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Received March 23, 2019; accepted June 22, 2019

ABSTRACT

An increased appreciation of the importance of optimizing drug-binding kinetics has lead to the development of various techniques for measuring the kinetics of unlabeled compounds. One approach is the competition-association kinetic binding method first described in the 1980s. The kinetic characteristics of the tracer employed greatly affects the reliability of estimated kinetic parameters, a barrier to successfully introducing these kinetic assays earlier in the drug discovery process. Using a modeling and Monte Carlo simulation approach, we identify the optimal tracer characteristics for determining the kinetics of the range of unlabeled ligands typically encountered during the different stages of a drug discovery program (i.e., rapidly dissociating, e.g., \( k_{\text{off}} = 10 \text{ minute}^{-1} \) low-affinity “hits” through to slowly dissociating e.g., \( k_{\text{off}} = 0.01 \text{ minute}^{-1} \) high-affinity “candidates”). For more rapidly dissociating ligands (e.g., \( k_{\text{off}} = 10 \text{ minute}^{-1} \)), the key to obtaining accurate kinetic parameters was to employ a tracer with a relatively fast off-rate (e.g., \( k_{\text{off}} = 1 \text{ minute}^{-1} \)) or, alternatively, to increase the tracer concentration. Reductions in assay start-time ≤1 second and read frequency ≤5 seconds significantly improved the reliability of curve fitting. Timing constraints are largely dictated by the method of detection, its inherent sensitivity (e.g., TR-FRET versus radiometric detection), and the ability to inject samples online. Furthermore, we include data from TR-FRET experiments that validate this simulation approach, confirming its practical utility. These insights into the optimal experimental parameters for development of competition-association assays provide a framework for identifying and testing novel tracers necessary for profiling unlabeled competitors, particularly rapidly dissociating low-affinity competitors.

Introduction

Historically, most drug discovery programs have relied upon equilibrium dissociation constant measurements when assessing the potential of future lead-drug candidates, rather than the kinetic parameters that comprise it (\( K_d = k_{\text{off}}/k_{\text{on}} \)). However, it is becoming more widely appreciated that optimizing the kinetics of drug binding (\( k_{\text{off}} \) and \( k_{\text{on}} \)) can enhance both compound efficacy and duration of action (Sykes et al., 2009; Guo et al., 2012; Copeland, 2016; Vauquelin, 2016). This has resulted in the introduction of novel methods to assess ligand binding kinetics at earlier stages in the drug discovery process, allowing the development of structure-kinetic relationships (Schmidtke et al., 2011; Georgi et al., 2018). A commonly applied method to assess the kinetics of unlabeled compounds is surface plasmon resonance (SPR) spectroscopy, but this is still not widely used for membrane proteins due to issues with protein purification and stability. To overcome this, competition-association kinetic binding assays using radioactive probes have been employed successfully to determine the kinetic values for both antagonists (Gillard and Chatelan, 2006; Slack et al., 2011; Fleck et al., 2012; Sykes et al., 2014, 2016) and agonists (Sykes et al., 2009; Sykes and Charlton, 2012) acting at a variety of G protein-coupled receptors at physiologic temperature and even more recently at non-G protein-coupled receptors targets (Yu et al., 2015).

Despite their undoubtable high sensitivity, the use of radioactive probes as tracers in kinetic competition binding assays presents a number of challenges. Classic radioactive binding requires the need for multiple washing steps to separate bound and free radioligand, adding complexity to the procedure and reducing throughput. Importantly, the wash step requirement also prevents any possibility of multiple single sample reads. More recently, scintillation

ABBREVIATIONS: HTRF, homogeneous time resolved fluorescence; PPHT-red, (±)-2-(n-phenethyl-n-propyl)amino-5-hydroxytetralin hydrochloride; 1-naphthalenol, 5,6,7,8-tetrahydro-6-[2-phenylethyl(propylamino)] derivative labeled with a red fluorescent probe; SPR, surface plasmon resonance; TR-FRET, time-resolved fluorescent resonance energy transfer.
proximity assays have been formulated that can be performed in homogeneous conditions, but signal-drift due to bead settling can complicate interpretation at early time points (Xia et al., 2016).

The emergence of time-resolved fluorescent resonance energy transfer (TR-FRET)-based methods (Schiele et al., 2015; Klein-Herenbrink et al., 2016) and bioluminescence based methods (Stoddart et al., 2018; Bouzo-Lorenzo et al., 2019) offers an alternative to radioactive binding assays and represents higher throughput methods to assess unlabeled ligand kinetics. With separation of bound and unbound label no longer necessary, multiple reads can be made from the same well (Emami-Nemini et al., 2013; Stoddart et al., 2015). Using the simple technique of TR-FRET we successfully characterized the binding of a series of dopamine D₂ specific agonists and clinically used antipsychotics employing a SNAP-tagged receptor labeled with terbium and a fluorescent tracer (Klein-Herenbrink et al., 2016; Sykes et al., 2017).

A key observation made during these studies was that the kinetic characteristics of the tracer had a profound effect on the reliability of the estimated kinetic parameters of unlabeled competitors (Klein-Herenbrink et al., 2016). This was most evident when using a slowly dissociating tracer to assess rapidly dissociating, low-affinity ligands. This represents a potential barrier to introducing successfully kinetic assays earlier in the drug discovery process (e.g., hit identification/ validation) as tracers need to be capable of determining the kinetics of low-affinity “hits” (in the micromolar range). Thus the aim of the current study was to use a modeling and simulation approach to identify the optimal tracer characteristics for determining the kinetics of a variety of unlabeled ligands typically encountered during different stages of a drug discovery program (i.e., low-affinity “hits” through to high-affinity “candidates”).

Using Monte Carlo simulations, we have assessed the ability of the global association method to accurately determine the kinetics of different model tracers. The global association method is a way of simultaneously fitting multiple tracer association curves (e.g., different tracer concentrations) to find best-fit $k_{\text{off}}$ and $k_{\text{on}}$ parameters across the whole dataset. Additionally, using Monte Carlo simulations we have explored the utility of these model tracers to predict the kinetic parameters of unlabeled ligands in competition-association binding mode. Furthermore, we have explored the influence of experimental design on the accuracy of kinetic parameter estimation, testing the impact of increasing tracer concentrations and investigating different plate reader configurations, particularly the influence of read frequency time and on-line (direct reagent injection into sample wells) versus off-line (reagent addition into plate prior to insertion into reader) reagent addition.

Finally we test the validity of this modeling approach through the detailed characterization of two dopamine D₂R specific fluorescent tracers, sipiperone-d2 and PPHT-red, assessing their ability to determine the kinetics of D₂R specific ligands with very varied kinetic characteristics, from the rapidly dissociating chlorpromazine to the very slowly dissociating butaclamol.

The analysis presented has important implications for the design of competition-kinetic approaches to assess unlabeled compound kinetics, providing a framework for the identification and testing of suitable tracers. In particular, we have identified the optimal tracer characteristics and experimental design for assessing low-affinity competitors, enabling the utilization of kinetics assays much earlier in the drug discovery process.

### Materials and Methods

**Determining the Kinetics of the Tracer for Use in Competition Kinetic Binding Studies**

Monte Carlo simulations are useful to investigate the behavior of a system under controlled situations and may be thought of as statistical experiments. They provide an estimation of variance for complex models, which ultimately helps researchers with experimental design and provides confidence in a particular experimental approach prior to testing. The underlying principle is to take a simulated dataset that is based on a set of “ideal” model parameters, add random error to the “dataset,” and then recreate the resulting dataset many times to obtain the parameters of interest. In this case the “dataset” comprises an XY table where X is time and Y is percentage of specific binding of the tracer. This process allows a frequency distribution to be built from the resulting parameter estimates, allowing an understanding of the associated error of each parameter estimate under the conditions employed (Christopoulos, 2001).

The competition association assay model (Motulsky and Mahan, 1984) relies on an accurate assessment of the kinetic properties of the labeled tracer, both the association rate constant ($k_{\text{on}}$ or $k_1$) and the dissociation rate constant ($k_{\text{off}}$ or $k_2$). Since the introduction of fitting software, such as GraphPad Prism (San Diego, CA), a popular experimental procedure to estimate the kinetics of the tracer is to monitor the binding of two or more different concentrations of tracer over time until equilibrium is reached; in doing so it is possible to calculate the $k_{\text{on}}$ and $k_{\text{off}}$ values of the tracer by simultaneously fitting all curves to generate global estimates for these rate parameters.

For the purposes of this study, tracer association was simulated using GraphPad Prism 6.0 employing the following equation, where $k_{\text{obs}}$ equals the observed rate of association and $k_{\text{on}}$ and $k_{\text{off}}$ are the association and dissociation rate constants, respectively, of the tracer:

$$k_{\text{obs}} = [L] + k_{\text{on}} + k_{\text{off}}$$

$$Y = Y_{\text{max}} \times (1 - \exp(-1 \times k_{\text{obs}} \times X))$$

In this globally fitted model of tracer binding, tracer concentrations ([L]) are fixed and $k_{\text{on}}$ and $k_{\text{off}}$ are shared parameters and independent of tracer concentration. Here, $Y$ is the level of receptor-bound tracer, $Y_{\text{max}}$ is the level of tracer binding at equilibrium, X is in units of time (e.g., minutes), and $k_{\text{obs}}$ (minute⁻¹) is the rate in which equilibrium is approached.

Tracer binding simulations were performed to assess the effect of on-line and off-line reagent addition on our ability to estimate accurately the kinetics of a variety of model tracers with varied kinetics parameters, consistent with the properties of compounds discovered in the various phases of the drug discovery cascade, the details of which are provided in Table 1. Tracer Monte Carlo simulations were performed using the following parameters: the $k_{\text{on}}$ and $k_{\text{off}}$ of each model tracer studied were allowed to vary, while the concentrations of tracer ([L]) employed were fixed at various multiples of the tracer’s affinity, specifically 30, 10, 3, 1, 0.3, 0.1 × $K_D$. Unless otherwise stated, the assay start time was fixed at either 1 second to mimic on-line addition of reagents via plate reader injectors or 30 seconds to mimic the delay in the time to read following off-line addition of membranes prior to insertion the assay plate into the plate reader. Read frequency (i.e., the time between each well read) was varied at 1, 5, 10, 20, and 60 seconds. Random error was added to the generated Y values by taking each theoretical (i.e., “correct”) value and...
Determining the Suitability of the Tracer for Competition Kinetic Binding Studies

The interactions of the tracer and unlabeled competitor with receptor is described by two differential equations that, when solved, yield a single equation describing the binding of the tracer as a function of time (Motulsky and Mahan, 1984). This model was used to simulate competition association curves to a variety of unlabeled compounds with different association-rate constants ($k_{on}$ or $k3$) and dissociation-rate constants ($k_{off}$ or $k4$), according to the following equations:

$$K_A = k_1[L] + k_2$$

$$K_S = k_3[I] + k_4$$

$$S = \sqrt{(K_A - K_S)^2 + 4k_1k_3[L][I]10^{-18}}$$

$$K_F = 0.5(K_A + K_S + S)$$

$$K_S = 0.5(K_A + K_S - S)$$

$$DIFF = K_F - K_S$$

$$Q = \frac{B_{max}K_L[L]10^{-9}}{DIFF}$$

(2)

where X = time (minutes), Y = specific binding (e.g., cpm or HTRF units), $k_1 = k_{on}$ tracer (molar$^{-1}$ minute$^{-1}$), $k_2 = k_{off}$ tracer (minute$^{-1}$), $[L] =$ concentration of tracer used (nanomolars), $[I] =$ concentration unlabeled ligand (nanomolars). Fixing the above parameters allows the following to be calculated: $k_3 =$ association-rate constant of unlabeled ligand (molar$^{-1}$ minute$^{-1}$), $k_4 = $ dissociation-rate constant of unlabeled ligand (minute$^{-1}$), $B_{max} = $ maximal specific binding of the system at equilibrium binding (e.g., cpm or HTRF units) (Motulsky and Mahan, 1984).

All competition kinetic association simulations were performed using the model tracers described in Table 1. These Monte Carlo simulation studies were designed to assess the effect of tracer kinetics on the ability of the Motulsky-Mahan model to distinguish compounds typically encountered in a drug discovery program with a wide range of kinetic off-rates ranging from the more rapidly dissociating (e.g., $k_{off} = 100$ minute$^{-1}$ representative of low-affinity fragments) to the more slowly dissociating (e.g., koff = 0.01 minute$^{-1}$ representative of a lead candidate compound).

Unless otherwise stated, the concentration of tracer ([L]) employed was fixed at 3 x $K_a$. The concentration of unlabeled competitor ([I]) was routinely fixed at various multiples of its own affinity, specifically 100, 30, 10, 3, 1 x $K_a$. Also taken into consideration were on-line versus off-line addition protocols and their effect on our ability to accurately estimate the kinetics of unlabeled compounds with varied kinetics parameters. Start time was fixed at either 1 second (to mimic on-line addition) or 30 seconds (to mimic off-line addition), and the read frequency was varied using either 1, 5, 10, 20, or 60 seconds unless otherwise stated.

Random error was added to the generated Y values by taking each theoretical (i.e., “correct”) value and adding to it a random number taken from a uniformly distributed population with an S.D. equal to one. The random error chosen for simulation was Gaussian absolute to directly reflect the pattern of error observed in our experimental data. The resulting simulated datasets were then individually fit to the competition association model (kinetics of competitive binding, Graph-Pad Prism 6.0) model and “Global (shared) parameters” including tracer $k_{off}$ and $k_{on}$ values were tabulated. In total, 200 simulations were performed per test condition.

To test the effect of tracer concentration on our ability to estimate the kinetic parameters of unlabeled compounds, we performed a further series of simulations, but this time we employed tracer concentrations ([L]) at 1 x, 10 x, and 30 x $K_a$. The concentrations of unlabeled competitor ([I]) were varied accordingly. This allowed us to test the effect of decreasing and increasing tracer competition on the accuracy of unlabeled compound kinetic determinations.

In another series of competition kinetic association simulations, the ability of a rapidly dissociating tracer to assess the kinetics of low affinity (1-1000 μM) unlabeled fragments was explored in both double and single concentration screening modes. All Monte Carlo simulations were performed using the following parameters: the $k_{on}$ and $k_{off}$ of the model tracer studied was fixed at 3 x $10^{-7}$ M$^{-1}$ minute$^{-1}$ and 10 minute$^{-1}$, respectively, with the concentration of tracer ([L]) employed fixed at 3 x $K_a$, while the concentrations of the fragments were fixed specifically at 10 and 100 μM (typical of a fragment screen where affinity of the fragments is unknown). Start time was fixed at 1 second (to mimic on-line addition), and a standard read frequency of 5 seconds was employed. The resulting simulated datasets were then individually fit to the competition association model and “Global (shared) parameters” including tracer $k_{off}$ and $k_{on}$ values were tabulated. In total, 200 simulations were performed per test condition. All the experiments described above assume that the Laws of Mass Action are obeyed and that ligand depletion does not occur under any of the conditions of tracer and competitor binding explored (Carter et al., 2007).

Monte Carlo Simulation Data Analysis

All Monte Carlo simulations were performed in Prism 6.0. In total, 200 simulations were performed per test condition using an associated error equivalent to an S.D. of one. “Global (shared) parameters” associated with eqs. 1 and 2 were recorded, and an outlier test (using an iterative Grubb’s test, α = 0.05) was performed on reported $k_{on}$ and $k_{off}$ values for tracers and unlabeled competitors. Outliers and ambiguous fits (those for which confidence intervals were extremely wide) were removed from the analysis before calculation of mean and % coefficient of variance (CV) values [i.e., (S.D./mean) x 100], which are indications of accuracy and variation around the expected input value. Mean fitting values were considered a reliably “good fit” if >90% of fits

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**TABLE 1**

Equilibrium affinity and kinetic properties of the tracer compounds used to construct Figs. 1–9 and Supplemental Figs. 1–7 and typical of the kinetic parameters of compounds discovered at all phases of the drug discovery screening cascade

<table>
<thead>
<tr>
<th>nM</th>
<th>Tracer $k_{off}$</th>
<th>Tracer $k_{on}$</th>
<th>Characteristic of Tracer</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.01</td>
<td>0.01</td>
<td>1 x 10$^9$</td>
<td>Very high-affinity candidate, very slow off</td>
</tr>
<tr>
<td>1</td>
<td>0.1</td>
<td>1 x 10$^8$</td>
<td>High-affinity candidate, slow off</td>
</tr>
<tr>
<td>100</td>
<td>1</td>
<td>1 x 10$^7$</td>
<td>Lead-like, fast off</td>
</tr>
<tr>
<td>300</td>
<td>10</td>
<td>3 x 10$^6$</td>
<td>Hit-like, very fast off</td>
</tr>
</tbody>
</table>
Fluorescent Ligand Binding Assays

Materials. PPHT ((±)-2-n-phenethyl-n-propylaminol-5-hydroxy-
tetralin hydrochloride;1-naphthalenol, 5,8,7,8-tetrahydro-6-(2-phenylethyl)
propyraminos) derivative labeled with a red fluorescent probe (PPHT-
red) was obtained from Cibisch Bioassays (Dagnolsur-Cesse, France).
Ninety-six-well polypropylene plates (Corning) were purchased from
Fisher Scientific UK (Loughborough, UK), and 384-well optiplate
plates were purchased from PerkinElmer (Beaconsfield, UK). GppNHp,
risperidone, chlorpromazine hydrochloride, and (+)-butaclamol used in
competition assays were obtained from Sigma-Aldrich (Poole, UK).
Dopemiderone and haloperidol hydrochloride used for competition
assays were obtained from Tocris Bioscience (Avonmouth, Bristol).
Perigole and ropinirole were kind gifts from Dr. Robert Lane (Monash
University, Melbourne, Australia), while bromocriptine was a kind gift
of Dr. Nicholas Holloway (Nottingham University, Nottingham, UK).

Determination of D2R fluorescent ligand binding kinetics. All
fluorescent binding experiments using PPHT-red and spiperone-d2 were
conducted in white 384-well Optiplate plates in assay binding
buffer, Hanks’ balanced salt solution containing 5 mM HEPES,
1% DMSO, and 0.02% pluronic acid pH 7.4, and 0.1 mM GppNHp.
GppNHp was included to remove the G protein-coupled population
of receptors that can result in two distinct populations of binding sites
in membrane preparations, since the Matkulsy–Mahan model is
only appropriate for ligands competing at a single site. In all cases,
nonspecific binding was determined in the presence of 10 μM
haloperidol.

To accurately determine association rate (k_on) and dissociation rate
(k_off) values, the observed rate of association (k_on) was calculated
using at least four different concentrations of either PPHT-red or
spiperone-d2. SNAP-tagged terbium-labeled human dopamine D2L
receptors expressed in CHO cell membranes (2 μg/well) were injected
into wells containing six different concentrations of the fluorescent
tracers PPHT-red (50–1.56 nM) or spiperone-d2 (10–0.3 nM) in a final
assay volume of 40 μL. A detailed description of the terbium labeling
procedure can be found in Sykes et al. (2017).

The degree of PPHT-red or spiperone-d2 bound to the receptor
was assessed at multiple time points by HTRF detection to allow
construction of association kinetic curves. The resulting data were
analyzed with the aid of the kinetic modeling (eq. 1) to derive a single
best-fit estimate for k_on and k_off. A detailed description of the terbium
labeling procedure can be found in Sykes et al. (2017).

Accurate Determination of Model Tracer Kinetic Parameters. To accurately calculate the kinetic on (k_on)
and off rates (k_off) of unlabeled competitor compounds, the
kinetic parameters of the tracer used must first be determined
by fitting tracer association data to a global kinetic model (see eq. 1). Simulations were performed for four model
tracers whose rates of dissociation differed up to 1000-fold (see Table 1).

For each tracer tested, a family of association kinetic curves
were constructed using six concentrations of each tracer,
ranging from ~30 to 0.1 × K_a, with each association curve
being monitored to the point of equilibrium, such that Y_max is
reached, allowing for the most accurate estimation of tracer
parameters, k_on and k_off (see Fig. 1, A and C).

To construct Fig. 1, A and C, using Monte Carlo simulations,
the k_off input value for the very rapidly dissociating tracer was
fixed at 3 × 10^-7 M^-1 min^-1 and k_off input value at
10 minute^-1. Figure 1, A and C, differ only in their initial
start time (representative of off-line injection) and 30 seconds
(representative of on-line addition), respectively, with read
frequency fixed every 10 seconds thereafter. What is immedi-
ately apparent under these simulation conditions is that the
equilibrium between a very rapidly dissociating tracer (k_off
10 minute^-1) and receptor is reached rapidly within the first
30 second, and, as a consequence, an accurate determination
of tracer association is improbable with a 30-second start time.
This is reflected in the graphical plots, Fig. 1, B and D,
showing the relationship between input tracer k_off and output
k_off withread frequency. With a start time of 1 second, tracer
kinetic determinations (data fits) are on the whole extremely
accurate, with a “good fit” achieved on >90% of the 200 fits (see
Fig. 1, B and C, and Supplemental Table 1). In contrast,
reliable determinations of tracer kinetics, as judged through
off-rate monitoring, following a 30-second start time repre-
sentative of off-line addition, are only possible when the off-
rate of the tracer is less than 1 minute^-1.

Also apparent is the effect of read frequency on the variation
in reported k_off values with shorter read frequencies, resulting

Non-specific binding was determined as the amount of HTRF signal
detected in the presence of haloperidol (10 μM) and was subtracted
from each time point, meaning that t = 0 was always equal to zero.
Each time point was conducted on the same 384-well plate incubated
at room temperature with orbital mixing (1 second of 100 rpm/cycle).

Data were globally fitted using eq. 2 to simultaneously calculate k_on
and k_off of the unlabeled compounds. Different competitor concentra-
tion ranges were chosen, as compounds with a long residence time
equilibrate more slowly, so a higher relative concentration is required to
ensure the experiments reach equilibrium within a reasonable time
frame (20 minutes), while still maintaining a good signal-to-noise.

Signal Detection

For the binding experiments described, signal detection was
performed on a Pherastrar FS (BMG Labtech, Offenburg, Germany)
using standard HTRF settings. The terbium donor was always excited
with up to 6 laser flashes at a wavelength of 337 nm. A kinetic TR-
FRET signal was collected at 5-second intervals at both 665 and 620
nm when using red acceptor. HTRF ratios were obtained by dividing
the acceptor signal (665 nm) by the donor signal (620 nm) and
multiplying this value by 10,000.

Results

Monte Carlo Simulations

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(i.e., >180/200 fits) were returned without being judged as either
outliers and/or ambiguous fits. Ambiguous fits are those for which
confidence intervals are extremely wide and therefore not reported
(http://www.graphpad.com/guides/prism/7/curve-fitting/index.htm).
Graphical representation of Monte Carlo data was also performed
in using R, and competition association data were fitted in Prism 6.0.

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in an improved quality of fit as highlighted by a reduction in the variability of the estimated output of tracer off-rate values. The results of Monte Carlo simulations using the kinetic association model equations are summarized in Supplemental Table 1. It must be emphasized that although our analysis focuses on the kinetic parameter $k_{\text{off}}$, the variability of $k_{\text{on}}$ of both tracer and competitor is also documented in Supplemental Tables 1-5.

**Competition Kinetic Binding between Tracer and Unlabeled Competitor Mimicking an Off-Line Addition Protocol**

To accurately determine the kinetics of unlabeled competitor compounds, it is conventional to construct a family of association kinetic curves using a fixed tracer concentration ($\sim 3 \times K_d$) and varying concentrations of the unlabeled compound with each association curve being monitored until equilibrium (see Fig. 2). In the case of the most slowly dissociating tracer, association curves were monitored for 180 minutes to reflect practical limitations. The resulting data were then fitted to the Motulsky–Mahan model, which describes the interaction between an unlabeled compound and a labeled tracer and allows us to calculate the $k_{\text{on}}$ and $k_{\text{off}}$ of the unlabeled compound (eq. 2).

Example results of Monte Carlo simulations using the Motulsky-Mahan model equation are shown in Fig. 2. In each case, the very rapidly dissociating tracer was employed; with input values of $k_{\text{on}}$ for the tracer fixed at $3 \times 10^7$ M$^{-1}$ minute$^{-1}$ and $k_{\text{off}}$ fixed at 10 minute$^{-1}$. Figure 2 differs only in respect of the kinetic properties of the unlabeled compound in competition with the tracer, with initial start times fixed at 1 second and with read frequency fixed every 10 seconds thereafter. Under these simulation conditions and in the presence of a rapidly dissociating unlabeled compound (with kinetic parameters; $k_{\text{off}}$ of 10 minute$^{-1}$, $k_{\text{on}}$ of $1 \times 10^4$ M$^{-1}$ minute$^{-1}$), equilibrium between the rapidly dissociating tracer, competitor compound, and receptor is reached rapidly within the first 30 seconds (see Fig. 2A).

In direct contrast in the presence of a more slowly dissociating competitor (with kinetic parameters; $k_{\text{off}}$ of 1 minute$^{-1}$, $k_{\text{on}}$ of $1 \times 10^7$ M$^{-1}$ minute$^{-1}$) then the time to equilibrium is markedly increased with a characteristic “overshoot” in the initial binding of the tracer (see Fig. 2B). The data contained in Fig. 2 represent a single simulation.
State of the art radioligand-based competition association binding assays are routinely formulated using only 12 time points (Sykes et al., 2010) employing either a single concentration of competitor (Sykes et al., 2014; Martella et al., 2017) or up to three concentrations of competitor (Sykes et al., 2009, 2010) and a tracer concentration in the range of 1–10× its own $K_d$. Current knowledge of FRET-based competition association binding assays is based on a small number of studies, which in general have employed an off-line addition protocol to improve experimental throughput but also to allow greater temperature control during the initial mixing step (Klein-Herrenbrink et al., 2016; Sykes et al., 2017; Sykes and Charlton, 2018; Sykes et al., 2018).

Figure 3 shows summary off-rate data from the Motulsky-Mahan model equation are summarized in Supplemental Tables 2–5. Figure 3 shows summary dissociation rate estimates markedly increases for the most rapidly dissociating unlabeled compounds examined (10–100 minute$^{-1}$), representing the limit of kinetic detection and is illustrative of the whole dataset, which is summarized in Fig. 3.

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State of the art radioligand-based competition association binding assays are routinely formulated using only 12 time points (Sykes et al., 2010) employing either a single concentration of competitor (Sykes et al., 2014; Martella et al., 2017) or up to three concentrations of competitor (Sykes et al., 2009, 2010) and a tracer concentration in the range of 1–10× its own $K_d$. Current knowledge of FRET-based competition association binding assays is based on a small number of studies, which in general have employed an off-line addition protocol to improve experimental throughput but also to allow greater temperature control during the initial mixing step (Klein-Herrenbrink et al., 2016; Sykes et al., 2017; Sykes and Charlton, 2018; Sykes et al., 2018).

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for this type of competition kinetic binding approach (see Supplemental Table 2). Also apparent was the influence of increasing read frequency on our ability to accurately predict the kinetics of unlabeled compounds with slower dissociation rates in the range of 0.01 minute$^{-1}$. Increased variation likely reflects the imprecise fitting of the tracer “overshoot,” which is reliant on early time points for accuracy. In contrast, a tracer with a $k_{\text{off}}$ value equal to 1 minute$^{-1}$ is seemingly suitable for the determination of off-rates in the range of $\sim 10$ minute$^{-1}$, but increasing variation in the mean is observed as the read frequency is increased from 1 to 60 seconds (see Fig. 4B). Practically this wider degree of variation in the mean will become especially apparent if the number of observations for a particular compound is kept low (i.e., $n \geq 4$). A different pattern is observed for a tracer with a $k_{\text{off}}$ value equal to 0.1 minute$^{-1}$, as shown in Fig. 4C, for compounds with rapid off-rates between 10 and 100 minute$^{-1}$ the number of ambiguous fits is very high (see Supplemental Table 4), and it is only compounds with a $k_{\text{off}}$ value of $\leq 1$ minute$^{-1}$ that can be measured without ambiguity at this concentration of tracer employed (i.e., $3 \times K_d$). Interestingly the appearance of ambiguity in fitting estimates precedes any inaccuracies in the fitting estimates themselves. Finally a kinetic tracer with a $k_{\text{off}}$ of 0.01 minute$^{-1}$ employed at a concentration of $3 \times K_d$ is only able to determine unambiguously (and accurately) the kinetics of compounds with $k_{\text{off}} \leq 1$ minute$^{-1}$ when the read frequency is shortened to 1 second (see Fig. 4D) and as a consequence is of limited value as a tracer for kinetic determinations of more rapidly dissociating compounds.

The figures that are derived from these Monte Carlo simulations are useful guides to enable investigators interested in compound kinetics and associated structure activity relationships to set up kinetic screens and choose appropriate tracers to profile the properties of unlabeled compounds. To illustrate some key points, Monte Carlo simulation results reproducing the experimentally observed effect of competitor off-rate on the competition profile observed with a rapidly dissociating tracer and a slowly dissociating tracer with varying start time are depicted in Fig. 5. Figure 5A shows competition between a fixed concentration ($3 \times K_d$) of a very rapidly dissociating tracer with the following kinetic parameters, $k_{\text{off}}$ of 1 minute$^{-1}$, $k_{\text{off}}$ of $3 \times 10^3$ M$^{-1}$ minute$^{-1}$ (A); a rapidly dissociating tracer with the following kinetic parameters, $k_{\text{off}}$ of 0.1 minute$^{-1}$, $k_{\text{off}}$ of $1 \times 10^3$ M$^{-1}$ minute$^{-1}$ (B); a slowly dissociating tracer with kinetic parameters, $k_{\text{off}}$ of 0.1 minute$^{-1}$, $k_{\text{off}}$ of $1 \times 10^2$ M$^{-1}$ minute$^{-1}$ (C); and a very slowly dissociating tracer with kinetic parameters, $k_{\text{off}}$ of 0.01 minute$^{-1}$, $k_{\text{off}}$ of $1 \times 10^2$ M$^{-1}$ minute$^{-1}$ (D). Blue symbols represent conditions that returned $>90\%$ reliable fits. Red symbols represent conditions that returned $<90\%$ reliable fits. In all cases tracer simulations were performed with an initial start time of 30 seconds representative of off-line addition of receptor to a reaction containing free tracer and unlabeled competitor, absolute values are the average of 200 simulations.

![Monte Carlo simulation results](image-url)
start time, representative of off-line addition. The increased start time has little effect on the ability to discriminate between the kinetic off-rates of these compounds primarily as the tracer itself is slow to reach equilibrium with the receptor.

Figure 5B shows competition between a fixed concentration (3 × $K_d$) of a rapidly dissociating tracer with kinetic parameters, $k_{\text{off}}$ of 10 minute$^{-1}$, $k_{\text{on}}$ of $3 \times 10^7$ M$^{-1}$ minute$^{-1}$ (A); a rapidly dissociating tracer with kinetic parameters, $k_{\text{off}}$ of 1 minute$^{-1}$, $k_{\text{on}}$ of $1 \times 10^8$ M$^{-1}$ minute$^{-1}$ (B); a slowly dissociating tracer with kinetic parameters, $k_{\text{off}}$ of 0.1 minute$^{-1}$, $k_{\text{on}}$ of $1 \times 10^7$ M$^{-1}$ minute$^{-1}$ (C); and a very slowly dissociating tracer with kinetic parameters $k_{\text{off}}$ of 0.01 minute$^{-1}$, $k_{\text{on}}$ of $1 \times 10^6$ M$^{-1}$ minute$^{-1}$ (D). Blue symbols represent conditions that returned >90% reliable fits. Red symbols represent conditions that returned <90% reliable fits. In all cases tracer simulations were performed with an initial start time of 1 second representative of injection of receptor to a reaction containing free tracer and unlabeled competitor; absolute values are the average of 200 simulations. $k_{\text{obs}}$, $k_{1/2}$ values (0.693/$k_{\text{obs}}$) for the tracers with dissociation rates of 10, 1, 0.1, and 0.01 minute$^{-1}$ were 1.1, 10.4 seconds and 1.7 and 17.3 minutes, respectively.

In line with previous kinetic studies (Klein-Herenbrink et al., 2016), the $k_{\text{off}}$ of unlabeled compounds was poorly estimated when the tracer $k_{\text{off}}$ is substantially slower than that of the unlabeled ligand. Increasing the $k_{\text{off}}$ of the tracer from 0.1 to 10 vastly improved the precision with which the values of $k_{\text{on}}$ and $k_{\text{off}}$ were estimated for the more rapidly dissociating compounds (as illustrated in Supplemental Tables 2–5). In contrast, $k_{\text{on}}$ had no impact on the accuracy of the parameter estimate reflective of the fact that changes in $k_{\text{on}}$ are completely compensated for by changes in the ligand concentration (data not shown), i.e., no change in $k_{\text{obs}}$.

**Effect of Tracer Concentration on Kinetic Determination**

The effect of tracer concentration on the accuracy of kinetic determinations was explored in a further series of Monte Carlo simulations. Tracer concentration was both lowered to 1 × $K_d$ and increased to 10 and 30 × $K_d$. We chose to highlight a tracer with kinetic properties commonly encountered in drug-receptor screening campaigns: high affinity (1 nM), fast on $1 \times 10^8$ M$^{-1}$ minute$^{-1}$, and slow off 0.1 minute$^{-1}$. The results obtained with this tracer in competition with compounds with kinetic off-rates ranging from 100 to 0.01 minute$^{-1}$ following an on-line addition protocol are shown in Fig. 6.
Simulations varying concentration following an on-line addition protocol were also performed for tracers with off-rates of 10, 1, and 0.01 as shown in Supplemental Figs. 1–3. $k_{obs}$ values (0.693/$k_{obs}$) for the tracers with dissociation rates of 10, 1, 0.1, and 0.01 minutes$^{-1}$ were 1.1, 10.4 seconds, 1.7, and 17.3 minutes, respectively, at tracer concentrations 3 × their respective $K_d$.

As a generality, an increase in ligand concentration was associated with an improvement in both the accuracy of kinetic determinations but also the number of fits deemed unambiguous and, in some cases, effectively rescued the ability of a tracer to determine the kinetics of a more rapidly dissociating competitor. For example, a competitor with an off-rate of 10 minutes$^{-1}$ competing with a tracer with an off rate of 0.1 minutes$^{-1}$ could be readily resolved when the tracer concentration was increased from 3 × $K_d$ (Fig. 6B) to 30 × $K_d$ (Fig. 6D). $k_{obs}$ values for a tracer with a dissociation rate of 0.1 minutes$^{-1}$ ranged from 3.5 to 0.2 minutes at concentrations of tracer 1–30 × $K_d$, respectively. Data for a competitor with an off-rate of 10 minutes$^{-1}$ competing with different concentrations of tracers with off-rates ranging from 0.01 to 10 minutes$^{-1}$ are summarized in Fig. 7.

Similarly the effect of tracer concentration was explored following an off-line addition protocol with the data summarized in Supplemental Figs. 4–7. In the case of a tracer with an off-rate of 0.1 minute$^{-1}$, increasing tracer concentration with the online method did not improve the goodness of fit when quantifying competitors with off-rates in the region of 10 minutes$^{-1}$, but tuning this particular experimental variable may still offer some benefit to researchers without access to detection instruments with injectors, reducing the overall error associated with kinetic determinations.

**Fragment Screening Using Competition-Kinetic Association Binding Assays**

Monte Carlo simulations were performed to recreate the competition profile observed with low-affinity fragments (31 in total with $K_d$ values ranging from 1 to 1000 μM). These competitor fragments with varied kinetic parameters ($k_{off}$ ranging from 3 to 200 minutes$^{-1}$ and $k_{on}$ from 3 × 10$^{6}$ to 2 × 10$^{8}$ M$^{-1}$ minute$^{-1}$) were simulated in competition with a fixed concentration (3 × $K_d$) of a rapidly dissociating tracer ($k_{off}$ of 10 minutes$^{-1}$, $k_{on}$ of 3 × 10$^{7}$ M$^{-1}$ minute$^{-1}$). Supplemental Fig. 8A shows the expected inhibition of tracer binding caused by a low-affinity competitor, fragment 25 (affinity of 0.25 mM, $k_{off}$ of 87.1 minutes$^{-1}$, $k_{on}$ of 5 × 10$^{5}$ M$^{-1}$ minute$^{-1}$) tested at two concentrations, 10 and 100 μM. Using a start time of 1 second and a read frequency of 5 seconds, we were able to estimate the affinity of such fragments with a high degree of accuracy as...
shown by the correlation of input $K_d$ with output $K_d$ (see Supplemental Fig. 8B). In contrast, as the $k_{off}$ of the fragment increases then the degree of error associated with the estimation of $k_{off}$ and $k_{on}$ is increased (as judged by the increase in the overall S.D. associated with the mean); however, the overall correlation of input and output kinetic parameters remains extremely good (see Supplemental Fig. 8, C and D). In all cases tracer simulations were performed with an initial start time of 1 second representative of injection of receptor to a reaction containing free tracer (L) and unlabeled competitor (I), absolute values are the average of 200 simulations. $k_{obs}$ values (0.693$/k_{off}$) for a tracer with a dissociation rate of 0.1 minute$^{-1}$, with increasing concentration of tracer 1, 3, 10 and 30 times $K_d$ were 3.5, 1.7, 0.6 and 0.2 minute respectively.

Fig. 6. Monte Carlo simulation results exploring the effect of tracer concentration and sample time on the accurate determination of competitor $k_{off}$ representative of an injection protocol. Effect of assay read frequency on measured $k_{off}$ of unlabeled competitor compounds with varied kinetics in competition with different concentrations of a slowly dissociating tracer with kinetic parameters, $k_{off}$ of 0.1 minute$^{-1}$, $k_{on}$ of $1 \times 10^8$ M$^{-1}$-minute$^{-1}$. Tracer concentrations were $1 \times K_d$ (A), $3 \times K_d$ (B), $10 \times K_d$ (C), and $30 \times K_d$ (D). Blue symbols represent conditions that returned >90% reliable fits. Red symbols represent conditions that <90% reliable fits. In all cases tracer simulations were performed with an initial start time of 1 second representative of injection of receptor to a reaction containing free tracer (L) and unlabeled competitor (I), absolute values are the average of 200 simulations. $k_{obs}$ values (0.693$/k_{off}$) for a tracer with a dissociation rate of 0.1 minute$^{-1}$, with increasing concentration of tracer 1, 3, 10 and 30 times $K_d$ were 3.5, 1.7, 0.6 and 0.2 minute respectively.

Experimental Validation of the Monte Carlo Simulation Approach

To validate the findings of these Monte Carlo simulations exploring the effect of tracer characteristics on kinetic determinations using the Motulsky-Mahan approach, we undertook an experimental study designed to explore the limits of a previously well characterized system, the dopamine D2L receptor using a TR-FRET-based system. In this study we chose to focus on a small number of compounds with diverse kinetics and employ the BMG pherastar FS (plate reader) injectors to make an assessment of tracer binding at the very earliest time points.

Determining the Kinetic of D2R Fluorescent Tracers

Representative kinetic association curves for PPHT-red and spiperone-d2 using a start time of ~1 second and a read frequency of 5 seconds are shown in Fig. 8, A and C, respectively. The kinetic parameters determined from these plots are presented in Table 2. PPHT-red dissociated rapidly from the dopamine D2R with a $t_{1/2}$ of 2.2 minutes (0.693$/k_{off}$), whereas spiperone-d2 dissociated more slowly ($t_{1/2}$ of 8.7 minutes). The removal of data points to reflect off-line addition (i.e., the first ~30 seconds) or increasing read frequencies
had little influence on the determination of the kinetic parameters of these two fluorescent tracers reflective of the fact that they are not so rapidly dissociating that the early time points are critical for an accurate estimation of their kinetic parameters. Figure 8, B and D, show the expected linear relationship between PPHT-red and spiperone-d2 concentration and the observed rate of association (or $k_{obs}$) and suggest that the Laws of Mass Action are observed.

Effect of Tracer Kinetics on Dopamine D2R Ligand Kinetic Determinations

To experimentally test the effect of tracer kinetics on $k_{off}$ estimates of unlabeled ligands, competition association experiments were performed for five dopamine D2 ligands: chlorpromazine, ropinerole, pergolide, domperidone, and butaclamol (Fig. 9). The associated errors and therefore the accuracy of kinetic determinations for rapidly dissociating compounds is highly dependent on the kinetic properties of the tracer employed to measure them. In contrast, the error associated with more slowly dissociating compounds is largely comparable and seemingly independent of the tracer's kinetic properties. This is illustrated in Fig. 9 and shows that the error associated with kinetic determinations made with the more slowly dissociating tracer spiperone-d2 is in general much larger than for the more rapidly dissociating tracer PPHT-red. This is evident from the spread of $k_{off}$ values on the x-axis (spiperone-d2) being wider than the spread of the $k_{off}$ values on the y-axis (PPHT-red). The errors associated with kinetic determinations of rapidly dissociating compounds following off-line addition of PPHT-red (Fig. 9A) is little changed compared with on-line addition (Fig. 9B); however, there was a tendency for the error to increase for the more slowly dissociating compounds and this situation is replicated for spiperone-d2 (see Fig. 9) and potentially represents the imprecise fitting of the tracer "overshoot."

Discussion

The growing awareness of the importance of optimizing drug-binding kinetics has led to a rapid increase in the development and utilization of assay systems capable of measuring the kinetics of unlabeled compounds. A popular format for investigating membrane-bound targets is the competition association binding assay, first described by Arányi (1980) and then popularized by Motulsky and Mahan (1984) and used extensively to characterize many different receptor systems (Gillard et al., 2002; Dowling and Charlton, 2006; Sullivan et al., 2006; Fleck et al., 2012; Sykes et al., 2014;
A key observation during this time has been that the accuracy of estimating kinetic parameters of rapidly dissociating molecules is poorer when using a slowly dissociating ligand as the tracer (Sykes and Charlton, 2012; Klein-Herenbrink et al., 2016; Bosma et al., 2019).

The purpose of the current study was to explore the effect of tracer kinetics on our ability to accurately estimate the kinetics of unlabeled compounds using the competition-association binding method. In particular, we were interested to test the limits of this model system in terms of its ability to assess the binding of very rapidly dissociating compounds likely to be representative of compounds identified in a screening campaign. In practical terms, we were also keen to investigate the importance of read frequency to assist with experimental design. To our knowledge, the current study is the first to fully explore these elements and provide clear guidance for the use of this assay format at all stages of the drug discovery process.

As shown in Fig. 1 and Supplemental Table 1, it was generally possible to derive accurate estimates (<2-fold difference) of the tracer kinetic parameters using the global association eq. 1 for simulated datasets. For the more rapidly dissociating tracers, the key to obtaining more accurate kinetic parameters was to reduce the assay start time and the read frequency. Our ability to control these parameters is largely instrument (reader) dependent. For example, assay start time is dependent upon the ability to inject sample while simultaneously reading from the same well. In addition, assay sensitivity determines the required sample acquisition time, which restricts minimal read frequency. Consequently the option to vary these parameters can be considered as critical factors in the process of determining unlabeled compound kinetics. This is readily illustrated in a comparison of Fig. 1, B

![Fig. 8. Determination of PPHT-red and spiperone-d2 kinetic binding parameters.](image)

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**TABLE 2**

Kinetic binding parameters of the tracers PPHT-red and spiperone-d2 determined from association binding studies using human dopamine D2L CHO cell membranes

<table>
<thead>
<tr>
<th>Tracer</th>
<th>Tracer $K_D$ (nM)</th>
<th>Tracer $k_{off}$ (min$^{-1}$)</th>
<th>Tracer $k_{on}$ (M$^{-1}$ min$^{-1}$)</th>
<th>No. of Observations</th>
</tr>
</thead>
<tbody>
<tr>
<td>PPHT-red</td>
<td>16.9 ± 1.1</td>
<td>0.32 ± 0.02</td>
<td>1.93 ± 0.21 $\times 10^7$</td>
<td>4</td>
</tr>
<tr>
<td>Spiperone-d2</td>
<td>0.62 ± 0.08</td>
<td>0.08 ± 0.01</td>
<td>1.30 ± 0.17 $\times 10^6$</td>
<td>4</td>
</tr>
</tbody>
</table>
Providing an initial start time of 1 second was used (representative of on-line injection), it was possible to determine accurately the kinetics of all model tracers with less than 10% variation around the mean. In contrast, employing a start time of 30 seconds prevented accurate kinetic determinations for the most rapidly dissociating tracer even when the read frequency was restricted to 1 second (Fig. 2, C and D), demonstrating the importance of on-line injection.

The situation with the Motulsky-Mahan eq. 2 for the estimation of unlabeled compound kinetics was a little more complicated. Although it was possible to derive accurate rate constant estimates for the majority of conditions, there were some combinations that failed to return reproducible estimates (Supplemental Tables 2–5). In general, employing a tracer concentration at $3C_2$, its $K_d$ was not possible to estimate reliably $k_{on}$ and $k_{off}$, where the dissociation of the tracer was ($\geq$10-fold) slower than the unlabeled competitor. For example, the accurate estimation of the rate constants of a rapidly dissociating ligand with $k_{off}$ of 100 minute$^{-1}$ requires a tracer that is also rapidly dissociating, i.e., in the range of 10 minute$^{-1}$. The initial read time was also critically important to determine the kinetics of unlabeled compounds. Where the simulations were started from 1 second, mimicking an on-line injection protocol, the kinetic parameters of unlabeled compounds were generally accurately estimated. In contrast, when measurements were started (sampled) 30 seconds after the beginning of the experiment, mimicking off-line addition, the estimates were poorer, particularly for rapidly dissociating ligands. This highlights the importance of early time points measured before equilibrium is established. This problem is exacerbated as read frequency is increased since definition on the tracer association curves are lost. In contrast, a short read frequency is associated with an increase in the number of data points, which is useful from an accuracy perspective (as illustrated by reduced % CV values; see Supplemental Table 2). However, short read frequency will have a negative impact on throughput, which is an important consideration especially when profiling hundreds to thousands of compounds during screening.

To date, SPR has been the main method for measuring kinetics of fragments at receptors; however, this technique is traditionally limited to artificially stabilized receptors (Shepherd et al., 2014). Thus the competitive binding model presented is an attractive alternative to SPR and should theoretically allow the investigator to reveal the kinetics of low-affinity fragments with off-rates in the order of 10 minute$^{-1}$ should an appropriate tracer be identified (see Supplemental Figs. 8 and 9).

In a previous paper exploring dopamine D$_2$R agonist kinetics, we were able to demonstrate the importance of tracer properties on our ability to determine the kinetics of rapidly dissociating ligands (Klein-Herenbrink et al., 2016). In the current study, we have further explored this observation employing an on-line injection protocol. In the previous study, [3H]-spiperone was unable to accurately determine the kinetics of the most rapidly dissociating D$_2$R agonists. This is likely...
the result of a number of factors, including its slower measured off-rate from the dopamine D2R, the relatively lower concentration of spiperone employed in the competition binding experiments, i.e., 3 × [3H]-spiperone (versus 10 × K_d spiperone-d2 in the current study) and the decision to employ an initial start time of 30 seconds.

As predicted by the Monte Carlo simulations, PPHT-red, a tracer with a relatively fast off-rate, was more reliable at determining the kinetic off-rates of the most rapidly dissociating antagonists and agonists tested in this study. In contrast, spiperone-d2 (10 × K_d), although adequate at determining the off-rates of the more slowly dissociating compounds, was prone to more variation in its determination of the off-rates of the most rapidly dissociating compounds. Importantly what this study clearly demonstrates is that a failure to demonstrate an accurate fit of the kinetics of a rapidly dissociating compound (10 minute^{-1}) can be overcome through the use of higher concentrations of a slowly dissociating tracer and the decision to employ a shortening start and read frequency (see Fig. 7). As one might predict, the use of higher tracer concentrations could not compensate for an increase in the assay start time, representative of an offline addition protocol (see Supplemental Fig. 6).

What is apparent from this study is that for a competitive binding approach to be used throughout the different phases of the drug discovery process, then the kinetics of the tracer need to be tailored to the appropriate properties of the unlabeled compounds. For low-affinity compounds, such as initial hits or fragments, a rapidly equilibrating tracer is required to accurately determine rate constants. This is fortunate as it opens the possibility to fluorescently label an early hit to serve as a tracer to discover and characterize new fragments. In this format, it would be necessary to read relatively small numbers of wells in rapid succession (i.e., a shortened read frequency) using a repeat on-line injection protocol to resolve the kinetics of the most rapidly dissociating fragments. Also apparent is that as we move further along the drug discovery pathway toward lead optimization, it may be beneficial to label a more slowly dissociating compound, allowing the off-line addition of membranes and increased sample throughput through extended read frequency. The decision to employ shorter read times is ultimately a compromise between throughput and the accurate resolution of kinetic parameters.

In summary, we have improved the current understanding of the Motulsky-Mahan approach, providing clear guidelines on the use of tracers to measure the kinetics of unlabeled competitors. Based on the detailed Monte Carlo approach presented, we propose the following four factors as being important considerations when formulating competition association binding experiments:

1. Tracer kinetics: fast off tracers in the region of 0.1–1 minute^{-1} appear to be critical for the determination of unlabeled compounds with rapid off kinetics in the region of 1–10 minute^{-1}.
2. Tracer concentration appears to play a crucial role in our ability to determine the kinetics of binding with greater accuracy and can even increase the range of compounds off rates that we can reliably measure.
3. Online injection capability proved critical to determine the off-rates of compounds and tracers that dissociate with off-rates in the region of 10 minute^{-1}.
4. Rapid read frequency dramatically improves the goodness of fit and reduces experimental variability and, like online injection, can rescue our ability to resolve the kinetics of the most rapidly dissociating compounds.

Experimental conditions can also be manipulated to enhance our ability to measure the kinetics of a particular tracer or competitor ligand. For example, reducing assay temperature will slow the off-rates of both the tracer and the competitor compound, a ploy that has been successfully used in the past to enable the determination of off-rates of more rapidly dissociating compounds using the off-line addition protocol (Contreras et al., 1986, Guo et al., 2012). It should be noted, however, that kinetic parameters calculated at lower temperatures are unlikely to reflect those in a physiologic system, significantly limiting their translational utility.

Overall, the findings in this paper highlight the importance of considering tracer kinetics and assay read start and read frequency when developing competition association assays. Notably, these simulations suggest that under the right conditions, the kinetic parameters of very low-affinity (millimolar) competitors can be measured, providing the opportunity for kinetic fragment-based receptor screens and the development of structure-kinetic relationships at all phases of the drug discovery cascade.

Acknowledgments
We thank Dr. Nick Holliday for his helpful suggestions during the preparation of this manuscript. In addition, we thank MBG Labtech Ltd. for their expert technical assistance and for personal sponsorship of David Sykes.

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Participated in research design: Sykes, Charlton.
Conducted experiments: Sykes.
Contributed new reagents or analytic tools: Jain.
Performed data analysis: Sykes, Jain.
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