general, practice is the best teacher to get a feeling for these assays and the expected output. Kinetic simulations are essential to understand the importance of reaction conditions and support assay design optimization.³⁴ In such simulations, one can freely change the parameters to visualize the effect on the output and validate that kinetic parameters found after data analysis correlate with the input values. This design precludes assay artifacts and human error, and also outputs the underlying concentrations of the different reaction species (e.g. unbound enzyme, enzyme-substrate complex), illustrating the relevance of the experimental assay conditions. Finally, kinetic simulations can validate if fitted experimental parameters correlate with the experimental read-out and aid the rational design of follow-up experiments by predicting the outcome.³⁵ Here, we use a set of customized kinetic simulation scripts based on numerical integration of the differential equations³⁶ to simulate the time-dependent product concentration as well as the underlying concentrations of various enzyme species (e.g. unbound, bound to inhibitor or substrate). Some concentrations are essentially constant under specific assay conditions, and treating these parameters as constants rather than variables reduces the computing/simulation time. An overview of our kinetic scripts and the assumptions on experimental assay conditions can be found in **Table 1**. Since understanding kinetics can be greatly facilitated by the ability to adjust reaction conditions and changing parameters without using expensive reagents, we have made interactive versions of these simulation scripts available free of charge at https://tinyurl.com/kineticsimulations. We encourage our readers to perform simulations with their own kinetic parameters to visualize how the underlying concentrations of enzyme species affect the detected read-out, and to get a feeling for realistic values and assay conditions. We selected one model inhibitor for each binding mode to generate the figures that exemplify the methods described (the kinetic parameters of each model inhibitor can be found in Table 2). All figures in this work can be recreated with the information in Table 1 and Table 2.

Our kinetic simulation scripts are tailored to competitive inhibition, where an intrinsically reactive inhibitor bearing an electrophilic warhead covalently targets a nucleophilic amino acid residue at the enzymatic substrate binding site, thus blocking substrate access.^{22, 33} Other covalent binding modes – e.g. prodrugs,³⁷ covalent allosteric inhibitors,³⁸ and multi-step mechanism-based inhibitors ³⁹⁻⁴⁰ – are outside the scope of this work, although the described experimental protocols can be useful in specific cases. For further instructions and detailed information on restrictions, we refer to the webpage itself. At the start of the simulations, we define the (pre)incubation time. The preincubation time is the elapsed time since the onset of enzyme inhibition by mixing enzyme and inhibitor, but before the onset of product formation: <u>after</u> substrate addition. In this work, we will distinguish between incubation and preincubation by using different symbols for preincubation t' (enzyme and inhibitor) and incubation *t* (enzyme, substrate and inhibitor) in all figures and equations to avoid confusion.

Reaction dynamics	Script	Simulat	Experimental restrictions	
$E+1 \xrightarrow{K_3} E1 \xrightarrow{K_{rest}} E1^*$	KinGen	Unbound inhibitor Unbound substrate Volume	$[1]_{0} = [1]_{t'} = [1]_{t}$ $[S]_{0} = [S]_{t}$ $V_{t'} = V_{t}$	$[I]_{0} > 10[E]_{0}$ $[S]_{0} > 10[E]_{0}$ $[P] < 0.1[S]_{0}$ $V_{sub} << V_{t'}$
$k_1 k_2$	KinSubDpl	Unbound inhibitor Volume	$[I]_0 = [I]_{t'} = [I]_t$ V _{t'} = V _t	$[I]_0 > 10[E]_0$ V _{sub} << V _t
k _{cat} E + P	KinVol	Unbound inhibitor Unbound substrate	$[I]_0 = [I]_t = (1 + (V_{sub}/V_t)) \times [I]_t$ $[S]_0 = [S]_t$	$[I]_0 > 10[E]_0$ [S]_0 > 10[E]_0 [P] < 0.1[S]_0
	KinInhDpl	Volume	$V_{t'} = V_t$	$V_{sub} \ll V_{t'}$
$ \begin{array}{c c} E_{\mathrm{log}} & EI_{\mathrm{log}} & EI_{\mathrm{log}} \\ \uparrow & K_{\mathrm{log}E} & \uparrow & K_{\mathrm{log}E} \\ \downarrow & K_{\mathrm{log}E} & I & K_{\mathrm{log}E} \\ & E + 1 & \overleftarrow{K_{4}} & EI & \overleftarrow{K_{\mathrm{log}E}} \\ & F & K_{\mathrm{log}} & E \\ & K_{\mathrm{log}} & K_{\mathrm{log}} \\ & K_{\mathrm{log}} & K_{\mathrm{log}} \\ & ES & \overleftarrow{K_{\mathrm{log}}} \\ & K_{\mathrm{cut}} \\ & E + P \end{array} $	KinDeg ^b	Unbound inhibitor Unbound substrate Volume	$[1]_{0} = [1]_{t'} = [1]_{t}$ $[S]_{0} = [S]_{t}$ $V_{t'} = V_{t}$	
	KinVolDeg ^b	Unbound inhibitor Unbound substrate	$[I]_0 = [I]_t = (1 + (V_{sub}/V_t)) \times [I]_t$ $[S]_0 = [S]_t$	[I] ₀ > 10[E] ₀ [S] ₀ > 10[E] ₀ [P] < 0.1[S] ₀

Table 1 Kinetic simulation scripts used in this work.^a

^a Available at <u>https://tinyurl.com/kineticsimulations</u>.^b First order spontaneous enzyme degradation/denaturation. [I]₀ = unbound inhibitor concentration at onset of inhibition, before (optional) enzyme binding. [I]_t = unbound inhibitor concentration during preincubation, after (optional) enzyme binding. [I]_t = unbound inhibitor concentration during incubation, after (optional) enzyme binding. [I]_t = unbound inhibitor concentration at onset of product formation, before enzyme binding. [S]_t = unbound substrate concentration during incubation, after (optional) enzyme binding and product formation. V_t = reaction volume during preincubation. V_{sub} = volume containing substrate. V_t = reaction volume during incubation (V_t = V_{sub} + V_t).

		Ing	out		Kinetic parameters ^a			
Reagent	k ₁ (M ⁻¹ s ⁻¹)	k ₂ (s ⁻¹)	k_{cat} (s ⁻¹)		κ (μΙ	d VI)	к_м (μМ)	k_{cat}/K_{M} (M ⁻¹ s ⁻¹)
Substrate S1	10 ⁸	100	1		1.0	1.00		9.9×10 ⁵
Ligand L1	10 ⁸	100	0		1.0	1.00		-
Inhibitor ^b	k ₃ (M ⁻¹ s ⁻¹)	k 4 (s ⁻¹)	k s (s ⁻¹)	k ₆ (s ⁻¹)	K _i (nM)	K _i * (nM)	K _I (nM)	$k_{\text{inact}}/K_{I} \text{ or } k_{\text{chem}}$ (M ⁻¹ s ⁻¹)
Α	10,000	0.0001	0	0	10	-	-	-
В	10 ⁸	10	0.001	0.0001	100	9.1	-	-
с	10 ⁸	10	0.001	0	100	-	100	10 ⁴
D	10,000	0	0	0	-	-	-	10 ⁴

Table 2 Kinetic parameters for simulated inhibitors used in this work to illustrate methods.

Reaction dynamics are illustrated in Table 1. ^aCalculated from microscopic rate constants in Table S1 and Figure S1D. ^b Mechanisms in Figure 1.

2.2. Inhibitor Binding Modes

Reversible noncovalent inhibitors inhibit enzymatic activity by formation of noncovalent EI complex in a single reaction step (**Figure 1A**). When the initial unbound inhibitor concentration is equal to inhibition constant K_i , the concentration of unbound enzyme E will be equal to the concentration of inhibitor-bound enzyme complex EI after steady-state equilibrium has been reached. For traditional fast-binding reversible inhibitors this equilibrium will be reached almost instantly, as association rate constant k_3 and dissociation rate constant k_4 are fast. In this work, the term 'reaction completion' relates to the endpoint of enzyme–inhibitor binding, which refers to reaching an equilibrium for reversible inhibitors (**Figure 1A** and **Figure 1B**) or reaching full inactivation for irreversible inhibitors (**Figure 1C** and **Figure 1D**). Contrary to classic fast-binding inhibitors, time-dependent or slow-binding inhibition is observed when the steady-state equilibrium or irreversible inactivation is reached relatively slowly on the assay timescale.⁴¹⁻⁴² Typically, this is observed for inhibitors with a covalent binding mode (**Figure 1B-D**), as formation of a covalent adduct is not an instantaneous process.

Reversible covalent adduct formation (**Figure 1B**) is a 2-step process consisting of (rapid) initial association to form noncovalent EI complex (*rapid equilibrium approximation*, discussed in more detail in *section 2.3*) preceding covalent EI* adduct formation. Covalent EI* adduct is at equilibrium with the noncovalent EI complex, as covalent adduct formation is reversible ($k_6 > 0$), with inhibition constant K_i reflecting the initial noncovalent E + I \leftrightarrow EI equilibrium and steady-state inhibition constant K_i* reflecting the steady-state (overall) E + I \leftrightarrow EI + EI* equilibrium. Development of reversible covalent inhibitors typically involves optimization of overall affinity (reflected in low K_i* values), preferably by slowing dissociation rates (**Figure 1B**). A slow off-rate (k_{off}) is favorable, as this is reciprocal with the drug-target residence time τ ($\tau = 1/k_{off}$), and a longer residence time has been linked to superior therapeutic potency.⁴³⁻⁴⁴ An overview of relevant kinetic parameters can be found in **Table S1**.

Inhibition is considered irreversible when its residence time exceeds the normal lifespan of the target enzyme.²² Dissociation from covalent EI* adduct is negligible, resulting in full enzyme engagement when reaction completion is reached for irreversible covalent inhibitors (**Figure 1C** and **Figure 1D**). The irreversible binding mode changes the obtainable kinetic parameters to rank inhibitor potency, as the biochemical IC_{50} may vary depending on the (pre)incubation time.^{3, 22} The potency of 2-step irreversible inhibitors that engage in an initial noncovalent enzyme–inhibitor complex EI prior to formation of covalent adduct EI* is driven by noncovalent affinity – reflected in inactivation constant K_1 – along with the maximum rate of inactivation k_{inact} (**Figure 1C**). Rate constant k_{inact}/K_1 is generally accepted as a more suitable measure of 2-step irreversible inhibitor potency, ^{3, 22, 37, 45} in an analogous fashion to k_{cat}/K_M reflecting the efficiency of enzymatic substrate conversion (a detailed comparison can be found in **Figure S1**). The binding mode becomes 1-step when noncovalent equilibrium is nonexistent, for example for highly reactive thiol-alkylating reagents,^{37, 46} with the parameter k_{chem} or $k_{obs}/[I]$ reflecting potency/efficiency (**Figure 1D**).

Drug development of irreversible covalent inhibitors is typically geared towards simultaneous improvement of the binding affinity (reflected in a lower K_I value) and faster covalent bond formation (reflected in a higher k_{inact} value) to generate irreversible covalent inhibitors



Figure 1 | Schematic overview of inhibitor binding modes.³⁹ E = unbound enzyme. I = unbound inhibitor. EI = noncovalent enzyme–inhibitor complex. EI* = covalent enzyme–inhibitor adduct. An overview of kinetic constants can be found in **Table S1**. (A) Classic 1-step reversible inhibition. Inhibitor potency ranking based on inhibition constant K_i (in M) or target residence time τ (in s). (B) Covalent 2-step reversible inhibition. Inhibitor potency ranking based on steady-state inhibition constant K_i^{*} (in M) for total E + I \leftrightarrow EI + EI* equilibrium or based on target residence time τ (in s). (C) Covalent 2-step irreversible inhibition (affinity label model). Inhibitor potency ranking based on inactivation efficiency: maximum rate of covalent adduct formation over inactivation constant k_{inact}/K_{I} (in M⁻¹s⁻¹). (D) Covalent 1-step irreversible inhibition (residue-specific reagent model). Inhibitor potency ranking based on inactivation efficiency: k_{chem} (in M⁻¹s⁻¹) = $k_{obs}/[I]$ (in M⁻¹s⁻¹).

with a high $k_{\text{inact}}/K_{\text{I}}$ value for the desired enzyme target, ^{45, 47} while minimizing the intrinsic reactivity with undesired enzymes such as GSH.⁴⁸⁻⁵⁰ Typical reported $k_{\text{inact}}/K_{\text{I}}$ values range from 10⁵-10⁷ M⁻¹s⁻¹ for kinase inhibitors, ^{45, 51-52} 10¹-10⁵ M⁻¹s⁻¹ for protease inhibitors, ^{28, 53-54} 10²-10⁴ M⁻¹s⁻¹ for other target classes, ⁵⁵⁻⁵⁷ to 10⁻²-10² M⁻¹s⁻¹ for covalent fragments. ⁵⁸⁻⁵⁹ Ranges of clinically relevant $k_{\text{inact}}/K_{\text{I}}$ values are highly dependent on the nucleophilicity of the targeted amino acid (cysteine typically being more reactive than serine) and concentration of naturally present competitors (e.g. ATP-competitive inhibitors need to overcome competition by ATP at physiological concentrations far exceeding the K_{MATP}).

2.3. Assumptions on Experimental Assay Conditions

Experimental conditions should meet certain criteria in order to use algebraic fitting methods. In this section, we focus on the assumptions on the experimental conditions that are embedded in algebraic equations to analyze time-dependent (covalent) inhibition (summarized in **Table 3**). Generally, these assumptions involve simplifying the enzyme–inhibitor binding reaction to a

single rate-determining step along with fixing inhibitor/substrate concentrations to a constant value. To use algebraic fitting, the experiment should meet all the required conditions outlined in this section. More complex systems – such as bisubstrate assays or other binding modes like allostery – violate one or more of these, and require a different method of fitting. For such systems, numerical integration with dedicated software packages – e.g. KinTek,⁶⁰ DynaFit⁶¹ – is recommended. These packages are very powerful, and can fit anything with good error even when the model does not reflect the biological situation.⁶² For these complex systems, it is crucial to ensure that the initial values are reasonable and the amount of (orthogonal) data is sufficient for the amount of parameters that are fitted. The first step, however, whether working with complex systems or reactions with a single rate-determining step, should always be optimization of the experimental conditions.

There are two distinct types of algebraic analysis: linear regression (fitting straight curves, compatible with commonly available software such as EXCEL) and nonlinear regression (fitting exponential curves, requiring sophisticated data fitting software). Linear regression was the predominant method to analyze kinetic data, but has now been surpassed by the more accurate nonlinear regression.⁶³ For our analyses, we use least-squares nonlinear regression with GraphPad Prism (RRID:SCR_002798), but other software packages are available too.⁶⁴ Please consult the detailed (online) guide on how to implement user-defined equations for nonlinear regression in GraphPad Prism.⁶⁵⁻⁶⁶

Michaelis-Menten enzyme kinetics. All experimental methods in this manuscript are based on enzyme activity assays with multiple turnovers per enzyme, with enzyme release after product formation. We assume that the uninhibited enzymatic substrate processing reaction $(E + S \leftrightarrow ES \rightarrow E + P)$ complies with Michaelis-Menten enzyme kinetics.^{35,64} The concentration of unbound substrate has to be constant ($[S]_t = [S]_0$) and not depleted by engagement in a (non)covalent complex ES ($[ES]_t < 0.1[S]_0$) or conversion into product. Therefore, substrate is added in a large excess over the enzyme ($[S]_0 > 10[E]_0$), and the uninhibited velocity of product formation (v^{ctrl}) is calculated over the linear part corresponding to less than 10% substrate conversion ($[P]_t < 0.1[S]_0$).⁶⁷ The signal corresponding to 10% substrate conversion can be

	Substrate concentration is constant Substrate is not depleted by complexation with enzyme	$[S]_t = [S]_0$ $[S]_0 > 10[E]_0$	
Michaelis-Menten enzyme kinetics	Uninhibited product formation is linear Product formation does not deplete substrate Enzyme activity is not affected by product inhibition The enzyme does not degrade significantly	$\begin{split} k_{\rm ctrl} &= 0 \\ [P]_t < 0.1[S]_0 \\ [P]_t << 0.1K_{\rm D,P} \\ k_{\rm deg} &= 0 \end{split}$	
Pseudo-first order kinetics	The unbound inhibitor concentration is constant Inhibitor is not depleted by complexation with enzyme	$[I]_{t} = [I]_{0}$ $[I]_{0} > 10[E]_{0}$	
Rapid equilibrium approximation	Noncovalent equilibrium is reached quickly for 2-step inhibitors Covalent adduct formation is rate-determining	k _{inact} << k ₄ k ₅ << k ₄	

 Table 3
 General assumptions on experimental assay conditions.

estimated from a product calibration/titration curve to avoid substrate depletion.⁶⁸⁻⁶⁹ The effect of substrate depletion can be investigated with the kinetic simulation script **KinSubDpl**. More complex enzymatic (bisubstrate) assays are outside of the scope of this work.³² However, the methods described herein could still be applicable under pseudo-single substrate (Hit-and-Run) conditions.

Enzyme stability. Unless otherwise noted, time-dependent decrease of enzyme activity is attributed solely to the presence of a (slow-binding) inhibitor. It is thus assumed that the enzyme activity is constant throughout the whole experiment, although this does not necessarily reflect the actual experimental situation. Recombinant enzymes do not have an eternal life; thus, timedependent loss of enzyme activity will inevitably occur due to spontaneous protein denaturation, degradation, or unfolding.⁷⁰ The Selwyn test is a relatively simple test to see if time-dependence of uninhibited enzyme activity is due to (spontaneous) enzyme inactivation.⁷¹ Spontaneous enzyme degradation/denaturation is similar to radioactive decay in a sense that inactivation is a first order reaction (*degradation rate* = $k_{degE} \times [E]$). Enzyme stability might be promoted by optimization of the assay buffer, and is less significant at shorter (pre)incubation times, but degradation cannot completely be avoided. Therefore, we included data analysis methods to account for spontaneous first order enzyme degradation/denaturation. Cannibalistic proteases follow a second order (auto)proteolysis rate (*degradation rate* = $k_{deeE} \times [E]^2$)⁷² and are as such outside of the scope of these methods. In simulations to illustrate the methods described herein (with kinetic simulation scripts **KinDeg** and **KinVolDeg**), we assumed that first order decay is uniform for all enzyme species ($k_{degE} = k_{degES} = k_{degEI} = k_{degEI^*}$) and combined the individual degradation rates into the enzyme degradation rate constant k_{deo} .

Constant uninhibited product formation velocity. The uninhibited controls should be linear for the whole measurement when analyzing time-dependent inhibition. There are various factors contributing to a slight time-dependent decrease of product formation velocity in the absence of inhibitor,³² thus violating this assumption. An overview of common troubleshooting options is listed in **Table 6** (located in *section 5*). As discussed above, substrate depletion ($[P] > 0.1[S]_0$) negatively influences the linearity over time, as does product inhibition $([P] > 0.1K_{D,P})$. Fortunately, this can be avoided by decreasing the enzyme concentration and/ or shortening the incubation time to reduce substrate turnover, thereby lowering the absolute and relative product concentration. Other factors can make the results look nonlinear – such as quenching of the fluorescent product signal by photobleaching.⁷³ This effect can be reduced by increasing the measurement interval and/or reducing the number of excitation cycles. Finally, optimization of assay conditions can minimize the effect of spontaneous loss of enzyme activity ($k_{deg} > 0$) but cannot be resolved completely. In this work, we will refer to the overall rate of nonlinearity in the uninhibited control (k_{obs} of [I] = 0) with the symbol k_{ctrl} , regardless of the underlying mechanism that causes the time-dependent decrease of product formation velocity.

Rapid equilibrium approximation. Algebraic analysis of (covalent) inhibition is based on the assumption that time-dependent inhibition is driven by a single rate-determining step. For 2-step covalent inhibitors (**Figure 1B** and **Figure 1C**), this means that the noncovalent $E + I \leftrightarrow EI$ equilibrium that precedes covalent EI^* adduct formation should be reached almost instantly after the onset of inhibition. After this rapid equilibrium, a much slower step

of covalent adduct formation follows ($k_{inact} \ll k_4$). Whether the noncovalent equilibrium indeed is reached rapidly is an intrinsic inhibitor property, and (kinase) inhibitors with a low-nM noncovalent potency are likely to violate this assumption: the association rate constant is diffusion-limited ($k_3 \le 10^9 \text{ M}^{-1}\text{s}^{-1}$), and thus k_4 must be relatively slow if $K_i \le 10^{-8} \text{ M}$.⁷⁴ Unfortunately, a slow initial, noncovalent step is not easily recognized from raw kinetic data, resulting in overestimation of the rate of inactivation k_{inact} and underestimation of the inactivation constant K_1 with algebraic rather than numerical data analysis. The inactivation constant K_1 approximates inhibition constant K_i ($K_1 \approx K_i$) when covalent bond formation is driven by the rate-determining conversion of noncovalent complex EI into covalent adduct EI* ($k_{inact} \ll k_4$) (**Figure 1C**), analogous to the Briggs–Haldane treatment of enzyme–substrate kinetics where $K_M \approx K_S$ if k_{cat} is rate-limiting.⁴⁶ Consequently, K_i and K_1 may have the same value, but they are not interchangeable, and it is as such recommended to report k_{inact}/K_1 rather than k_{inact}/K_i .

Pseudo-first order reaction kinetics without inhibitor depletion. Algebraic analysis of (covalent) inhibition is typically based on the assumption that the unbound inhibitor concentration is a constant value $([I]_t = [I]_0)$ unaffected by enzyme binding.³⁵ This assumption is only valid when the inhibitor is present in large excess with respect to the enzyme $([I]_0 > 10[E]_0)$ at reaction initiation. The enzyme occupancy after reaching the noncovalent equilibrium is driven solely by the excess inhibitor concentration relative to the (apparent) inhibition constant K_i^{app} : $[EI]_{eq}/[E]_0 = 1/(1 + (K_i^{app}/[I]))$. The effect of inhibitor depletion can be investigated with the kinetic simulation script **KinInhDpl**. Violation of this assumption results in an appreciable reduction of the remaining population of unbound inhibitor upon complexation with enzyme. Consequently, the inhibitor occupancy at equilibrium no longer reflects the apparent inhibition constant K_i^{app} because the equilibrium is now driven by both enzyme and inhibitor concentration (**Figure 2A**). Algebraic correction for inhibitor depletion ($[I]_t < [I]_0$) to find the equilibrium constant K_i is often performed for 1-step reversible inhibitors displaying tight-



Figure 2 | Consequences of inhibitor depletion. Simulated with **KinInhDpl** for 50 nM inhibitor **C** with 5 nM enzyme $([I]_0 = 10[E]_0)$ or 50 nM enzyme $([I]_0 = [E]_0)$. (A) Inhibitor depletion (*blue line*) results in lower noncovalent equilibrium occupancy $[EI]_{eq}$ and slower reaction rates, resulting in longer incubation time to reach full inactivation than for excess inhibitor (*black line*). (B) First order reaction conditions with constant half-life t¹/₂ when inhibitor is present in excess (*left*). Second order reaction conditions with variable half-life t¹/₂ and longer overall reaction time when inhibitor is depleted (*right*).

binding behavior (with low inhibitor concentrations because K_i^{app} approaches [E^{total}]), by fitting the (steady-state) equilibrium product formation velocity to (variants of) Morrison's quadratic equation that treat the inhibitor concentration as a variable rather than a constant value.⁷⁵⁻⁷⁶ However, these equations are only compatible with inhibitors with a reversible binding mode after equilibrium has been reached, and are thus not suitable for irreversible inhibition.

Binding of inhibitor to enzyme is, in principle, a second order reaction: the association rate depends on the concentration of unbound enzyme as well as unbound inhibitor, which both decrease upon formation of association product EI. Towards the end of the reaction, the reaction rate is significantly slower when less of the unbound components are left. Algebraic analysis of second order (ir)reversible association curves is complicated (data not included, simulated with simulation script KinInhDpl) – even for inhibitors with a 1-step binding mode - thus, it is strongly advised to analyze second order reactions of 2-step (ir)reversible inhibitors by numeric integration.⁷⁷ However, as mentioned above, unbound inhibitor concentrations remain more or less constant during the reaction if the inhibitor is present in excess at reaction initiation $([I]_0 > 10[E]_0)$. Consequently, the second order binding reaction of enzyme and inhibitor behaves like a first order reaction when the inhibitor is present in excess: pseudo-first order reaction kinetics.⁷⁷ The time-dependent association reaction for a (pseudo-)first order reaction has a constant half-life t¹/₂, and the progress curves can be fitted to standard onephase exponential association equations (Figure 2B, *left*), as will be discussed in more detail in section 2.4. Second order kinetic association reactions require a longer overall time to reach reaction completion of the enzyme-inhibitor binding reaction (inactivation or equilibrium) with a variable half-life t^{1/2} (Figure 2B, right), because the association reaction rate slows down when the remaining unbound inhibitor concentration decreases. For 2-step (ir)reversible inhibitors, the time-dependent reduction in covalent reaction rate is a direct consequence of the decreasing noncovalent occupancy upon inhibitor depletion. The rate-determining step of covalent adduct formation is preceded by noncovalent EI complex formation, and is thus limited by noncovalent occupancy, which decreased over time.

2.4. Time-Dependent Inhibitor Potency

Methods to analyze time-dependent inhibitors are based on the fact that it takes time to reach completion, and we use this information to obtain kinetic parameters. Under pseudo-first order conditions based on a single rate-determining step,⁷⁷ inhibitor binding follows an exponential one-phase association reaction ³⁵ from the rapid initial binding (*rapid equilibrium approximation*) to (slowly) reaching a plateau at <u>reaction completion</u>: equilibrium for reversible inhibitors (**Figure 3A**, *left*) or inactivation for irreversible inhibitors (**Figure 3A**, *right*). The incubation time to reaction completion is infinite, but after five half-lives ($t = 5t\frac{1}{2}$) the reaction progress is at 97%, which is generally sufficient to be considered reaction completion. Reaction half-life $t\frac{1}{2}$ is inversely related to observed reaction rate k_{obs} : $t\frac{1}{2} = LN(2)/k_{obs}$.⁷⁷ k_{obs} is the experimental reaction rate for reaction progress from initial binding to reaction completion under the specific assay conditions. Inhibitor concentration as well as competing substrate concentration are major contributors to the observed reaction rate k_{obs} . The experimental k_{obs} value can be obtained by fitting the time-dependent binding/occupancy curve to exponential one-phase association **Equation I** (shown in **Figure 3B**) from initial to final enzyme occupancy.



Figure 3 | Time-dependent inhibition and reaction completion. Simulated with KinGen for 1 pM enzyme with substrate S1. (A) Time-dependent enzyme occupancy simulated for 50 nM 1-step reversible inhibitor A (left) or 2-step irreversible inhibitor **C** (right) in presence of 100 nM substrate **S1**. Each half-life t_{λ} , the occupancy increases by 50% (of the remaining span). After 5t%, occupancy is at 97% of its maximum (equilibrium concentration [EI]_{e0} or total enzyme concentration [E] $_0$) and generally considered as reaction completion. Half-life t% is inversely related with observed reaction rate k_{obs} (under pseudo-first order conditions). (B) Bounded exponential association Equation I from initial occupancy (rapid equilibrium) to final occupancy (reaction completion). (C) Progress curve of time-dependent product formation for enzyme inhibition in panel A. Product formation velocity (slope, in AU/s), reflecting the (remaining) enzyme activity decreases until reaction completion is reached (steady-state equilibrium or inactivation). (D) Exponential association Equation II from initial velocity v_i (rapid equilibrium) to final velocity v_s (reaction completion). (E) k_{obs} curves in absence (black, [S] = 0) or presence (gray, [S] = 2K_M) of competing substrate. Apparent values are not yet corrected for substrate competition. (F) 2-step irreversible covalent inhibitors display 1-step behavior at non-saturating inhibitor concentrations ($[I] \le 0.1 K_{\tau}$). Fit straight line with Y-intercept = k_{ctrl} to obtain $k_{\text{chem}} = (k_{\text{inact}}/K_{\text{I}})$ from the linear slope. (G) 2-step irreversible covalent inhibitors display 1-step behavior at saturating inhibitor concentrations ($[I] > 10K_1$). Distinguish from non-saturating inhibitor concentrations in panel F: Y-intercept > k_{ctrl} when fitting a straight line to the k_{obs} curve.

Biochemical inhibitor potency is seldom assessed by direct observation of enzyme complex/ adduct. Typically, enzyme inhibition is indirectly assessed in *in vitro* assays with a detectable read-out for product formation as a measure of (remaining) enzyme activity. Consequently, reversible enzyme inhibition may have reached the enzyme–inhibitor binding equilibrium (*reaction completion*), but not all enzyme is occupied (unless [I] >> K_i^{app}) so the remaining fraction of unbound enzyme continues to convert substrate into product (**Figure 3C**, *left*). The reaction is no longer accurately reflected by **Equation I** (shown in **Figure 3B**), as product concentration at reaction initiation does not reflect the initial binding equilibrium, and product concentration does not reach a plateau after reaching the noncovalent equilibrium (*reaction completion*) for reversible inhibitors. Therefore, time-dependent product formation is fitted to exponential one-phase association **Equation II** (shown in **Figure 3D**) to obtain observed reaction rate k_{obs} from initial to final product formation velocity. For irreversible inhibitors, the initial velocity v_i reflects the (remaining) enzyme activity after rapid noncovalent association, and final velocity $v_s = 0$ as this reflects full enzyme inactivation (**Figure 3C**, *right*).

Typically, substrate competition assays are run at various inhibitor concentrations, and the concentration-dependent k_{obs} is fitted to obtain kinetic parameters (Figure 3E). In this work, equations and simulations are tailored to competitive binding of inhibitor and substrate.^{22, 64} Consequently, the observed reaction rate k_{obs} in the presence of competing substrate is slower, and apparent kinetic constants (marked with ^{app}) need to be corrected for substrate competition to reflect the kinetic inhibitor potency. Unless otherwise noted, nonlinearity in the uninhibited control k_{ctrl} (k_{obs} of [I] = 0) is assumed to be 0. The relation between k_{obs} and inhibitor concentration holds important information on the inhibitor binding mechanism. A linear k_{obs} increase with inhibitor concentration is a hallmark of a 1-step binding mode, as reaction rates are only limited by experimental factors such as solubility. Plots of k_{obs} against 2-step inhibitor concentrations are hyperbolic, as the experimental covalent EI* association rate is limited by EI occupancy, which reaches its maximum (k_{inact} or k_5) at saturating inhibitor concentration, as shown in **Figure 3E**: $[I] > 10K_1$ for 2-step irreversible inhibitors or $[I] > 10K_1$ for 2-step reversible inhibitors. An exception to this general observation is inhibitors with a 2-step binding mode that will display a linear relationship when assessed at all non-saturating inhibitor concentrations (Figure 3F) or all saturating inhibitor concentrations (Figure 3G).³⁷ These 1-step binding behaviors can be distinguished from the Y-intercept (Y0 = k_{crt} for $[I] \ll K_i^{app}$ and Y0 > k_{ctrl} for $[I] \gg K_i^{app}$ along with the noncovalent inhibition of enzyme activity ($v_i = v^{ctrl}$ for [I] << K_i^{app}) and $v_i < v^{ctrl}$ for [I] >> K_i^{app}).

3. Theoretical Framework for Experimental Methods and Data Analysis

We will discuss four methods in this work, that all have accompanying data analysis procedures tailored to the inhibitor binding mode (summarized in **Table 4**). In this section, we will first give an overview of the general conceptual background and assay design considerations for each assay method. Subsequent data analysis is tailored to a specific inhibitor binding mode, and we will illustrate the 'ideal' situation with kinetic simulations to guide interpretation of results. Furthermore, pointers on identification of deviations such as nonlinearity in the uninhibited control ($k_{ctrl} > 0$) will be given along with algebraic corrections or troubleshooting options to

resolve issues. The stepwise assay protocols and accompanying data analysis protocols can be found in *section 4*. A practical comment on the nomenclature used: we use the word 'fit' for nonlinear fits of raw data (in e.g. GraphPad Prism, as part of the *Data Analysis Protocols* in *section 4*) and 'calculate' to denote that we calculate parameters from experimental values (in EXCEL, as part of the *Sample Calculations* in *section 4*).

Methods I and *II* are based on incubation time-dependent enzyme inhibition (**Figure 4**). Here, substrate and inhibitor are mixed, and the reaction is initiated by addition of enzyme: i.e. simultaneous onset of product formation and enzyme inhibitor. *Methods III* and *IV* are based on enzyme inhibition after preincubation. Here, enzyme is preincubated with inhibitor before substrate addition. Two major factors contribute to selection of the appropriate experimental method for your enzymatic inhibition assay: the available enzyme activity assay and the inhibitor binding mode (the most important considerations are summarized for each method and inhibitor binding mode in **Table 4**). Recombinant enzyme inhibition is assessed in an *in vitro* enzyme activity assay with detectable product formation.⁷⁸⁻⁷⁹ This can be a continuous read-out for enzymatic processing of fluorogenic substrates (e.g. fluorescence intensity, FRET) or be a stopped/quenched assay that may require a secondary development/quenching or separation step to detect the formed product (or remaining substrate) such as LCMS-based

Assay Method	Data Analysis Protocol	Binding Mode	Read-out and Assay Conditions ^a	Obtainable Kinetic Parameters	Comments/Remarks	Ref	
	1A 2-step IRREV		Continuous	k _{inact} K _I k _{inact} /K _I	Continuous detection of enzyme activity. Optimization of reaction conditions to minimize assay artefacts can be challenging		
1B 1 1C 1D		1-step IRREV	Continuous	Continuous k _{chem} arteracts can be challenging but rewards with the simplest experimental procedure. Method I is generally favored: mos			
	18	2-step IRREV	Continuous [I] << K _i ^{app}	$k_{\text{inact}}/\text{K}_{\text{I}}$	informative in a single measurement. Favored for very potent inhibitors because competing substrate is always present		
	1C	2-step REV	Continuous k _{ctrl} << k ₆	K _i *	Method I is disfavored over Method III as algebraic correction for spontaneous loss of enzyme activity is NOT possible for 2-step REV inhibitors.		
	1D	2-step IRREV	Continuous [P] _t > 0.1[S] ₀ [S] << 0.1K _M	k _{inact} K _I k _{inact} /K _I	Algebraic correction for substrate depletion.	80	
П	2	2-step IRREV	Continuous, Quenched k _{ctri} = 0	$k_{ m inact} \ { m K_{I}} \ { m K_{I}} \ { m k}_{ m inact} / { m K_{I}}$	Incubation time-dependent potency. Compatible with quenched assays but is sensitive to spontaneous loss of enzyme activity.	81	

Table 4 Concise Summary of Methods.

Table 4 continues on the next page

Assay Method	Data Analysis Protocol	Binding Mode	Read-out and Assay Conditions ^a	Obtainable Kinetic Parameters	Comments/Remarks	Ref	
111	3 + 3Ai	2-step IRREV	Continuous, Quenched [S] << K _M V _{sub} << V _t ,	k _{inact} K _I k _{inact} /K _I			
	3 + 3Aii	2-step IRREV	Continuous, Quenched [S] << K _M V _{sub} << V _t .	k _{inact} K _I k _{inact} /K _I	enzyme activity after preincubation, with detection of inhibition by El and El* formed during preincubation. Experimental assays are relatively time-consuming.		
	3 + 3Bi	1-step IRREV	Continuous, Quenched [S] << K _M V _{sub} << V _t ,	k _{chem}	Favored for inhibitors with low potency and disfavored for very potent (tight-binding) inhibitors as preincubation is performed in absence of competing substrate. Data Analysis Protocols 3Ai and 3Bi are favored for comparison of		
		2-step IRREV	Continuous, Quenched [S] << K _M V _{sub} << V _t , [I] << K ₁	$k_{\rm inact}/{\rm K_I}$			
	3 + 3Bii	1-step IRREV	Continuous, Quenched [S] << K _M V _{sub} << V _t .	k _{chem} k _{obs} /I	multiple inhibitors on a single target. Data Analysis Protocols 3Aii and 3Bii are favored for selectivity evaluation of a single inhibitor on multiple	-	
		2-step IRREV	Continuous, Quenched [S] << K _M V _{sub} << V _t , [I] << K _I	$k_{\rm inact}/{\rm K_I}$	targets.		
	3 + 3C	2-step REV	Continuous, Quenched [S] << K _M V _{sub} << V _t .	K _i *	Favored over <i>Method I</i> for 2-step REV: normalization enables algebraic correction for spontaneous loss of enzyme activity.		
IV	4 + 4Ai	2-step IRREV	Continuous, Quenched [S] >> K _M V _{sub} >> V _t ,	k _{inact} K _I k _{inact} /K _I	Enzyme activity after preincubation,		
	4 + 4Aii	2-step IRREV	Continuous, Quenched [S] >> K_M $V_{sub} >> V_t$	$k_{ m inact} \ { m K_{I}} \ { m K_{I}} \ { m k_{inact}} / { m K_{I}}$	with detection of inhibition by covalent adduct EI* formed during preincubation (comparable to direct detection): dilution into (excess)		
	4 + 4Bi	1	1-step IRREV	Continuous, Quenched [S] >> K _M V _{sub} >> V _t ,	k _{chem}	substrate promotes noncovalent El dissociation. Favored for inhibitors with low	
		2-step IRREV	Continuous, Quenched [S] >> K _M V _{sub} >> V _t . [I] << K _I	k _{inact} /Κ _Ι	 (noncovalent) affinity, or to study the contribution of covalent bond formation. Data Analysis Protocols 4Ai and	83	
	4 + 4Bii	1-step IRREV	Continuous, Quenched [S] >> K _M V _{sub} >> V _t ,	k _{chem} k _{obs} /l	4Bi are favored for comparison of multiple inhibitors on a single target. Data Analysis Protocols 4Aii and 4Bii are favored for selectivity evaluation of a single inhibitor on multiple		
		2-step IRREV	Continuous, Quenched $[S] >> K_M$ $V_{sub} >> V_{t'}$ $[I] << K_I$	k _{inact} /K _I k _{obs} /I	targets.		

Table 4 Concise Summary of Methods. (continued)

^a General assay conditions for all methods unless otherwise specified: [I] > 10[E]. [S] > 10[E]. $[P]_t < 0.1[S]_0$.



Figure 4 | Schematic overview of experimental protocols to analyze covalent inhibitor potency included in this work. Incubation time-dependent enzyme inhibition in *Method I* and *II*. Preincubation time-dependent enzyme inhibition in *Method III* and *IV*. Data analysis is tailored to 2-step irreversible inhibition (shown in **Figure 1C**), 1-step irreversible inhibition (shown in **Figure 1D**), or 2-step reversible inhibition (shown in **Figure 1B**).

assays, conversion of radiolabeled substrate, and commercial assay technologies including ADP-GloTM (Promega) ATP consumption/ADP production assays, HTRF[®] KinEASETM (Cisbio) and Z'-LYTE (Invitrogen) phosphorylation assays, and Amplex[®] Red (Invitrogen) hydrogen peroxide/peroxidase assays.⁷⁸⁻⁷⁹ *Method I* is only compatible with homogeneous enzymatic assays that allow continuous read-out, such as cleavage of fluorogenic reporter peptides by proteases. *Methods II-IV* are also compatible with quenched/stopped assays with development step prior to read-out.

METHOD I: Progress Curve Analysis of Substrate Association Competition

Progress curve analysis is an established method for kinetic analysis of slow-binding inhibitors based on continuous detection of product formation after the substrate processing/product formation reaction has been initiated by addition of enzyme to a mixture of inhibitor and substrate (**Figure 5A**). A single measurement at each inhibitor concentration is sufficient, which is convenient when comparing the potency of multiple inhibitors on the same target. However, this method requires the availability of an activity assay format with a continuous read-out, thereby limiting the substrates that can be used. Additionally, assay optimization for progress curve analysis is labor intensive: it is not uncommon to perform multiple pilot experiments to find suitable concentrations of substrate, enzyme, and inhibitor that ensure linear product formation in the uninhibited control (consult **Table 6** in *section 5* for troubleshooting).

For 'slow-binding' inhibitors, the slope of time-dependent product formation exponentially decreases from initial product formation velocity v_i (rapid noncovalent inhibition) to the final product formation velocity v_s (reaction completion) (Figure 5B).⁴¹ The progress curve of timedependent product formation (as detected signal F_t in AU) is fitted to a general exponential inhibitor association Equation II (shown in Figure 5C) to obtain the observed rate of reaction completion k_{obs} (in s⁻¹) from initial velocity v_i (in AU/s) to final velocity v_s (in AU/s). A 1-step or 2-step binding mode can be identified by (visual) inspection of the initial velocity (Figure 5B). The value of initial velocity v_i is inhibitor concentration-dependent for 2-step (ir)reversible inhibitors that form a rapid (noncovalent) equilibrium ($v_i < v^{ctrl}$) because the noncovalent enzyme-inhibitor complex already inhibits the enzyme activity (rapid equilibrium approximation). Similarly, the value of initial velocity v_i is equal to the uninhibited velocity v^{ctrl} in lieu of a rapid initial binding step, as can be observed for 2-step (ir)reversible inhibitors at non-saturating concentrations ([I] $\ll K_i^{app}$) and 1-step (ir)reversible inhibitors ($v_i < v^{ctrl}$). Irreversible inhibitors are expected to reach 100% inhibition at reaction completion for all inhibitor concentrations, provided inhibitor is present in large excess and the reaction does not exceed the dynamic enzyme lifetime. Therefore, the final velocity v_s is restrained to full inhibition ($v_s = 0$) for 2-step irreversible inhibitors (*Data Analysis 1A*) and 1-step irreversible inhibitors (Data Analysis 1B). A 2-step reversible inhibitor will reach a reversible steady-state equilibrium ($v_s \ge 0$) upon reaction completion (*Data Analysis 1C*). Be aware that the product formation progress curve is not only linear for fast-binding inhibitors but will also appear linear for slow-binding inhibitors if reaction completion is much slower than the time course of the



Figure 5 | Method I: Progress curve analysis of substrate association competition. Simulated with KinGen for 1 pM enzyme and 100 nM substrate S1. (A) The reaction between enzyme, inhibitor, and substrate is initiated by addition of enzyme. Product formation is monitored continuously to detect the time-dependent enzyme activity. Simulated for 50 nM inhibitor B (top) and inhibitor C (bottom). Enzyme inhibition increases with time-dependent formation of covalent EI* until reaching reaction completion. Initially, total enzyme occupancy [EI + EI*] reflects the rapid noncovalent equilibrium $[EI]_{eq}$. At reaction completion (t > 5t%), total enzyme occupancy EI + EI* reflects the steady-state equilibrium (reversible) or inactivation (irreversible). (B) Typical progress curves for enzyme activity in presence of time-dependent inhibitors. Time-dependent product formation decreases exponentially from initial velocity v_i (dashed green line) to the steady-state velocity v_c (dashed purple line) at reaction completion (t > 5t½), with v^{ctrl} = linear product formation in uninhibited control (gray line). $v_i = v^{\text{ctrl}}$ for 1-step IRREV inhibitors and for 2-step IRREV inhibitors when $[I] << K_i^{app}$. Simulated for 50 nM 1-step REV inhibitor A, 2-step REV inhibitor B, 1-step IRREV inhibitor D, and 2-step IRREV inhibitor C. (C) General exponential association Equation II to fit progress curves of time-dependent inhibition. Parameters are constrained depending on the inhibitor binding mode. $v_s = 0$ for irreversible inhibition: inactivation at reaction completion. $v_i = v^{\text{ctrl}}$ for 1-step inhibition: noncovalent complex is not significant at non-saturating inhibitor concentrations. The uninhibited controls are also fitted to obtain reference values for uninhibited velocity v^{ctrl} and the rate of nonlinearity in the uninhibited control k_{ctrl} . F_t = time-dependent signal resulting from product formation. F_0 = Y-intercept = background signal at reaction initiation. vi = initial product formation velocity. vs = final/steady-state product formation velocity. t = incubation time after enzyme addition. kobs = observed rate of time-dependent inhibition from initial v_i to final v_c .

assay ($t \ll t\frac{1}{2}$). Importantly, the noncovalent equilibrium is assumed to be reached instantly for 2-step inhibitors (*rapid equilibrium approximation*). An algebraic solution to analyze irreversible 2-step inhibitors violating the rapid equilibrium approximation is available as a preprint.⁷⁴

It is crucial to have linear product formation in the uninhibited control (F^{ctrl}), as progress curve fitting for time-dependent (ir)reversible inhibition relies on the assumption that uninhibited product formation is absolutely linear. This ideal situation is often not feasible to achieve

experimentally, as there are many factors contributing to a slight time-dependent decrease of product formation velocity in the uninhibited control, and not all of them are resolvable (common troubleshooting options are listed in **Table 6**, *section 5*). Algebraic correction for nonlinearity in the uninhibited control k_{ctrl} caused by spontaneous enzyme degradation/ denaturation is possible for irreversible inhibitors (*Data Analysis 1A-B*). Furthermore, it is also possible to perform an algebraic correction for substrate depletion for 2-step irreversible inhibitors (*Data Analysis 1D*).⁸⁴ Ultimately, numerical integration is the preferred method in complex systems where multiple events contribute to the observed nonlinearity.

Data Analysis 1A: 2-Step Irreversible Covalent Inhibition.

Data obtained for 2-step irreversible inhibitors with *Method I* (Figure 6A) is processed with *Data Analysis Protocol 1A*. The progress curve of time-dependent product formation of each inhibitor concentration is fitted to exponential Equation II (shown in Figure 5C) constraining final velocity to 100% inhibition ($v_s = 0$) at reaction completion (Figure 6B). The inhibitor concentration-dependent observed rate of inactivation k_{obs} reflects the rate from initial velocity v_i (rapid noncovalent equilibrium) to final velocity v_s (inactivation at reaction completion). The plot of inhibitor concentration-dependent k_{obs} (Figure 6C) reaches maximum rate of inactivation k_{inact} in the presence of saturating inhibitor concentration ($[I] >> K_I^{app}$) with the Y-intercept at 0 when the progress curve in absence of inhibitor is strictly linear. Importantly, the inhibitor concentration that results in half-maximum enzyme inactivation ($k_{obs} = \frac{1}{2}k_{inact}$) has to be corrected for competition by the substrate during incubation but maximum rate of inactivation k_{inact} is unaffected.

A linear plot of inhibitor concentration-dependent k_{obs} (with Y-intercept = k_{ctrl}) and an initial velocity independent of inhibitor concentration ($v_i = v^{ctrl}$) are indicative of a 1-step binding mechanism: the inhibitor concentration is not saturating ($[I] \le 0.1K_1^{app}$ and $[I] \le 0.1K_i^{app}$). This can be resolved by increasing the inhibitor concentration, reducing the substrate concentration, or processing the data with *Data Analysis Protocol 1B*. Inhibitors with a high noncovalent potency ($[I] >> K_i^{app}$) might exhibit tight-binding behavior: complete inactivation is reached at reaction initiation ($v_i = 0$), even at the lowest inhibitor concentration, without violating the pseudo-first order reaction conditions ($[I]_0 \ge 10[E]_0$). This can be resolved by lowering the inhibitor concentration, but only if the assay robustness is sufficient to also lower the enzyme concentration, and/or by increasing the concentration of competing substrate, thus increasing the apparent inhibitors with inhibitor depletion ($[I]_0 \le 10[E]_0$) is not compatible with 2-step inhibition.⁴¹ Numeric fitting is a possibility to fit progress curves with depletion of substrate as well as inhibitor.⁸⁰ Alternatively, tight-binding 2-step irreversible covalent inhibition can be assessed with *Method IV* if covalent adduct formation is relatively slow.

Spontaneous enzyme degradation/denaturation (**Figure 6D**) causes a nonlinearity in the uninhibited control ($k_{ctrl} > 0$) that violates the assumption that time-dependence in the inhibitor-treated samples is a direct effect of the inhibitor (**Figure 6E**). The first order enzymatic degradation rate contributes to k_{obs} independent of inhibitor concentration ($k_{degE} = k_{degEI} = k_{degEI}$). Consequently, the Y-intercept of the k_{obs} against inhibitor concentration plot now corresponds to observed rate k_{ctrl} in absence of inhibitor, and k_{max} is higher



Figure 6 | <u>Data Analysis 1A</u>: Progress curve analysis for 2-step irreversible covalent inhibition. Simulated with **KinGen** (panel A-C) or **KinDeg** (panel D-F) for 2-step IRREV inhibitor **C** with 1 pM enzyme and 100 nM substrate **S1**. (A) Schematic enzyme dynamics during incubation. (B) Time-dependent product formation in absence of inhibitor *F*^{ctrl} or in presence of inhibitor. The progress curve for each inhibitor concentration is fitted individually to **Equation II** (shown in **Figure 5C**) (constraining $v_s = 0$) to obtain the observed rate of inactivation k_{obs} . (C) Inhibitor concentration-dependent k_{obs} reaches k_{inact} at saturating inhibitor concentration constant K_1^{app} . (D) Schematic enzyme dynamics during incubation with spontaneous loss of enzyme activity. Simulated with $k_{degE} = k_{degEI} = 0.0003 \text{ s}^{-1}$. (E) Time-dependent product formation in absence of inhibitor *F*^{ctrl} is not linear because $k_{ctrl} > 0$. The progress curves for each inhibitor concentration and uninhibited control are fitted individually to **Equation II** (shown in **Figure 5C**) (constraining $v_s = 0$) to obtain the observed rates of inactivation k_{obs} . (F) Inhibitor concentration-dependent k_{obs} with spontaneous enzyme degradation increases with k_{ctrl} , but the span from k_{min} (= k_{ctrl}) to k_{max} (= $k_{inact} + k_{ctrl}$) still equals k_{inact} . Fit with algebraic correction for nonlinearity (*black line*, $k_{ctrl} > 0$). Ignoring the nonlinearity (*gray line*, constrain $k_{ctrl} = 0$) results in underestimation of K_1^{app} (overestimation of potency) and overestimation of k_{inact} .

 $(k_{\text{max}} = k_{\text{inact}} + k_{\text{ctrl}})$ (**Figure 6F**). Performing a simple algebraic correction for the observed nonlinearity due to spontaneous enzyme degradation results in good estimates for k_{inact} and K_1^{app} (**Figure 6F**). Ignoring the nonlinearity in the uninhibited control by restraining $k_{\text{ctrl}} = 0$ implies that all time-dependent loss of enzyme activity should be attributed to inhibitor-mediated inactivation, resulting in an underestimation of inactivation constant K_1^{app} (overestimation of potency) and overestimation of k_{inact} . This effect is less pronounced when spontaneous enzyme degradation is much slower than the maximum rate of covalent adduct formation $(k_{\text{inact}} >> k_{\text{ctrl}})$. It is important to note that stabilization of the enzyme species by (noncovalent) inhibitor binding also decreases the contribution of k_{ctrl} to the observed rate k_{obs} at saturating inhibitor concentrations ($k_{\text{max}} = k_{\text{inact}}$). This impairs the accuracy of the algebraic correction

Data Analysis 1A: 2-step IRREV

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unless k_{ctrl} is relatively small (k_{max} approaches k_{inact} if $k_{inact} >> k_{ctrl}$). This algebraic correction does not accurately correct for nonlinearity due to substrate depletion ([P]_t > 0.1[S]₀): substrate depletion is dependent on the total product formation and does not (significantly) contribute to k_{max} at saturating inhibitor concentration because enzyme inhibition reduces the total amount of product formed ($k_{max} = k_{inact}$). Please consult *Data Analysis 1D* for algebraic correction of nonlinearity due to substrate depletion.

Data Analysis 1B: 1-Step Irreversible Covalent Inhibition.

Data obtained for 1-step irreversible inhibitors with *Method I* (Figure 7A) is processed with *Data Analysis Protocol 1B*. The progress curve of time-dependent product formation of each inhibitor concentration is fitted to exponential Equation II (shown in Figure 5C) constraining final velocity to inactivation ($v_s = 0$) at reaction completion (Figure 7B). The initial velocity v_i equals the uninhibited product formation velocity ($v_i = v^{\text{ctrl}}$), as noncovalent inhibitor binding



Figure 7 | <u>Data Analysis 1B</u>: Progress curve analysis for 1-step irreversible covalent inhibition. Simulated with **KinGen** (panel A-C) or **KinDeg** (panel D-F) for 1-step IRREV inhibitor **D** with 1 pM enzyme and 100 nM substrate **S1.** (**A**) Schematic enzyme dynamics during incubation. (**B**) Time-dependent product formation in absence of inhibitor F^{ctrl} or in presence of inhibitor. The progress curve for each inhibitor concentration is fitted individually to **Equation II** (shown in **Figure 5C**) (constraining $v_s = 0$) to obtain the observed rate of inactivation k_{obs} . $v_l = v^{ctrl}$ for 1-step irreversible inhibitors and 2-step irreversible inhibitor s at non-saturating concentrations ([I] << K_i^{app}). (**C**) Inhibitor concentration-dependent k_{obs} increases linearly with inhibitor concentration, with k_{chem}^{app} as the slope. (**D**) Schematic enzyme dynamics during incubation with spontaneous loss of enzyme activity. Simulated with $k_{degE} = k_{degEI} = 0.0003 \text{ s}^{-1}$. (**E**) Time-dependent product formation in absence of inhibitor F^{ctrl} is not linear because $k_{ctrl} > 0$. The progress curves for each inhibitor concentration and uninhibited control are fitted individually to **Equation II** (shown in **Figure 5C**) (constraining $v_s = 0$) to obtain the observed rates of inactivation k_{obs} . (**F**) Inhibitor concentration-dependent k_{obs} with spontaneous enzyme degradation/denaturation increases by k_{ctrl} . Fit with algebraic correction for nonlinearity (*black line*, $k_{ctrl} > 0$) or ignoring nonlinearity (*gray line*, constrain $k_{ctrl} = 0$). Ignoring the nonlinearity (assuming Y-intercept = 0) results in overestimation of k_{chem}^{app} (steeper slope).

does not contribute to enzyme inhibition by 1-step irreversible inhibitors. A linear plot of inhibitor concentration-dependent k_{obs} is indicative of a 1-step binding mechanism with k_{chem}^{app} as the slope (**Figure 7C**). A 2-step irreversible covalent inhibitor also has a linear k_{obs} against inhibitor concentration plot at non-saturating concentrations ([I] $\leq 0.1K_1^{app}$): $k_{chem}^{app} = k_{inact}/K_1^{app}$. The slope has to be corrected for substrate competition to obtain the inactivation constant k_{chem} . Substrate will occupy a fraction of the unbound enzyme to reach the noncovalent E + S \leftrightarrow ES equilibrium (how much depends on [S]/K_M), thus reducing the unbound enzyme concentration. It may seem counterintuitive to correct for substrate competition, as the pseudo-first order rate of covalent adduct formation ($k_{obs} = k_{chem}^{app} \times [I]$) does not seem to involve unbound enzyme (provided inhibitor is present in large excess), but formation of EI* is limited by the available unbound enzyme at that moment and it is not possible to form covalent adduct EI* when competing substrate blocks access to the enzyme active site.

It is important to have linear product formation in the uninhibited control ($k_{ctrl} = 0$) or to perform an algebraic correction for nonlinearity in the uninhibited control ($k_{ctrl} > 0$) caused by spontaneous first order enzyme degradation/denaturation (**Figure 7D-F**). Failure to correct for the contribution of enzyme degradation when fitting the observed rate of inactivation k_{obs} against inhibitor results in overestimation of k_{chem}^{app} (**Figure 7F**, *gray line*). The contribution of nonlinearity k_{ctrl} becomes less pronounced at elevated inhibitor concentrations as k_{ctrl} becomes significantly smaller than k_{obs} ($k_{ctrl} << k_{chem}^{app} \times [I]$). (De)stabilization of enzyme upon inhibitor binding (k_{degEI^*}) does not affect k_{obs} , as EI* formation is already irreversible, thus removing the species from the available pool of catalytic enzyme. To our knowledge, methods to algebraically correct for substrate depletion have not been reported.

Data Analysis 1C: 2-Step Reversible Covalent Inhibition.

Data obtained for 2-step reversible inhibitors with Method I (Figure 8A) is processed with Data Analysis Protocol 1B. The progress curve of time-dependent product formation of each inhibitor concentration (Figure 8B) is fitted to exponential Equation II (shown in Figure 5C). The inhibitor concentration-dependent observed rate for reaction completion k_{obs} reflects the rate from initial velocity v_i (rapid noncovalent equilibrium) to final velocity v_s (slow steadystate equilibrium). Contrary to irreversible inhibition, steady-state velocity v_s is not constrained to inactivation ($v_s > 0$) because the reversible steady-state equilibrium is reached at reaction completion. Maximum rate of reaction completion k_{max} is reached in the presence of saturating inhibitor concentration ([I] >> K_i^{app}), and the covalent association rate constant k_5 is obtained from the span between k_{\min} and k_{\max} (Figure 8C). Interestingly, the Y-intercept k_{\min} is equal to covalent dissociation rate constant k_6 ; therefore, the k_{obs} of uninhibited control (k_{ctrl}) is excluded from the fit. Steady-state inhibition constant K_i^{*app} can be calculated from the fitted values of K_i^{app} , k_5 , and k_6 , but this is not the preferred approach, as a small error in k_6 has huge implications for the calculation of K_i^* . Other methods such as jump dilution assays generate more reliable estimates of k_6 , which is especially important for very potent 2-step reversible covalent inhibitors: relatively small k_6 -values cannot accurately be estimated from the Y-intercept.^{29, 33} Generally, more reliable estimates of the apparent steady-state inhibition constant K_i^{*app} are generated from the dose-response curve of steady-state velocity v_s against inhibitor concentration (Figure 8D).



Figure 8 Data Analysis 1C: Progress curve analysis for 2-step reversible covalent inhibition. Simulated with **KinGen** for 2-step REV inhibitor **B** with 1 pM enzyme and 100 nM substrate **S1**. (**A**) Schematic enzyme dynamics during incubation. (**B**) Time-dependent product formation in absence of inhibitor F^{ctrl} or in presence of inhibitor. The progress curve for each inhibitor concentration is fitted individually to **Equation II** (shown in **Figure 5C**) to obtain the observed rate of inactivation k_{obs} and steady-state velocity v_s . (**C**) Inhibitor concentration-dependent k_{obs} equals k_{max} at saturating inhibitor concentration ($k_{max} = k_s + k_6$) and approaches k_6 in absence of inhibitor ($k_{min} = k_6$). Half-maximum $k_{obs} = k_{min} + \frac{1}{2}(k_{max} - k_{min}) = k_6 + \frac{1}{2}k_5$ is reached when inhibitor concentration equals the apparent inhibition constant K_i^{app} . Steady-state inhibition constant K_i^{app} has to be calculated from the fitted values of k_5 , k_6 and K_i^{app} , thus being very sensitive to errors and (non)linearity in the uninhibited background (illustrated in **Figure 9**). (**D**) Steady-state inhibition constant K_i^{*app} is equal to the IC₅₀ of steady-state velocity v_s .

Contrary to irreversible covalent inhibitors that become more potent with a faster k_{inact} , reversible covalent inhibitors are more potent if they have a longer residence time τ , which is driven by a slow dissociation rate k_6 (Figure 1B).⁴³⁻⁴⁴ It is crucial to have strictly linear product formation in the uninhibited control ($k_{ctrl} = 0$) because it is not possible to perform an algebraic correction for spontaneous enzyme degradation/denaturation (Figure 9A-B). Unfortunately, potent reversible covalent inhibitors are likely to violate this condition. Violation of this assumption ($k_{ctrl} > 0$) can be identified by fitting the uninhibited product formation F^{ctrl} to Equation II (shown in Figure 5C): initial velocity v_i^{ctrl} should not be larger than steady-state v_s^{ctrl} . The consequence of nonlinearity in the uninhibited control is 'contamination' of reaction rate k_{obs} and final velocity v_s (based on the reversible reaction to reach steady-state equilibrium: $v_s > 0$) with the rate of enzyme degradation k_{ctrl} (based on an inactivation reaction: $v_s = 0$). Y-intercept approaching k_{ctrl} instead of k_6 even though the uninhibited control is not included in the fit is an indication that spontaneous enzyme degradation dominates k_{obs} at low inhibitor concentrations (Figure 9C). This 'red flag' should not be ignored, as it will result in over/underestimation of kinetic parameters. To our knowledge, models to perform an



Figure 9 | <u>Data Analysis 1C</u>: Incompatibility with spontaneous loss of enzyme activity (2-step REV). Simulated with **KinDeg** for 2-step REV inhibitor **B** with 1 pM enzyme, 100 nM substrate **S1**, and $k_{ctrl} = k_{degEI} = k_{degEI} = k_{degEI}$. **(A)** Schematic enzyme dynamics during incubation with spontaneous loss of enzyme activity due to degradation/denaturation. **(B)** Time-dependent product formation in absence of inhibitor *F*^{ctrl} is not linear because $k_{ctrl} > 0$. The progress curve for each inhibitor concentration is fitted individually to **Equation II** (shown in **Figure 5C**) to obtain the observed rate of inactivation k_{obs} and steady-state velocity v_s . Simulated for $k_{ctrl} = 0.00003 \text{ s}^{-1}$. **(C)** Inhibitor concentration-dependent k_{obs} is driven by spontaneous enzyme degradation at low inhibitor concentrations, thus lowering the Y-intercept (k_{min} approaches k_{ctrl}). Ignoring the nonlinearity in the uninhibited control k_{ctrl} results in poor fits with underestimation of k_6 , even if k_{ctrl} is slower than k_6 . Simulated for $k_{ctrl} = 0.00003 \text{ s}^{-1}$ (*Icitf*), $k_{ctrl} = 0.00003 \text{ s}^{-1}$ (*indidle*) and $k_{ctrl} = 0.0003 \text{ s}^{-1}$ (*iright*). **(D)** Final velocity v_s has been 'contaminated' by the contribution of irreversible inactivation to the time-dependent inhibition, and approaches $v_s = 0$ at low inhibitor concentrations. Final velocity v_s no longer reflects the steady-state equilibrium: IC₅₀ is larger than K_i^{*app} (underestimation of steady-state potency) unless k_{ctrl} is much smaller than k_6 .

algebraic correction have not been reported. Calculating steady-state inhibition constant K_i^* from final velocity v_s also results in an underestimation of the steady-state potency because the contribution of spontaneous enzyme degradation to final velocity v_s is dominant at low inhibitor concentrations (**Figure 9D**). Underestimation of the steady-state potency of reversible covalent inhibitors that have a relatively slow k_6 is more severe than for the less potent counterpart with a faster k_6 . We were able to find reasonable estimates of K_i^* when the contribution of nonlinearity was significantly smaller than covalent adduct dissociation ($k_{ctrl} << k_6$). Preincubation time-

dependent inhibition (*Method III*) is a more suitable method to analyze 2-step reversible inhibition affected by enzyme instability: it is possible to algebraically correct for enzyme instability in this method (*Data Analysis 3C*).

<u>Data Analysis 1D</u>: Algebraic Correction for Substrate Depletion (2-Step Irreversible Covalent Inhibition).

Scientists from BioKin and Pfizer derived an algebraic model for 2-step irreversible covalent inhibitors to correct for nonlinearity caused by substrate depletion in *Method I* (**Figure 10A**).⁸⁴ Substrate depletion causes a nonlinearity in the uninhibited control because the unbound substrate concentration is no longer constant ($[S]_t < [S]_0$) when a significant fraction of the substrate has been converted into product ($[P]_t > 0.1[S]_0$). The contribution of substrate depletion to the progress curve is directly related to the enzyme activity, as >10% substrate conversion is more likely to be exceeded when enzyme activity is high (**Figure 10B**). Algebraic correction is performed by globally fitting all progress curves in presence of inhibitor (**Figure 10C**) to **Equation III** (shown in **Figure 10D**) with shared values for k_{inact} and K_1^{app} . Substrate depletion should be the only factor contributing to nonlinearity, because the uninhibited control is not included in the global fit. Violation of this (and other) assumption requires data analysis by numerical solving.⁸⁰

The authors demonstrate their algebraic model to correct for substrate depletion with the EGFR inhibitor afatinib in a homogeneous kinase activity assay. A bisubstrate kinase activity



Figure 10 | Data Analysis 1D: Algebraic correction for substrate depletion in progress curve analysis for 2-step irreversible covalent inhibition. Simulated with **KinSubDpl** for 2-step IRREV inhibitor **C** with 100 pM enzyme and 10 nM substrate **S1**. (**A**) Enzyme dynamics. Algebraic correction for substrate depletion is restricted to a Hit-and-Run model (E + S \rightarrow E + P) for product formation. (**B**) Substrate depletion ([P]_t > 0.1[S]₀) results in a decrease of product formation in the uninhibited control (*solid line*) compared to product formation, assuming substrate conversion does not affect product formation rates (*dashed line*, simulated with **KinGen**). The contribution of substrate depletion to nonlinearity increases with higher enzyme activity (less inhibition). (**C**) Time-dependent product formation in the absence of inhibitor *F*^{ctrl} or in presence of inhibitor with time-dependent loss of enzyme activity due to substrate depletion. Inhibitor-treated progress curves are globally fitted to **Equation III** with shared values for k_{inact} and K_{I}^{app} . (**D**) **Equation III**. Algebraic model to correct for substrate depletion at low substrate concentrations.⁸⁰ F_0 = Y-intercept = background signal at reaction initiation (in AU). r_p = product coefficient for detected signal *F* per formed product [P] (in AU/M). k_{sub} = reaction rate constant for Hit-and-Run model of enzymatic product formation E + S \rightarrow E + P (in M⁻¹s⁻¹).

assay is different from our simulations with a single substrate, but this algebraic model can be applied in both systems: product formation in single-substrate as well as bisubstrate reactions can be simplified to a Hit-and-Run model (E + S \rightarrow E + P) with rate constant $k_{sub} = k_{cat}/K_M$ as long as the substrate concentration is far below its K_M ([S] < 0.1 K_M) (**Figure 10A**). The accuracy of k_{inact} and K_1 was very good with low substrate concentrations ([S] \leq 0.01 K_M). A slightly higher substrate concentration ([S] \geq 0.1 K_M) resulted in underestimation of k_{inact} and overestimation of K₁, but a good estimation of overall second order inactivation rate constant k_{inact}/K_1 . Importantly, a calibration/titration should be performed prior to data analysis to determine product coefficient r_p (in AU/M) that transforms the detected signal F_t into product concentration [P]_t.⁶⁸⁻⁶⁹

METHOD II: Incubation Time-dependent Potency IC₅₀(t)

The observed potency of irreversible inhibitors increases with longer (pre)incubation time, as more enzyme is irreversibly bound. In this method, sometimes dubbed 'the Krippendorff method', the time-dependence of potency $IC_{50}(t)$ is utilized to directly find the relevant kinetic parameters for 2-step irreversible covalent inhibition. Contrary to progress curve analysis (*Method I*), this method is compatible with quenched/stopped assays that require a development/separation/quenching step before read-out, as continuous measurement of product formation is not required (but optional).

The incubation time-dependent potency $IC_{50}(t)$ reflects the inhibitor concentration resulting in a 50% decrease of cumulative product formed F_t during incubation compared to cumulative product formed in the uninhibited control F^{ctrl} . Enzymatic product formation is initiated by enzyme addition without preincubation of enzyme and inhibitor (**Figure 11A**). Fractional cumulative product formation F_t/F^{ctrl} decreases with longer incubation times (**Figure 11B**). Importantly, this does not reflect the current enzyme activity because read-out F_t reflects that



Figure 11 | Method II: Incubation time-dependent potency $IC_{50}(t)$. Simulated with **KinGen** for 50 nM 2-step IRREV inhibitor **C** with 1 pM enzyme and 100 nM substrate **S1**. (**A**) The reaction between enzyme, inhibitor, and substrate is initiated by addition of enzyme. Enzyme inhibition increases with time-dependent formation of covalent EI* until reaching reaction completion. (**B**) Read-out of cumulative product formation (reflected in signal F_t) in presence of 2-step irreversible covalent inhibitor relative to product formed the uninhibited control (F^{ctrl}) decreases upon longer incubation. (**C**) Cumulative product F_t (blue line) is 'contaminated' with product formed prior to reaching 100% inhibition even if the current enzyme activity (green line) is fully inhibited.

the cumulative product formed during incubation will be 'contaminated' with product that was formed before full inhibition was reached. Consequently, incubation time-dependent potency $IC_{50}(t)$ calculated from the fractional product formation F_t/F^{ctrl} against inhibitor concentration will increase with longer incubation times (for slow-binding inhibitors), but will underestimate the potency compared to the values based on the current enzyme activity $[E+ES]_t/[E]_0$ (**Figure 11C**). $IC_{50}(t)$ does not approach K_i^{*app} (2-step reversible inhibition) or $\frac{1}{2}[E]_0$ (irreversible inhibition) at infinite incubation times. An implicit algebraic model based on multipoint $IC_{50}(t)$ values has been derived for 2-step irreversible covalent inhibitors (*Data Analysis 2*).⁸¹ Additionally, a two-point $IC_{50}(t)$ method for 2-step irreversible covalent inhibitors have been reported in a preprint.⁸⁵ To our knowledge, algebraic methods to calculate K_i^{*app} for 2-step reversible covalent inhibitors from (endpoint) $IC_{50}(t)$ values have not been reported.

Data Analysis 2: 2-Step Irreversible Covalent Inhibition.

Krippendorff and co-workers report an algebraic model to calculate k_{inact} and K_1 of irreversible covalent inhibitors from the incubation time-dependent potency $IC_{50}(t)$ after multiple incubation times (**Figure 12A**).⁸¹ Detection of cumulative product formation after several incubation times is compatible with continuous assays, but more importantly also with



Figure 12 | <u>Data Analysis 2</u>: Incubation time-dependent potency $IC_{50}(t)$ for 2-step irreversible covalent inhibition. Simulated with **KinGen** for 2-step IRREV inhibitor **C** with 1 pM enzyme and 100 nM substrate **S1**. (**A**) Schematic enzyme dynamics during incubation. (**B**) Time-dependent cumulative product formation in absence of inhibitor F^{ctrl} or in presence of inhibitor F_t is detected with longer measurement intervals compatible with quenched assays. (**C**) Incubation time-dependent potency $IC_{50}(t)$ reflects the inhibitor concentration that reduces cumulative product formation during incubation by 50% compared to the uninhibited control. (**D**) Incubation time-dependent potency $IC_{50}(t)$ against incubation time is fitted to **Equation IV**. $IC_{50}(0)$ approaches apparent noncovalent inhibition constant K_1^{app} but $IC_{50}(0)$ is never included in the fit because product formation does not start until initiation of the incubation ($F_0 = F^{\text{ctrl}} = 0$). (**E**) Implicit algebraic **Equation IV** derived by Krippendorff *et al.*⁸¹

stopped/quenched assays that require a development step to visualize product formation (**Figure 12B**). Incubation time-dependent potency $IC_{50}(t)$ is calculated for each incubation time from fractional product formation F_t/F^{ctrl} (**Figure 12C**) and plotted against the incubation time (**Figure 12D**). Finally, the authors derived *implicit* algebraic **Equation IV** (shown in **Figure 12E**) to calculate k_{inact} and K_1 from the incubation time-dependent potency $IC_{50}(t)$. This method is restricted to substrate-competitive irreversible (multi-step) covalent inhibitors: k_{inact} and K_1 do not have a biological meaning for reversible inhibitors or for 1-step covalent inhibitors. This method requires software (e.g. GraphPad Prism) that allows fitting a model defined by an implicit equation (where Y appears on both sides of the equal sign).

Product formation in the uninhibited control should be strictly linear ($k_{ctrl} = 0$): normalization of cumulative product formation (F_t/F^{ctrl}) does not correct for spontaneous loss of enzyme activity or substrate depletion. It is relatively easy to miss violations of this assumption because nonlinearity in the uninhibited control ($k_{ctrl} > 0$) is not evident from visual inspection of the dose-response curves. Violation of this assumption results in a significant underestimation of k_{inact} and K_1 values, also when nonlinearity is relatively small ($k_{ctrl} << k_{inact}$) (**Figure 13A**). Another



Figure 13 | <u>Data Analysis 2</u>: Experimental/Assay Restrictions. **(A)** Enzyme degradation/denaturation simulated with **KinDeg** for 2-step IRREV inhibitor **C** with 1 pM enzyme, 100 nM substrate **S1**, and $k_{ctrl} = k_{degEI} = k_{de$

important assumption is that the onset of product formation and enzyme inhibition occur simultaneously: inhibition and product formation are both initiated by addition of enzyme, without preincubation of enzyme and inhibitor prior to substrate addition. Unfortunately, numerous publications refer to preincubation of enzyme and inhibitor as 'incubation', resulting in the understandable but incorrect fitting of preincubation time-dependent potency IC₅₀(t') to the Krippendorff model (Figure 13B).⁸⁵ Preincubation time-dependent potency $IC_{50}(t')$ is calculated from product formation velocity $v_{t'}$, reflecting the enzyme activity after preincubation rather than cumulative product formation F_t/F^{ctrl} . Enzyme activity $v_{t'}$ is not 'contaminated' by product formed prior to read-out because product formation is initiated after preincubation. Furthermore, substrate does not compete with inhibitor for enzyme binding during preincubation. Fitting $IC_{50}(t')$ values to the Krippendorff model resulted in an overestimation of k_{inact} and an overestimation of the overall inactivation potency $k_{\text{inact}}/K_{\text{I}}$ (Figure 13B). This method is not compatible with ligand binding competition assays (such as the LanthaScreen kinase binding assay) where inhibitor binding competes with ligand (tracer) binding to form enzyme-ligand complex EL as the detectable product (Figure 13C). The enzyme-ligand equilibrium after incubation in presence of inhibitor reflects the current inhibitor competition and is unaffected by binding equilibria prior to read-out (not cumulative). Furthermore, unbound enzyme is not released after formation of product EL, thereby limiting the product formation to a single turnover per enzyme. Fitting $IC_{50}(t)$ values obtained in ligand binding assays to the Krippendorff model result in overestimation of k_{inact} and/or unstable parameters (Figure 13C).

METHOD III: Preincubation Time-Dependent Inhibition Without Dilution

Preincubation of enzyme and inhibitor prior to initiation of product formation by addition of substrate is an established method for kinetic analysis of slow-binding (ir)reversible (covalent) inhibitors.^{41, 82} In the benchmark protocol by Ito and co-workers, a low substrate concentration $([S] \leq K_M)$ is added in a relatively small volume $(V_{sub} \leq V_{t'})$ to keep the noncovalent enzymeinhibitor $E + I \leftrightarrow EI$ equilibrium intact. However, (partial) disruption of the noncovalent equilibrium does not affect the accuracy of preincubation experiments for irreversible inhibition, as is illustrated by Method IV. Product formation is inhibited by formation of EI and EI* during preincubation in absence of competing substrate (Figure 14A). Preincubation timedependent product formation velocity v_t reflects the total inhibition by noncovalent as well as covalent inhibitor binding, and is calculated after a relatively short incubation time ($t \ll t'$) to minimize additional (time-dependent) inhibition of enzyme activity during incubation resultant from enzyme-inhibitor complex/adduct formation during incubation (Figure 14B). Enzyme activity after preincubation in the presence of time-dependent inhibitors v_{t} decreases exponentially from rapid (initial) equilibrium K_i^{app} (Y-intercept: v_i) to reach a plateau at reaction completion (t' > 5t¹/₂), corresponding to the steady-state equilibrium ($v_s > 0$) or inactivation $(v_s = 0)$ (Figure 14C). Observed rate of reaction completion k_{obs} (from enzyme activity without preincubation v_i to final enzyme activity v_s) is obtained by fitting to bounded exponential decay **Equation v** (shown in **Figure 14D**). Importantly, this equation fits enzyme activity v_t rather than directly fitting product signal F.



Figure 14 | Method III: Preincubation time-dependent inhibition without dilution. Simulated with KinGen for 1 pM enzyme and 100 nM substrate S1. (A) Enzyme is preincubated with inhibitor to form noncovalent complex EI and covalent adduct EI* in absence of competing substrate, followed by addition of substrate. Addition of a low substrate concentration in a small volume to avoid disruption of the noncovalent E + I ↔ EI equilibrium. Simulated for 50 nM 2-step IRREV inhibitor C with preincubation t' = 1800 s. (B) Preincubation time-dependent enzyme activity $v_{t'}$ is obtained from the slope of (initial) linear product formation velocity with a short incubation time t relative to preincubation t' to minimize ΔEI^* formation after substrate addition. This measurement is performed separately for each preincubation time, thus requiring more material than incubation time-dependent inhibition protocols with continuous product read-out. Simulated for 50 nM 2-step IRREV inhibitor C with preincubation t' = 0-1800 s. (C) Enzyme activity v_{μ} of time-dependent inhibitors decreases exponentially from rapid (initial) equilibrium K_i^{app} (Y-intercept = enzyme activity without preincubation v_i) to reaching reaction completion (t' > 5t%): inactivation for irreversible inhibitors ($v_s = 0$) and steady-state equilibrium K_i^{sapp} for reversible inhibitors ($v_s > 0$). Enzyme activity without preincubation v, equals the uninhibited enzyme activity v^{ctrl} for 1-step inhibitors and for 2-step inhibitors at non-saturating concentration ([I] $<< K_i^{app}$). Simulated for 50 nM 1-step REV inhibitor A, 2-step REV inhibitor B, 1-step IRREV inhibitor D, and 2-step IRREV inhibitor C. (D) General bounded exponential decay Equation v to fit preincubation time-dependent enzyme activity $v_{t'}$ against preincubation time t'. Parameters are constrained depending on the inhibitor binding mode. $v_s = 0$ for irreversible inhibition: inactivation at reaction completion. $v_i = v^{ctrl}$ for 1-step inhibition: noncovalent complex is not significant at non-saturating inhibitor concentrations. v_t = preincubation time-dependent enzyme activity. v_i = enzyme activity based without preincubation. v_s = enzyme activity after preincubation based on reaching reaction completion (t' > 5t%). t' = preincubation time of enzyme and inhibitor before substrate addition. k_{nbc} = observed rate of time-dependent inhibition from initial v_i to final v_s .

Preincubation assays are generally disfavored because their experimental execution requires more material and is more laborious than substrate competition assays with continuous read-out (*Method I* and *II*). Here, substrate has to be added after the indicated preincubation time, thus requiring multiple individual measurements for each inhibitor concentration. However, preincubation experiments are still favored when reaction completion is too slow for detection during the normal time course of a substrate competition assay ($t \ll t_2$ in *Method I*): substrate competition reduces the (covalent) reaction rate and inhibitor solubility limits the maximum inhibitor concentration. Instead, preincubation is performed in the absence of competing substrate, thus reaching the maximum reaction rate at a low inhibitor concentration. Therefore, preincubation experiments are frequently conducted for compounds that display 1-step irreversible inhibition behavior because they have a poor noncovalent affinity, such as covalent fragments.⁸ Additionally, preincubation times can exceed the maximum incubation time of progress curve analysis, which is limited by linear product formation ($[P]_t > 0.1[S]_0$), as the onset of product formation does not start until preincubation is completed.

This method is less suitable for enzymatic assays with a relatively slow uninhibited product formation velocity v^{ctrl} , as assay sensitivity might be insufficient to produce enough product signal F_t during a short incubation time. Reaction completion (t' > 5t¹/₂) and/or full inhibition $(v_{t'} = 0)$ should not be reached before the first (shortest) preincubation time because it will be impossible to detect time-dependent changes in enzyme activity. This can be resolved by increasing the measurement interval (shorter dt'), reduction of the inhibitor concentration, or selection of a different experimental protocol. This method is compatible with 2-step irreversible inhibition (Data Analysis 3A) and 1-step irreversible inhibition (Data Analysis 3B), but also with (2-step) reversible inhibition (Data Analysis 3C). Algebraic analysis by linear regression to obtain k_{obs} from the (initial) linear slope of LN(enzyme activity) against preincubation time t' is still frequently reported. This is probably because linear regression is part of benchmark protocols for kinetic analysis of preincubation time-dependent enzyme inactivation.⁸²⁻⁸³ It is important to note that these benchmark protocols were published before dedicated data analysis software for nonlinear regression was available.⁶³ Visualization of this 'linear' relationship is possible by plotting the enzyme activity against preincubation time t' on a semilog scale (illustrated in Figure S2).

Data Analysis 3A: 2-Step Irreversible Covalent Inhibition.

Data obtained for 2-step irreversible inhibitors (Figure 15A) is processed with Data Analysis Protocol 3, followed by Data Analysis Protocol 3Ai or 3Aii. Time-dependent product formation is fitted to a straight line for each inhibitor concentration to obtain the enzyme activity after preincubation v_{t} from the linear (initial) slope (Figure 15B, *left*). It is important that the incubation time be relatively short ($t < 0.1t\frac{1}{2}$) to minimize artifacts caused by significant formation of covalent adduct EI* after substrate addition (Δ EI*) because v_r should reflect the enzyme activity at the end of preincubation. As a rule of thumb, incubation time t should be much shorter than the shortest preincubation time t'. A short incubation time may result in insufficient product formation for a robust signal, which can be resolved by increasing the incubation time and obtaining enzyme activity v_{r} from the initial velocity of the exponential association progress curve, provided that the assay is compatible with progress curve analysis (continuous read-out) (Figure 15B, right). Enzyme activity after preincubation v_t is fitted to bounded exponential decay Equation v (shown in Figure 14D) (constraining $v_s = 0$) for each inhibitor concentration to obtain the observed rate of reaction completion k_{obs} from enzyme activity without preincubation (Y-intercept at v_i) to reaching the final enzyme inactivation (plateau at $v_s = 0$) (Figure 15C). Enzyme activity without preincubation in presence of inhibitor v_i is lower than the uninhibited enzyme activity v^{ctrl} for 2-step (ir)reversible inhibitors, because v_i reflects the rapid noncovalent equilibrium (K_i^{app}) after substrate addition.⁴¹ The plot of inhibitor concentration-dependent k_{obs} reaches maximum rate of inactivation k_{inact} in presence of saturating inhibitor concentration ([I] >> K₁) with the Y-intercept at $k_{ctrl} = 0$ when uninhibited enzyme activity v^{ctrl} is independent of preincubation time (**Figure 15D**). Inhibitor concentrations should correspond with the inhibitor concentration <u>during preincubation</u> (rather than after substrate addition). Correction of inactivation constant K₁ for substrate competition is not necessary because preincubation is performed in absence of substrate. The rapid noncovalent $E + I \leftrightarrow EI$ equilibrium does not significantly contribute to inhibition at non-saturating inhibitor concentrations ([I] << K_i^{app}), resulting in 1-step binding behavior (illustrated in **Figure 3F**). This will be apparent from the observation that initial velocity v_i is independent of inhibitor concentration ($v_i = v^{ctrl}$) along with a linear plot of k_{obs} against [I]. This is resolved either by increasing the inhibitor concentration or performing *Data Analysis 3B*. Increasing the substrate concentration can resolve issues with assay sensitivity associated with short incubation times, as this will result in a higher product signal. However, substrate concentration in a relatively large volume ($V_{sub} > 0.1V_t$) and/or addition of a competitive substrate concentration



Figure 15 | Data Analysis 3A: Preincubation time-dependent inhibition without dilution for 2-step irreversible covalent inhibition. Simulated with **KinGen** for 2-step IRREV inhibitor **C** with 1 pM enzyme and 100 nM substrate **S1.** (A) Schematic enzyme dynamics during preincubation in absence of substrate and during incubation after substrate addition. (**B**) Time-dependent product formation after preincubation in absence of inhibitor F^{ctrl} or in presence of inhibitor (t' = 1800 s). *Left:* Enzyme activity after preincubation $v_{t'}$ is obtained from the linear slope if the incubation time is relatively short (t < t'): gray area is excluded from the fit. *Right:* Enzyme activity after preincubation $v_{t'}$ is obtained from the initial velocity of the exponential association progress curve of each inhibitor concentration. (**C**) Preincubation time-dependent enzyme activity $v_{t'}$ is fitted to **Equation v** (shown in **Figure 14D**) (constraining $v_s = 0$) for each inhibitor concentration to obtain observed rates of inactivation k_{obs} . Alternatively, $v_{t'}$ can be normalized to a fraction of the uninhibited enzyme activity v^{trl} (**D**) Inhibitor concentration-dependent k_{obs} reaches k_{inact} at saturating inhibitor concentration ($k_{max} = k_{inact}$). Half-maximum $k_{obs} = \frac{y_s}{k_{inact}}$ is reached when inhibitor concentration constant K_1 . $v_{t'}$ reflects the enzyme activity after preincubation in absence of competing substrate.

 $([S] > 0.1K_M)$ causes (partial) disruption of the reversible equilibrium, although this does not affect the accuracy of k_{obs} for irreversible inhibitors. In fact, disruption of the noncovalent complex can be employed to detect covalent adduct formation of 2-step irreversible inhibitors that exhibit tight-binding behavior resulting from very potent noncovalent inhibition,⁷⁵⁻⁷⁶ as will be discussed in *Method IV*.

Uninhibited enzyme activity v^{ctrl} decreases when preincubation is long enough for significant spontaneous enzyme degradation (t' >> 0.1t¹/₂) (**Figure 16A-B**). A simple algebraic correction for spontaneous enzyme degradation results in good estimates for k_{inact} and K₁ if all enzyme species have the same first order enzymatic degradation rate ($k_{\text{degE}} = k_{\text{degEI}} |$ (**Figure 16C**). Alternatively, normalizing the enzyme activity $v_{t'}$ to uninhibited enzyme activity $v_{t'}^{\text{ctrl}}$ at each preincubation time corrects for enzyme degradation (**Figure 16D**), and k_{obs} obtained from normalized enzyme activity $v_{t'}/v^{\text{ctrl}}$ results in good estimates of k_{inact} and K₁ without further correction (**Figure 16E**).



Figure 16 | Data Analysis 3A: Corrections for spontaneous loss of enzyme activity (2-step IRREV). Simulated with **KinDeg** for 2-step IRREV inhibitor **C** with 1 pM enzyme, 100 nM substrate **S1**, and $k_{degE} = k_{degEI} = 0.0003 \text{ s}^{-1}$. (A) Schematic enzyme dynamics during preincubation in absence of substrate and during incubation after substrate addition with spontaneous enzyme degradation/denaturation. (B) Uninhibited enzyme activity after preincubation v_t^{ctrl} is not linear. Preincubation time-dependent enzyme activity v_t is fitted to **Equation V** (shown in **Figure 14D**) (constraining $v_s = 0$) for each inhibitor concentration to obtain observed rates of inactivation k_{obs} , as well as fitting uninhibited activity v_t^{ctrl} to obtain the rate of nonlinearity k_{ctrl} . (C) Inhibitor concentration-dependent k_{obs} with spontaneous enzyme degradation increases with k_{ctrl} but the span from $k_{min} (= k_{ctrl})$ to $k_{max} (= k_{inact} + k_{ctrl})$ still equals k_{inact} . Fit with algebraic correction for nonlinearity (*black line*, $k_{ctrl} > 0$). Ignoring the nonlinearity (*gray line*, constrain $k_{ctrl} = 0$) results in underestimation of K_{I} (overestimation of potency) and overestimation of k_{inact} . (D) Normalized enzyme activity v_t^{ctrl} is fitted to **Equation V** (shown in **Figure 14D**) (constraining $v_s = 0$) for each inhibitor concentration by fitting normalized enzyme activity v_t/v^{ctrl} and does not require further corrections.
Data Analysis 3B: 1-Step Irreversible Covalent Inhibition.

Data obtained for 1-step irreversible inhibitors (**Figure 17A**) is processed with *Data Analysis Protocol 3*, followed by *Data Analysis Protocol 3Bi* or *3Bii*. Time-dependent product formation is fitted to a straight line for each inhibitor concentration to obtain the enzyme activity after preincubation $v_{t'}$ from the linear slope (**Figure 17B**, *left*). Incubation must be short enough to minimize formation of covalent adduct EI* after substrate addition ($t << t^{1/2}$); otherwise k_{chem} will be overestimated. Similar to *Data Analysis 3A*, preincubation-dependent enzyme activity $v_{t'}$ can also be obtained from the initial velocity of the exponential association progress curve, provided that the read-out is continuous (**Figure 17B**, *right*). Enzyme activity after preincubation $v_{t'}$ (**Figure 17C**) is fitted to bounded exponential decay **Equation v** (shown in **Figure 14D**) to obtain observed rate of reaction completion k_{obs} from uninhibited enzyme activity without preincubation (Y-intercept at $v_i = v^{ctrl}$) to reaching the final enzyme inactivation (constraining $v_s = 0$). Inhibited enzyme activity without preincubation is equal to uninhibited enzyme activity ($v_i = v^{ctrl}$), as rapid noncovalent inhibitor binding does not contribute to



Figure 17 | Data Analysis 3B: Preincubation time-dependent inhibition without dilution for 1-step irreversible covalent inhibition. Simulated with **KinGen** for 1-step IRREV inhibitor **D** with 1 pM enzyme and 100 nM substrate **S1.** (A) Schematic enzyme dynamics during preincubation in absence of substrate and during incubation after substrate addition. (B) Time-dependent product formation after preincubation in absence of inhibitor F^{ctrl} or in presence of inhibitor (t' = 1800 s). *Left:* Enzyme activity after preincubation v_t is obtained from the linear slope if the incubation time is relatively short (t < t'): gray area is excluded from the fit. *Right:* Enzyme activity after preincubation v_t is obtained from the linear slope if the incubation time is relatively short (t < t'): gray area is excluded from the fit. *Right:* Enzyme activity after preincubation v_t is obtained from the linear slope if the incubation time is relatively short (t < t'): gray area is excluded from the fit. *Right:* Enzyme activity after preincubation v_t is obtained from the initial velocity of the exponential association progress curve of each inhibitor concentration. (**C**) Preincubation time-dependent enzyme activity v_t is fitted to **Equation v** (shown in **Figure 14D**) (constraining $v_s = 0$) for each inhibitor concentration to obtain observed rates of inactivation k_{obs} . $v_i = v^{ctrl}$ for 1-step irreversible inhibitors and 2-step irreversible inhibitors at non-saturating concentrations ([I] << K_i^{app}). Alternatively, v_t can be normalized to a fraction of the uninhibited enzyme activity v^{ctrl} . (**D**) Inhibitor concentration-dependent k_{obs} increases linearly with inhibitor concentration, with k_{chem} as the slope. v_t reflects the enzyme activity after preincubation in absence of competing substrate.

enzyme inhibition by 1-step irreversible inhibitors. The slope of the linear plot of k_{obs} against inhibitor concentration <u>during preincubation</u> is equal to k_{chem} (Figure 17D), which should not be corrected for substrate competition as preincubation is performed in absence of competing substrate. Substrate addition in a relatively large volume ($V_{sub} > 0.1V_t$) and/or addition of a competitive substrate concentration ([S] > $0.1K_M$) does not significantly affect the accuracy of k_{obs} because 1-step irreversible inhibition does not involve a rapid noncovalent equilibrium that can be disrupted (also see *Method IV*). Increasing the substrate concentration can resolve issues with assay sensitivity: higher substrate concentration results in a higher product concentration after the same incubation time ($v^{ctrl} = V_{max}[S]/([S]+K_M)$), which in turn will result in a better signal to noise ratio.

Uninhibited enzyme activity v^{ctrl} decreases with longer preincubation due to spontaneous enzyme degradation (Figure 18A-B). This especially affects assays where preincubation is long



Figure 18 | Data Analysis 3B: Corrections for spontaneous loss of enzyme activity (1-step IRREV). Simulated with **KinDeg** for 1-step IRREV inhibitor **D** with 1 pM enzyme, 100 nM substrate **S1**, and $k_{degE} = k_{degES} = k_{degEI} = 0.0003 s^{-1}$. (**A**) Schematic enzyme dynamics during preincubation in absence of substrate and during incubation after substrate addition with spontaneous enzyme degradation/denaturation. (**B**) Uninhibited enzyme activity after preincubation v_t^{ctrl} is not linear: $k_{ctrl} > 0$. Preincubation time-dependent enzyme activity v_t is fitted to **Equation V** (shown in **Figure 14D**) (constraining $v_s = 0$ and shared value for $v_i =$ uninhibited enzyme activity without preincubation v_0^{ctrl}) for each inhibitor concentration to obtain observed rates of inactivation k_{obs} , as well as fitting uninhibited activity v_t^{ctrl} to obtain the rate of nonlinearity k_{ctrl} . (**C**) Inhibitor concentration-dependent k_{obs} with spontaneous enzyme degradation/denaturation increases by k_{ctrl} . Fit with algebraic correction for nonlinearity (*black line*, $k_{ctrl} > 0$) or ignoring nonlinearity (gray line, constrain $k_{ctrl} = 0$). Ignoring the nonlinearity v_t/v^{ctrl} is fitted to **Equation V** (shown in **Figure 14D**) (constraining $v_s = 0$ and Y-intercept = $v_i/v_0^{ctrl} = 1$) for each inhibitor concentration increases by k_{ctrl} . Fit with algebraic correction for nonlinearity (*black line*, $k_{ctrl} > 0$) or ignoring nonlinearity (gray line, constrain $k_{ctrl} = 0$). Ignoring the nonlinearity (assuming Y-intercept = 0) results in overestimation of k_{chem} (steeper slope). (**D**) Normalized enzyme activity v_t/v^{ctrl} is fitted to **Equation V** (shown in **Figure 14D**) (constraining $v_s = 0$ and Y-intercept = $v_i/v_0^{ctrl} = 1$) for each inhibitor concentration to obtain corrected observed rates of inactivation k_{obs} . (**E**) Inhibitor concentration-dependent k_{obs} has been corrected for enzyme degradation/denaturation by fitting normalized enz

enough for significant enzyme degradation (t' > 0.1t½). Algebraic correction for spontaneous enzyme degradation ($k_{degE} = k_{degES}$) in the secondary k_{obs} plot is relatively simple (**Figure 18C**). Alternatively, correction for enzyme degradation is performed by normalizing enzyme activity v_t to uninhibited enzyme activity v_t^{ctrl} at each preincubation time (**Figure 18D-E**). Stabilization of enzyme upon inhibitor binding ($k_{degEI*} < k_{degE}$) does not affect k_{obs} , as EI* formation is already irreversible thus removing the species from the available pool of catalytic enzyme.

Data Analysis 3C: 2-step Reversible Covalent Inhibition.

Data obtained for 2-step reversible inhibitors (Figure 19A) is processed with *Data Analysis Protocol 3*, followed by *Data Analysis Protocol 3C*. Time-dependent product formation is



Figure 19 | Data Analysis 3C: Preincubation time-dependent inhibition without dilution for 2-step reversible covalent inhibition. Simulated with KinGen for 2-step REV inhibitor B with 1 pM enzyme and 100 nM substrate S1. (A) Schematic enzyme dynamics during preincubation in absence of substrate and during incubation after substrate addition. (B) Time-dependent product formation after preincubation in absence of inhibitor F^{ctrl} or in presence of inhibitor (t' = 1800 s). Enzyme activity after preincubation v, is obtained from the linear slope if the incubation time is relatively short ($t \ll t'$): gray area is excluded from the fit. Alternatively, enzyme activity after preincubation v_{r} is obtained from the initial velocity of the exponential association progress curve of each inhibitor concentration. (C) Preincubation time-dependent enzyme activity $v_{t^{\prime}}$ is fitted to Equation V (shown in Figure 14D) for each inhibitor concentration to obtain observed rates of inactivation k_{obs} and steady-state velocity v_s (plateau > 0). Alternatively, v_t can be normalized to a fraction of the uninhibited enzyme activity v^{ctrl} . (D) Inhibitor concentration-dependent k_{obs} equals k_{max} at saturating inhibitor concentration ($k_{max} = k_s + k_6$) and approaches k_6 in absence of inhibitor ($k_{min} = k_6$). Half-maximum $k_{obs} = k_{min} + \frac{1}{2}(k_{max} - k_{min}) = k_6 + \frac{1}{2}k_5$ is reached when inhibitor concentration equals the inhibition constant K_i. Steady-state inhibition constant K_i^{*} calculated from the fitted values of k_{s} , k_{b} , and K_{i} is thus very sensitive to errors and (non)linearity in the uninhibited background (illustrated in Figure 9C). No correction: v_t reflects the enzyme activity after preincubation in absence of competing substrate. (E) Steady-state inhibition constant K_i^* corresponds with the IC_{50} of steady-state velocity v_{s} obtained by fitting the dose-response curve to the Hill equation.³³ No correction: v_{t} reflects the enzyme activity after preincubation in absence of competing substrate.

fitted to a straight line for each inhibitor concentration to obtain the enzyme activity after preincubation v_t from the linear slope (**Figure 19B**). Again, it is important that the incubation time be much shorter than the shortest preincubation time t' ($t \ll t'$), but enzyme activity $v_{t'}$ can also be calculated from the initial velocity of the exponential association progress curve, provided that the assay is compatible with progress curve analysis (continuous read-out). Enzyme activity after preincubation $v_{t'}$ is fitted to bounded exponential decay **Equation v** (shown in **Figure 14D**) for each inhibitor concentration to obtain observed rate of reaction completion k_{obs} from rapid noncovalent equilibrium (Y-intercept at $v_i < v^{ctrl}$) to slowly reaching steady-state equilibrium (plateau at $v_s > 0$) (**Figure 19C**). Enzyme activity without preincubation in presence of inhibitor v_i is lower than the uninhibited enzyme activity v^{ctrl} for 2-step (ir)reversible inhibitors because v_i reflects the rapid noncovalent equilibrium (K_i^{app}) after substrate addition.⁴¹ Contrary to irreversible inhibition, the plateau ($v_s > 0$) does not approximate enzyme inactivation but reaches the steady-state equilibrium (K_i^*) instead.

Steady-state inhibition constant K_i^* can be calculated from the fitted values of K_i , k_5 and k_6 (Figure 19D), but this is not the preferred approach as a small error in k_6 has huge implications for the calculation of K_i* (as illustrated in Figure 9C). Generally, more reliable estimates of the steadystate inhibition constant K_i^{*} are generated from dose-response curves of steady-state velocity $v_{\rm s}$ against inhibitor concentration <u>during preincubation</u> (Figure 19E). Steady-state inhibition constant K_i^* reflects the reversible $E + I \leftrightarrow EI + EI^*$ equilibrium that can be disrupted by substrate addition in a relatively large volume ($V_{sub} > 0.1V_{t'}$) and/or addition of a competitive substrate concentration ([S] > $0.1K_{\rm M}$). Simulations with high substrate concentration ([S] = $10K_{\rm M}$) show that the IC₅₀ of the dose-response curve for steady-state velocity v_s was slightly higher than steady-state inhibition constant K_i^* , but still significantly lower than K_i^{*app} , as covalent dissociation will not be significant provided the incubation time is significantly shorter than the dissociation half-life (t << t $\frac{1}{2}$ _{diss}). Altogether, fitting exponential association rather than increasing the substrate concentration is the desired solution to resolve issues with assay sensitivity associated with short incubation times. Alternatively, reasonable estimates of the steady-state inhibition constant Ki* were obtained from the endpoint preincubation timedependent potency $IC_{50}(t')$ with minimal substrate competition ([S] << K_M) and preincubation times exceeding the required time to reach reaction completion at all inhibitor concentrations $(t' > 5t^{1/2}).$

As mentioned before, spontaneous loss of enzyme activity (**Figure 20A**) due to first order degradation and/or denaturation of enzyme species ($k_{degE} = k_{degES} = k_{degEI}$) results in a preincubation time-dependent decrease of uninhibited enzyme activity v^{ctrl} (**Figure 20B**). The biggest advantage of *Method III* (*Data Analysis 3C*) over *Method I* (*Data Analysis 1C*) is that it is possible to perform an algebraic correction for the enzyme instability in kinetic analysis of 2-step reversible covalent inhibitors with *Data Analysis 3C*. Enzyme activity v_t is normalized to uninhibited enzyme activity v_t^{ctrl} at each preincubation time (**Figure 20C**), and the normalized enzyme activity after preincubation v_t/v^{ctrl} is fitted to bounded exponential decay **Equation v** (shown in **Figure 14D**) for each inhibitor concentration to obtain observed rate of reaction completion k_{obs} and steady-state velocity v_s . Kinetic analysis of k_{obs} (**Figure 20D**) and steady-state velocity v_s (**Figure 20E**) against inhibitor concentration during preincubation result in



Figure 20 | Data Analysis 3C: Corrections for spontaneous loss of enzyme activity (2-step REV). Simulated with **KinDeg** for 2-step REV inhibitor **B** with 1 pM enzyme, 100 nM substrate **S1**, and $k_{degE} = k_{degEI} = 0.0003 \text{ s}^{-1}$. (A) Schematic enzyme dynamics during preincubation in absence of substrate and during incubation after substrate addition with spontaneous enzyme degradation. (B) Uninhibited enzyme activity after preincubation v_t^{ctrl} is not linear. Fitting preincubation time-dependent enzyme activity v_t to **Equation v** (shown in **Figure 14D**) for each inhibitor concentration gives observed rates of inactivation k_{obs} as well as the rate of nonlinearity k_{ctrl} for uninhibited enzyme degradation if enzyme activity is not normalized. (C) Enzyme activity v_t is normalized to the uninhibited enzyme activity v_t^{ctrl} after each preincubation time before fitting to **Equation v** (shown in **Figure 14D**). (D) Inhibitor concentration-dependent k_{obs} has been corrected for enzyme degradation/denaturation by fitting normalized enzyme activity v_t^{ctrl} and does not require further corrections (even if $k_{ctrl} > k_6$). (E) Steady-state velocity v_s has been corrected for enzyme degradation/denaturation by fitting normalized enzyme activity v_t , v_{trl}^{ctrl} and does not require further corrections (even if $k_{ctrl} > k_6$). (E) Steady-state velocity v_s has been corrected for enzyme degradation by fitting normalized enzyme activity v_t , v_{trl}^{ctrl} and does not require further corrections (even if $k_{ctrl} > k_6$). Final velocity v_s obtained from uncorrected v_t is 'contaminated' by the contribution of irreversible inactivation to the time-dependent inhibition, and does not result in accurate estimates of steady-state inhibition constant K_i^* (illustrated in **Figure 9D**).

good estimates of the kinetic parameters without further correction, even when k_{ctrl} is faster than the covalent dissociation rate k_6 ($k_{ctrl} > k_6$). We strongly advise that enzyme activity be normalized prior to analysis of reversible covalent inhibition even when k_{ctrl} is not directly obvious from the uninhibited control v_t^{ctrl} .

METHOD IV: Preincubation Time-Dependent Inhibition With Dilution/Competition

Preincubation time-dependent inhibition with dilution and/or competition is a variant of *Method III* reported for kinetic analysis of irreversible covalent inhibitors (**Figure 21**).⁸³ Enzyme and inhibitor are preincubated in absence of competing substrate to form noncovalent EI complex and covalent EI* adduct, followed by dilution in a 10-100× larger volume ($V_{sub} >> V_{t'}$)

and/or addition of a high concentration of competing substrate ($[S] >> K_M$) (Figure 21A). The inhibitor concentration after substrate addition is far below the equilibrium concentration $([1]_t << 0.1K_i^{app})$, thereby inducing dissociation of inhibitor from the noncovalent enzymeinhibitor complex EI and quenching the formation of covalent EI* during incubation $(\Delta[EI^*]_t = 0)$. The approach is two-pronged: either dilution (reducing [I]_t) or saturating substrate concentration (increasing K_1^{app} and decreasing k_{chem}^{app}) can be sufficient as long as covalent EI* adduct formation is fully guenched, for example by dissociation of noncovalent EI complex. Preincubation time-dependent product formation velocity v_t reflects the inhibition by covalent EI* adduct formed during preincubation, and is calculated from the linear slope of product formation (Figure 21B). Enzyme activity v_t decreases exponentially from 0% covalent adduct without preincubation (Y-intercept = v^{ctrl}) to reach a plateau at 100% covalent adduct upon reaction completion (t' > $5t\frac{1}{2}$) for irreversible covalent inhibitors (Figure 21C). Observed rate of reaction completion k_{obs} (from 0-100% inhibition) is obtained by fitting to bounded exponential decay Equation VI (shown in Figure 21D). This is a simplified version of Equation v (shown in Figure 14D) in Method III (constraining $v_e = 0$) because we only consider 2-step irreversible inhibition (Data Analysis 4A) and 1-step irreversible inhibition (Data Analysis 4B). Reversible (2-step) covalent inhibition with a slow rate of covalent dissociation k_6 (t¹/_{2 diss} = LN(2)/ k_6) can be analyzed with preincubation dilution assays using the initial product formation velocity after rapid/jump dilution^{29, 33} but will not be discussed here because the (slow) dissociation of covalent EI* adduct may complicate algebraic analysis.

Generally, preincubation assays are disfavored because their experimental execution requires more material and measurements than incubation assays with continuous read-out. However, as already mentioned in Method III, preincubation methods are favored for inhibitors that have a slow covalent reaction rate and/or a poor noncovalent affinity. Additionally, dilution in excess substrate can resolve issues for enzyme assays that do not generate enough product for a robust signal (slow v^{ctrl}), as the maximum incubation time to calculate $v_{t'}$ is not limited by formation of EI* during incubation (Δ [EI*]_{*t*} = 0): incubation time can be longer than preincubation time. It is important to mention that there is still a limit to the incubation time: competition and/or dilution cannot fully mitigate the covalent adduct formation reaction, but it can be reduced to a negligible rate during the incubation. Finally, this method allows the assessment of covalent adduct formation potency without contamination by reversible inhibition. This can be beneficial in the analysis of 2-step covalent inhibitors that exhibit tightbinding behavior (customary for kinase inhibitors that have to compete with ATP): very potent noncovalent affinity 'shields' or 'contaminates' the rate of covalent adduct formation in the other protocols but not in this method, as detection is based solely on inhibition by covalent EI* adduct. However, the enzyme concentration during incubation is much lower than during preincubation, and inhibitor has to be present in excess during preincubation (pseudo-first order conditions), thus limiting the inhibitor concentration to higher concentrations than with other methods, which might be impractical. Be aware that dilution in (excess) substrate will change the absolute enzyme/inhibitor concentrations from preincubation to incubation, and make sure to calculate the desired enzyme concentration during incubation accordingly. Reaction completion ($v_{t'} < 0.1v^{\text{ctrl}}$) should not be reached before the first (shortest) preincubation time



Figure 21 | Method IV: Preincubation time-dependent inhibition with dilution/competition. Simulated with KinVol for 100 pM enzyme and 50 nM inhibitor C (before dilution) in $V_{t'}$ = 1, and 10 μ M substrate S1 in V_{sub} = 99: corresponding with 100-fold dilution in excess substrate ($[S] = 10K_{M}$). (A) Enzyme is preincubated with inhibitor to form noncovalent complex EI and covalent adduct EI* in absence of competing substrate, followed by dilution in excess substrate. Initial noncovalent EI complex forms rapidly ($[I]_{t'}/K_i = 0.5$) but fully dissociates upon dilution in a large volume ($V_{sub} >> V_{t'}$) and/or addition of a high concentration of competing substrate ([S] > K_{M}), as the E + I \leftrightarrow EI equilibrium has shifted towards fully unbound enzyme ([I],/K_i^{app} << 0.1). (B) Preincubation timedependent enzyme activity v_{t} is obtained from the (linear) slope of product formation velocity. Dilution in excess substrate quenches EI* formation after substrate addition (Δ EI* = 0), thus enabling longer incubation times compared to Method III. This measurement must be performed separately after each preincubation time. (C) Enzyme activity v_{t} , decreases exponentially from 0% covalent adduct (Y-intercept = enzyme activity without preincubation v_i) to 100% covalent adduct ($v_s = 0$). Enzyme activity without preincubation v_i equals the uninhibited enzyme activity v^{ctrl} for 1-step as well as 2-step irreversible inhibitors: dilution in excess substrate should induce full dissociation of noncovalently bound inhibitor ([I], << 0.1Ki app), and covalent adduct does not form instantly. (D) Bounded exponential decay Equation VI to fit preincubation time-dependent enzyme activity v., after dilution in (excess) competing substrate against preincubation time t' for irreversible 1-step and 2-step inhibition. This is a simplified version of Equation v (shown in Figure 14D): constraining $v_c = 0$ (inactivation at reaction completion). v_i = enzyme activity without preincubation = uninhibited enzyme activity v^{ctrl} because covalent adduct has not yet been formed and noncovalent complex has been disrupted by dilution in excess substrate. v_t = preincubation time-dependent enzyme activity reflecting covalent EI* adduct formed. t' = preincubation time of enzyme and inhibitor before substrate addition. k_{nhs} = observed rate of time-dependent inhibition from initial v_i to final v_s .

because it will be impossible to detect time-dependent changes in enzyme activity. This can be resolved by increasing the measurement interval (shorter dt') or reducing the inhibitor concentration whenever possible. This method is less suitable for inhibitors with a very fast covalent adduct formation k_{inact} because preincubation is performed in absence of competing substrate (thus allowing the maximum rate of covalent adduct formation possible at this inhibitor concentration).

Data Analysis 4A: 2-Step Irreversible Covalent Inhibition.

Data obtained for 2-step irreversible inhibitors (**Figure 22A**) is processed with *Data Analysis Protocol 4*, followed by *Data Analysis Protocol 4Ai* or *4Aii*. Kinetic analysis of enzyme activity with dilution/competition after preincubation in the presence of a 2-step covalent inhibitor is similar to data analysis of preincubation without dilution/competition (*Data Analysis 3A*), with the exception that longer incubation times are possible to calculate

Data Analysis 4A: 2-step IRREV



Figure 22 | Data Analysis 4A: Preincubation time-dependent inhibition with dilution/competition for 2-step irreversible covalent inhibition. Simulated with **KinVol** for 2-step IRREV inhibitor **C** with 100 pM enzyme in $V_t = 1$ ([E^{total}]_t = 10, [E^{total}]_t = 1), and 10 µM substrate **S1** ([S] = 10K_M) in $V_{sub} = 99$. (**A**) Schematic enzyme dynamics during preincubation in absence of substrate and during incubation after dilution in excess substrate. (**B**) Time-dependent product formation after preincubation (t' = 1800 s) in absence of inhibitor *F*^{ctrl} or in presence of various inhibitor concentrations. Enzyme activity after preincubation $v_{t'}$ is obtained from the linear slope. (**C**) Preincubation time-dependent enzyme activity $v_{t'}$ is fitted to **Equation VI** (shown in **Figure 21D**) for each inhibitor concentration with global shared value for v_i ($v_i = v^{ctrl}$) to obtain observed rates of inactivation k_{obs} . Alternatively, $v_{t'}$ can be normalized to a fraction of the uninhibited enzyme activity v^{ctrl} . (**D**) Half-maximum $k_{obs} = \frac{1}{2}k_{inact}$ is reached when inhibitor concentration during preincubation equals the inactivation constant K_I. No correction for substrate competition: v_t reflects the remaining unbound/noncovalent enzyme activity after preincubation in absence of competing substrate.

enzyme activity v_t from the slope (**Figure 22B**), and enzyme activity without preincubation v_i should be equal to the uninhibited enzyme activity v^{ctrl} (**Figure 22C**). Contrary to *Method III*, this does not imply that the inhibitors show 1-step behavior: it merely confirms that extensive dilution/substrate competition successfully induced inhibitor dissociation from noncovalent EI complex to unbound enzyme. It is essential to plot the rate of covalent adduct formation k_{obs} against the inhibitor concentration <u>during preincubation</u> (**Figure 22D**) to obtain kinetic parameters: k_{obs} is based on the formation of EI* during preincubation, and the inhibitor concentration during preincubation is much higher than the inhibitor concentration after dilution in substrate ($[I]_{t'} >> [I]_t$). Insufficient dilution/competition will partially disrupt noncovalent EI complex, resulting in a time-dependent decrease of enzyme activity due to formation of EI* after substrate addition, and deviation from $v_i = v^{\text{ctrl}}$, as noncovalent complex EI contributes to inhibition without preincubation. Increasing substrate concentration and/or



Data Analysis 4A: 2-step IRREV

Figure 23 | Data Analysis 4A: Corrections for spontaneous loss of enzyme activity (2-step IRREV). Simulated with **KinVolDeg** for 2-step IRREV inhibitor **C** with 100 pM enzyme in $V_{t'} = 1$ ([E^{total}]_t = 100, [E^{total}]_t = 1), and 10 μ M substrate **S1** ([S] = 10K_M) in $V_{sub} = 99$, and $k_{degE} = k_{degE5} = k_{degE1} = 0.0003 \text{ s}^{-1}$. (**A**) Schematic enzyme dynamics during preincubation in absence of substrate and during incubation after dilution in excess substrate with spontaneous enzyme degradation/denaturation. (**B**) Uninhibited enzyme activity after preincubation v_t^{ctrl} decreases with longer preincubation. Enzyme activity $v_{t'}$ is fitted to **Equation VI** (shown in **Figure 21D**) for each inhibitor concentration during preincubation with globally shared value for v_i ($v_i = v_0^{ctrl}$) to obtain observed rates of inactivation k_{obs} , as well as fitting uninhibited activity v_t^{ctrl} to obtain the rate of nonlinearity k_{ctrl} . (**C**) Inhibitor concentration-dependent k_{obs} with spontaneous enzyme degradation increases with k_{ctrl} but the span from k_{min} (= k_{ctrl}) to k_{max} (= $k_{inact} + k_{ctrl}$) still equals k_{inact} . Fit with algebraic correction for nonlinearity (*black line*, $k_{ctrl} > 0$). Ignoring the nonlinearity (**g***ay line*, constrain $k_{ctrl} = 0$) results in underestimation of K_I (overestimation of potency) and overestimation of k_{inact} . (**D**) Normalized enzyme activity $v_t' v_0^{ctrl}$ is fitted to **Equation VI** (shown in **Figure 21D**) for each inhibitor concentration during preincubation (constrain $v_i v_0^{ctrl} = 1$) to obtain corrected observed rates of inactivation k_{obs} . (**E**) Inhibitor concentration-dependent k_{obs} has been corrected for enzyme degradation by fitting normalized enzyme activity $v_t' v_{ctrl}^{ctrl}$ and does not require further corrections.

dilution in a larger volume might resolve this. Alternatively, enzyme activity with partial disruption of noncovalent EI analyzed with *Data Analysis Protocol 3A* still results in reliable estimates of k_{obs} . Please note that, although detection based only on covalent adduct formation allows analysis of 2-step inhibitors displaying tight-binding behavior (very high noncovalent affinity resulting in full inhibition at all inhibitor concentrations), these inhibitor concentrations are saturating if they comply with the rapid equilibrium approximation ($K_i \approx K_1$); thus, it would only be possible to determine the lower limit of k_{inact} and the upper limit of K_1 (see **Figure 3G**).

Correction for enzyme (in)stability during preincubation (**Figure 23A**) by correcting for the rate of spontaneous degradation k_{ctrl} has been reported for dilution experiments with irreversible covalent inhibitors (**Figure 23B-C**).⁸⁶ Alternatively, enzyme activity after preincubation $v_{t'}$ can be normalized to the uninhibited enzyme activity after preincubation $v_{t'}$ (**Figure 23D-E**).

Data Analysis 4B: 1-Step Irreversible Covalent Inhibition.

Data obtained for 1-step irreversible inhibitors (**Figure 24A**) is processed with *Data Analysis Protocol 4*, followed by *Data Analysis Protocol 4Bi* or *4Bii*. Kinetic analysis of enzyme activity with dilution/competition after preincubation in presence of a 1-step covalent inhibitor is almost identical to data analysis of preincubation without dilution in excess substrate (*Data Analysis 3B*), with the exception that enzyme activity $v_{t'}$ can now be calculated from the slope after longer incubation times (**Figure 24B**). It is essential to plot the rate of covalent adduct formation k_{obs} (**Figure 24C**) against the inhibitor concentration <u>during preincubation</u> (**Figure 24D**) to obtain kinetic parameters: k_{obs} is based on the formation of EI* during preincubation, and the inhibitor concentration during preincubation will be much higher than the inhibitor concentration after dilution in substrate ($[I]_{t'} >> [I]_t$). Dilution/ competition does not disrupt any noncovalent EI complex, as this is nonexistent for 1-step inhibitors, but the rate of covalent adduct formation k_{obs} should be negligible after dilution in



Figure 24 | Data Analysis 4B: Preincubation time-dependent inhibition with dilution/competition for 1-step irreversible covalent inhibition. Simulated with **KinVol** for 1-step IRREV inhibitor **D** with 100 pM enzyme in $V_t = 1$ ([E^{total}]_t = 10, [E^{total}]_t = 1), and 10 µM substrate **S1** ([S] = 10K_M) in $V_{sub} = 99$. (**A**) Schematic enzyme dynamics during preincubation in absence of substrate and during incubation after dilution in excess substrate. (**B**) Time-dependent product formation after preincubation (t' = 1800 s) in absence of inhibitor *F*^{ctrl} or in presence of various inhibitor concentrations. Enzyme activity after preincubation v_t is obtained from the linear slope. (**C**) Preincubation time-dependent enzyme activity v_t is fitted to **Equation VI** (shown in **Figure 21D**) for each inhibitor concentration with global shared value for $v_i (v_i = v^{ctrl})$ to obtain observed rates of inactivation k_{obs} . Alternatively, v_t can be normalized to a fraction of the uninhibited enzyme activity v^{ctrl} . Inhibitor concentration-dependent k_{obs} increases linearly with inhibitor concentration during preincubation, with k_{chem} as the slope. No correction for substrate competition: $v_{t'}$ reflects the remaining unbound enzyme activity after preincubation in the absence of competing substrate.

excess substrate, to prevent formation of covalent EI*. Insufficient dilution and/or competition $(\Delta[EI^*]_t > 0)$ can result in time-dependent decrease of enzyme activity due to formation of EI* after substrate addition. Increasing substrate concentration and/or dilution in a larger volume might resolve this if necessary, but simply performing analysis with *Data Analysis Protocol 3B* also results in reliable estimates of k_{obs} . Inhibitor concentrations that reach reaction completion during the shortest preincubation time should be excluded from the fit (highest concentration in **Figure 24C**) as these fits are not reliable.

Correction for enzyme (in)stability (**Figure 25A**) using the rate of spontaneous degradation k_{ctrl} has been reported for dilution experiments with irreversible covalent inhibitors (**Figure 25B-C**).⁸⁶ Alternatively, enzyme activity after preincubation v_t can be normalized to the uninhibited enzyme activity after preincubation v_t^{ctrl} (**Figure 25D-E**).



Figure 25 | Data Analysis 4B: Corrections for spontaneous loss of enzyme activity (1-step IRREV). Simulated with **KinVolDeg** for 1-step IRREV inhibitor **D** with 100 pM enzyme in $V_{t'} = 1$ ([E^{total}]_t = 100, [E^{total}]_t = 1), and 10 μ M substrate **S1** ([S] = 10K_M) in $V_{sub} = 99$, and $k_{degE} = k_{degES} = k_{degEI} = 0.0003 s^{-1}$. (**A**) Schematic enzyme dynamics during preincubation in absence of substrate and during incubation after dilution in excess substrate with spontaneous enzyme degradation/denaturation. (**B**) Uninhibited enzyme activity after preincubation v_t^{ctrl} decreases with longer preincubation with globally shared value for v_i ($v_i = v_0^{ctrl}$) to obtain observed rates of inactivation k_{obs} , along with fitting uninhibited activity v_t^{ctrl} to obtain the rate of nonlinearity k_{ctrl} . (**C**) Inhibitor concentration-dependent k_{obs} with spontaneous enzyme degradation/denaturation increases by k_{ctrl} . Fit with algebraic correction for nonlinearity (*black line*, $k_{ctrl} > 0$) or ignoring nonlinearity (*gray line*, constrain $k_{ctrl} = 0$). Ignoring the nonlinearity (assuming Y-intercept = 0) results in overestimation of k_{chem} (steeper slope). (**D**) Normalized enzyme activity $v_t'v^{ctrl}$ is fitted to **Equation VI** (shown in **Figure 21D**) for each inhibitor concentration divenge network v_{obs} , we degradation **VI** (shown in **Figure 21D**) for each inhibitor concentration-dependent k_{obs} solution $v_i/v_0^{ctrl} = 0$) results in overestimation of k_{chem} (steeper slope). (**D**) Normalized enzyme activity $v_t'v^{ctrl}$ is fitted to **Equation VI** (shown in **Figure 21D**) for each inhibitor concentration during preincubation (constrain $v_i/v_0^{ctrl} = 1$) to obtain corrected observed rates of inactivation k_{obs} . (**E**) Inhibitor concentration-dependent k_{obs} has been corrected for enzyme degradation by fitting normalized enzyme activity v_t'/v^{ctrl} and does not require further corrections.

4. Experimental Protocols

Assay Protocol I. Progress Curve Analysis of Substrate Association Competition

The protocol below provides a generic set of steps to accomplishing this type of measurement.

Materials

- 1× Assay/reaction buffer supplemented with co-factors and reducing agent
- Active enzyme, 4× solution in assay buffer
- Substrate with continuous read-out, 4× solution in assay buffer
- Positive control: vehicle/solvent as DMSO stock, or 2% solution in assay buffer
- Negative control: known inhibitor or alkylating agent as DMSO stock, or 2× solution in assay buffer
- Inhibitor: as DMSO stock, or serial dilution of 2× solution in assay buffer with 2% DMSO
- 384-well low volume microplate with nonbinding surface (e.g. Corning 3820 or 4513) for incubation and read-out
- Optical clear cover/seal (e.g. Perkin Elmer TopSeal-A Plus, #6050185, or Corning 6575 Universal Optical Sealing Tape,
- or Duck Brand HP260 Packing Tape)
- 1.5 mL (Eppendorf) microtubes to prepare stock solutions
- Optional: 96-well microplate to prepare serial dilution of inhibitor concentration
- Microplate reader equipped with appropriate filters to detect product formation (e.g. CLARIOstar microplate reader)
- Optional: Automated (acoustic) dispenser (e.g. Labcyte ECHO 550 Liquid Handler acoustic dispenser)

Exemplary assay concentrations

	incubation t		
	[stock]	Volume	[conc] _t
Enzyme	4 nM	5 μL	0.99 nM
Inhibitor	20 nM	10.2 μL	10.10 nM
Substrate	4 µM	5 μL	0.99 μM
Total		20.2 μL	

Before you start, optimize assay conditions in the uninhibited control to ensure compliance with assumptions and restrictions for progress curve analysis - most importantly linear product formation in the uninhibited control for the duration of the experiment $(k_{ctrl} = 0)$ – by activating the enzyme before reaction initiation (e.g. preincubation with reducing agent for proteases, or ATP for kinases and ligases), testing the enzyme activity on the (fluorogenic) substrate in absence of inhibitor, and adjusting the enzyme and substrate concentration $([S]_0 > 10[E]_0)$ to reach maximum 10% substrate conversion at the end of the measurement window ($[P]_{,} < 0.1[S]_{0}$). Further optimization typically involves tuning the reader settings for optimal sensitivity, measurement of a calibration curve for product concentration,68-69 and calculation of the Z'-score from the uninhibited and inhibited controls (ideally 8 replicates)⁸⁷ in a separate experiment to validate that enough product is formed for a good signal/noise ratio (Z' > 0.5) at the end of the measurement. Consult Table 6 in section 5 for common optimization and troubleshooting options. The read-out of product formation must be homogeneous/continuous. Product formation of substrates with a less sensitive read-out (e.g. fluorescence polarization) may generate a relatively low product signal relative to the unprocessed substrate, and substrate depletion is unavoidable to generate a sufficient Z'-score.⁸⁷ Algebraic analysis of 2-step irreversible inhibition with substrate depletion ($[P]_{t} < 0.1[S]_{0}$) can be performed with Data Analysis Protocol 1D after completion of steps 2-6 of Assay Protocol I.

1. Add inhibitor or control to each well with the uninhibited control for full enzyme activity containing the same volume vehicle/solvent instead of inhibitor (we use DMSO in this protocol). Add a constant volume of serially diluted inhibitor in assay buffer supplemented with DMSO (e.g. $10.2 \,\mu$ L of 2× solution containing 2% DMSO) or add inhibitor and controls by (acoustic) dispensing of the pure DMSO stocks, with DMSO backfill to a constant volume (e.g. $0.2 \,\mu$ L), followed by addition of assay buffer to each well (e.g. $10 \,\mu$ L) and gentle shaking (300 rpm) to homogenize the solution.

Typically, measurements are performed in triplicate (or more replicates) with at least 8 inhibitor concentrations. Inhibitor concentrations might need optimization, but a good starting point is $0.1-10 \times |C_{50}$; the highest inhibitor concentration should correspond to maximum 90% initial (noncovalent) inhibition ($v_i > 0.1v^{ctri}$), as it can be difficult to accurately detect the increase from 90% to 100% inhibition.

 Add substrate in assay buffer to each well (e.g. 5 μL of 4× solution) and homogenize the solutions by gentle shaking (300 rpm).

The order of substrate or inhibitor addition is not important *per se*, as long as enzyme is the last reagent to be added, and DMSO stocks are added prior to buffered (aqueous) solutions. Optionally, gently centrifuge the plate (1 min at 1000 rpm) to ensure that assay components are not stuck at the top of the well.

3. Add active enzyme in assay buffer to each well (e.g. 5 μ L of 4× solution), with minimal delay between addition to the first and the last well. Optionally, gently centrifuge the plate (1 min at 1000 rpm) if bubbles are formed (especially for buffers containing surfactants), as these will induce assay artifacts, and to ensure assay components are in solution together rather than stuck to the wall at the top of the well.

Manual addition of enzyme solution and physically moving the plate to the plate reader introduces a delay that may slightly affect the accuracy of the measurement, as it can be variable (depending on the total number of wells, distance to the machine and walking pace of the researcher). This should not be significant if the delay is short compared to the total reaction time, but it can affect the outcome in the data analysis when t0 is actually 1-2 min. One method to monitor the delay between reaction initiation (onset of product formation and inhibition) and the start of product detection is *step 6* is evaluation of the Y-intercept values (as discussed in **Table 6**, *section 5*). Alternatively, enzyme addition with an injector built into the plate reader minimizes the delay between reaction initiation (onset of product formation and inhibition) and starting the measurement.

4. Seal the wells by applying an optical clear cover.

Continuous kinetic measurements are subject to assay artifacts such as drift due to evaporation. In our experience, application of an optical clear cover/seal prior to measurement improves the assay robustness and resolves significant aberrant nonlinearity unrelated to enzyme activity.

5. Measure product formation in microplate reader by detection of the product read-out.

A typical assay measurement window is 60-240 min, with a measurement interval of 1-2 min. The inhibitor-binding reaction does not have to reach completion (100% inhibition for irreversible inhibitors, equilibrium for reversible inhibitors) within this window, but data will be more reliable when completion is reached before the end of the measurement (see also **Figure 5B**).

6. Proceed to Data Analysis Protocols to calculate the appropriate kinetic parameters for each covalent binding mode: Data Analysis Protocol 1A for 2-step irreversible inhibitors, Data Analysis Protocol 1B for 1-step irreversible inhibitors, Data Analysis Protocol 1C for 2-step reversible inhibitors, or Data Analysis Protocol 1D for 2-step irreversible inhibitors with substrate depletion.

	Data Analysis Protocol				
EXP Conditions	2-step IRREV	1-step IRREV	2-step REV		
$k_{\rm ctrl} = 0$	1A	1B	1C		
$k_{\text{degE}} > 0$	1A	1B	-		
$[P]_t > 0.1[S]_0$	1D	-	-		

Data Analysis Protocol 1A for 2-Step IRREV Inhibition

Processing of raw data obtained with Assay Protocol I for 2-step irreversible covalent inhibitors.

1. Plot signal F against incubation time t

Plot signal (in AU) on the Y-axis against incubation time (in s) on the X-axis for each inhibitor concentration and the controls (Figure 6B). Product formation in the uninhibited control *F*^{ctrl} should be linear. Consult **Table 6** (section 5) for

troubleshooting of nonlinearity of the uninhibited control. Optionally, perform background correction to correct for assay artifacts such as bleaching and drift that cause a negative final velocity ($v_s < 0$ AU/s) in the fully inhibited control. This correction can be subtraction of the background in presence of substrate (and inhibitor) but absence of enzyme, or subtraction of the fully inhibited control. Consult the guidelines of your data fitting software for instructions on background corrections (e.g. GraphPad Prism).⁸⁸

2. Fit signal F_t against t to obtain k_{obs}

Fit signal F_t against incubation time t to **Equation II** (Figure 6B/E). Constrain final velocity $v_s = 0$ for backgroundcorrected product formation, or $v_s =$ value for full inhibition control. A lack of initial noncovalent complex ($v_i = v^{ctrl}$) is indicative of 1-step binding behavior.

$$F_t = v_s t + \frac{v_i - v_s}{k_{obs}} \left[1 - e^{-k_{obs} t} \right] + F_0$$
(II)

Nonlinear regression of user-defined explicit equation Y = (vs*X) + ((vi-vs)/kobs)*(1-EXP(-kobs*X))) + Y0 with $Y = signal F_t$ (in AU), X = incubation time t (in s), and $vs = final slope v_s$ (in AU/s, constrained) to find Y0 = Y-intercept F_0 = background signal at t = 0 (in AU), $vi = initial slope v_i$ (in AU/s) and kobs = observed reaction rate k_{obs} (in s⁻¹).

3. Plot k_{obs} against [I]

Plot the mean and standard deviation of k_{obs} (in s⁻¹) on the Y-axis against inhibitor concentration (in M) after reaction initiation by enzyme addition (in the final solution) on the X-axis (**Figure 6C/F**). The plot of k_{obs} against [1] should reach a maximum k_{obs} at saturating inhibitor concentration. Note that a linear curve is indicative of 1-step binding behavior at non-saturating inhibitor concentrations ([I] << 0.1K_1^{app}: see **Figure 3F**) with $v_i = v^{ctri}$ (low initial inhibition). Proceed to *step 4* of *Data Analysis Protocol 1B* after it has been validated that the linear curve is not resultant from saturating inhibitor concentrations ([I] >> $10K_1^{app}$: see **Figure 3G**) as identified by $v_i << v^{ctri}$ (significant initial inhibition), by repeating the measurement with a higher competitive substrate concentration (increase K_1^{app}) and/or lower inhibitor concentration.

4. Fit k_{obs} against [I] to obtain k_{inact} and K_{I}^{app}

Fit k_{obs} against inhibitor concentration to **Equation VII** to obtain maximum inactivation rate constant k_{inact} and apparent inactivation constant K_1^{app} . Constrain $k_{ctrl} = k_{obs}$ of the uninhibited control (**Figure 6F**). Calculate inactivation constant K_1 and irreversible covalent inhibitor potency k_{inact}/K_1 with *Sample Calculation 1&2*.

$$k_{\rm obs} = k_{\rm ctrl} + \frac{k_{\rm inact} [I]}{K_{\rm I}^{\rm app} + [I]}$$
(VII)

Nonlinear regression of user-defined explicit equation Y=YO+((kmax*X)/((Klapp)+X)) with $Y = observed reaction rate k_{obs} (in s^{-1}), X = inhibitor concentration (in M), and YO = rate of nonlinearity in uninhibited control <math>k_{ctrl}$ (in s^{-1} , constrained) to find kmax = maximum reaction rate k_{max} (in s^{-1}) and Klapp = Apparent inactivation constant K_{L}^{app} (in M).

5. EXTRA: Plot and fit v, against [I] to obtain K, app

Inhibition constant K₁ can be calculated from the initial velocity v_i (obtained in *step 3*), reflecting the rapid (initial) noncovalent enzyme–inhibitor equilibrium. Plot the mean and standard deviation of v_i (in AU/s) on the Y-axis against inhibitor concentration on the X-axis (similar to **Figure 8D**). Fit v_i against [1] to four-parameter nonlinear regression Hill **Equation VIII** to obtain apparent inhibition constant K_i^{app.33} Constrain the top to the uninhibited v_i (maximum velocity = v^{tril}) and the bottom to the fully inhibited v_i (minimum velocity = v₁^{min}. For (background-)corrected product formation v_i^{min} = 0). Calculate inhibition constant K_i with *Sample Calculation 3*.

$$v_{i} = v_{i}^{\min} + \frac{v^{\text{ctrl}} - v_{i}^{\min}}{1 + \left(\frac{[I]}{K_{i}^{\text{app}}}\right)^{h}}$$
(VIII)

Nonlinear regression of four-parameter dose-response equation $Y=Bottom+(Top-Bottom)/(1+(X/IC50)^HillSlope)$ with Y = initial product formation velocity v, (in AU/s), X = inhibitor concentration (in M), Bottom = velocity in fully inhibited control v^{min} (in AU/s, constrained), and Top = maximum velocity in uninhibited control v^{ctd} (in AU/s, constrained) to find Hillslope = Hill coefficient h (unitless) and IC50 = apparent inhibition constant K^{app} (in M).

6. Optional: Validate experimental kinetic parameters with kinetic simulations

Proceed to *Kinetic Simulations* 1 to compare the experimental progress curves to the progress curves simulated with scripts **KinGen** and **KinDeg** (using experimental rate constant $k_{inact} = k_5$) to confirm that the calculated kinetic constants are in accordance with the experimental data.

Data Analysis Protocol 1B for 1-Step IRREV Inhibition

Processing of raw data obtained with Assay Protocol I for 1-step irreversible covalent inhibitors and 2-step irreversible inhibitors at non-saturating inhibitor concentrations ([I] $\leq 0.1K_i^{app}$).

1. Plot signal F against incubation time t

Plot signal (in AU) on the Y-axis against incubation time (in s) on the X-axis for each inhibitor concentration and the controls (**Figure 7B**). Product formation in the uninhibited control F^{ctrl} should be linear. Consult **Table 6** (section 5) for troubleshooting of nonlinearity of the uninhibited control. Optionally, perform background correction to correct for assay artifacts such as bleaching and drift that cause a negative final velocity ($v_s < 0$ AU/s) in the fully inhibited control. This correction can be subtraction of the background in presence of substrate (and inhibitor) but absence of enzyme, or subtraction of the fully inhibited control. Consult the guidelines of your data fitting software for instructions on background corrections (e.g. GraphPad Prism).⁸⁸

2. Fit F_t against t to obtain k_{obs}

Fit signal F_t against incubation time t to **Equation II (Figure 7B/E)**. Constrain final velocity $v_s = 0$ for background-corrected product formation, or $v_s =$ value for full inhibition control. Initial velocity v_i should be a shared value because noncovalent inhibition does not significantly contribute to the initial inhibition for inhibitors displaying 1-step behavior.

$$F_t = v_s t + \frac{v_i - v_s}{k_{obs}} \left[1 - e^{-k_{obs}t} \right] + F_0$$
(II)

Nonlinear regression of user-defined explicit equation Y = (vs*X) + ((vi-vs)/kobs)*(1-EXP(-kobs*X))) + Y0 with $Y = signal F_t$ (in AU), X = incubation time t (in s), and $vs = final slope v_s$ (in AU/s) to find Y0 = Y-intercept F_0 = background signal at t = 0 (in AU), vi = initial slope v_i (in AU/s, shared value), and kobs = observed reaction rate k_{obs} (in s^{-1}).

3. Plot k_{obs} against [I]

Plot the mean and standard deviation of k_{obs} (in s⁻¹) on the Y-axis against inhibitor concentration (in M) after reaction initiation by enzyme addition (in the final solution) on the X-axis (**Figure 7C/F**). The plot of k_{obs} against inhibitor concentration [I] is linear for 1-step irreversible inhibitors and for 2-step irreversible inhibitors at non-saturating inhibitor concentrations ([I] << 0.1K_i^{app}).

4. Fit k_{obs} against [I] to obtain k_{chem}^{app}

Fit k_{obs} against inhibitor concentration to **Equation IX** to obtain apparent inhibitor potency k_{chem}^{app} from the linear slope. Constrain Y-intercept $k_{ctrl} = k_{obs}$ of the uninhibited control (**Figure 7F**). Calculate k_{chem} reflecting inhibitor potency for 1-step irreversible covalent inhibition with *Sample Calculation 4*. Calculate k_{inact}/K_1 for 2-step irreversible inhibitors at non-saturating inhibitor concentrations ([I] $\leq 0.1K_1^{app}$) with *Sample Calculation 5 and 6*.

$$k_{\rm obs} = k_{\rm ctrl} + k_{\rm chem}^{\rm app}$$
 [I]

(IX)

Nonlinear regression of straight line Y=YIntercept+Slope*X with Y = observed reaction rate k_{obs} (in s⁻¹), X = inhibitor concentration (in M), and YIntercept = rate of nonlinearity in uninhibited control k_{ctrl} (in s⁻¹, constrained) to find Slope = apparent inactivation rate constant k_{chem}^{app} (in M⁻¹s⁻¹).

5. Optional: Validate experimental kinetic parameters with kinetic simulations

Proceed to *Kinetic Simulations* 1 to compare the experimental progress curves to the progress curves simulated with scripts **KinGen** and **KinDeg** (using experimental rate constant $k_{chem} = k_3$) to confirm that the calculated kinetic constants are in accordance with the experimental data.

Data Analysis Protocol 1C for 2-Step REV Inhibition

Processing of raw data obtained with Assay Protocol I for 2-step reversible covalent inhibitors.

1. Plot signal F against incubation time t

Plot signal (in AU) on the Y-axis against incubation time (in s) on the X-axis for each inhibitor concentration and the controls (**Figure 8B**). Product formation in the uninhibited control *F*^{ctri} should be linear. Consult **Table 6** (*section 5*) for troubleshooting of nonlinearity of the uninhibited control. Optionally, perform background correction to correct for assay artifacts such as bleaching and drift that cause a negative final velocity ($v_s < 0$ AU/s) in the fully inhibited control. This correction can be subtraction of the background in the presence of substrate (and inhibitor) but absence of

enzyme, or subtraction of the fully inhibited control. Consult the guidelines of your data fitting software for instructions on background corrections (e.g. GraphPad Prism).⁸⁸

2. Fit F_t against t to obtain k_{obs} and v_s

Fit signal F_t against incubation time t to **Equation II (Figure 8B)** to obtain final product formation velocity v_s and the observed reaction rate k_{obs} from initial equilibrium v_i to steady-state equilibrium v_s . Do not constrain initial velocity v_i or final velocity v_s . Also fit the progress curve of the uninhibited control (F^{ctri}) to validate that product formation is strictly linear ($v_i^{ctri} = v_s^{ctri}$), because algebraic correction for nonlinearity in the uninhibited control is not possible (illustrated in **Figure 9**). The observed rate k_{obs} reflects the exponential reaction rate from initial noncovalent equilibrium (v_i) to final steady-state equilibrium (v_s).

$$F_{t} = v_{s}t + \frac{v_{i} - v_{s}}{k_{obs}} \left[1 - e^{-k_{obs}t}\right] + F_{0}$$
(II)

Nonlinear regression of user-defined explicit equation Y = (vs*X) + ((vi-vs)/kobs)*(1-EXP(-kobs*X))) + Y0 with $Y = signal F_t$ (in AU) and X = incubation time t (in s) to find Y0 = Y-intercept F_0 = background signal at t = 0 (in AU), vi = initial slope v_i (in AU/s), vs = final slope v_s (in AU/s), and kobs = observed reaction rate k_{obs} (in s^{-1}).

3. Plot and fit v_s against [I] to obtain K_i^{*app}

Apparent steady-state inhibition constant K_i^{*app} can be calculated from the final velocity v_s (obtained in the previous step) reflecting enzyme activity after reaching the steady-state inhibitor equilibrium (*reaction completion*). Plot the mean and standard deviation of v_s (in AU/s) on the Y-axis against inhibitor concentration (in M) on the X-axis and fit to four-parameter nonlinear regression Hill **Equation x** to obtain apparent steady-state inhibition constant K_i^{*app} (**Figure 8D**).³³ Constrain the top to uninhibited velocity v^{ctri} (maximum velocity) = v_s^{max}) and the bottom to the fully inhibited $v_s (v_s^{min}$, minimum velocity). For (background-)corrected product formation, $v_s^{min} = 0$. Accurate values are only obtained when uninhibited product formation is strictly linear ($k_{ctri} = 0$) or when the rate of spontaneous inactivation k_{ctri} is much smaller than the covalent dissociation k_6 (**Figure 9D**). Validate that v_s is not driven by spontaneous enzyme degradation ($k_{ctri} < k_6$) by also fitting without constraints for v_s^{max} . Calculate steady-state inhibition constant K_i^* with *Sample Calculation* 7.

$$v_{s} = v_{s}^{\min} + \frac{v^{\text{ctrl}} - v_{s}^{\min}}{1 + \left(\frac{[I]}{K_{i}^{*app}}\right)^{h}}$$
(x)

Nonlinear regression of four-parameter dose-response equation $Y=Bottom+(Top-Bottom)/(1+(X/IC50)^HillSlope)$ with $Y = final product formation velocity <math>v_s$ (in AU/s, X = inhibitor concentration (in M), Bottom = velocity in fully inhibited control v_s^{min} (in AU/s, constrained) and Top = maximum velocity in uninhibited control v_s^{crit} (in AU/s, constrained) to find Hillslope = Hill coefficient h (unitless) and IC50 = apparent steady-state inhibitor constant V_s^{map} (in AU).

4. Optional: Plot and fit k_{obs} against [I] to obtain K_i^{app} , k_5 , and k_6

This is an optional data processing step to obtain kinetic parameters by fitting to the observed rate k_{obs} (obtained in *step 2* of *Data Analysis 1C*), and is used to validate K_1^{*app} values found in the previous step, to check if nonlinearity in the uninhibited control k_{ctrl} affects the fit, and/or to generate experimental k_5 and k_6 values to use in kinetic simulations. Plot the mean and standard deviation of k_{obs} (in s⁻¹) on the Y-axis against inhibitor concentration (in M) on the X-axis (Figure 8C). Exclude k_{obs} of uninhibited control (k_{ctrl}) from the fit. Fit k_{obs} against inhibitor concentration to Equation XI to obtain rate constants for the covalent association k_5 and covalent dissociation k_6 , as well as apparent noncovalent inhibition constant K_1^{app} reflecting the rapid (initial) noncovalent equilibrium. Use the inhibitor concentration after reaction initiation by enzyme addition (in the final solution). Accurate values are only obtained when uninhibited product formation is strictly linear ($k_{ctrl} = 0$). Y-intercept approaching k_{ctrl} despite the uninhibited control not being included in the fit is a red flag that should not be ignored, as this is indicative of spontaneous enzyme degradation rather than k_6 dominating k_{obs} at low inhibitor concentrations, for which algebraic corrections are not available (Figure 9C). Calculate noncovalent K₁^{*} with *Sample Calculation 8*. Optionally, perform step 6 of Data Analysis 1A to obtain apparent noncovalent inhibition constant K₁^{**} perform the initial velocity v_1 (obtained in the fit 2 of Data Analysis 1A to obtain apparent noncovalent inhibition constant K₁^{**} with Sample Calculation 8 of Data Analysis 1A to obtain apparent noncovalent inhibition constant K₁^{**} perform step 2 of Data Analysis 1A to obtain apparent noncovalent inhibition constant K₁^{**} obtain apparent noncovalent inhibition constant K₁^{**} with Sample formal to perform step 2 of Data Analysis 1A.

$$k_{\rm obs} = k_6 + \frac{k_5 [I]}{K_i^{\rm app} + [I]}$$
(XI)

Nonlinear regression of user-defined explicit equation Y=YO+((kmax*X)/((Kiapp)+X)) with Y = observed reaction rate k_{obs} (in s⁻¹) and X = inhibitor concentration (in M) to find YO = covalent dissociation rate constant k_6 (in s⁻¹), kmax = covalent association rate constant k_5 (in s⁻¹) and Kiapp = Apparent inhibition constant K_{ipp}^{ipp} (in M).

5. Optional: Validate experimental kinetic parameters with kinetic simulations

Proceed to *Kinetic Simulations 1* to compare the experimental progress curves to the progress curves simulated with scripts **KinGen** and **KinDeg** to confirm that the calculated kinetic constants are in accordance with the experimental data. Experimental estimates of k_s and k_s are generated in the previous step of this protocol.

Data Analysis Protocol 1D for 2-Step IRREV Inhibition with Substrate Depletion

Processing of raw data obtained with Assay Protocol I for 2-step irreversible covalent inhibitors with nonlinearity in the uninhibited control resultant from substrate depletion $([P]_r < 0.1[S]_0)$.

Before you start, validate compliance with essential assay reaction conditions such as the Hit-and-Run model. This algebraic correction for substrate depletion has additional requirements for assay conditions, ⁸⁴ and is only compatible with 2-step irreversible inhibition (**Figure 10**). Validate that the product formation reaction complies with the Hit-and-Run model $E + S \rightarrow E + P$ (shown in **Figure 10A**): substrate concentration must be far below the K_M ($[S]_0 < 0.1K_M$) to calculate the pseudo-first order reaction rate constant for enzymatic product formation $k_{sub} = k_{cat}/K_M$ ($M^{-1}s^{-1}$). Observed nonlinearity in the uninhibited control should be fully attributed to substrate depletion (**Figure 10B**). Convert the maximum signal F^{ctrl} (in AU) into product concentration experiment. ⁶⁸⁻⁶⁹ Validate that the total substrate conversion to product exceeds 10% of the initial substrate concentration ($[Pctrl]_t > 0.1[S]_0$), and that substrate depletion is the only factor that contributes to the observed nonlinearity: uninhibited product formation should be linear when incubation times are shorter ($[P]_t < 0.1[S]_0$) or enzyme concentration is lower. Alternatively, perform kinetic analysis by numeric solving if one or more assumptions are violated.⁸⁰

$$[\mathbf{P}]_t = \frac{F_t - F_0}{r_{\mathbf{p}}}$$

Calculate: Pt=(Ft-F0)/rp with Pt = product concentration at the end of the incubation $[P]_t$ (in A), Ft = signal in uninhibited control at the end of the incubation time F_t (in AU), F0 = substrate background signal F_0 (in AU) and rp = product coefficient r_p (in AU/M product).

1. Plot signal F against incubation time t

Plot signal (in AU) on the Y-axis against incubation time (in s) on the X-axis for each inhibitor concentration (Figure 10C). Label the columns with the inhibitor concentration (in M).

2. Perform background correction

Correct for assay artifacts such as fluorescence bleaching and drift that cause a declining signal in the fully inhibited control. This correction can be subtraction of the time-dependent background in absence of enzyme but in presence of substrate (and inhibitor), or subtraction of the fully inhibited control. Consult the guidelines of your data fitting software for instructions on background corrections (e.g. GraphPad Prism).⁸⁸

3. Globally fit F_t against t to obtain k_{inact} and K_T^{app}

Globally fit the progress curves of time-dependent signal F_t for all inhibitor concentrations to **Equation III (Figure 10C)**. Consult the guidelines of your data fitting software for instructions on user-defined (implicit) equations.⁶⁶ Exclude the dataset of the fully inhibited control from the fit. Constrain $[E]_0$ (in M), $[S]_0$ (in M), and $[I] = [I]_0$ (in M) to their theoretical values. Originally, $[I]_0$ was locally optimized,⁸⁰ but we used fixed values of $[I]_0$ in GraphPad Prism. Constrain product coefficient r_p (in AU/M product) to the value determined in a separate product calibration experiment. Constrain k_{inact} , K_1 , and k_{sub} to a shared value that must be greater than 0 for all datasets and provide initial values that are in the anticipated range. Note that **Equation III** is in agreement with **equation C.16** in Appendix C of the original publication,⁸⁴ but $[I]_0$ and k_{inact} were unintentionally displaced in **equation 3** in the main text of this publication. Calculate inactivation constant K_1 and irreversible covalent inhibitor potency k_{inact}/K_1 with *Sample Calculations 1* and 2.

$$\begin{split} F_t &= F_0 + r_p \left[S \right]_0 \left\{ 1 - e^{-\beta \left(1 - e^{-\alpha t} \right)} \\ \alpha &= \frac{k_{\text{inact}} \left[I \right]}{K_1^{\text{app}} + \left[I \right]} \\ \beta &= \left(\frac{\left[E \right]_0 k_{\text{sub}}}{k_{\text{inact}}} \right) \left(\frac{K_1^{\text{app}}}{\left[I \right]} \right) \end{split}$$

(III)

Nonlinear regression of user-defined explicit equation:

a=kinact*I0/(I0+KIapp)
b=(E0*ksub/kinact)*(KIapp/I0)
P=S0*(1-exp(-b*(1-exp(-a*X))))
Y=Y0+(rp*P)

with Y = time-dependent signal F_t (in AU), X = incubation time t (in s), rp = product coefficient r_p (AU/M product, constrained), E0 = maximum unbound enzyme concentration at reaction initiation [E]₀ (in M, constrained), S0 = maximum unbound substrate concentration at reaction initiation [S]₀ (in M, constrained) and I0 = maximum unbound inhibitor concentration [I] (in M, column value) to find globally shared values for ksub = product formation rate constant $K_{sub} = k_{cat}/K_{M}$ (in M⁻¹s⁻¹, shared), kinact = maximum rate of inactivation k_{imact} (in s⁻¹, shared) and Klapp = apparent inactivation constant K_{a}^{avp} (in M, shared).

4. Optional: Validate experimental kinetic parameters with kinetic simulations

Proceed to *Kinetic Simulations* 1 to compare the experimental progress curves to the progress curves simulated with script **KinSubDpl** (using experimental rate constant $k_{inact} = k_s$) to confirm that the calculated kinetic constants are in accordance with the experimental data.

Assay Protocol II. Incubation Time-Dependent Potency IC₅₀(t)

The below protocol provides a generic set of steps to accomplishing this type of measurement.

Materials

- 1× Assay/reaction buffer supplemented with co-factors and reducing agent
- Active enzyme, 4× solution in assay buffer
- Competitive substrate with continuous or quenched read-out, 4× solution in assay buffer
- Positive control: vehicle/solvent as DMSO stock, or 2% solution in assay buffer
- Negative control: known inhibitor or alkylating agent as DMSO stock, or 2× solution in assay buffer
- Inhibitor: as DMSO stock, or serial dilution of 2× solution in assay buffer with 2% DMSO
- Optional: Development/quenching solution
- 384-well low volume microplate with nonbinding surface (e.g. Corning 3820 or 4513) for incubation/read-out
- Optical clear cover/seal (e.g. Perkin Elmer TopSeal-A Plus, #6050185, Corning 6575 Universal Optical Sealing Tape or Duck Brand HP260 Packing Tape) for <u>continuous</u> read-out, or a general microplate cover/lid (e.g. Corning 6569 Microplate Aluminum Sealing Tape) for <u>noncontinuous</u> read-out
- 1.5 mL (Eppendorf) microtubes to prepare stock solutions
- Optional: 96-well microplate to prepare serial dilution of inhibitor concentration
- Optional: Microtubes to perform incubations (e.g. Eppendorf Protein Lobind Microtubes, #022431018)
- Microplate reader equipped with appropriate filters to detect product formation (e.g. CLARIOstar microplate reader)
- Optional: Automated (acoustic) dispenser (e.g. Labcyte ECHO 550 Liquid Handler acoustic dispenser)

Exemplary assay concentrations

	incubation t		
	[stock]	Volume	[conc] _t
Enzyme	4 nM	5 μL	0.99 nM
Inhibitor	20 nM	10.2 μL	10.10 nM
Substrate	4 µM	5 μL	0.99 μM
Total		20.2 μL	

Before you start, optimize assay conditions in the uninhibited control to ensure compliance with assumptions and restrictions as outlined for Assay Protocol I (see also Figure 13). It is crucial to ensure that uninhibited product formation is linear with incubation time for the duration of the measurement: no enzyme degradation ($k_{deg} = 0$) or other factors contributing to a nonlinearity in product formation in the uninhibited control ($k_{ctri} = 0$) are allowed, as correction for nonlinearity is not possible in Data Analysis Protocol 2. This method is compatible with homogeneous (continuous) assays but also with assays that require a development/quenching step to visualize formed product.

 Add inhibitor or control (e.g. 0.2 μL) and assay buffer (e.g. 10 μL) to each well with the uninhibited control for full enzyme activity containing the same volume of vehicle/solvent instead of inhibitor as outlined in step 1 of Assay Protocol I.

Typically, measurements are performed in triplicate (or more replicates) with at least 8 inhibitor concentrations spanning the $IC_{50}(t)$. Inhibitor concentrations might need optimization, but a good starting point is $[I] = 0.1-5 \times IC_{50}(t)$ at the shortest incubation time *t*. Alternatively, larger-volume incubations can be performed in (Eppendorf) Protein Lobind microtubes, from which aliquots are transferred to a microplate after the indicated incubation time. Whether incubation in tube or plate is performed is a matter of personal preference, compatibility with lab equipment and automation, and convenience of dispensing small volumes.

 Add substrate in assay buffer to each well (e.g. 5 μL of 4× solution) and homogenize the solutions by gentle shaking (1 min at 300 rpm).

The order of substrate or inhibitor addition is not important *per se*, as long as DMSO stocks are added prior to buffered (aqueous) solutions and the enzyme is the last reagent to be added, to avoid unintentional preincubation. Inhibitor binding mode must be competitive with substrate. Optionally, gently centrifuge the plate or microtubes (1 min at 1000 rpm) to ensure assay components are not stuck at the top of the well.

3. Add active enzyme in assay buffer to each well (e.g. 5 μL of 4× solution) or tube as outlined in step 3 of Assay Protocol I.

The accuracy of the measurement improves if the incubation time is monitored precisely.

- 4. Seal the wells by applying an (optical clear) cover or lid, or close the caps of microtubes to prevent evaporation of assay components during incubation.
- Optional: Transfer aliquots (e.g. 20 µL) from the reaction mixture to the microplate after each time point; if incubation is performed in large volumes (in Protein Lobind microtubes or 96-well NBS plate) rather than incubation of replicates in a 384-well microplate.
- 6. *Quenching:* Add development solution to the reaction mixture in the microplate to quench the product formation reaction for assay formats that require a development/quenching step to visualize formed product.

Incubation time *t* is the elapsed time between reaction initiation by enzyme addition (*step 3*) and (optional) quenching of the enzyme activity by addition of development/quenching solution (*step 6*).

Measure formed product after incubation by detection of the product read-out in microplate reader.

Follow manufacturer advice on waiting time after addition of development solution before read-out. A typical assay measurement window is >2 hours, measuring cumulative product formation every 5-30 min (Figure 11B). The best results are obtained when inhibitor concentrations cover at least 50% of the DRC at all incubation times and there is a significant decrease from the earliest to the last $IC_{so}(t)$ value (Figure 12D).

8. Proceed to Data Analysis Protocol 2 to calculate relevant kinetic parameters for 2-step irreversible covalent inhibition.

	Data Analysis Protocol			
EXP Conditions	2-step IRREV	1-step IRREV	2-step REV	
k _{ctrl} = 0	2	-	-	

Data Analysis Protocol 2 for 2-Step IRREV Inhibition

Processing of raw data obtained with Assay Protocol I or Assay Protocol II for 2-step irreversible covalent inhibitors.

1. Plot signal F against incubation time t

Plot cumulative signal (in AU) on the Y-axis against incubation time (in s) on the X-axis for each inhibitor concentration and for the controls (**Figure 12B**). Label the columns with the inhibitor concentration (in M). It is not possible to algebraically correct for spontaneous loss of enzyme activity. Validate that the product formation in the uninhibited control F^{ctrl} is linear ($v_i = v_s$) by performing *steps 1-3* of *Data Analysis Protocol 1A* with $k_{obs} = k_{ctrl}$. Consult **Table 6** (*section 5*) for troubleshooting of nonlinearity of the uninhibited control.

2. Perform background correction

Correct for assay artifacts such as fluorescence bleaching and drift that cause a declining signal in the fully inhibited control. This correction can be subtraction of the time-dependent background in absence of enzyme but in presence of substrate (and inhibitor), or subtraction of the fully inhibited control. Consult the guidelines of your data fitting software for instructions on background corrections (e.g. GraphPad Prism).⁸⁸

3. Transpose to plot signal F against inhibitor concentration [I]

For each incubation time, transpose the X and Y values to plot signal F_t (in AU) on the Y-axis against inhibitor concentration (in M) on the X-axis. Also include product formation in the uninhibited control F^{ctri} ([1] = 0).

4. Normalize F_t/F^{ctrl}

Normalize F_t to lowest value = 0 and highest value = uninhibited product formation F^{ctrl} to obtain fractional product formation in presence of inhibitor F_t/F^{ctrl} . Consult the guidelines of your data fitting software for instructions on data normalization to the positive and negative controls.⁸⁸

5. Plot and fit F_t/F^{ctrl} against [I] to obtain the incubation time-dependent potency IC₅₀(t)

Plot the dose-response curve of fractional signal F_t/F^{ctrl} against inhibitor concentration (in M), and fit to four-parameter nonlinear regression Hill **Equation XII** to obtain the incubation time-dependent potency IC₅₀(t) (**Figure 12C**).³³ Use the inhibitor concentration <u>during</u> incubation: after reaction initiation by enzyme addition but before the (optional) addition of development solution (*Assay Protocol II, step 3*).

$$\frac{F_t}{F^{\text{ctrl}}} = \frac{1}{1 + \left(\frac{\text{IC}_{50}(t)}{[\text{I}]}\right)^h} \tag{XII}$$

Nonlinear regression of four-parameter dose-response equation $Y=Bottom+(Top-Bottom)/(1+(IC50/X)^HillSlope)$ with $Y = fractional product signal <math>F_c/F^{ctrl}$ (unitless), $X = inhibitor concentration [I] (in M), Bottom = normalized fully inhibited product signal = 0 (unitless, constrained), and Top = normalized uninhibited product signal <math>F_c^{trl}/F_c^{ctrl} = 1$ (unitless) and IC50 = incubation time-dependent potency ($C_{S0}(t)$ (in M).

6. Plot and fit $IC_{50}(t)$ against t to obtain k_{inact} and K_{I}

Plot the mean and standard deviation of $IC_{50}(t)$ (in M) on the Y-axis against incubation time t (in s) on the X-axis (Figure 12D). The rate of covalent bond formation at saturating inhibitor concentration k_{inact} and inactivation constant K_I are obtained by solving implicit Equation IV (shown in Figure 12E).⁸¹ Use the substrate concentration <u>during</u> incubation (*Assay Protocol II, step 3*): after reaction initiation by enzyme addition but before (optional) addition of development/ quenching solution. It is important that the Michaelis constant K_M be accurate for the reaction conditions (buffer, temperature, substrate), as this value is directly used to correct inactivation constant K_I for substrate competition. Consult the guidelines of your data-fitting software (e.g. GraphPad Prism)⁶⁵ for instructions on solving implicit equations (where Y appears on both sides of the equal sign). Proceed to *Sample Calculation 2* to calculate irreversible covalent inhibitor potency k_{inact}/K_I with propagation of error.

$$IC_{50}(t) = K_{I}\left(1 + \frac{[S]_{0}}{K_{M}}\right)\left(\frac{2 - 2e^{-\eta k_{inact}t}}{\eta k_{inact}t} - 1\right) with \eta = \frac{IC_{50}(t)}{K_{I}\left(1 + \frac{[S]_{0}}{K_{M}}\right) + IC_{50}(t)}$$
(IV)

Nonlinear regression of user-defined implicit equation: Y = (KI*(1+(S/KM)))*(((2-(2*EXP(-(Y/((KI*(1+(S/KM)))+Y))*kinact*X)))/((Y/((KI*(1+(S/KM)))+Y))*kinact*X))-1), with Y = incubation time-dependent potency [C₂₆(!) (in M), X = incubation time t (in s), S = maximum unbound substrate concentration at reaction initiation [S]₀ (in M, constrained), and KM = Michaelis constant K_M (in M, constrained) to find kinact = inactivation rate constant K_M (in S⁻¹) and KI = inactivation constant K₁ (in M).

7. Optional: Validate experimental kinetic parameters with kinetic simulations

Proceed to *Kinetic Simulations 1* to compare the experimental read-out to the product formation simulated with scripts **KinGen** and **KinDeg** (using experimental rate constant $k_{inact} = k_5$) to confirm that the calculated kinetic constants are in accordance with the experimental data and found $IC_{sn}(t)$ values.

Assay Protocol III. Preincubation Time-Dependent Inhibition Without Dilution

The protocol below provides a generic set of steps to accomplishing this type of measurement.

Materials

- 1× Assay/reaction buffer supplemented with co-factors and reducing agent
- Active enzyme, 2× solution in assay buffer
- Substrate with continuous or quenched read-out, 11× solution in assay buffer
- Positive control: vehicle/solvent as DMSO stock, or 2% solution in assay buffer
- Negative control: known inhibitor or alkylating agent as DMSO stock, or 2× solution in assay buffer
- Inhibitor: as DMSO stock, or serial dilution of 2× solution in assay buffer with 2% DMSO
- Optional: Development/quenching solution
- 1.5 mL (Eppendorf) microtubes to prepare stock solutions
- 384-well low volume microplate with nonbinding surface (e.g. Corning 3820 or 4513) for preincubation/read-out
- General microplate cover/lid (e.g. Corning 6569 Microplate Aluminum Sealing Tape) if preincubation is conducted in a microplate
- Optional: 96-well microplate to prepare serial dilution of inhibitor concentration
- Optional: Microtubes to perform preincubations (e.g. Eppendorf Protein Lobind Microtubes, #022431018)
- Microplate reader equipped with appropriate filters to detect product formation (e.g. CLARIOstar microplate reader)
- Optional: Automated (acoustic) dispenser (e.g. Labcyte ECHO 550 Liquid Handler acoustic dispenser)

Exemplary assay concentrations

	preincubation t'		incubation t			
	[stock]	Volume	[conc] _{t'}	[stock]	Volume	[conc] _t
Enzyme	2 nM	10 µL	0.99 nM	-	-	0.90 nM
Inhibitor	20 nM	10.2 μL	10.10 nM	-	-	9.19 nM
Substrate	-	-	-	11 µM	2 µL	0.99 μM
Total		20.2 μL			22.2 μL	

Before you start, optimize assay conditions in the uninhibited control to ensure compliance with assumptions and restrictions, as outlined in section 2.3 and in Assay Protocol I. Consult **Table 6** (section 5) for common optimization and troubleshooting options. Specific adjustments for Method III are that substrate concentration should be relatively low $([S]_0 << K_M)$ to minimize disruption of the noncovalent $E+1 \leftrightarrow EI$ equilibrium or reduction of reaction rates by competition (illustrated in **Figure 14A**); adjustment of the enzyme concentration might be required to ensure that maximum 10% of the substrate is processed during the read-out $([P]_t < 0.1[S]_0)$ and product formation is linear in the uninhibited control. Furthermore, incubation time **t** must be relatively short to minimize additional time-dependent enzyme inhibition after substrate addition. As a rule of thumb, incubation must be much shorter than the shortest preincubation (t << t'), unless the product formation read-out is continuous (more details in Data Analysis Protocol 3, step 3). Validate that enough product is formed for a good signal/noise ratio (Z' > 0.5) by calculating the Z'-score from the uninhibited and inhibited controls (ideally 8 replicates) in a separate experiment.⁸⁷ This method is compatible with homogeneous (continuous) assays but also with assays that require a development/quenching step to visualize formed product. Note that this protocol was designed for preincubation and read-out in a 384-well microplate.

 Add inhibitor or control (e.g. 0.2 μL) and assay buffer (e.g. 10 μL) to each well with the uninhibited control for full enzyme activity containing the same volume vehicle/solvent instead of inhibitor as outlined in step 1 of Assay Protocol I.

Gently shake to mix DMSO with the aqueous buffer. Typically, measurements are performed in triplicate (or more replicates) with at least 8 inhibitor concentrations for at least 5 preincubation times. Inhibitor concentrations might need optimization, but a rational starting point is to use inhibitor concentrations below $5 \times IC_{50}$ at the shortest preincubation time t': inhibition is expected to improve in a time-dependent manner and the best results are obtained when full inhibition is not achieved already at the shortest preincubation time. Alternatively, larger-volume preincubations (e.g. >200 µL) can be performed in (Eppendorf) microtubes from which aliquots (e.g. 20.2μ L) are transferred to a microplate after the indicated preincubation time. Whether preincubation is performed in a tube or microplate is a matter of personal preference, compatibility with lab equipment and automation, and convenience of dispensing small volumes.

 Add active enzyme in assay buffer to each well (e.g. 10 μL of 2× solution) or tube to start preincubation of enzyme with inhibitor and homogenize the solution by gently shaking (1 min at 300 rpm). Alternatively, dispensing the enzyme at a high flow rate will also mix the components.

The order of enzyme and inhibitor addition is not important *per se*, as long as DMSO stocks are added prior to buffered (aqueous) solutions. Inhibitor must be present in excess during preincubation $([I]_0 > 10[E]_0)$. Optionally, gently centrifuge the plate or microtubes (1 min at 1000 rpm) to ensure assay components are not stuck at the top of the well.

- Seal the wells with a cover or lid, and close the caps of microtubes to prevent evaporation of assay components during preincubation.
- Optional: Transfer aliquots (e.g. 20.2 μL) from the reaction mixture to the microplate after completion of preincubation if performed in larger volumes.
- Add substrate in assay buffer (e.g. 2 µL of 11× solution) to (at least) three replicates after preincubation time t'.

Typically, preincubation can run anywhere from several minutes to hours depending on the enzyme stability and anticipated inhibitor potency, with superior accuracy if the preincubation time is monitored precisely. Substrate should be added in a negligible volume ($V_{sub} < 0.1V_t$) to minimize disruption of the noncovalent equilibria by dilution ($V_t = V_t$.) (Figure 14A). Because at steady-state the equilibrium can be disrupted by dilution in too much competitive substrate, keep the substrate volume V_{sub} and substrate concentration low ($[S]_0 < 0.1K_M$) for successful analysis of 2-step reversible inhibitors (*Data Analysis 3C*). Optionally, homogenize the solutions by gentle shaking (300 rpm) and centrifuge the plate or microtubes (1 min at 1000 rpm) to ensure that assay components are not stuck at the top of the well.

 Quenching: Add development solution to the reaction mixture in the microplate to quench the product formation reaction if read-out of product formation requires a development/quenching step to visualize formed product after incubation time t.

Follow manufacturer advice on waiting time after addition of development solution before read-out. Incubation time t is the elapsed time between onset of product formation by substrate addition (*step 5*) and addition of development/ quenching solution (*step 6*). A possible advantage to the use of a quenched assay is the possibility to store the samples after addition of quenching/development solution (*step 6*) and measure product formation (*step 7*) in all samples after completion of the final preincubation rather than performing multiple separate measurements (after each preincubation time).

7. Measure formed product after incubation by detection of the product read-out in microplate reader.

Incubation time (after substrate addition) is relatively short ($t \ll LN(2)/k_{obs}$) to minimize additional (time-dependent) inhibition of enzyme activity during incubation (illustrated in **Figure 14B**).

8. Repeat steps 4-7 of Assay Protocol III for at least another four preincubation times.

Preincubation time t' is the elapsed time between onset of inhibition by mixing enzyme and inhibitor (*step 2*) and addition of substrate (*step 5*). A typical preincubation assay consists of multiple hours of measuring enzyme activity every 5-30 min, depending on enzyme stability and inhibitor reaction rates. Best results are obtained if the incubation time t used to calculate enzyme activity is kept constant at all preincubation times.

9. Proceed to *Data Analysis Protocol 3* to convert the raw experimental data into preincubation time-dependent enzyme activity.

Data Analysis Protocol 3 for all binding modes

Processing of raw experimental data obtained with Assay Protocol III.

1. Plot signal F against incubation time t

Plot signal F (in AU) on the Y-axis against the incubation time (in s) on the X-axis for each inhibitor concentration and for the controls (Figure 14B). Do this separately for each preincubation time. Proceed to step 3 of this protocol for continuous read-out assays that require a longer incubation time to produce enough product for a good signal/noise ratio.

2. Fit F_t against t to obtain $v_{t'}$

Fit signal *F* against incubation time *t* to **Equation XIII** (Figure 15B/Figure 17B, *left*) to obtain preincubation timedependent product formation velocity v_{t} from the linear slope. Consult **Table 6** (*section 5*) for troubleshooting if product formation is not linear.

$F_t = F_0 + v_{t'} t$

(XIII)

Nonlinear regression of straight line **Y=YIntercept+Slope*X** with Y = signal F_t (in AU) and X = incubation time t (in s) to find YIntercept = background signal at reaction initiation F_0 (in AU) and Slope = preincubation time-dependent product formation velocity v_t (in AU/s).

3. Alternative for continuous: Fit F_t against t to obtain $v_{t'}$

This is an alternative method to obtain v_t from the initial velocity for assays with a continuous read-out, using the initial velocity in progress curve analysis (*Method I*). Fit signal F_t against incubation time t to exponential association **Equation XIV** (Figure 15B/Figure 17B, *right*) to obtain preincubation time-dependent product formation velocity v_t from the initial velocity. This resolves issues with low signal/noise ratios for continuous read-out assays where v_t is not linear (due to additional covalent modification during the incubation) by allowing longer incubation times to produce sufficient signal.

$$F_t = v_s t + \frac{v_t - v_s}{k} \left[1 - e^{-kt}\right] + F_0$$
(XIV)

Nonlinear regression of user-defined explicit equation Y = (vs*X) + ((vi-vs)/kobs)*(1-EXP(-kobs*X))) + Y0 with $Y = signal F_t$ (in AU) and X = incubation time t (in s) to find Y0 = Y-intercept F_0 = background signal at t = 0 (in AU), $v = initial slope = preincubation time-dependent product formation velocity <math>v_t$ (in AU/s), $v = final slope <math>v_s$ (in AU/s) and kobs = nonlinearity reaction rate k (in s^{-1}).

4. Proceed to Data Analysis Protocols to obtain the appropriate kinetic parameters for each covalent binding mode: Data Analysis Protocol 3Ai or 3Aii for 2-step irreversible inhibitors, Data Analysis Protocol 3Bi or 3Bii for 1-step irreversible inhibitors, and Data Analysis Protocol 3C for 2-step reversible inhibitors.

Selection of a data analysis method for inhibitors with an irreversible binding mode depends on the desired visual representation as well as personal preference. Generally, *Data Analysis Protocols 3Ai* and *3Bi* have less data processing/ manipulation and are more informative for comparison of various inhibitors on a single enzyme target, as they are compatible with assessment of inhibitor potency simultaneous with visual assessment of time-dependent enzyme stability k_{ctrl} (**Figure 16B** and **Figure 18B**). *Alternative Data Analysis Protocols 3Aii* and *3Bii* involve normalization of the enzyme activity that aids visual assessment of inhibitory potency of a single inhibitor on multiple enzyme targets (that might have a variable stability) (**Figure 16D** and **Figure 18D**).

	Data Analysis Protocol					
EXP Conditions	2-step IRREV 1-step IRREV 2-step REV					
$k_{\rm ctrl} = 0$	3Ai	3B	3C			
$k_{\text{degE}} > 0$	3Ai or 3Aii	3Bi or 3Bii	3C			

Data Analysis Protocol 3Ai for 2-Step IRREV Inhibition

Processing of experimental data obtained with *Assay Protocol III* that has been processed according to *Data Analysis Protocol 3* for 2-step irreversible covalent inhibitors.

1. Plot v_t against preincubation time t' for each inhibitor concentration

Plot the mean and standard deviation of $v_{t'}$ (in AU/s) on the Y-axis against preincubation time t' (in s) on the X-axis for each inhibitor concentration and the uninhibited control (Figure 15C). Validate that inhibitor concentrations are not too high: inhibition should be less than 100% at the shortest t' for at least six inhibitor concentrations. Check whether the uninhibited enzyme activity is independent of preincubation time ($v_0^{ctrl} = v_t^{ctrl}$, Figure 15C): an algebraic correction for enzyme instability ($k_{ctrl} > 0$, Figure 16B) can be performed in step 4 of this protocol by accounting for nonlinearity in the uninhibited control in the secondary k_{obs} plot (Figure 16C). Alternatively, proceed to Alternative Data Analysis Protocol 3Aii to correct for enzyme instability ($v_0^{ctrl} > v_t^{ctrl}$) by normalization of the enzyme activity v_t/v_t^{ctrl} (Figure 16D-E).

2. Fit $v_{t'}$ against preincubation time t' to obtain k_{obs}

Fit the mean and standard deviation of v_t against preincubation time t' (Figure 15C/Figure 16B) to Equation v. Constrain v_s = value in fully inhibited control to obtain the observed reaction rate k_{obs} from initial velocity v_i (Y-intercept) to full inactivation (Plateau = 0). A lack of initial noncovalent complex ($v_i = v_0^{ctrl}$) is indicative of 1-step binding behavior.

$$v_{t'} = v_s + (v_i - v_s) e^{-k_{obs}t'}$$

(V)

Nonlinear regression of exponential one-phase decay equation Y = (YO-Plateau) * EXP(-k*X) + Plateau with Y = preincubation timedependent product formation velocity v_t (in AU/s), X = preincubation time t' (in s), and Plateau = final velocity in the fully inhibited control v_s (in AU/s, constrained) to find YO = Y-intercept = initial velocity v_t (in AU/s) and k = observed reaction rate k_{obs} (in s^{-1}).

3. Plot k_{obs} against [I]

Plot the mean and standard deviation of k_{obs} (in s^{-1}) on the Y-axis against inhibitor concentration (in M) <u>during</u> <u>preincubation</u> (*before* addition of substrate) on the X-axis (**Figure 15D**/**Figure 16C**). The plot of k_{obs} against [I] should reach a maximum k_{obs} at saturating inhibitor concentration. Note that a linear curve is indicative of 1-step binding behavior at non-saturating inhibitor concentrations ([I] << 0.1K₁^{app}: see **Figure 3F**) with $v_i = v_0^{ctri}$ (shared Y-intercept in the previous step). Proceed to *step 4* of *Data Analysis Protocol 3Bi* after it has been validated that the linear curve is not resultant from saturating inhibitor concentrations ([I] >> 10K₁^{app}: see **Figure 3G**) as identified by $v_i << v_0^{ctri}$, by repeating the measurement with lower inhibitor concentrations.

4. Fit k_{obs} against [I] to obtain k_{inact} and K_{I}

Fit k_{obs} against inhibitor concentration <u>during preincubation</u> to **Equation xv** to obtain maximum inactivation rate constant k_{inact} and inactivation constant K_1 . Constrain $k_{ctrl} = k_{obs}$ of the uninhibited control (**Figure 16C**). Inactivation constant K_1 does not have to be corrected for substrate competition because preincubation is conducted in absence of competing substrate. Calculate irreversible covalent inhibitor potency k_{inact}/K_1 with propagation of error with *Sample Calculation 2*.

$$k_{\rm obs} = k_{\rm ctrl} + \frac{k_{\rm inact} \left[l \right]}{K_{\rm I} + \left[l \right]} \tag{xv}$$

Nonlinear regression of user-defined explicit equation Y=Y0+((kmax*X)/((KI)+X)) with Y = observed reaction rate k_{obs} (in s⁻¹), X = inhibitor concentration during preincubation (in M), and Y0 = rate of nonlinearity in uninhibited control k_{ctrl} (in s⁻¹, constrained) to find kmax = maximum reaction rate k_{inact} (in s⁻¹) and KI = Inactivation constant K₁ (in M).

5. Optional: Validate experimental kinetic parameters with kinetic simulations

Proceed to *Kinetic Simulations 1* to compare the experimental read-out to the product formation simulated with scripts **KinGen** and **KinDeg** (using experimental rate constant $k_{inact} = k_5$) to confirm that the calculated kinetic constants are in accordance with the experimental data. Also perform simulations with **KinVol** and **KinVolDeg** to confirm that addition of substrate does not significantly affect the noncovalent interactions.

Alternative Data Analysis Protocol 3Aii for 2-Step IRREV Inhibition

Processing of experimental data obtained with *Assay Protocol III* that has been processed according to *Data Analysis Protocol 3* for 2-step irreversible covalent inhibitors.

1. Plot $v_{t'}$ against preincubation time t' for each inhibitor concentration

Plot the mean and standard deviation of $v_{t'}$ (in AU/s) on the Y-axis against preincubation time t' (in s) on the X-axis for each inhibitor concentration and the uninhibited control (**Figure 15C/Figure 16B**). Validate that inhibitor concentrations are not too high: inhibition should be less than 100% at the shortest t' for at least six inhibitor concentrations.

2. Normalize $v_{t'}$ to obtain $v_{t'}/v^{ctrl}$

Normalize v_t of each inhibitor concentration and the controls to lowest value = 0 (or full inhibition control) and highest value = uninhibited product formation v_t^{ctrl} to obtain normalized enzyme activity v_t/v^{ctrl} . Perform this correction separately for each preincubation time.

3. Plot and fit $v_{t'}/v^{ctrl}$ against preincubation time t' to obtain k_{obs}

Plot the mean and standard deviation of $v_{t'}/v^{ctrl}$ on the Y-axis against preincubation time t' (in s) on the X-axis (**Figure 16D**). Fit to exponential decay **Equation XVI** to obtain k_{obs} from initial velocity v_i/v_0^{ctrl} to full inactivation (Plateau = 0). A lack of initial noncovalent complex ($v_i/v_0^{ctrl} = 1$) is indicative of 1-step binding behavior.

$$\left(\frac{v_{t'}}{v_{t'}^{ctrl}}\right) = \left(\frac{v_i}{v_0^{ctrl}}\right) e^{-k_{obs}t'} \tag{XVI}$$

Nonlinear regression of exponential one-phase decay equation Y=(YO-Plateau)*EXP(-k*X)+Plateau with Y = normalized preincubation time-dependent product formation velocity v_c/v^{crt} (unitless), X = preincubation time t' (in s), and Plateau = normalized final velocity $v_s/v_s^{crt} = 0$ (unitless, constrained) to find Y0 = Y-intercept = normalized initial velocity v_i/v_0^{crt} (unitless) and k = observed reaction rate k_{obs} (in s⁻¹).

Plot k_{obs} against [I]

Plot the mean and standard deviation of k_{obs} (in s⁻¹) on the Y-axis against inhibitor concentration (in M) <u>during</u> <u>preincubation</u> (*before* addition of substrate) on the X-axis (**Figure 16E**). The plot of k_{obs} against [I] should have a Y-intercept = 0 and reach a maximum k_{obs} at saturating inhibitor concentration. Note that a linear curve is indicative of 1-step binding behavior at non-saturating inhibitor concentrations (II] << 0.1K₁^{app}: see **Figure 3F**) with $v_1 = v_0^{ctrl}$ (shared Y-intercept = 1 in the previous step). Proceed to *step 5* of *Data Analysis Protocol 3Bii* after it has been validated that the linear curve is not resultant from saturating inhibitor concentrations (II) >> 10K₁^{app}: see **Figure 3G**) as identified by $v_1 < v_0^{ctrl}$ (shared Y-intercept = 0 in the previous step), by repeating the measurement with lower inhibitor concentrations.

5. Fit k_{obs} against [I] to obtain k_{inact} and K_{I}

Fit k_{obs} against inhibitor concentration <u>during preincubation</u> to **Equation XVII** to obtain maximum inactivation rate constant k_{inact} and inactivation constant K_1 (**Figure 16E**). Do <u>not</u> correct for enzyme instability ($k_{ctrl} > 0$), as this correction has already been performed by normalizing v_t to v_t/v^{ctrl} in *step 2* of this protocol. Inactivation constant K_1 does not have to be corrected for substrate competition because preincubation is conducted in absence of competing substrate. Calculate irreversible covalent inhibitor potency k_{inart}/K_1 with propagation of error with *Sample Calculation 2*.

$$k_{\rm obs} = \frac{k_{\rm inact}[I]}{K_{\rm I} + [I]} \tag{XVII}$$

Nonlinear regression of user-defined explicit equation Y=Y0+((kmax*X)/((KI)+X)) with Y = observed reaction rate k_{obs} (in s⁻¹), X = inhibitor concentration during preincubation (in M), and Y0 = 0 (in s⁻¹, constrained) to find kmax = maximum reaction rate k_{inact} (in s⁻¹) and KI = Inactivation constant K₁ (in M).

6. Optional: Validate experimental kinetic parameters with kinetic simulations

Proceed to *Kinetic Simulations 1* to compare the experimental read-out to the product formation simulated with scripts **KinGen** and **KinDeg** (using experimental rate constant $k_{inact} = k_s$) to confirm that the calculated kinetic constants are in accordance with the experimental data. Also perform simulations with **KinVol** and **KinVolDeg** to confirm that addition of substrate does not significantly affect the noncovalent interactions.

Data Analysis Protocol 3Bi for 1-step IRREV Inhibition

Processing of experimental data obtained with *Assay Protocol III* that has been processed according to *Data Analysis Protocol 3* for 1-step irreversible covalent inhibitors and 2-step irreversible inhibitors at non-saturating inhibitor concentrations ([I] $\leq 0.1K_1$).

1. Plot $v_{t'}$ against preincubation time t' for each inhibitor concentration

Plot the mean and standard deviation of $v_{t'}$ (in AU/s) on the Y-axis against preincubation time t' (in s) on the X-axis for each inhibitor concentration and the uninhibited control (Figure 17C). Validate that inhibitor concentrations are not too high: inhibition should be less than 100% at the shortest t' for at least six inhibitor concentrations. Check whether the uninhibited enzyme activity is independent of preincubation time ($v_0^{\text{ctrl}} = v_t^{\text{ctrl}}$, Figure 17C): an algebraic correction for enzyme instability ($k_{\text{ctrl}} > 0$, Figure 18B) can be performed in step 4 of this protocol by accounting for nonlinearity in the uninhibited control in the secondary k_{obs} plot (Figure 18C). Alternatively, proceed to Alternative Data Analysis Protocol 3Bii to correct for enzyme instability ($v_0^{\text{ctrl}} > v_t^{\text{ctrl}}$) by normalization of the enzyme activity v_t/v_t^{ctrl} (Figure 18D-E).

2. Fit $v_{t'}$ against preincubation time t' to obtain k_{obs}

Fit the mean and standard deviation of v_t against preincubation time t' (Figure 17C/Figure 18B) to Equation v. Constrain v_s = value in fully inhibited control to obtain the observed reaction rate k_{obs} from initial velocity v_i (Y-intercept) to full inactivation (Plateau = 0). A lack of initial noncovalent complex ($v_i = v_0^{ctrl}$) is indicative of 1-step binding behavior.

$$v_{t'} = v_s + (v_i - v_s) e^{-k_{obs}t}$$

(V)

Nonlinear regression of exponential one-phase decay equation Y = (YO-Plateau) *EXP(-k*X)+Plateau with Y = preincubation time $dependent product formation velocity <math>v_i$. (in AU/s), X = preincubation time t' (in s), and Plateau = final velocity in the fully inhibited control v_s (in AU/s, constrained) to find YO = Y-intercept = initial velocity v_i = uninhibited initial velocity v_0^{ctrl} (in AU/s, shared value), and k = observed reaction rate k_{obs} (in s^{-1}).

3. Plot k_{obs} against [I]

Plot the mean and standard deviation of k_{obs} (in s⁻¹) on the Y-axis against inhibitor concentration (in M) <u>during</u> <u>preincubation</u> (*before* addition of substrate) on the X-axis (**Figure 17D/Figure 18C**). The plot of k_{obs} against inhibitor concentration [I] is linear for 1-step irreversible inhibitors and for 2-step irreversible inhibitors at non-saturating inhibitor concentrations ([I] << 0.1K₁).

4. Fit k_{obs} against [I] to obtain k_{chem}

Fit k_{obs} against inhibitor concentration <u>during preincubation</u> to **Equation XVIII** to obtain inhibitor potency k_{chem} from the linear slope. Constrain Y-intercept $k_{ctrl} = k_{obs}$ of the uninhibited control (**Figure 18C**). Inhibitor potency k_{chem} does <u>not</u> have to be corrected for substrate competition because preincubation is conducted in absence of competing substrate. Calculate k_{inact}/K_1 for 2-step irreversible inhibitors at non-saturating inhibitor concentrations ([I] $\leq 0.1K_1$) with *Sample Calculation* 9.

$k_{\rm obs} = k_{\rm ctrl} + k_{\rm chem} [I]$

(XVIII)

Nonlinear regression of straight line Y=YIntercept+Slope*X with Y = observed reaction rate k_{obs} (in s^{-1}), X = inhibitor concentration during preincubation (in M) and YIntercept = rate of nonlinearity in uninhibited control k_{ctri} (in s^{-1} , constrained) to find Slope = inactivation rate constant k_{chem} (in $M^{-1}s^{-1}$).

5. Optional: Validate experimental kinetic parameters with kinetic simulations

Proceed to *Kinetic Simulations 1* to compare the experimental read-out to the product formation simulated with scripts **KinGen** and **KinDeg** (using experimental rate constant $k_{chem} = k_3$) to confirm that the calculated kinetic constants are in accordance with the experimental data. Also perform simulations with **KinVol** and **KinVolDeg** to confirm that addition of substrate does not significantly affect the reaction rates by dilution and/or competition.

Alternative Data Analysis Protocol 3Bii for 1-step IRREV Inhibition

Processing of experimental data obtained with *Assay Protocol III* that has been processed according to *Data Analysis Protocol 3* for 1-step irreversible covalent inhibitors and 2-step irreversible inhibitors at non-saturating inhibitor concentrations ($[I] \le 0.1K_1$).

1. Plot $v_{t'}$ against preincubation time t' for each inhibitor concentration

Plot the mean and standard deviation of v_t (in AU/s) on the Y-axis against preincubation time t' (in s) on the X-axis for each inhibitor concentration and the uninhibited control (Figure 17C/Figure 18B). Validate that inhibitor concentrations are not too high: inhibition should be less than 100% at the shortest t' for at least six inhibitor concentrations.

2. Normalize $v_{t'}$ to obtain $v_{t'}/v^{ctrl}$

Normalize v_t of each inhibitor concentration and the controls to lowest value = 0 (or full inhibition control) and highest value = uninhibited product formation v_t^{ctrl} to obtain normalized enzyme activity v_t/v^{ctrl} . Perform this correction separately for each preincubation time.

3. Plot and fit $v_{t'}/v^{ctrl}$ against preincubation time t' to obtain k_{obs}

Plot the mean and standard deviation of $v_{t'}/v^{ctrl}$ on the Y-axis against preincubation time t' (in s) on the X-axis (Figure 18D). Fit to exponential decay Equation XIX to obtain k_{obs} from initial velocity v_i/v_0^{ctrl} to full inactivation (Plateau = 0). A lack of initial noncovalent complex ($v_i/v_0^{ctrl} = 1$) is indicative of 1-step binding behavior.

$$\left(\frac{v_{t'}}{v_{t'}^{trl}}\right) = e^{-k_{obs}t'} \tag{XIX}$$

Nonlinear regression of exponential one-phase decay equation Y=(Y0-Plateau)*EXP(-k*X)+Plateau with Y = normalized preincubation time-dependent product formation velocity v_c/v^{ctri} (unitless), X = preincubation time t' (in s), Y0 = Y-intercept = normalized initial velocity $v_c/v^{ctri}_{cri} = 0$ (unitless, constrained), and Plateau = normalized final velocity $v_s/v^{ctri}_s = 0$ (unitless, constrained) to find k = observed reaction rate k_{bb} (in s⁻¹).

4. Plot k_{obs} against [I]

Plot the mean and standard deviation of k_{obs} (in s⁻¹) on the Y-axis against inhibitor concentration (in M) <u>during</u> <u>preincubation</u> (*before* addition of substrate) on the X-axis (**Figure 18E**). The plot of k_{obs} against inhibitor concentration [I] is linear for 1-step irreversible inhibitors and for 2-step irreversible inhibitors at non-saturating inhibitor concentrations ([I] << 0.1K₁).

5. Fit k_{obs} against [I] to obtain k_{chem}

Fit k_{obs} against inhibitor concentration <u>during preincubation</u> to **Equation xx** to obtain inhibitor potency k_{chem} from the linear slope (**Figure 18E**). Do <u>not</u> correct for enzyme instability ($k_{ctrl} > 0$), as this correction has already been performed by normalizing v_t : to v_t/v^{ctrl} in *step 2* of this protocol. Inhibitor potency k_{chem} does not have to be corrected for substrate competition because preincubation is conducted in absence of competing substrate. Calculate k_{inact}/K_1 for 2-step irreversible inhibitors at non-saturating inhibitor concentrations ([I] $\leq 0.1K_1$) with *Sample Calculation 9*. Alternatively, inhibitor potency k_{chem} or k_{inact}/K_1 can be directly calculated from a single k_{obs} and [I] with *Sample Calculation 10*.

$$k_{obs} = k_{chem}$$
 [I]

(XX)

Nonlinear regression of straight line Y=YIntercept+Slope*X with Y = observed reaction rate k_{obs} (in s^{-1}), X = inhibitor concentration during preincubation (in M), and YIntercept = 0 (in s^{-1} , constrained) to find Slope = inactivation rate constant k_{chem} (in $M^{-1}s^{-1}$).

6. Optional: Validate experimental kinetic parameters with kinetic simulations

Proceed to *Kinetic Simulations 1* to compare the experimental read-out to the product formation simulated with scripts **KinGen** and **KinDeg** (using experimental rate constant $k_{chem} = k_3$) to confirm that the calculated kinetic constants are in accordance with the experimental data. Also perform simulations with **KinVol** and **KinVolDeg** to confirm that addition of substrate does not significantly affect the reaction rates by dilution and/or competition.

Data Analysis Protocol 3C for 2-Step REV Inhibition

Processing of experimental data obtained with *Assay Protocol III* that has been processed according to *Data Analysis Protocol 3* for 2-step reversible covalent inhibitors.

1. Plot $v_{t'}$ against preincubation time t' for each inhibitor concentration

Plot the mean and standard deviation of v_t (in AU/s) on the Y-axis against preincubation time t' (in s) on the X-axis for each inhibitor concentration and the uninhibited control (Figure 19C/Figure 20B). Validate that inhibitor concentrations are not too high: inhibition should be less than 100% at the shortest t' for at least six inhibitor concentrations. Enzyme activity is never truly independent of preincubation time ($v_0^{ctrl} > v_t^{ctrl}$) and kinetic analysis of reversible inhibitors is very sensitive to small deviations (as illustrated in Figure 9). Therefore, correction for enzyme instability is always performed by normalization of the enzyme activity v_t/v_t^{ctrl} in the next step (illustrated in Figure 20).

2. Normalize $v_{t'}$ to obtain $v_{t'}/v^{ctrl}$

Normalize $v_{t'}$ of each inhibitor concentration and the controls to lowest value = 0 (or full inhibition control) and highest value = uninhibited product formation $v_{t'}^{ctrl}$ to obtain normalized enzyme activity $v_{t'}/v^{ctrl}$ (Figure 20C). Perform this correction <u>separately</u> for each preincubation time.

3. Plot and fit $v_{t'}/v^{ctrl}$ against preincubation time t' to obtain k_{obs} and v_s/v_s^{ctrl}

Plot the mean and standard deviation of v_t / v^{ctrl} on the Y-axis against preincubation time t' (in s) on the X-axis (**Figure 20C**). Fit to exponential decay **Equation XXI** to obtain k_{obs} from initial velocity v_i / v_0^{ctrl} reflecting rapid noncovalent equilibrium (Y-intercept $v_i / v_0^{\text{ctrl}} \le 1$) to the final velocity v_s / v_s^{ctrl} reflecting steady-state equilibrium (Plateau $v_s / v_s^{\text{ctrl}} \ge 0$).

$$\left(\frac{v_{t'}}{v_{t'}^{\text{ctrl}}}\right) = \left(\frac{v_{s}}{v_{s}^{\text{ctrl}}}\right) + \left(\frac{v_{i}}{v_{0}^{\text{ctrl}}} - \frac{v_{s}}{v_{s}^{\text{ctrl}}}\right)e^{-k_{\text{obs}}t'}$$
(XXI)

Nonlinear regression of exponential one-phase decay equation $\mathbf{Y}=(\mathbf{Y}\mathbf{0}-\mathbf{P}\mathbf{lateau})*\mathbf{EXP}(-\mathbf{k}*\mathbf{X})+\mathbf{P}\mathbf{lateau}$ with Y = normalized preincubation time-dependent product formation velocity v_t/v^{ctrl} (unitless) and X = preincubation time t^* (in s) to find $Y\mathbf{0} = Y$ -intercept = normalized initial velocity v_t/v_0^{ctrl} (unitless), Plateau = normalized final velocity v_s/v_s^{ctrl} (unitless), and k = observed reaction rate k_{obs} (in s⁻¹).

4. Plot and fit v_s/v_s^{ctrl} against [I] to obtain K_i^*

Steady-state inhibition constant K_i^* can be calculated from v_s/v_s^{ctrl} (obtained in the previous step) reflecting remaining fractional enzyme activity after reaching the steady-state inhibitor equilibrium (*reaction completion*) (Figure 20E). Plot the mean and standard deviation of v_s/v_s^{ctrl} on the Y-axis against inhibitor concentration (in M) <u>during preincubation</u> (*before* addition of substrate) on the X-axis (Figure 20E), and fit the dose-response curve to four-parameter nonlinear regression Hill Equation XXII to obtain steady-state inhibition constant $K_i^{*,33}$ The maximum product formation velocity at reaction completion corresponds with the uninhibited enzyme activity $v_s^{ctrl}/v_s^{ctrl} = 1$ and minimum velocity $v_s^{min}/v_s^{ctrl} = 0$ for (background-)corrected enzyme activity in the full inhibition control. Steady-state equilibrium constant K_i^* does <u>not</u> have to be corrected for substrate competition because preincubation is conducted in absence of competing substrate.

$$\left(\frac{v_s}{v_s^{\text{ctrl}}}\right) = 1 + \frac{1}{1 + \left(\frac{[I]}{K_i^*}\right)^h}$$
(XXII)

Nonlinear regression of four-parameter dose-response equation $Y=Bottom+(Top-Bottom)/(1+(X/IC50)^HillSlope)$ with Y = fractional steady-state product formation velocity v_s/v_s^{ctrl} (unitless), X = inhibitor concentration during preincubation (in M), Bottom = velocity in fully inhibited control $v_s^{imn}/v_s^{ctrl} = 0$ (unitless, constrained), and Top = uninhibited enzyme activity $v_s/v_s^{ctrl} = 1$ (unitless, constrained) to find Hillslope = Hill coefficient h (unitless) and IC50 = steady-state inhibition constant K⁺_s (in M).

5. Optional: Plot and fit k_{obs} against [I] to obtain K_i, k_5 , and k_6

This is an optional data processing step to obtain kinetic parameters by fitting to the observed rate k_{obs} (obtained in *step 3* of *Data Analysis Protocol 3C*), and can be used to validate K_i^* values found in the previous step or to find values for k_s and k_6 to use in kinetic simulations (next step in this protocol). Plot the mean and standard deviation of k_{obs} (in s^{-1}) on the Y-axis against inhibitor concentration <u>during preincubation</u> (in M) on the X-axis (**Figure 20D**). Exclude the uninhibited control ($k_{ctri} = 0$ for normalized enzyme activity) from the fit because Y-intercept = k_6 rather than k_{ctri} . Fit k_{obs} against inhibitor concentration to **Equation XXIII** to obtain rate constants for the covalent association k_5 and covalent dissociation k_6 as well as noncovalent inhibition constant K_i reflecting the rapid (initial) noncovalent equilibrium. Noncovalent equilibrium constant K_i does <u>not</u> have to be corrected for substrate competition because preincubation

(XXIII)

is conducted in absence of competing substrate. Proceed to Sample Calculation 8 to calculate steady-state inhibition constant K_i^* from experimental values of K_i , k_5 , and k_6 .

$$k_{\rm obs} = k_6 + \frac{k_5 \left[I \right]}{K_i + \left[I \right]}$$

Nonlinear regression of user-defined explicit equation $Y=Y0+((kmax^*X)/((Ki)+X))$ with Y = observed reaction rate k_{obs} (in s^{-1}) and X = inhibitor concentration during preincubation (in M) to find Y0 = covalent dissociation rate constant k_6 (in s^{-1}), kmax = covalent association rate constant k_5 (in s^{-1}), and Ki = inhibition constant K_i (in M).

6. Optional: Validate experimental kinetic parameters with kinetic simulations

Proceed to *Kinetic Simulations 1* to compare the experimental read-out to the product formation simulated with scripts **KinGen** and **KinDeg** to confirm that the calculated kinetic constants are in accordance with the experimental data. Also perform simulations with **KinVol** and **KinVolDeg** to confirm that addition of substrate does not significantly affect the noncovalent interactions/equilibria or reaction rates by dilution and/or competition. Experimental estimates of k_s and k_c are generated in the previous step of this protocol.

Assay Protocol IV. Preincubation Time-Dependent Inhibition With Dilution/Competition

The protocol below provides a generic set of steps to accomplish this type of measurement.

Materials

- 1× Assay/reaction buffer supplemented with co-factors and reducing agent
- Active enzyme, 200× solution in assay buffer
- Substrate with continuous or quenched read-out, 1× solution in assay buffer
- · Positive control: vehicle/solvent as DMSO stock, or 2% solution in assay buffer
- Negative control: known inhibitor or alkylating agent as DMSO stock, or 200× solution in assay buffer
- Inhibitor: as DMSO stock, or serial dilution of 200× solution in assay buffer with 2% DMSO
- · Optional: Development/quenching solution
- 1.5 mL (Eppendorf) microtubes to prepare stock solutions
- 384-well low volume microplate with nonbinding surface (e.g. Corning 3820 or 4513) for preincubation
- Microplate cover/lid (e.g. Corning 6569 Microplate Aluminum Sealing Tape) to seal 384-well plate during preincubation
- 96-well low volume microplate with nonbinding surface (e.g. Corning 3650 or 3820) for quenching and read-out
- Optional: 96-well microplate to prepare serial dilution of inhibitor concentration
- Optional: Microtubes to perform preincubations (e.g. Eppendorf Protein Lobind Microtubes, #022431018)
- Optional: 384-well low volume microplate with nonbinding surface (e.g. Corning 3820 or 4513) for read-out
- · Microplate reader equipped with appropriate filters to detect product formation (e.g. CLARIOstar microplate reader)
- Optional: Automated (acoustic) dispenser (e.g. Labcyte ECHO 550 Liquid Handler acoustic dispenser)

Exemplary assay concentrations

		incubation t'			incubation t	
	[stock]	Volume	[conc] _{t'}	[stock]	Volume	[conc] _t
Enzyme	200 nM	10 µL	99 nM	-	1 μL	1.0 nM
Inhibitor	2000 nM	10.2 μL	1010 nM	-	1 µL	10 nM
Substrate	-	-	-	10 µM	198 µL	9.9 μM
Total		20.2 μL			200 µL	

Before you start, optimize assay conditions in the uninhibited control to ensure compliance with assumptions and restrictions, as outlined in *Assay Protocol I.* Consult **Table 6** (*section 5*) for common optimization and troubleshooting options.

Specific adjustments for Method IV

Substrate should be added in a large volume ($V_{sub} >> V_{t'}$) and/or at a high concentration ([S]₀ >> K_M) to guench time-dependent enzyme inhibition (as illustrated in Figure 21). Enzyme concentration after dilution [Etotal], should be adjusted to correspond to maximum 10% substrate conversion until the end of the incubation in the uninhibited control ($[P]_r < 0.1[S]_0$), and substrate should be present in excess $([S]_0 > 10[E^{total}]_t)$. Preincubation-dependent enzyme activity should be calculated from initial, linear product formation after substrate addition. Validate that enough product is formed for a good signal:noise ratio (Z' > 0.5) by calculating the Z'-score from the uninhibited and inhibited controls (ideally 8 replicates) in a separate experiment.⁸⁷ This method is compatible with homogeneous (continuous) assays but also with assays that require a development/quenching step to visualize formed product. Note that preincubation in very small volumes (<10 µL) is not representative/reliable and the volume after 100-fold dilution in substrate will often exceed the maximum well volume of assay plates. Therefore, preincubation is typically performed in a larger volume (tube or plate) from which aliquots are removed at the end of the preincubation. In this protocol, we perform incubations in triplicate (20 µL per replicate) in a 384-well plate, from which 2 µL aliquots are removed and guenched in 198 µL substrate in a 96-well plate that is also used for read-out. Optionally, it is possible to then transfer 20 µL to a 384-well plate for read-out, but multiple transfers of assays solutions will introduce errors. Alternatively, preincubation can be performed in microtubes or a 96-well plate.

 Add inhibitor or control (e.g. 0.2 μL) and assay buffer (e.g. 10 μL) to each well with the uninhibited control for full enzyme activity containing the same volume vehicle/solvent instead of inhibitor, as outlined in step 1 of Assay Protocol III.

Gently shake to mix DMSO with the aqueous buffer. Typically, measurements are performed in triplicate (or more replicates) with at least 8 inhibitor concentrations for at least 5 preincubation times. Inhibitor concentrations might need optimization, but a rational starting point is to use inhibitor concentrations below $5 \times IC_{50}$ at the shortest preincubation time t': inhibition is expected to improve in a time-dependent manner, and the best results are obtained when full inhibition is not achieved already at the shortest preincubation time. Whether preincubation is performed in a tube or microplate is a matter of personal preference, compatibility with lab equipment and automation, and convenience of dispensing small volumes.

 Add active enzyme in assay buffer to each well (e.g. 10 μL of 200× solution) or tube to start preincubation of enzyme with inhibitor and homogenize the solution by gently shaking (1 min at 300 rpm). Alternatively, dispensing the enzyme at a high flow rate will also mix the components.

The order of enzyme and inhibitor addition is not important *per se*, as long as DMSO stocks are added prior to buffered (aqueous) solutions. Inhibitor must be present in excess during preincubation $([I]_0 > 10[E]_0)$. Optionally, gently centrifuge the plate or microtubes (1 min at 1000 rpm) to ensure assay components are not stuck at the top of the well.

- Seal the wells with a cover or lid, and close the caps of microtubes to prevent evaporation of assay components during preincubation.
- 4. Remove a single aliquot in volume Vt (e.g. 2 µL) from the reaction mixture, and transfer to a 96-well microplate already containing a large volume (volume V_{sub}) of substrate (e.g. 198 µL of 1× solution in assay buffer) after preincubation time t'.

Substrate should be added in a large volume (V_t << V_t) and/or at a high concentration ([S] >> K_M) to quench timedependent addition enzyme inhibition during incubation by dilution ([I]_t << [I]_t.) or competition (increasing K₁^{app} or decreasing k_{chem}^{app}). Dilution to inhibitor concentration far below the equilibrium concentration ([I]_t << K_i^{app}) promotes dissociation of noncovalently bound inhibitor after substrate addition (illustrated in **Figure 21A**). The accuracy of the measurement improves if the preincubation time is monitored precisely. Optionally, homogenize the solutions by gentle shaking (300 rpm) and centrifuge the plate (1 min at 1000 rpm) to ensure assay components are not stuck at the top of the well.

 Quenching: Add development solution to the reaction mixture in the microplate to quench the product formation reaction if read-out of product formation requires a development/quenching step to visualize formed product after incubation time t.

Follow manufacturer advice on waiting time after addition of development solution before read-out. Incubation time t is the elapsed time between onset of product formation by substrate addition (*step 4*) and addition of development/ quenching solution (*step 5*). A possible advantage to the use of a quenched assay is the ability to store the samples

after addition of quenching/development solution (*step 5*) and measure product formation (*step 6*) in all samples after completion of the final preincubation rather than performing multiple separate measurements (after each preincubation time).

6. Optional: Transfer aliquot (e.g. 20 µL) to a 384-well microplate for read-out.

Typically, the total volume after dilution in substrate solution ($V_t = V_{sub} + V_{t'}$) exceeds the maximum well volume of a 384-well microplate. Transfer an appropriate amount of reaction mixture (at least two technical replicates) to a microplate. This step can be skipped if read-out is performed in a 96-well plate.

7. Measure formed product after incubation by detection of the product read-out in reader.

Incubation time *t* (after substrate addition) is arbitrary as long as product formation is linear in uninhibited as well as inhibited samples (Figure 21B).

8. Repeat steps 4-7 of Assay Protocol IV for at least another four preincubation times.

Preincubation time t' is the elapsed time between onset of inhibition by mixing enzyme and inhibitor (*step 2*) and addition of substrate (*step 4*). A typical preincubation assay is multiple hours measuring enzyme activity every 5-30 min, depending on enzyme stability and inhibitor reaction rates. Best results are obtained if the incubation time t used to calculate enzyme activity is kept constant at all preincubation times.

9. Proceed to *Data Analysis Protocol 4* to convert the raw experimental data into preincubation time-dependent enzyme activity.

Data Analysis Protocol 4 for 1-step and 2-step IRREV Inhibition

Processing of raw experimental data obtained with Assay Protocol IV for irreversible inhibitors.

1. Plot signal F against incubation time t

Plot signal *F* (in AU) on the Y-axis against the incubation time (in s) on the X-axis for each inhibitor concentration and for the controls (Figure 22B, Figure 24B). Do this separately for each preincubation time.

2. Fit F_t against t to obtain $v_{t'}$

Fit signal F_t against incubation time t to Equation XIII (Figure 22B, Figure 24B) to obtain preincubation time-dependent product formation velocity $v_{t'}$ from the linear slope. Linear product formation is indicative of effective disruption of additional covalent modification during incubation by dilution in excess substrate (Figure 21A). If product formation is not linear: consult Table 6 (section 5) for troubleshooting or change to Data Analysis Protocol 3.

$F_t = F_0 + v_{t'} t$

(XIII)

Nonlinear regression of straight line **Y=YIntercept+Slope***X with Y = signal F_t (in AU) and X = incubation time t (in s) to find YIntercept = background signal at reaction initiation F_0 (in AU) and Slope = preincubation time-dependent product formation velocity v_r (in AU/s).

 Proceed to Data Analysis Protocols to obtain the appropriate kinetic parameters for each covalent binding mode: Data Analysis Protocol 4Ai or 4Aii for 2-step irreversible inhibitors and Data Analysis Protocol 4Bi or 4Bii for 1-step irreversible inhibitors

Selection of a data analysis protocols for inhibitors with an irreversible binding mode depends on the desired visual representation as well as personal preference. Generally, *Data Analysis Protocols 4Ai* and *4Bi* have less data processing/ manipulation and are more informative for comparison of various inhibitors on a single enzyme target, as they are compatible with assessment of inhibitor potency simultaneous with visual assessment of time-dependent enzyme stability k_{ctrl} (Figure 23B-C, Figure 25B-C). *Alternative Data Analysis Protocols 4Aii* and *4Bii* involve normalization of the enzyme activity that aids visual assessment of inhibitory potency of a single inhibitor on multiple enzyme targets (that might have a variable stability) (Figure 23D-E, Figure 25D-E).

_	Data Analysis Protocol		
EXP Conditions	2-step IRREV	1-step IRREV	2-step REV
$k_{\rm ctrl} = 0$	4Ai or 4Aii	4Bi or 4Bii	-
$k_{\text{degE}} > 0$	4Ai or 4Aii	4Bi or 4Bii	-

Data Analysis Protocol 4Ai for 2-step IRREV Inhibition

Processing of experimental data obtained with *Assay Protocol IV* that has been processed according to *Data Analysis Protocol 4* for 2-step irreversible inhibitors.

1. Plot $v_{t'}$ against preincubation time t' for each inhibitor concentration

Plot the mean and standard deviation of v_t (in AU/s) on the Y-axis against preincubation time t' (in s) on the X-axis for each inhibitor concentration and the uninhibited control (Figure 22C). Validate that inhibitor concentrations are not too high: inhibition should be less than 100% at the shortest t' for at least six inhibitor concentrations. Check whether the uninhibited enzyme activity is independent of preincubation time ($v_0^{\text{ctrl}} = v_t^{\text{ctrl}}$): an algebraic correction for enzyme instability ($k_{\text{ctrl}} > 0$, Figure 23B) can be performed in *step* 4 of this protocol by accounting for nonlinearity in the uninhibited control in the secondary k_{obs} plot (Figure 23C). Alternatively, proceed to Alternative Data Analysis Protocol 4Bii to correct for enzyme instability ($v_t^{\text{ctrl}} > v_t^{\text{ctrl}}$) by normalization of the enzyme activity v_t/v_t^{ctrl} (Figure 23D-E).

2. Fit $v_{t'}$ against preincubation time t' to obtain k_{obs}

Fit the mean and standard deviation of v_t against preincubation time t' (Figure 22C/Figure 23B) for each inhibitor concentration to bounded exponential decay Equation VI with shared value for initial velocity v_i to obtain the observed reaction rate k_{obs} from initial velocity v_i (V-intercept) to full inactivation (v_s in fully inhibited control). A lack of initial noncovalent complex ($v_i = v_0^{ctrl}$) is indicative of effective disruption of noncovalent interactions by dilution in excess substrate. Validate this by fitting without constraints for v_i . Proceed to *Data Analysis Protocol 3Ai* if deviations ($v_i < v_0^{ctrl}$) are observed.

$$v_{t'} = v_0^{\text{ctrl}} e^{-k_{\text{obs}}t}$$

(VI)

Nonlinear regression of exponential one-phase decay equation Y = (Y0-Plateau) *EXP(-k*X)+Plateau with Y = preincubation time $dependent product formation velocity <math>v_t$. (in AU/s), X = preincubation time t' (in s) and Plateau = final velocity $v_a = 0$ or v_a^{ctri} in fully inhibited control (in AU/s, constrained) to find Y0 = Y-intercept = initial velocity v_i = uninhibited velocity v_0^{ctri} (in AU/s, shared value) and k = observedreaction rate k_{obs} (in s⁻¹).

3. Plot k_{obs} against [I]

Plot the mean and standard deviation of k_{obs} (in s^{-1}) on the Y-axis against inhibitor concentration (in M) <u>during</u> <u>preincubation</u> (before addition of substrate) on the X-axis (**Figure 22D**/**Figure 23C**). The plot of k_{obs} against [I] should reach a maximum k_{obs} at saturating inhibitor concentration. Note that a linear curve is indicative of 1-step binding behavior at non-saturating inhibitor concentrations ([I] << 0.1K₁^{app}: see **Figure 3F**) with $v_i = v_0^{ctrl}$ (shared Y-intercept in the previous step). Proceed to step 4 of Data Analysis Protocol 4Bi after it has been validated that the linear curve is not resultant from saturating inhibitor concentrations ([I] >> 10K₁^{app}: see **Figure 3G**) as identified by $v_i << v_0^{ctrl}$, by repeating the measurement with lower inhibitor concentrations.

Fit k_{obs} against [I] to obtain k_{inact} and K_I

Fit k_{obs} against inhibitor concentration <u>during preincubation</u> to **Equation xv** to obtain maximum inactivation rate constant k_{inact} and inactivation constant K_1 . Constrain $k_{ctrl} = k_{obs}$ of the uninhibited control (**Figure 23C**). Inactivation constant K_1 does <u>not</u> have to be corrected for substrate competition because preincubation is conducted in absence of competing substrate. Calculate irreversible covalent inhibitor potency k_{inact}/K_1 with propagation of error with *Sample Calculation 2*.

$$k_{\rm obs} = k_{\rm ctrl} + \frac{k_{\rm inact} \left[I \right]}{K_{\rm I} + \left[I \right]} \tag{xv}$$

Nonlinear regression of user-defined explicit equation Y=Y0+((kmax*X)/((KI)+X)) with Y = observed reaction rate k_{obs} (in s⁻¹), X = inhibitor concentration during preincubation (in M), and Y0 = rate of nonlinearity in uninhibited control k_{ctrl} (in s⁻¹, constrained) to find kmax = maximum reaction rate k_{inact} (in s⁻¹), and KI = Inactivation constant K₁ (in M).

5. Optional: Validate experimental kinetic parameters with kinetic simulations

Proceed to *Kinetic Simulations 1* to compare the experimental read-out to the product formation simulated with scripts **KinVol** and **KinVolDeg** (using experimental rate constant $k_{inact} = k_5$) to confirm that the calculated kinetic constants are in accordance with the experimental data.

Alternative Data Analysis Protocol 4Aii for 2-step IRREV Inhibition

Processing of experimental data obtained with *Assay Protocol IV* that has been processed according to *Data Analysis Protocol 4* for 2-step irreversible inhibitors.

1. Plot $v_{t'}$ against preincubation time t' for each inhibitor concentration

Plot the mean and standard deviation of v_t (in AU/s) on the Y-axis against preincubation time t' (in s) on the X-axis for each inhibitor concentration and the uninhibited control (**Figure 23B**). Validate that inhibitor concentrations are not too high: inhibition should be less than 100% at the shortest t' for at least six inhibitor concentrations.

2. Normalize $v_{t'}$ to obtain $v_{t'}/v^{ctrl}$

Normalize v_t of each inhibitor concentration and the controls to lowest value = 0 (or full inhibition control) and highest value = uninhibited product formation v_t^{ctrl} to obtain normalized enzyme activity v_t/v^{ctrl} . Perform this correction separately for each preincubation time.

3. Plot and fit $v_{t'}/v^{ctrl}$ against preincubation time t' to obtain k_{obs}

Plot the mean and standard deviation of v_t / v^{ctrl} on the Y-axis against preincubation time t' (in s) on the X-axis (**Figure 23D**). Fit to exponential decay **Equation XIX** to obtain k_{obs} from initial velocity v_i / v_0^{ctrl} to full inactivation (Plateau = 0). A lack of initial noncovalent complex ($v_i = v_0^{\text{ctrl}}$) is indicative of effective disruption of noncovalent interactions by dilution in excess substrate. Validate this by fitting without constraints for v_i . Proceed to *Data Analysis Protocol 3Aii* if deviations ($v_i < v_0^{\text{ctrl}}$) are observed.

$$\left(\frac{v_{t'}}{v_{t'}^{\text{ctrl}}}\right) = e^{-k_{\text{obs}}t'} \tag{XIX}$$

Nonlinear regression of exponential one-phase decay equation Y=(YO-Plateau)*EXP(-k*X)+Plateau with Y = normalized preincubation time-dependent product formation velocity v_r/v_c^{ctrl} (unitless), X = preincubation time t' (in s), YO = Y-intercept = normalized initial velocity $v_r/v_c^{ctrl} = 1$ (unitless, constrained), and Plateau = normalized final velocity $v_s/v_s^{ctrl} = 0$ (unitless, constrained) to find k = observed reaction rate k_{bb} (in s⁻¹).

4. Plot k_{obs} against [I]

Plot the mean and standard deviation of k_{obs} (in s⁻¹) on the Y-axis against inhibitor concentration (in M) <u>during</u> <u>preincubation</u> (*before* addition of substrate) on the X-axis (**Figure 23E**). The plot of k_{obs} against [I] should reach a maximum k_{obs} at saturating inhibitor concentration. Note that a linear curve is indicative of 1-step binding behavior at non-saturating inhibitor concentrations ([I] < 0.1K₁^{app}: see **Figure 3F**) with $v_1 = v_0^{ctrl}$ (shared Y-intercept = 1 in the previous step). Proceed to *step 5* of *Data Analysis Protocol 4Bii* after it has been validated that the linear curve is not resultant from saturating inhibitor concentrations ([I] >> 10K₁^{app}: see **Figure 3G**) as identified by $v_1 < v_0^{ctrl}$ (shared Y-intercept = 0 in the previous step), by repeating the measurement with lower inhibitor concentrations.

5. Fit k_{obs} against [I] to obtain k_{inact} and K_I

Fit k_{obs} against inhibitor concentration <u>during preincubation</u> to **Equation XVII** to obtain maximum inactivation rate constant k_{inact} and inactivation constant K_1 (**Figure 23E**). Do not correct for enzyme instability ($k_{ctrl} > 0$), as this correction has already been performed by normalizing v_t . Inactivation constant K_1 does <u>not</u> have to be corrected for substrate competition because preincubation is conducted in absence of competing substrate. Calculate irreversible covalent inhibitor potency k_{mact}/K_1 with propagation of error with *Sample Calculation 2*.

$$k_{\rm obs} = \frac{k_{\rm inact}\left[I\right]}{K_{\rm I} + \left[I\right]} \tag{XVII}$$

Nonlinear regression of user-defined explicit equation Y=Y0+((kmax*X)/((KI)+X)) with Y = observed reaction rate k_{obs} (in s⁻¹), X = inhibitor concentration during preincubation (in M), and Y0 = 0 (in s⁻¹, constrained) to find kmax = maximum reaction rate k_{inact} (in s⁻¹) and KI = Inactivation constant K₁ (in M).

6. Optional: Validate experimental kinetic parameters with kinetic simulations

Proceed to *Kinetic Simulations 1* to compare the experimental read-out to the product formation simulated with scripts **KinVol** and **KinVolDeg** (using experimental rate constant $k_{inact} = k_5$) to confirm that the calculated kinetic constants are in accordance with the experimental data.

Data Analysis Protocol 4Bi for 1-step IRREV Inhibition

Processing of experimental data obtained with *Assay Protocol IV* that has been processed according to *Data Analysis Protocol 4* for 1-step irreversible covalent inhibitors and 2-step irreversible inhibitors at non-saturating inhibitor concentrations ($[I] \le 0.1K_1$).

1. Plot $v_{t'}$ against preincubation time t' for each inhibitor concentration

Plot the mean and standard deviation of v_t (in AU/s) on the Y-axis against preincubation time t' (in s) on the X-axis for each inhibitor concentration and the uninhibited control (Figure 24C). Validate that inhibitor concentrations are not too high: inhibition should be less than 100% at the shortest t' for at least six inhibitor concentrations. Check whether the uninhibited enzyme activity is independent of preincubation time ($v_0^{\text{ctrl}} = v_t^{\text{ctrl}}$): an algebraic correction for enzyme instability ($k_{\text{ctrl}} > 0$, Figure 25B) can be performed in *step* 4 of this protocol by accounting for nonlinearity in the uninhibited control in the secondary k_{obs} plot (Figure 25C). Alternatively, proceed to Alternative Data Analysis Protocol 4Bii to correct for enzyme instability ($v_0^{\text{ctrl}} > v_t^{\text{ctrl}}$) by normalization of the enzyme activity v_t , v_t^{ctrl} (Figure 25D-E).

2. Fit $v_{t'}$ against preincubation time t' to obtain k_{obs}

Fit the mean and standard deviation of v_t against preincubation time t' (Figure 24C/Figure 25B) for each inhibitor concentration to bounded exponential decay Equation VI. Constrain initial velocity v_i to a shared value to obtain observed reaction rate k_{obs} from initial velocity v_i (Y-intercept) to full inactivation ($v_s = 0$ or value in fully inhibited control).

$$v_{t'} = v_0^{\text{ctrl}} e^{-k_{\text{obs}}t}$$

(VI)

Nonlinear regression of exponential one-phase decay equation Y = (YO-Plateau) *EXP(-k*X)+Plateau with Y = preincubation time $dependent product formation velocity <math>v_t$ (in AU/S), X = preincubation time t' (in s), and Plateau = final velocity $v_e = 0$ or v_e^{ctri} in fully inhibited control (in AU/S, constrained) to find YO = Y-intercept = initial velocity v_i = uninhibited velocity v_0^{ctri} (in AU/S, shared value) and k = observedreaction rate k_{obs} (in 5⁻¹).

3. Plot k_{obs} against [I]

Plot the mean and standard deviation of k_{obs} (in s⁻¹) on the Y-axis against inhibitor concentration (in M) <u>during</u> <u>preincubation</u> (*before* addition of substrate) on the X-axis (**Figure 24D/Figure 25C**). The plot of k_{obs} against inhibitor concentration [I] is linear for 1-step irreversible inhibitors and for 2-step irreversible inhibitors at non-saturating inhibitor concentrations ([I] << 0.1K₁).

4. Fit k_{obs} against [I] to obtain k_{chem}

Fit k_{obs} against inhibitor concentration <u>during preincubation</u> (in M) to **Equation XVIII** to obtain inhibitor potency k_{chem} from the linear slope. Constrain Y-intercept to k_{ctrl} : k_{obs} of the uninhibited control (**Figure 25C**). Inhibitor potency k_{chem} does <u>not</u> have to be corrected for substrate competition because preincubation is conducted in absence of competing substrate. Calculate k_{inact}/K_1 for 2-step irreversible inhibitors at non-saturating inhibitor concentrations ([I] $\leq 0.1K_1$) with Sample Calculation 9.

$k_{\rm obs} = k_{\rm ctrl} + k_{\rm chem} [I]$

(XVIII)

Nonlinear regression of straight line Y=YIntercept+Slope*X with Y = observed reaction rate k_{obs} (in s^{-1}), X = inhibitor concentration during preincubation (in M), and YIntercept = rate of nonlinearity in uninhibited control k_{ctrl} (in s^{-1} , constrained) to find Slope = inactivation rate constant k_{chem} (in $M^{-1}s^{-1}$).

5. Optional: Validate experimental kinetic parameters with kinetic simulations

Proceed to *Kinetic Simulations 1* to compare the experimental read-out to the product formation simulated with scripts **KinVol** and **KinVolDeg** (using experimental rate constant $k_{chem} = k_3$), to confirm that the calculated kinetic constants are in accordance with the experimental data.

Alternative Data Analysis Protocol 4Bii for 1-step IRREV Inhibition

Processing of experimental data obtained with *Assay Protocol IV* that has been processed according to *Data Analysis Protocol 4* for 1-step irreversible covalent inhibitors and 2-step irreversible inhibitors at non-saturating inhibitor concentrations ($[I] \le 0.1K_1$).

1. Plot $v_{t'}$ against preincubation time t' for each inhibitor concentration

Plot the mean and standard deviation of v_t (in AU/s) on the Y-axis against preincubation time t' (in s) on the X-axis for each inhibitor concentration and the uninhibited control (Figure 25B). Validate that inhibitor concentrations are not too high: inhibition should be less than 100% at the shortest t' for at least six inhibitor concentrations.

2. Normalize $v_{t'}$ to obtain $v_{t'}/v^{ctrl}$

Normalize v_t of each inhibitor concentration and the controls to lowest value = 0 (or full inhibition control) and highest value = uninhibited product formation v_t^{ctrl} to obtain normalized enzyme activity v_t'/v^{ctrl} . Perform this correction separately for each preincubation time.

3. Plot and fit $v_{t'}/v^{ctrl}$ against preincubation time t' to obtain k_{obs}

Plot the mean and standard deviation of $v_{t'}/v^{ctri}$ on the Y-axis against preincubation time t' (in s) on the X-axis (**Figure 25D**). Fit to exponential decay **Equation XIX** to obtain k_{obs} from initial velocity v_i/v_0^{ctri} (shared value) to full inactivation (Plateau = 0).

$$\left(\frac{v_{t'}}{v_{t'}^{\text{ctrl}}}\right) = e^{-k_{\text{obs}}t'} \tag{XIX}$$

Nonlinear regression of exponential one-phase decay equation Y = (YO-Plateau)*EXP(-k*X) + Plateau with Y = normalizedpreincubation time-dependent product formation velocity v_t/v^{ctrl} (unitless), X = preincubation time t' (in s), Plateau = normalized final velocity $v_s/v_s^{ctrl} = 0$ (unitless, constrained), and YO = Y-intercept = normalized initial velocity $v_t/v_0^{ctrl} = 1$ (unitless, constrained) to find $k = observed reaction rate <math>k_{obs}$ (in s^{-1}).

4. Plot k_{obs} against [I]

Plot the mean and standard deviation of k_{obs} (in s⁻¹) on the Y-axis against inhibitor concentration (in M) <u>during</u> <u>preincubation</u> (*before* addition of substrate) on the X-axis (**Figure 25E**). The plot of k_{obs} against inhibitor concentration [I] is linear for 1-step irreversible inhibitors and for 2-step irreversible inhibitors at non-saturating inhibitor concentrations ([I] << 0.1K₁).

5. Fit k_{obs} against [I] to obtain k_{chem}

Fit k_{obs} against inhibitor concentration <u>during preincubation</u> to **Equation XX** to obtain inhibitor potency k_{chem} from the linear slope (**Figure 25E**). Do not correct for enzyme instability ($k_{ctrl} > 0$), as this correction has already been performed by normalizing v_t to v_t / v^{ctrl} in *step 2* of this protocol. Inhibitor potency k_{chem} does <u>not</u> have to be corrected for substrate competition because preincubation is conducted in absence of competing substrate. Calculate k_{inact}/K_I for 2-step irreversible inhibitors at non-saturating inhibitor concentrations ([I] $\leq 0.1K_I$) with propagation of error with *Sample Calculation 9*. Alternatively, inhibitor potency k_{chem} or k_{inact}/K_I can be directly calculated from a single k_{obs} and [I] with *Sample Calculation 10*.

$$k_{\rm obs} = k_{\rm chem} [I]$$

(XX)

Nonlinear regression of straight line $\mathbf{Y} = \mathbf{YIntercept} + \mathbf{Slope} * \mathbf{X}$ with $\mathbf{Y} = \text{observed}$ reaction rate k_{obs} (in s^{-1}), $\mathbf{X} = \text{inhibitor}$ concentration during preincubation (in M), and YIntercept = 0 (in s^{-1} , constrained) to find Slope = inactivation rate constant k_{chem} (in $M^{-1}s^{-1}$).

6. Optional: Validate experimental kinetic parameters with kinetic simulations

Proceed to *Kinetic Simulations 1* to compare the experimental read-out to the product formation simulated with scripts **KinVol** and **KinVolDeg** (using experimental rate constant $k_{chem} = k_3$), to confirm that the calculated kinetic constants are in accordance with the experimental data.

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4.1. Sample Calculations

The values obtained in the data analysis protocols have to be converted into relevant inhibition parameters. These are fairly straightforward linear calculations and can be performed with more basic software like Microsoft EXCEL. For each equation, the values on the right-hand side of the equal sign are known, so it becomes a linear calculation to obtain the parameter on the left-hand side. The calculations are listed in order of appearance in the manuscript. We have outlined the key assumptions and a little background on the used variables for improved readability and direct applicability following the protocols.

Sample Calculation 1. Calculate K₁ from K₁^{app}

Apparent inactivation constant K_1^{app} (in M) found in *Data Analysis Protocols 1A* or *1D* for competitive 2-step irreversible inhibitors is corrected for substrate competition to obtain inactivation constant K_1 (in M), with propagation of error. Use substrate concentration [S] (in M) after reaction initiation and K_M (in M) as determined for these specific assay conditions (buffer, temperature, enzyme, substrate). Proceed to *Sample Calculation 2* to calculate k_{inart}/K_1 .

$$K_{I} = \frac{K_{I}^{app}}{\left(1 + \frac{[S]}{K_{M}}\right)} \qquad \sigma_{K_{I}} = \sqrt{\left(\frac{1}{1 + \frac{[S]}{K_{M}}}\right)^{2} \sigma_{K_{I}^{app}}^{2} + \left(-\frac{K_{I}^{app} K_{M}}{\left(K_{M} + [S]\right)^{2}}\right)^{2} \sigma_{[S]}^{2} + \left(\frac{K_{I}^{app} [S]}{\left([S] + K_{M}\right)^{2}}\right)^{2} \sigma_{K_{M}}^{2}}$$

<u>Sample Calculation 2.</u> Calculate k_{inact}/K_I from k_{inact} and K_I

Irreversible covalent inhibitor potency k_{inact}/K_{I} (in $M^{-1}s^{-1}$) is calculated from k_{inact} (in s^{-1}) and K_{I} (in M) values found in *Data Analysis Protocols 1A*, *1D*, *2*, *3Ai*, *3Aii*, *4Ai* or *4Aii* and *Sample Calculation 1* for 2-step irreversible inhibitors, with propagation of error.

$$\left(\frac{k_{\text{inact}}}{K_{\text{I}}}\right) = \frac{k_{\text{inact}}}{K_{\text{I}}} \qquad \sigma_{\frac{k_{\text{inact}}}{K_{\text{I}}}} = \left(\frac{k_{\text{inact}}}{K_{\text{I}}}\right) \sqrt{\left(\frac{\sigma_{k_{\text{inact}}}}{k_{\text{inact}}}\right)^{2} + \left(\frac{\sigma_{K_{\text{I}}}}{K_{\text{I}}}\right)^{2}}$$

Sample Calculation 3. Calculate K_i from K_i^{app}

Apparent inhibition constant K_i^{app} (in M) found in *Data Analysis Protocols 1A*, *1C*, *3Ai*, *3Aii* or *3C* for competitive 2-step (ir)reversible inhibitors is corrected for substrate competition to obtain inhibition constant K_i (in M) for the initial noncovalent equilibrium.⁸⁹ Use substrate concentration [S] (in M) after reaction initiation and K_M (in M) as determined for these specific assay conditions (buffer, temperature, enzyme, substrate). Inhibition constant K_i for 2-step irreversible inhibitors if covalent bond formation is rate-limiting (rapid equilibrium assumption).

$$K_{i} = \frac{K_{i}^{app}}{\left(1 + \frac{[S]}{K_{M}}\right)} \qquad \sigma_{K_{i}} = \sqrt{\left(\frac{1}{1 + \frac{[S]}{K_{M}}}\right)^{2} \sigma_{K_{i}^{app}}^{2} + \left(-\frac{K_{i}^{app} K_{M}}{(K_{M} + [S])^{2}}\right)^{2} \sigma_{[S]}^{2} + \left(\frac{K_{i}^{app} [S]}{([S] + K_{M})^{2}}\right)^{2} \sigma_{K_{M}}^{2}}$$

<u>Sample Calculation 4.</u> Calculate k_{chem} from k_{chem}^{app}

Apparent inhibitor potency k_{chem}^{app} (in M⁻¹s⁻¹) found in *Data Analysis Protocol 1B* for competitive 1-step irreversible inhibitors is corrected for substrate competition to obtain inhibition potency k_{chem} (in M⁻¹s⁻¹) with propagation of error. Use substrate concentration [S] (in M) after reaction initiation and K_M (in M) as determined for these specific assay conditions (buffer, temperature, enzyme, substrate).

$$k_{\rm chem} = k_{\rm chem}^{\rm app} \left(1 + \frac{[S]}{K_{\rm M}}\right) \qquad \sigma_{k_{\rm chem}} = \sqrt{\left(1 + \frac{[S]}{K_{\rm M}}\right)^2 \sigma_{k_{\rm chem}^{\rm app}}^2 + \left(\frac{k_{\rm chem}^{\rm app}}{K_{\rm M}}\right)^2 \sigma_{[S]}^2 + \left(-\frac{k_{\rm chem}^{\rm app}[S]}{K_{\rm M}^2}\right)^2 \sigma_{K_{\rm M}}^2}$$
<u>Sample Calculation 5.</u> Calculate k_{inact}/K_{I}^{app} from k_{chem}^{app}

The linear slope k_{chem}^{app} (in M⁻¹s⁻¹) found in *Data Analysis Protocol 1B* for 2-step irreversible inhibitors equals k_{inact}/K_{I}^{app} when all inhibitor concentrations are non-saturating ([I] $\leq 0.1K_{I}^{app}$). It is not possible to obtain individual values of k_{inact} and K_{I} from a linear graph, but it is possible to estimate the upper and lower limits: K_{I}^{app} is much larger than the highest inhibitor concentration if this concentration is non-saturating ($K_{I}^{app} >> [I]_{max}$). An unchanged slope upon constraining the Y-intercept k_{ctrl} (*step 5* in *Data Analysis Protocol 1B*) to the experimental value for the uninhibited control validates that all inhibitor concentrations are non-saturating (see **Figure 3G**). Proceed to *Sample Calculation 6* to calculate k_{inart}/K_{I} .

$$k_{\rm chem}^{\rm app} = \left(\frac{k_{\rm inact}}{{\rm K}_{\rm I}^{\rm app}}\right)$$

<u>Sample Calculation 6.</u> Calculate k_{inact}/K_I from k_{inact}/K_I^{app}

Apparent inactivation potency k_{inact}/K_i^{app} (in M⁻¹s⁻¹) found in *Data Analysis Protocols 1A* or 1D, or calculated in *Sample Calculation 5* for competitive 2-step irreversible inhibitors is corrected for substrate competition to obtain k_{inact}/K_i (in M) with propagation of error. Use substrate concentration [S] (in M) after reaction initiation and K_M (in M) as determined for these specific assay conditions (buffer, temperature, enzyme, substrate).

$$\frac{k_{\text{inact}}}{K_{\text{I}}} = \left(\frac{k_{\text{inact}}}{K_{\text{I}}^{\text{app}}}\right) \left(1 + \frac{[S]}{K_{\text{M}}}\right) \quad \sigma_{\underline{k_{\text{inact}}}} = \sqrt{\left(1 + \frac{[S]}{K_{\text{M}}}\right)^2 \sigma_{\underline{k_{\text{inact}}}^2}} + \left(\frac{\left(\frac{k_{\text{inact}}}{K_{\text{I}}^{\text{app}}}\right)}{K_{\text{M}}}\right)^2 \sigma_{[S]}^2 + \left(-\frac{\left(\frac{k_{\text{inact}}}{K_{\text{I}}^{\text{app}}}\right)[S]}{K_{\text{M}}^2}\right)^2 \sigma_{\underline{k_{\text{M}}}^2}$$

Sample Calculation 7. Calculate K_i^{*} from K_i^{*app}

Apparent steady-state inhibition constant K_i^{*app} (in M) found in *Data Analysis Protocols 1C* or 3C for competitive 2-step reversible covalent inhibitors is corrected for substrate competition to obtain steady-state inhibition constant K_i^* (in M). Use substrate concentration [S] (in M) after reaction initiation and K_M (in M) as determined for these specific assay conditions (buffer, temperature, enzyme, substrate).

$$K_{i}^{*} = \frac{K_{i}^{*app}}{\left(1 + \frac{[S]}{K_{M}}\right)} \qquad \sigma_{K_{i}^{*}} = \sqrt{\left(\frac{1}{1 + \frac{[S]}{K_{M}}}\right)^{2} \sigma_{K_{i}^{*app}}^{2} + \left(-\frac{K_{i}^{*app} K_{M}}{\left(K_{M} + [S]\right)^{2}}\right)^{2} \sigma_{[S]}^{2} + \left(\frac{K_{i}^{*app} [S]}{\left([S] + K_{M}\right)^{2}}\right)^{2} \sigma_{K_{M}^{*2}}^{2}}$$

<u>Sample Calculation 8.</u> Calculate K_i^* from K_i , k_5 , and k_6

Steady-state inhibition constant K_i^* (in M) of 2-step reversible inhibitors can be calculated from experimental values of K_i (in M), k_5 (in s⁻¹), and k_6 (in s⁻¹) found with *Data Analysis Protocols 1C* or *3C*, and *Sample Calculation 3*. Reliable (relatively) small k_6 -values can only be obtained with more sensitive methods such as rapid dilution assays.^{29, 33} The uninhibited control must be strictly linear ($k_{ctrl} = 0$) for values found with *Data Analysis Protocol 1C*. This calculation is not the preferred method to obtain K_i^* due to its sensitivity to (experimental) errors in k_6 and contribution of k_{ctrl} : values obtained in *Data Analysis Protocol 1C* or *3C* and *Sample Calculation 7* should generally be considered as more reliable.

$$\mathbf{K}_{i}^{*} = \frac{\mathbf{K}_{i}}{\left(1 + \frac{k_{5}}{k_{6}}\right)} \qquad \qquad \mathbf{\sigma}_{\mathbf{K}_{i}^{*}} = \sqrt{\left(\frac{1}{1 + \frac{k_{5}}{k_{6}}}\right)^{2} \mathbf{\sigma}_{\mathbf{K}_{i}^{2}} + \left(-\frac{\mathbf{K}_{i} k_{6}}{\left(k_{6} + k_{5}\right)^{2}}\right)^{2} \mathbf{\sigma}_{k_{5}^{2}}^{2} + \left(\frac{\mathbf{K}_{i} k_{5}}{\left(k_{5} + k_{6}\right)^{2}}\right)^{2} \mathbf{\sigma}_{k_{6}^{2}}^{2}}$$

<u>Sample Calculation 9.</u> Calculate k_{inact}/K_I from k_{chem}

The linear slope k_{chem} (in M⁻¹s⁻¹) found in *Data Analysis Protocols 3Bi*, *3Bi*, *4Bi* or *4Bii* for 2-step irreversible inhibitors equals k_{inact}/K_I when all inhibitor concentrations are non-saturating ([I] $\leq 0.1K_I$). It is not possible to obtain individual values of k_{inact} and K_I from a linear graph, but it is possible to estimate the upper and lower limits: K_I is much larger than the highest inhibitor concentration if this concentration is non-saturating ($K_I >> [I]_{max}$). An unchanged slope upon constraining the Y-intercept k_{ctrl} to the experimental value for the uninhibited control in *step 4* of *Data Analysis Protocols 3Bi* and *4Bi* validates that all inhibitor concentrations are non-saturating (see **Figure 3F**) rather than saturating (see **Figure 3G**).

$$k_{\rm chem} = \left(\frac{k_{\rm inact}}{{\rm K}_{\rm I}}\right)$$

<u>Sample Calculation 10.</u> Calculate k_{chem} or k_{inact}/K_{I} from k_{obs} and [I]

Divide the k_{obs} -value (in s⁻¹) obtained in Alternative Data Analysis Protocols 3Bii or 4Bii by its corresponding inhibitor concentration (in M) to calculate irreversible inhibitor potency k_{chem} (in $M^{-1}s^{-1}$) or k_{inact}/K_{I} (in $M^{-1}s^{-1}$). This calculation is only accurate for normalized k_{obs} values (unaffected by contribution of k_{ctrl}), in absence of competing substrate, and (only applicable for 2-step irreversible inhibitors) at non-saturating inhibitor concentration.

$$k_{\text{chem}} = \frac{k_{\text{obs}}}{[1]}$$
$$\left(\frac{k_{\text{inact}}}{K_{1}}\right) = \frac{k_{\text{obs}}}{[1]}$$

4.2. Kinetic simulations

The figures illustrating the protocols in this work are generated using kinetic simulation scripts. These scripts are available online and can be used to validate the obtained kinetic parameters or help in optimizing your assay. On a more educational level, these scripts can show what your assay result could look like when using wildly different parameters to obtain more insight into how these affect your assay. A tutorial on how to perform kinetic simulations can be found on the website of our kinetic simulation scripts. Note: loading this page for the first time can take up to 5 minutes, and involves an automatic redirection (reloading) to the Landing Page.

Materials

- Kinetic Simulation Script (<u>https://tinyurl.com/kineticsimulations</u>)
- Software to open csv files (e.g. EXCEL)
- Data fitting software (e.g. GraphPad Prism)
- Experimental values found in Data Analysis Protocols 1-4

Kinetic Simulation 1. Validation of Experimental Values

Perform kinetic simulations to validate that calculated kinetic parameters are in accordance with experimental RAW data. Estimate microscopic rate constants from reported (literature) values, or use association rate constants $k_1 = k_3 = 10^{6} \cdot 10^{9} \text{ M}^{-1} \text{s}^{-1}$ (rapid noncovalent association) to calculate the dissociation rate constants from the experimental equilibrium constants: $k_4 = K_1 \times k_3$ (**Table S1**) and $k_2 = (K_M \times k_1) - k_{cat}$ (**Figure S1**). Ideally, also simulate the HTS reaction conditions to validate that the calculated kinetic constants give rise to the experimental inhibition or IC_{50}^{-35}

Kinetic Simulation 2. Rational Design of Validation Assays

Perform kinetic simulations with the calculated kinetic parameters to rationalize assay conditions for subsequent validation assays such as the minimum/maximum (pre)incubation times for reversibility assays or MS detection of the covalent adduct (formulas can be found in **Table 5**).

Reaction conditions for linearity in uninhibited control	$E + S \leftrightarrow ES \to E + P$
Maximum enzyme concentration $[E]_0$ for 10% substrate conversion during incubation in the uninhibited control $([S]_0 \ge 10[E]_0)$	$[E]_0 = \frac{0.1 ([S] + K_M)}{k_{\text{cat}} t} = \frac{[S] + K_M}{10 k_{\text{cat}} t}$
Maximum incubation time t for 10% substrate conversion in the uninhibited control ([S] ₀ \ge 10[E] ₀)	$t = \frac{[S] + K_{M}}{10 k_{cat} [E]_{0}}$
Reaction conditions for covalent occupancy	$E + I \leftrightarrow E I \to E I^*$
Expected covalent occupancy ${\rm [EI^*]}_{\mathfrak{r}}/{\rm [E]}_0$ after preincubation t^{\star} (in absence of competitor)	$\frac{[\text{EI*}]_{t'}}{[\text{E}]_0} = 1 - e^{-k_{\text{obs}}t'}$
Minimum preincubation time t' (in absence of competitor) to reach covalent occupancy $[EI^*]_t/[E]_0 \ge 0.6$	$t' = \frac{LN\left(1 - \frac{[EI^*]_{t'}}{[E]_0}\right)}{k_{obs}} = \frac{LN(1 - 0.6)}{k_{obs}}$

Table 5		Calcul	ations	to	rational	ize	assay	conditions.
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5. Troubleshooting

Like with any experimental method, our described methods will also require the necessary optimization. Since data analysis depends heavily on the experimental input, it is very important to optimize assay conditions, rather than trying to apply data corrections, to obtain reliable kinetic parameters. As the assay conditions will vary widely, depending on the enzyme used,⁷⁹ we can only give general pointers on the optimization of the assay conditions (**Table 6**). Luckily, many model substrates come with a satisfactory user manual or are described in extensive methods papers.⁶⁸⁻⁶⁹ These resources generally state reagents required for the reaction (e.g. fresh reducing agent, for cysteine-based catalysis) or additives that stabilize the read-out (such as BSA or Tween-20; to prevent aspecific aggregation). The control for full inhibition of (catalytic) cysteines is typically a thiol-alkylating reagent such as iodoacetamide (IAc) or N-methylmaleimide (NEM), or a known inhibitor.

As the assay performance is essential to get reliable fits, we recommend focusing on potential experimental problems before looking into issues with fitting. A great guide for general assay optimization can be found online in the assay guidance manual of the NCATS.⁹⁰ A comprehensive troubleshooting table with potential solutions that deal with various issues causing a troublesome read-out can be found on the next pages (**Table 6**). For the first half of **Table 6**, these solutions are generally related to the assay conditions and can generally be executed in the optimization stage. The latter half of **Table 6** is more geared towards after the data analysis of an initial experiment. The problems and accompanying solutions deal more with the experimental setup: how much inhibitor or substrate one needs to add becomes more apparent after these first data points. Some solutions, like changing inhibitor or substrate concentrations, can be simulated with our set of interactive kinetic simulation scripts. For better understanding and help in optimizing, we recommend simulating these conditions with our scripts to see what would happen when changing the concentrations.

Problem	Possible cause	Solutions
	Enzyme is not active (enough)	Increase [E] (not always possible with very potent inhibitors) Increase [S] to increase absolute maximum signal Optimize buffer components Switch to a substrate that is processed faster Activate enzyme with fresh reagents (e.g. DTT, ATP) in single-use aliquots Minimize freeze/thaw cycles
Difference positive and negative control is not significant	Signal product is not significant compared to substrate	Change fluorophore/read-out Optimize buffer components
	Negative control or inhibitor does not inhibit	Change to reported (specific) inhibitor Use thiol-alkylating reagent (e.g. NEM, IAc) for cysteines Use no-enzyme as negative control Increase concentration of inhibitor Make fresh dilution/aliquots of inhibitor solution
. ,	DMSO in positive control acts as inhibitor	<u>Validate:</u> compare enzyme activity with/without DMSO Reduce DMSO to max. 1% of final solution
	Machine settings/ sensitivity	Check if [P] is within the sensitivity range of used machine Optimize gain settings for [P] = $0-20\%$ [S] ₀ Check if correct wavelengths/settings are selected
	Pipetting error	Frequently replace pipette tips to avoid contamination of positive control with inhibitor (from negative control) Avoid well-to-well contamination by using an automated dispenser
Nonlinear uninhibited product formation curve <i>F</i> ^{ctrl}	Substrate depletion $([P]_t > 0.1[S]_0)$	Decrease [E] Increase [S] Shorter incubation time
	Spontaneous inactivation of enzyme $(k_{deg} > 0)$	Optimize buffer conditions for stability Use nonbinding surface plates Shorter incubation time
	Drift/evaporation	Cover/seal plate with optical clear cover Shorter incubation time
	Pre-steady state kinetics (lag phase)	Increase [S] to reach E + S ↔ ES equilibrium faster Preincubate enzyme with reducing agent/ATP
	Solution is not homogeneous	Introduce mixing step before addition of final component
	Fluorescence bleaching/ quenching	Optimize excitation conditions (e.g. lower no. of flashes) Longer measurement intervals/less measurements
Linear inhibited progress curve F _t	Inhibition is not time-dependent (or k _{obs} is too slow)	Longer (pre)incubation time (t > 0.1t½) Increase [I] Reduce [S] to decrease competition Activate enzyme with fresh reagents (e.g. DTT, ATP) Validate with different enzyme batch/construct
Full initial inhibition for all [l] (v _i = 0)	Noncovalent affinity is too potent ([I] >> K_i^{app})	Reduce [I] Higher [S] to increase competition (higher K _i ^{app}) Use method based on covalency (<i>Method IV</i> or direct detection)
	<pre>k_{obs} is too fast for detection/resolvable range (inhibition is not slow- binding)</pre>	Shorter minimal (pre)incubation time Higher [S] to increase competition (slower k_{obs}) Reduce [I] (slower k_{obs})

 Table 6
 I Troubleshooting and Optimization Experimental Assay Conditions.

Table 6 continues on the next page

Problem	Possible cause	Solutions		
F ₀ is not constant	Delay between enzyme addition and read-out	Reduce [E] (less substrate conversion during delay) Correcting t = 0 for actual time after addition Use injector in plate reader <u>Validate</u> row effect: change lay-out of plate (first well has higher F_0 than last well, but containing same components) and reduce number of samples in one measurement.		
	Fluorescence interference inhibitor	Validate: check F0 for inhibitor (no substrate and enzyme), substrate (no enzyme) and substrate and inhibitor (no enzyme) Exclude high [I] Background subtraction (subtract values substrate/inhibitor without enzyme from enzyme/substrate/inhibitor signal)		
	Pipetting error substrate	Check for bubbles when pipetting Use low-binding tips		
k _{obs} values are low compared to uninhibited control k _{ctrl}	Enzyme is unstable (high k _{ctri})	Optimize assay conditions to improve linearity of uninhibited control (lower $k_{\rm ctrl}$) Use preincubation protocol (<i>Method III & IV</i>): higher $k_{\rm obs}$ without competition		
	Enzyme is not reactive (low k _{obs})	Optimize buffer conditions to increase enzyme reactivity Add (fresh) reagents (e.g. DTT, ATP) in single-use aliquots Validate with different enzyme batch/construct Too many freeze/thaw cycles		
	Low inhibitor concentration ([I] << K _i ^{app})	Decrease [S] to reduce competition Increase [I] Use preincubation protocol <i>(Method III & IV)</i> : higher k _{obs} without competition		
	Slow reaction k _{obs}	Reduce [S] (less competition) Longer (pre)incubation time (t > 0.1t½) Use preincubation protocol (<i>Method III & IV</i>): higher k _{obs} without competition Optimize buffer conditions to increase enzyme reactivity		
k _{obs} vs. [I] is linear	Inhibitor has 1-step binding mode	Validate: Y-intercept = k_{ctrl} in k_{obs} vs. [I] plot Validate: $v_l = v_0^{ctrl}$ in [P], vs. t or v_t vs. t' plots Increase [I] to exclude 2-step [I] << K_1^{app} Decrease [S] to exclude 2-step [I] << K_1^{app}		
	2-step IRREV inhibitor is non-saturating ([I] << K _I ^{app})	Validate: Y-intercept = k_{ctrl} in k_{obs} vs. [I] plot Validate: $v_1 = v_0^{ctrl}$ in $[P]_t$ vs. t or v_t vs. t' plots Fit k_{obs} vs. [I] to linear function for combined value k_{inact}/K_1 Increase [I] Decrease [S] to reduce competition (lower K_1^{app}) Use preincubation protocol (Method III & IV): no competition		
	2-step IRREV inhibitor is saturating ([I] >> K _I ^{app})	Validate: Y-intercept > k_{ctrl} in k_{obs} vs. [I] plot Validate: $v_l < v_0^{ctrl}$ in $[P]_t$ vs. t or v_t vs. t' plots Decrease [I] Increase [S] to increase competition (higher K_1^{app})		
k _{obs} decreases with increasing [I]	Inhibitor concentration beyond resolvable range: noncovalent affinity is too potent ([I] >> K _i ^{app})	Optimize [I] range ($v_i = 0.1-0.9 \times v^{ctrl}$) Increase [S] (increase competition to increase K_i^{app}) Exclude unlikely values from fit		
	Incorrect formula to calculate k _{obs}	<u>Validate</u> if correct equation is used to determine k_{obs} ; reversible covalent/irreversible covalent, 1-step/2-step etc.		

Table 6	Troubleshooting and	Optimization	Experimental	Assav Conditions	(continued)
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6. Conclusion

The background of covalent inhibition kinetics and critical parameters for enzyme activity assays can be found in *section 2*. It is recommended to refer to this section before setting up your kinetic inhibition experiments as well as the core references by Copeland to get a general background on enzyme activity assays.³²⁻³³ We would like to reiterate that good experimental performance is essential for obtaining reliable parameters for your covalent inhibitor. Our kinetic simulation scripts can help validate the found values by 'rerunning' the experiment without human error or experimental artifacts. Not only will this give insight into the reliability of your assay, but it can also help to improve the assay setup and can show what wildly different values of concentrations would do for the read-out. In fact, figures in this manuscript have been created this way, and can as such be reproduced. Keep in mind that these are simulations, and real-life examples will always deviate due to machine artifacts or pipetting errors. Nevertheless, with a working activity assay and these instructions in hand, adequate analysis of covalent inhibitors should be very feasible.

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Data and Materials Availability

All figures in this work can be recreated with the information in **Table 1** and **Table 2**. Interactive versions of the simulation scripts are available free of charge at <u>https://mybinder.org/v2/gh/</u> <u>sroet/Elma/main?labpath=Landing%20Page.ipynb</u> (Full URL).

Author Contributions

<u>E.M.</u>: Conceptualization, Formal Analysis, Investigation, Visualization, Writing – Original Draft, Writing – Review & Editing. <u>S.R.</u>: Conceptualization, Resources, Software, Writing – Review & Editing. <u>R.Q.K.</u>: Supervision, Validation, Writing – Original Draft, Writing – Review & Editing. <u>M.P.C.M.</u>: Project Administration, Supervision, Writing – Review & Editing.

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8. Supporting Information

8.1. Symbols

E	Unbound enzyme
1	Unbound inhibitor
EI	Noncovalent enzyme–inhibitor complex
EI*	Covalent enzyme–inhibitor adduct
S	Unbound substrate
ES	Noncovalent enzyme-substrate complex
Р	(Detectable) product
<i>k</i> ₁	Second order association rate constant for $E + S \leftrightarrow ES$ reaction (in $M^{-1}S^{-1}$)
k ₂	First order dissociation rate constant for $E + S \leftrightarrow ES$ equilibrium (in s^{-1})
к ₃	Second order association rate constant for $E + I \leftrightarrow EI$ reaction (in M ⁻¹ s ⁻¹)
K ₄	First order dissociation rate constant for $E + 1 \leftrightarrow EI$ equilibrium (in s ⁻¹)
к ₅	First order association rate constant for EL \rightarrow EL* reaction (in s ⁻¹)
к ₆ Е	First order dissociation rate constant for EI \leftrightarrow EI equilibrium (in s) Detected signal reflecting product formation in presence of inhibitor after incubation t (in AU)
F ^{ctrl}	Detected signal reflecting product formation in the uninhibited control (in AII)
, F-	Background signal at reaction initiation (in All)
r _o	Product coefficient for detected signal per formed product (in AU/M)
V:	Initial product formation velocity in presence of inhibitor (in AU/s)
V.	Steady-state/final product formation velocity in presence of inhibitor (in AU/s)
V _{t'}	Product formation velocity after preincubation t' (in AU/s)
v ^{ctrl}	Product formation velocity in the uninhibited control (in AU/s)
$v_{t'}^{ctrl}$	Product formation velocity in the uninhibited control after preincubation t' (in AU/s)
v ₀ ^{ctrl}	Product formation velocity in the uninhibited control without preincubation: t'= 0 (in AU/s)
t	Incubation time after onset of product formation (in s)
ť'	Preincubation time after onset of enzyme inhibition (in s)
t½	Half-life for reaction progress (in s)
t½ _{diss}	Halt-life for dissociation reaction (in s)
τ	larget residence time (in s)
K _{obs}	Observed reaction rate constant (In s ⁻¹)
K _{max}	Maximum reaction rate constant at saturating inhibitor concentration for 2-step inhibition (in S^{-1})
k _{inact}	Indulvation rate constant for poplingarity or loss of enzyme activity in uninhibited control (in s^{-1})
k	Enzyme degradation rate constant for $F \rightarrow Fdeg$ (in s ⁻¹)
k .	Product formation rate constant for $E \rightarrow E + P$ (in s ⁻¹) at saturating substrate concentration
Keub	Reaction rate constant for $E + S \rightarrow E + P$ (in $M^{-1}s^{-1}$) (= k_{ev}/K_{M} if [S] << 0.1K_M)
k _{chom}	Reaction rate constant for $E + I \rightarrow EI^*$ of 1-step IRREV inhibitors (in $M^{-1}s^{-1}$)
k _{off}	Overall dissociation rate constant from bound to unbound enzyme $EI + EI^* \rightarrow E + I$ (in s ⁻¹)
K	Inhibition/dissociation constant (in M) for noncovalent E + I \leftrightarrow EI equilibrium of 2-step inhibition
K ^{app}	Apparent noncovalent inhibition constant (in M): with substrate competition
K [*]	Steady-state inhibition constant (in M) for E + I \leftrightarrow EI + EI* equilibrium of 2-step REV inhibition
K ^{*app}	Apparent steady-state inhibition constant (in M): with substrate competition
K	Inactivation constant for $E + I \rightarrow EI^*$ (in M) of 2-step IRREV inhibition
KIabb	Apparent inactivation constant (in M): with substrate competition
K _M	Michaelis-Menten constant for $E + S \rightarrow E + P$ (in M)
$K_{\text{inact}}/K_{\text{I}}$	Inactivation efficiency: reaction rate constant for $E + I \rightarrow EI^{+}$ of 2-step IRREV inhibitors (in M ⁻ S ⁻)
$1C_{50}$	Inhibitor concentration resulting in half-maximum inhibition after incubation time t (in M)
[Ftotal]	Combined total concentration of all anyme species ($F^{\text{total}} = F + F + $
[E]_	Unbound enzyme concentration at reaction initiation (before binding to inhibitor/substrate)
[1]	Unbound inhibitor concentration at onset of inhibition (before binding to enzyme)
[S]_	Unbound substrate concentration at onset of product formation (before binding to enzyme)
[EI]	Noncovalent El concentration at (steady-state) equilibrium
[X] ₀	Concentration of component X at reaction initiation (before binding to other reaction components)
$[X]_t$	Concentration of component X at incubation time t
[X] _{t'}	Concentration of component X at preincubation time t'
V _t	Incubation reaction volume containing enzyme, inhibitor and substrate $(V_t = V_t + V_{sub})$
V _{t'}	Preincubation reaction volume containing enzyme and inhibitor
V _{sub}	Volume containing substrate

8.2. Kinetic Parameters and Microscopic Rate Constants

	1-step REV	2-step REV	2-step IRREV	1-step IRREV
к _i (М)	$\frac{k_4}{k_3}$	$\frac{k_4}{k_3}$	$rac{k_4}{k_3}$	-
к _i * (М)	-	$\frac{k_4}{k_3 + \frac{k_3 k_5}{k_6}}$	-	-
К ₁ (М)	_	_	$\frac{k_4 + k_{\text{inact}}}{k_3}$	-
$k_{\rm off}$ (s ⁻¹)	k_4	$\frac{k_4 k_6}{k_4 + k_5 + k_6}$	0	0
k _{obs} (s ⁻¹)	$k_4 + k_3[I]$	$k_6 + \frac{k_5 \left[I \right]}{\left(\frac{k_4}{k_3} \right) + \left[I \right]}$	$k_{6} + \frac{k_{\text{inact}}\left[I\right]}{\left(\frac{k_{4} + k_{\text{inact}}}{k_{3}}\right) + [I]}$	k _{chem} [I]
Inactivation rate constant (M ⁻¹ s ⁻¹)	_	_	$\frac{k_{\text{inact}}}{K_{\text{I}}} = \frac{k_3 k_{\text{inact}}}{k_4 + k_{\text{inact}}}$	k _{chem}
t½ (s)	$\frac{\text{LN}(2)}{k_4 + k_3[1]}$	$\frac{\text{LN (2)}}{k_6 + \frac{k_5 [I]}{\left(\frac{k_4}{k_3}\right) + [I]}}$	$\frac{\left(\frac{k_4 + k_{\text{inact}}}{k_3} + [I]\right) \text{LN (2)}}{k_{\text{inact}}[I]}$	$\frac{\text{LN (2)}}{k_{\text{chem}}[1]}$
t¼ _{diss} (s)	$\frac{\mathrm{LN}\left(2\right)}{k_{4}}$	$\frac{(k_3 + k_5 + k_6) \ln(2)}{k_4 k_6}$	0	0
τ (s)	$\frac{1}{k_4}$	$\frac{\left(k_3+k_5+k_6\right)}{k_4\ k_6}$	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	0

 Table S1
 Kinetic parameters and microscopic rate constants for pseudo-first order reactions.



8.3. Similarities Michael-Menten Enzyme Kinetics and 2-step IRREV Inhibition

Figure S1 | Similarities and differences between MM enzyme kinetics and 2-step IRREV inhibition. Simulated for substrate **S1** with **KinSubDpl** (*left*) and 2-step IRREV inhibitor **C** with **KinGen** (*right*). **(A)** Binding mode. **(B)** Time-dependent product formation at $[S] = K_M$ (*left*) or $[I] = K_1$ (*right*). The biggest difference here is that a linear or an exponential model is fitted. **(C)** Secondary plot fits a bounded one-phase exponential equation. The maximum is driven by the rate of the second (irreversible) step and relevant kinetic parameters are derived from the maximum and the concentration at half-maximum. **(D)** Similarities in parameters for kinetic description of substrate hydrolysis for enzymes complying with Michaelis-Menten kinetics, and inactivation potency of 2-step IRREV inhibitors. K_0 = dissociation constant for noncovalent $E + S \leftrightarrow ES$ equilibrium (M). k_1 = association rate constant for $E + S \leftrightarrow ES$ equilibrium (M⁻¹s⁻¹). k_2 = dissociation rate constant for $E + S \leftrightarrow ES$ equilibrium (S⁻¹). k_{aat} = rate constant for irreversible $ES \rightarrow E + P$ reaction (s⁻¹). K_M = Michaelis constant for substrate hydrolysis (M). K_1 = inhibition constant for noncovalent $E + I \leftrightarrow EI$ equilibrium (M⁻¹s⁻¹). k_4 = dissociation rate constant for $E + I \leftrightarrow EI$ equilibrium (S⁻¹). k_{max} = maximum initial velocity (M/s). k_{obs} = [I]-dependent observed rate constant for 2-step inactivation (M). v_0 = [S]-dependent initial velocity (M/s). V_{max} = maximum initial velocity (M/s). k_{obs} = [I]-dependent observed rate constant for reaction from [EI*] = 00% (s⁻¹).

8.4. Linear vs. Nonlinear Regression

Linear regression methods are sometimes used to analyze preincubation-dependent enzyme inhibition (*Method III and IV*) because linear regression does not require dedicated graphical software, and benchmark protocols promoting linear regression originate from a time that computation with nonlinear regression was not readily available.⁸²⁻⁸³ Here, observed reaction rate k_{obs} is calculated from the linear slope of the natural logarithm of the percentage remaining enzyme activity against preincubation time (**Figure S2A**). The straight line enables relatively simple visual inspection of the fit. For readers that prefer this visual output, we recommend fitting the data by nonlinear regression but plotting the preincubation time-dependent enzyme activity on a semilog scale (**Figure S2B**).



Figure S2 | Comparison of linear and nonlinear regression illustrated with the data of inhibitor **C** in *Method III* (original data in **Figure 15C**). (**A**) Linear regression. Reaction rate k_{obs} is obtained from the negative slope of the natural logarithm of percentage enzyme activity against preincubation time t'. LN(enzyme activity) = LN(100% × v_t/v^{ctrl}). Negative values for LN(enzyme activity) corresponding with enzyme activity below 1% are excluded as assay sensitivity is normally insufficient to accurately distinguish 99% from 99.9% inhibition though this error will significantly affect the linear fit of k_{obs} . (**B**) Nonlinear regression. Plot preincubation time-dependent enzyme activity against preincubation time on a semilog scale for visual inspection of the fit.