A live cell NanoBRET binding assay allows the study of ligand-binding kinetics to the adenosine A_3 receptor

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Abstract

There is a growing interest in understanding the binding kinetics of compounds that bind to G protein-coupled receptors prior to progressing a lead compound into clinical trials. The widely expressed adenosine A_3 receptor (A_3AR) has been implicated in a range of diseases including immune conditions, and compounds that aim to selectively target this receptor are currently under development for arthritis. Kinetic studies at the A_3AR have been performed using a radiolabelled antagonist, but due to the kinetics of this probe, they have been carried out at 10 °C in membrane preparations. In this study, we have developed a live cell NanoBRET ligand binding assay using fluorescent A_3AR antagonists to measure kinetic parameters of labelled and unlabelled compounds at the A_3AR at physiological temperatures. The kinetic profiles of four fluorescent antagonists were determined in kinetic association assays, and it was found that XAC-ser-tyr-X-BY630 had the longest residence time (RT = 288 ± 62 min) at the A_3AR. The association and dissociation rate constants of three antagonists PSB-11, compound 5, and LUF7565 were also determined using two fluorescent ligands (XAC-ser-tyr-X-BY630 or AV039, RT = 6.8 ± 0.8 min) as the labelled probe and compared to those obtained using a radiolabelled antagonist ([\(^3\)H]PSB-11, RT = 44.6 ± 3.9 min). There was close agreement in the kinetic parameters measured with AV039 and [\(^3\)H]PSB-11 but significant differences to those obtained using XAC-S-ser-S-tyr-X-BY630. These data indicate that selecting a probe with the appropriate kinetics is important to accurately determine the kinetics of unlabelled ligands with markedly different kinetic profiles.

Keywords Adenosine A_3 receptor · Binding kinetics · Residence time · NanoBRET

Introduction

The nucleoside adenosine is a ubiquitous signalling molecule which modulates cellular responses to stress. Upon cellular stress caused by mechanical, inflammatory or hypoxic stressors, high concentrations of ATP are released [1, 2] which are rapidly hydrolysed to adenosine [3]. Adenosine signals via four related family A G protein-coupled receptors (GPCRs) [4] of which the adenosine A_3 receptor (A_3AR) is thought to play an important role in the control of infection and related inflammation due to its expression on immune cells [5]. It has also been proposed to play a cardio- [6, 7] and neuroprotective [8, 9] role. In addition, the A_3AR receptor has been shown to be expressed in a variety of cancer cell lines, and since the tumour micro-environment is often hypoxic, it may play a role in tumour progression [10]. This has led to an interest in developing molecules that target the A_3AR in the treatment of immune conditions and cancer [11].

In recent years, many molecules targeting GPCRs, including those acting at adenosine receptors [12], have exhibited a lack of efficacy in clinical trials [13]. The optimization of molecules for clinical trials has historically focused on developing a candidate with high affinity and selectivity for the target receptor, and these parameters are often measured at equilibrium in model cell systems over-expressing the receptor of interest [14, 15]. Within the last 10 years, it has become clear that measuring the binding properties of a molecule at
equilibrium may not be the most effective way to determine potential in vivo efficacy and that determining the duration of protein-drug interactions may be a better predictor of in vivo action [16, 17]. As the equilibrium dissociation constant \( (K_d) \) of a ligand at a GPCR is a function of the ligand’s association \( (k_{on}) \) and dissociation rate \( (k_{off}) \) constants \( (K_d = k_{off}/k_{on}) \), molecules that have the same measured \( K_d \) at equilibrium can have markedly different \( k_{on} \) and \( k_{off} \) rate constants. To address this, Copeland et al. (2006) introduced the concept residence time (RT), which is the reciprocal of the dissociation rate constant \( (RT = 1/k_{off}) \), as a measure of the duration of drug-target complex formation [18]. Depending on the clinical setting, different RTs may be required. For example, drugs with a long RT are preferred when an extended duration of action is required and can reduce administration to once a day. This has been demonstrated for the M3 muscarinic receptor antagonist tiotropium and the \( \beta_2 \) adrenoceptor agonist olodaterol which both have long RTs and are once a day bronchodilators used to treat chronic obstructive pulmonary disease [14, 19, 20]. Short RT ligands are advantageous when the rapid, i.e. seconds to minutes, replacement of the drug by an endogenous ligand is crucial to avoid long-term side effects which has been proposed to be the case for drugs targeting the dopamine D2 receptor [17, 21, 22].

A variety of techniques exist to measure the kinetic parameters of molecules binding to the protein target of interest. As GPCRs are integral membrane proteins, biophysical techniques such as surface plasmon resonance which require purified protein can be challenging and often require mutagenesis to stabilise the receptor and allow it to be purified within lipid micelles or nanodiscs [23, 24]. Techniques that do not require the receptor to be purified include those that use radiolabelled or fluorescently labelled ligands [25, 26]. One limitation of these techniques is that they can only directly measure the kinetic parameters of the labelled compound. To overcome this, the methodology proposed by Motulsky and Mahan is often employed [27]. This technique measures the association kinetics of a labelled ligand in the presence of an unlabelled ligand and through knowledge of the kinetic parameters of the labelled ligand both the association and dissociation rate constants of the unlabelled ligand can be calculated. This technique is widely used with radiolabelled ligands and has more recently been successfully applied in combination with fluorescently labelled ligands in resonance energy transfer techniques such as time-resolved resonance energy transfer (TR-FRET) and bioluminescence resonance energy transfer (BRET) [28, 29].

For the A3AR, two recent studies have used radioligands to determine the kinetic parameters of unlabelled ligands [30, 31]. In the study by Xia et al., however, the radiolabelled antagonist (\(^{3}H\)PSB-11) had to be used at low temperatures (10 °C) to slow the association rate sufficiently to give enough resolution to accurately determine the kinetic binding parameters of unlabelled ligands [30]. Due to this limitation and the inherent issue of throughput associated with radioligands [25], there is a need to develop additional methods to measure kinetic parameters. One way to potentially overcome these issues is through the use of fluorescently labelled ligand which have increased throughput, and previous studies have suggested that the fluorescent A3AR antagonist CA200645 has a slower association rate than \(^{3}H\)PSB-11 [32]. For the A3AR, a number of fluorescently labelled antagonist probes have been developed based on two structurally distinct antagonists which have been shown to retain high affinity for the receptor [32–34]. Therefore, in this study, we have developed a live-cell BRET-based kinetic binding assay for the A3AR using four different fluorescently labelled antagonists. This assay has then been compared to the radioligand binding assay described in Xia et al. [30] for the determination of the kinetic binding parameters of unlabelled ligands.

**Material and methods**

**Materials**

Foetal calf serum (FCS) was obtained from PAA Laboratories (Wokingham, UK). Furimazine was purchased from Promega (Southampton, UK). Bicinchoninic acid protein assay kit and white 96-well microplates were obtained from Thermo Fisher Scientific (Waltham, MA, USA). GF/B filter plates and Microscint-O were from PerkinElmer (Groningen, The Netherlands). CA200645 was obtained from HelloBio (Bristol, UK). The synthesis of AV039 was described in Vernall et al. as compound 19 [34], while the synthesis of XAC-S-ser-S-tyr-X-BY630 (compound 27) and XAC-S-tyr-S-tyr-X-BYFL (compound 28) was described in Vernall et al. 2013 [33]. PSB-11 and MRS1220 were purchased from Tocris Bioscience (Bristol, UK), and NECA was obtained from Sigma-Aldrich (Zwijndrecht, The Netherlands). \(^{3}H\)-8-Ethyl-4-methyl-2-phenyl-(8H)-imidazo[2,1-i]-purin-5-one (\(^{3}H\)PSB-11) was kindly donated by Prof. C.E. Müller (University of Bonn, Germany) and its synthesis described in Müller et al. [35]. 1-Benzyl-8-methoxy-1H,3H-pyrido[2,1-f]purine-2,4-dione (compound 5) synthesis was described in Priego et al. as compound number 3 [36] and referred in Xia et al. [37] as compound number 5, while LUF7565 synthesis was described in Xia et al. as compound 27 [30]. All other chemicals and reagents were obtained from Sigma-Aldrich (Gillingham, UK).

**Cell culture and membrane preparation**

Generation of human embryonic kidney 293 (HEK293) cells stably expressing the human A3AR tagged at the N-terminus...
with NanoLuc (Nluc-A3AR) is described in Stoddart et al. [38], and these cells were used throughout this study. Nluc-A3AR HEK293 cells were maintained in Dulbecco’s modified Eagle’s medium containing 10% FCS and 2 mM L-glutamine at 37 °C, 5% CO2.

For membrane preparation, Nluc-A3AR HEK293 cells were grown to confluence in 500-cm² dishes. Normal growth media was replaced with ice-cold PBS, and the cells were removed from the dish using a cell scraper. The cells were then transferred to a 50-ml tube and centrifuged at 250×g for 5 min. The supernatant was discarded, and the resulting pellets were stored at −80 °C. Thawed pellets were resuspended in ice-cold PBS and homogenised using an IKA RW16 overhead stirrer fitted to an IKA T10 Ultra-Turrax disperser in 10 × 5 s bursts at 15,000 rpm. After removal of unbroken cells and nuclei by centrifugation at 1200×g for 10 min, the supernatant was centrifuged at 41,415×g for 30 min to obtain the membrane pellet. The pellet was then resuspended in ice-cold PBS and homogenised by 20 passes using a Kartell serrated pestle and a borosilicate glass homogeniser mortar fitted to an IKA RW16 overhead stirrer set to 1000 rpm. Finally, protein concentration was determined using a bicinchoninic acid protein assay [37] and membranes stored at −80 °C until needed.

**NanoBRET binding assays**

HEK293 cells stably expressing human Nluc-A3AR were seeded in normal growth medium 24 h prior to experimentation in white 96-well microplates coated with poly-D-lysine (100 μg/ml poly-D-lysine in PBS, 30 min room temperature, then washed in normal growth medium prior to use). Immediately before experimentation, media was replaced with HEPES buffered saline solution (HBSS; 145 mmol/L NaCl, 5 mmol/L KCl, 1.7 mmol/L CaCl2, 1 mmol/L MgSO4, 10 mmol/L HEPES, 2 mmol/L sodium pyruvate, 1.5 mmol/L NaHCO3, 10 mmol/L D-glucose, pH 7.4). For saturation and competition binding assays, the required concentration of fluorescent ligand and competing ligand was added simultaneously and incubated for 1 h (for AV039) or 3 h (for XAC-S-ser-S-tyr-X-BY630) at 37 °C (no CO2). After 1 h, 10 μM furimazine was added to each well and the plate incubated for 5 min at 37 °C before reading. Prior to all kinetic experiments, the medium was replaced by HBSS containing 10 μM furimazine and incubated at room temperature in the dark for 15 min to allow the luminescence signal to reach equilibrium [29]. For association kinetic experiments, following the furimazine incubation, the required concentration of fluorescent ligand in the presence or absence of 10 μM MRS1220 was added simultaneously and the plate read immediately once per minute for up to 3 h at 37 °C. For kinetic competition association assays, after furimazine incubation, either XAC-S-ser-S-tyr-X-BY630 (20 nM) or AV039 (40 nM) were added simultaneously with the required concentration of unlabelled ligand or 10 μM MRS1220 to determine non-specific binding and read once per minute for the indicated times at 37 °C. For all NanoBRET experiments, fluorescence and luminescence were read using a PHERAstar FS plate reader (BMG Labtech, Ortenberg, Germany). Filtered light emissions were measured at 460 nm (80 nm band pass) and at > 610 nm (long pass) for the BY630 labelled ligands and at 450 nm (80 nm band pass) and > 550 nm (long pass) for the BYFL labelled ligand. The raw NanoBRET ratio was calculated by dividing the fluorescence emission (610 or 550 nm) by the luminescence emission (460 or 450 nm).

**Radioligand binding assays**

Prior to all experiments, membranes from HEK293 membranes expressing human Nluc-A3AR were diluted to 20 μg/well in a total volume of 100 μL/well assay buffer (50 mM Tris-HCl, 5 mM MgCl2, supplemented with 0.01% CHAPS and 1 mM EDTA, pH 7.4) and homogenised using an IKA T10 Ultra-Turrax disperser in 3 × 5 s bursts at 15,000 rpm. For equilibrium displacement assays, membranes were placed in 96-well microplates at 10 °C and the required concentration of competing agonist was added in the presence of a final concentration of ~10 nM [3H]PSB-11 with nonspecific binding determined in the presence of 100 μM NECA (final concentration). For association assays, Nluc-A3AR HEK293 membranes were placed at 10 °C in 96-well microplates and the amount of radioligand bound to the receptor was measured at different time points during a total incubation of 120 min. For dissociation experiments, HEK293 membranes expressing human Nluc-A3AR were incubated for 120 min with ~10 nM [3H]PSB-11 at 10 °C prior the addition of 10 μM PSB-11 (final concentration) at various time points during a further 120 min. The competition association assays were initiated by the addition of Nluc-A3AR HEK293 membranes at different time points for a total of 240 min to a total volume of 100 μL/well of assay buffer at 10 °C with ~10 nM [3H]PSB-11 in the absence or presence of a single concentration (2 × Kd) of competing A3AR ligands. For all experiments, incubation was terminated by rapid filtration performed on 96-well GF/B filter plates using a PerkinElmer Filtermate-harvester (PerkinElmer, Groningen, The Netherlands). After drying the filter plate at 50 °C for 30 min, the filter-bound radioactivity was determined by scintillation spectrometry using a 2450 MicroBeta2 Plate Counter (PerkinElmer, Boston, MA). In addition, the exact concentration of [3H]PSB-11 used in each experiment was determined by scintillation spectrometry and this concentration was used in the data analysis.
Data analysis

All experimental data were analysed using Prism7 (GraphPad Software, San Diego, CA).

NanoBRET total and non-specific saturation binding curves were fitted simultaneously using the following equation:

\[
\text{BRET ratio} = \frac{B_{\text{max}}[B]}{[B]} + \left(\frac{1}{K_D} + \left(M [B]\right) + C\right)
\]

where \(B_{\text{max}}\) is the maximal specific binding achieved, \([B]\) is the concentration of fluorescent ligand, \(K_D\) is the equilibrium dissociation constant, \(M\) is the slope of the non-specific binding component and \(C\) is the intercept with the Y-axis.

Equilibrium competition binding curves were fitted with the following equation:

\[
K_1 = \frac{\text{IC}_{50}}{1 + \frac{[L]}{K_D}}
\]

where \([L]\) is the concentration of \(^3\)HPSB11, AV039 or XAC-S-ser-S-tyr-X-BY630 and \(K_D\) is the equilibrium dissociation constant of the labelled ligand (9.9 nM for \(^3\)HPSB-11, 14.5 nM for XAC-S-ser-S-tyr-X-BY630 and 32.5 nM for AV039 as determined in saturation binding assay performed as part of this study). The IC_{50} is obtained as follows:

\[
\% \text{inhibition of specific binding} = \frac{100[A]}{[A] + \text{IC}_{50}}
\]

where \([A]\) is the concentration of unlabelled competing drug and \(\text{IC}_{50}\) is the molar concentration of this competing ligand required to inhibit 50% of the specific binding of the concentration \([L]\) of the labelled ligand.

For NanoBRET association kinetic data, non-specific binding was determined in wells with fluorescent ligand plus 10 μM MRS1220 and this was obtained for each concentration of fluorescent ligand at each time point. This was subsequently subtracted from total binding at the equivalent time point. The data were simultaneously fit to the following equations:

\[
Y = Y_{\text{max}} \left(1 - e^{-k_{\text{obs}}t}\right)
\]

\[
k_{\text{on}} = \frac{k_{\text{obs}} - k_{\text{off}}}{[L]}
\]

where \(Y_{\text{max}}\) equals the level of binding at infinite time (\(t\)), \(k_{\text{obs}}\) is the rate constant for the observed rate of association at a particular concentration of \(L\), \([L]\) is the ligand concentration in molar, \(k_{\text{off}}\) is the dissociation rate constant of the ligand in per minute and \(k_{\text{on}}\) is the association rate constant in per molar per minute. From this, the equilibrium dissociation constant (\(K_D\)) is determined as follows:

\[
K_D = \frac{k_{\text{off}}}{k_{\text{on}}}
\]

The binding kinetics of unlabelled ligands was quantified using the competition association assay based on the theoretical framework by Motulsky and Mahan [27]. NanoBRET and radioligand data were fitted into the competition association model using ‘kinetics of competitive binding’ to determine association and dissociation rate constants of the unlabelled compounds:

\[
K_A = \frac{k_1[L] \cdot 10^{-9} + k_2}{k_3[L] \cdot 10^{-9} + k_4}
\]

\[
S = \sqrt{(K_A - K_B)^2 + 4k_1k_3L}\cdot I \cdot 10^{-18}
\]

\[
K_F = 0.5(K_A + K_B + S)
\]

\[
K_S = 0.5(K_A + K_B - S)
\]

\[
Q = \frac{\text{B}_{\text{max}}^2 \cdot k_1 \cdot L \cdot 10^{-9}}{K_F \cdot K_S}
\]

\[
Y = \frac{k_4(K_F - K_S)}{K_F \cdot K_S} + \frac{k_4 - k_F}{K_F} e^{(-K_F \cdot X)} \cdot \frac{k_4 - K_S}{K_S} e^{(-K_S \cdot X)}
\]

where \(k_1\) is the \(k_{\text{on}}\) of the labelled ligand (M⁻¹ min⁻¹), \(k_2\) is the \(k_{\text{off}}\) of the labelled ligand (min⁻¹), \(L\) is the concentration of the labelled ligand in nanomolar, \(I\) is the concentration of the unlabelled competitor (nM), \(X\) is the time (min) and \(Y\) is the specific binding of the labelled ligand (NanoBRET ratio or DPM).

\(k_1\) and \(k_2\) values were generated from association kinetic experiments. \(k_3\), \(k_4\) and \(B_{\text{max}}\) were then calculated, where \(k_3\) represents the \(k_{\text{on}}\) (M⁻¹ min⁻¹) of the unlabelled ligand, \(k_4\) stands for the \(k_{\text{off}}\) (min⁻¹) of the unlabelled ligand and \(B_{\text{max}}\) equals the total binding (NanoBRET ratio or DPM). The residence time (RT) was calculated as the reciprocal of the ligand dissociation rate constant as follows:

\[
RT = \frac{1}{k_{\text{off}}}
\]

Statistical significance was determined by Student’s unpaired t test where \(p < 0.05\) was considered statistically significant throughout this study.

Results

NanoBRET binding profile of fluorescent probes at Nluc-A3AR in live cells

The four fluorescent ligands used in this study have previously been shown to display high affinity for the A3AR, and their structures are shown in Fig. 1 [32–34]. Three of the ligands are based on the non-selective adenosine receptor antagonist
xanthine amine congener (XAC). Two of these XAC linked compounds contain a serine, tyrosine dipeptide (both in the S configuration) to link the pharmacophore to the fluorophore which is either BODIPY630/650 (excitation max 630 nm, emission max 650 nm) to give XAC-S-ser-S-tyr-BY630 (Fig. 1a, or BODIPY-FL (excitation max 503 nm, emission max 512 nm) to give XAC-S-ser-S-tyr-BYFL (Fig. 1b) [33]. The third XAC-based compound, CA200645 (Fig. 1c), contains a polyamide linker (β-alanine, β-alanine) connected to the BY630 fluorophore [32]. AV039 (Fig. 1d) is based on the A3AR selective antagonist 1,2,4-triazolo[4,3-a]quinoxalin-1-one linked to BY630 and has been shown to retain A3AR selectivity [34].

Initially, we determined the affinity of each of these fluorescent antagonists at NanoLuc tagged A3AR (Nluc-A3AR) in the NanoBRET assay. An advantage of NanoBRET compared to radioligand binding experiments is the large concentration range of fluorescent ligands over which binding can be measured, and to represent this accurately, binding is fitted as a sigmoidal curve with BRET ratio values plotted versus the log concentration of fluorescent ligand. In all the cases, we obtained a saturable curve clearly dependent on the concentration of the fluorescent ligands (Fig. 2). Non-specific binding was determined by co-incubation with 10 μM of the unlabelled high affinity A3AR antagonist MRS1220, and all four fluorescent

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**Fig. 1** Structures of the A3AR fluorescent (a-d) and unlabelled (e-g) antagonists used in this study.
ligands exhibited very low levels of non-specific NanoBRET in line with previously published NanoBRET data [29, 38]. The point on the curve which gives the 50% of the binding was then taken as $K_D$. The $pK_D$ values calculated for the four fluorescent antagonists are shown in Table 1. All four ligands displayed high affinities for Nluc-A3AR which were consistent with literature values, and the range of $pK_D$ values were found to span less than 1 log unit ($pK_D$ 7.78–8.11).

To determine the kinetic parameters of fluorescent ligands at the A3AR, Nluc-A3AR expressing HEK293 cells were initially treated with the Nanoluc substrate, furimazine (10 μM), for 15 min prior to the fluorescent ligand to allow the luminescence signal to stabilise as described previously [29]. Increasing concentrations of fluorescent ligands were then added and the change in NanoBRET monitored at 37 °C over time, with specific binding determined in the presence of the high affinity A3AR antagonist MRS1220 (10 μM) for every time point (Fig. 3). The kinetic parameters obtained from globally fitting the association curves showed that all four fluorescent ligands had similar association rates within a factor of two of each other (Table 1). As expected, the range of dissociation rates for A V039, CA200645 and XAC-S-ser-S-tyr-X-BY630 were also within a factor of two of each other, leading to similar RTs. The dissociation rate for XAC-S-ser-S-tyr-X-BY630, however, was over ten times slower than that of the other three fluorescent ligands. This produced a markedly different RT for XAC-S-ser-S-tyr-X-BY630 which was of the order of hours (4.8 h) compared to the much more rapid values.

![Fig. 2 Determination of the binding affinity of four fluorescent antagonists at human A3AR. NanoBRET saturation binding curves obtained by treating Nluc-A3AR HEK293 cells with increasing concentrations of the fluorescent antagonists AV039 (a), CA200645 (b), XAC-S-ser-S-tyr-X-BYFL (c) or XAC-S-ser-S-tyr-X-BY630 (d) in the absence (black circles) or presence (red squares) of 10 μM MRS1220 for 1 h at 37 °C. The data shown are mean ± SEM and are representative of five independent experiments each performed in triplicate.](image)

| Compound            | $pK_D$  | $k_{on}$ ($\times 10^6$) (M$^{-1}$ min$^{-1}$) | $k_{off}$ (min$^{-1}$) | RT (min) | $n$ | $pK_D$  | $n$ |
|---------------------|--------|-----------------------------------------------|------------------------|----------|-----|--------|-----|
| AV039               | 7.35 ± 0.10* | 3.67 ± 0.62                                   | 0.15 ± 0.02            | 6.8 ± 0.8 | 4   | 8.11 ± 0.10* | 5   |
| CA200645            | 7.58 ± 0.04* | 2.58 ± 0.05                                   | 0.069 ± 0.006          | 14.8 ± 1.1 | 5   | 7.80 ± 0.07* | 5   |
| XAC-S-ser-S-tyr-X-BY630 | 8.57 ± 0.15 | 1.66 ± 0.36                                   | 0.0043 ± 0.0009        | 288 ± 62  | 5   | 8.51 ± 0.07 | 5   |
| XAC-S-ser-S-tyr-X-BYFL | 7.75 ± 0.14 | 3.95 ± 0.73                                   | 0.069 ± 0.006          | 14.9 ± 1.1 | 4   | 7.78 ± 0.05 | 5   |

Saturation $pK_D$ values were calculated directly from the saturation binding curves of the fluorescent ligands binding to Nluc-A3AR HEK293 cells. The kinetic parameters, $k_{on}$, $k_{off}$ and $pK_D$ values, were obtained by monitoring the NanoBRET signal over time of various concentrations of fluorescent ligand in HEK293 Nluc-A3AR cells at 37 °C. The residence time (RT) was calculated as the reciprocal of the $k_{off}$ values from each individual experiment. All values represent mean ± SEM from $n$ separate experiments performed in triplicate.

* $p < 0.05$, kinetic versus equilibrium saturation $pK_D$ values according to unpaired Student’s $t$ test.
(min) obtained for the other three compounds. Additionally, it can be noted that the pK_D values from saturation studies of XAC-S-ser-S-tyr-X-BY630 and XAC-S-ser-S-tyr-X-BYFL are comparable to those obtained from the kinetic data, whereas for CA200645 and AV039, the saturation pK_D values are significantly higher than the kinetic pK_D (Table 1, p < 0.05, unpaired t test).

**Radioligand binding characterisation of fluorescent probe**

To confirm the differences in the kinetics observed with AV039 and XAC-S-ser-S-tyr-X-BY630, radioligand equilibrium and kinetic binding assays in membranes from HEK293 cells stably expressing Nluc-A3AR were undertaken. Initially, the binding affinity and kinetic parameters of the radiolabelled A3AR antagonist [3H]PSB-11 at Nluc-A3AR were resolved from its association and dissociation curves at 10 °C (Fig. 4a, b; Table 2) and were similar to those obtained previously with the wild-type A3AR [30]. Next, the ability of XAC-S-ser-S-tyr-X-BY630 and AV039 to compete at equilibrium with specific [3H]PSB-11 binding to the Nluc-A3AR was examined (Fig. 4c) and the pKi for each of these compounds determined (Table 2). The affinities obtained in the radioligand binding assay for both fluorescent probes were lower than those obtained in the NanoBRET experiments (Table 1), which may reflect differences in affinity values determined in whole cells compared to cell membranes, and also the impact of temperature (10 °C versus 37 °C), as previously observed for the histamine H1 receptor [29].

The association and dissociation rates of XAC-S-ser-S-tyr-X-BY630 and AV039 were determined in kinetic association assays where the specific binding of [3H]PSB-11 in the presence and absence of a single concentration (2 × IC50) of the fluorescent ligand was monitored over time and the resulting data fitted to the model proposed by Motulsky and Mahan [27]. As expected from the kinetic behaviour observed in the NanoBRET assay, the association curve of [3H]PSB-11 obtained in the presence of AV039 quickly reached an equilibrium plateau (Fig. 4d) indicating that AV039 has a similar or faster kinetic profile compared to [3H]PSB-11. This was borne out in the kinetics parameters obtained from fitting the data to the Motulsky and Mahan model (Table 2). In comparison, XAC-S-ser-S-tyr-X-BY630 induced an 'overshoot' in [3H]PSB-11 specific binding followed by a steady decrease (Fig. 4e) which is characteristic of long RT compounds. As such, both the association and dissociation rate constants of this fluorescent ligand are more than 10 times slower than those of AV039, whilst the pK_D values are very similar. As a consequence, it was confirmed that XAC-S-ser-S-tyr-X-BY630 had a long RT (over 7.5 h) compared to AV039 and [3H]PSB-11.
Determination of the affinity and kinetic constants of unlabelled ligands

To extend the use of the NanoBRET binding assay, it is necessary to use this setup to determine the kinetics of unlabelled ligands using the Motulsky and Mahan approach. To demonstrate its utility, it is important that the NanoBRET assay can distinguish between fast and slow unlabelled compounds. First, it was important to select unlabelled compounds with the desired kinetic profile. Using \(^{3}H\)PSB-11 as the tracer, the kinetic parameters of two previously described high-affinity A\(_3\)AR antagonists, compound 5 [36] (Fig. 1e) and LUF7565 [30] (Fig. 1f) were determined at Nluc-A\(_3\)AR. Competition binding experiments indicated compound 5 and LUF7565 membranes at 10 °C to generate competition binding curves. Data were normalised to the maximal specific \(^{3}H\)PSB-11 labelling in each experiment. Data points are combined mean ± SEM from five separate experiments performed in duplicate. The data shown are representative examples from four (d) and five (e) independent experiments performed in duplicate with each data point shown.
gave a concentration-dependent inhibition of the specific binding of [3H]PSB-11 with LUF7565 displaying higher affinity for Nluc-A3AR than compound 5 (Fig. 5a; Table 2).

The association and dissociation rate constants of compound 5

| Kinetics | Equilibrium competition |
|----------|-------------------------|
| pKD     | k<sub>on</sub> (× 10<sup>6</sup> M<sup>−1</sup> min<sup>−1</sup>) | k<sub>off</sub> (min<sup>−1</sup>) | RT (min) | n | pKi | n |
| [3H]PSB-11 | 8.04 ± 0.08 | 2.75 ± 0.52 | 0.023 ± 0.002 | 44.6 ± 3.9 | 5 | ND | ND |
| AV039    | 7.63 ± 0.08 | 3.27 ± 0.47 | 0.076 ± 0.009 | 14.0 ± 1.6 | 5 | 7.65 ± 0.02 | 5 |
| XAC-S-ser-S-tyr-X-BY630 | 7.74 ± 0.08* | 0.14 ± 0.04 | 0.0023 ± 0.0002 | 447 ± 44 | 4 | 7.21 ± 0.06* | 5 |
| compound 5 | 7.53 ± 0.13 | 6.61 ± 1.76 | 0.198 ± 0.045 | 6.5 ± 2.1 | 4 | 7.35 ± 0.02 | 5 |
| LUF7565  | 9.11 ± 0.04* | 8.17 ± 0.95 | 0.0065 ± 0.0014 | 168 ± 25 | 4 | 8.99 ± 0.02* | 5 |

pKi values were calculated from inhibition of equilibrium [3H]PSB-11 binding to Nluc-A3AR HEK293 membranes at 10 °C. The kinetic parameters, k<sub>on</sub>, k<sub>off</sub> and pKD values of PSB-11 were obtained from association and dissociation curves of 10 nM [3H]PSB-11 in Nluc-A3AR HEK293 membranes at 10 °C. Radioligand dissociation was initiated by the addition of 10 μM unlabeled PSB-11. The kinetic parameters, k<sub>on</sub>, k<sub>off</sub> and pKD values of AV039, XAC-S-ser-S-tyr-X-BY630, compound 5 and LUF7565 were obtained by competition association with ~ 10 nM [3H]PSB-11 at 10 °C. The RT was calculated as the reciprocal of the k<sub>off</sub> values from each individual experiment. All values represent mean ± SEM from n separate experiments performed in duplicate.

ND not determined

*p < 0.05, kinetic pKD versus equilibrium competition pKi values according to unpaired Student’s t test.

Fig. 5 Radioligand binding characterisation of unlabelled ligands at human A<sub>3</sub>AR. a HEK293 membranes expressing human Nluc-A3AR were treated with increasing concentrations of compound 5 (red triangles) and LUF7565 (green squares), and 10 nM [3H]PSB-11 at 10 °C for 2 h and levels of [3H]PSB-11 binding monitored by scintillation counting. Data were normalised to the maximal specific [3H]PSB-11 labelling in each experiment. Data points are combined mean ± SEM from five separate experiments performed in duplicate. b. Association of [3H]PSB-11 on Nluc-A3AR HEK293 membranes at 10 °C in the absence (control, black circles) or presence of a single concentration (2 × K<sub>i</sub> value) of and LUF7565 (b) or compound 5 (c). The data shown are representative examples from four independent experiments performed in duplicate with each data point shown.
and LUF7565 were then determined from the decrease in the specific binding of [3H]PSB-11 in the presence of a single concentration (2 × IC50) of the unlabelled ligands. The competition association curve of compound 5 (Fig. 5c) exhibited the kinetic profile of a short RT ligand, while LUF7565 induced an ‘overshoot’ in the [3H]PSB-11-specific binding (Fig. 5b) which is the characteristic profile of a long RT compound, and this was confirmed from the calculated kinetic parameters (Table 2). Radioligand kinetic affinity was comparable to that obtained by radioligand displacement for both the unlabelled ligands.

The selective A3AR fluorescent antagonist AV039 was chosen as a short RT probe to evaluate the affinity and kinetic parameters of compound 5, LUF7565 alongside those of PSB-11. Initially, the affinity of these three antagonists was determined in an equilibrium competition assay using 20 nM AV039 in HEK 293 cells stably expressing the Nluc-A3AR (Fig. 6a; Table 3). As expected, all three ligands exhibited high affinity for the A3AR, confirming the data obtained using [3H]PSB-11. To further investigate the properties of these compounds, their binding kinetic parameters were determined using AV039 as the probe. The time-dependent decrease in the association binding of AV039 induced by the addition of different concentrations of PSB-11, compound 5 and LUF7565 was monitored at 37 °C (Fig. 6b–d) to determine the kinetic parameters of these unlabelled compounds. The competition association curves of PSB-11 (Fig. 6b) and compound 5 (Fig. 6c) exhibited the typical kinetic profile of a short RT ligand, while LUF7565 (Fig. 6d) induced a very pronounced initial overshoot in binding of AV039 followed by a decrease which is characteristic of a long RT compound. As expected from the shape of the competition association curves, LUF7565 was found to have the slowest dissociation rate and consequently the longest RT (Table 3). The kinetic pKD of the short RT compounds PSB-11 and compound 5 was similar to the equilibrium pKᵢ, while for the slow dissociation compound LUF7565, the kinetic pKD was significantly higher than pKᵢ (p < 0.05, unpaired t test).

As XAC-S-ser-S-tyr-X-BY630 was found to have a considerably longer RT than AV039 and [3H]PSB-11, it was used to investigate the effect of the RT and binding kinetic parameters of the labelled probe on the measured kinetic parameters of unlabelled ligands. Firstly, it was confirmed that PSB-11, compound 5 and LUF7565 inhibited the specific XAC-S-ser-S-tyr-X-BY630 binding in a concentration-dependent manner in Nluc-A3AR HEK 293 cells (Fig. 7a; Table 3). It was noted that the affinity obtained for all the compounds was around half a log unit lower than the pKᵢ value obtained with AV039.
but exhibited the same trend, with PSB-11 displaying the highest affinity at A3AR and compound 5 the least potent.

The concentration-dependent decrease in the association binding of XAC-S-ser-S-tyr-X-BY630 induced by the addition of different concentrations of PSB-11, compound 5 and LUF7565 allowed us to determine the association and dissociation rates of these unlabelled compounds when using a long RT compound as the probe. The competition association curves obtained in the presence of all three unlabelled ligands showed a continuous increase in XAC-S-ser-S-tyr-X-BY630 signal parallel to that observed with the fluorescent ligand only (Fig. 7b–d). Only a small initial overshoot in binding was observed in the presence of 100 and 300 nM of LUF7565. The kinetic parameters obtained for all ligands are presented in Table 3. The kinetic pKD for compound 5 and PSB-11 was similar to that obtained at equilibrium, while LUF7565’s kinetic pKD was slightly higher than the equilibrium pKi. In comparison to the kinetic parameters obtained with AV039, the k\text{on} values were lower when using XAC-S-ser-S-tyr-X-BY630 as the probe for all three unlabelled compounds. For PSB-11 and compound 5, an increase in RT was observed with XAC-S-ser-S-tyr-X-BY630 in comparison to that with AV039, whereas a decrease in the RT of LUF7565 was seen.

**Discussion**

It is becoming increasingly clear that a ligand’s binding kinetics are a crucial part of its pharmacology which can play a role in its success in the clinic [16, 18, 20]. In this study, we set out to develop an assay to determine the kinetic parameters of molecules binding to the A3AR using NanoBRET in living cells and at physiological temperatures. We also compared the values obtained to those obtained in a more classical radioligand binding assay that used cell membrane preparations and was performed at much lower temperatures (10 °C). We found that kinetic parameters of unlabelled ligands obtained in both assays were comparable when using a labelled probe with similar kinetics (AV039 versus [3H]PSB-11). Moreover, when a long RT probe (XAC-S-ser-S-tyr-X-BY630) was used, it was difficult to resolve the differences in kinetic parameters of a fast and slow unlabelled compound. This indicates that the kinetic profile of the labelled probe can influence the range of k\text{on} and k\text{off} values that can be accurately determined.

For many years, the toolbox available for studying binding kinetics at GPCRs was limited to the use of radioligand binding assays [14]. With the increasing range of fluorescent ligands available for the study of GPCRs alongside more suitable proteins for use in energy transfer techniques (such as NanoLuc and SNAP), there has been increased interest in adapting these techniques to measure ligand-binding kinetics.
Here, we have developed a BRET-based kinetic assay for the A3AR. This is particularly important for the A3AR, as historically there has only been a radiolabelled agonist commercially available, and radiolabelled antagonists, such as the ones used in this study, have to be custom synthesised [35]. The A3AR offered a unique opportunity for this since multiple fluorescent ligands based on different parent molecules have been described in the literature and one of these ligands is commercially available [32–34]. Three of these fluorescent ligands measured in this study were based on the non-selective adenosine receptor antagonist XAC. Two of these ligands, CA200645 and XAC-S-ser-S-tyr-X-BYFL, were found to have similar $k_{on}$ and $k_{off}$ rate constants, whereas XAC-S-ser-S-tyr-X-BY630 had a similar value for $k_{on}$ but a much slower $k_{off}$ rate constant. In the development of fluorescent ligands with improved imaging properties, incorporation of a peptide linker between the pharmacophore and fluorophore was found to prevent the ligand from crossing the cell membrane in a non-specific manner [29, 33]. Molecular modelling has suggested that this linker region makes additional contacts at the top of the transmembrane domains of the receptor and that the fluorophore is buried within the lipid environment of the plasma membrane [33]. This has also been observed in molecular modelling of fluorescent ligands binding to the histamine H1 receptor [29].

Other studies that have developed fluorescent ligands have also found that the fluorophore can influence the affinity of the resulting molecule [39, 40]. It is therefore clear that the fluorophore portion of fluorescent ligand is not a passive part of the molecule but can influence both the affinity and binding kinetics. The chemical structures of XAC-S-ser-S-tyr-X-BY630 and XAC-S-ser-S-tyr-X-BYFL are very similar and only differ in the structure of the fluorophore. It has previously been shown with the $\beta_2$-adrenoceptor that the interaction of ligands with the membrane influences the observed association rates [41]. The interaction of a ligand with a membrane can create a micro-environment which can alter the observed kinetic profiles [42]. It is possible that the BODIPY fluorophores interact differently with the cell membrane to increase the RT of XAC-S-ser-S-tyr-X-BY630 compared to XAC-S-ser-S-tyr-X-BYFL. It is worth noting, however, that the mere presence of a BY630 fluorophore is not a determinant for a longer RT compound since both CA200645 and AV039 contain the same fluorophore.

The availability of a radiolabelled antagonist, $[^3H]$PSB-11, allowed us to compare the kinetic parameters of unlabelled ligands, measured using either long or short RT fluorescent ligand in the NanoBRET assay, with those obtained in a more traditional radioligand binding assay. We found that the $k_{on}$ rate constants measured with all three unlabelled ligands were performed in triplicate.

**Fig. 7** Characterisation of the effect of unlabelled ligands on the binding of XAC-S-ser-S-tyr-X-BY630 to Nluc-A3AR using NanoBRET. a Nluc-A3AR cells were treated with 40 nM XAC-S-ser-S-tyr-X-BY630 and increasing concentrations of PSB-11 (black circles), compound 5 (red triangles) and LUF7565 (green squares) for 3 h at 37 °C and competition binding curves generated. Data were normalised to the maximal BRET signal in each experiment. Data points are combined mean ± SEM from five (PSB-11) or six (compound 5 and LUF7565) separate experiments.
all within a similar range (Table 2) although the rank order within data sets was not the same. For example, LUF7565 was found to have the fastest association rate when either [3H]PSB-11 or XAC-S-ser-S-tyr-X-BY630 were used as the probe, whereas LUF7565 had the slowest association rate when measured against AV039. Although here we have only compared the kinetic parameters of three compounds, this is similar to previous studies comparing the kinetic parameters of GnRH ligands determined using radioligand binding and time-resolved FRET (TR-FRET) where no correlation between association rates was found in the two assay setups [43].

In comparison to the tight range of $k_{\text{off}}$ values, there was a wider range of measured $k_{\text{off}}$ values. For both [3H]PSB-11 and AV039, there was between a 30- and 94-fold difference, respectively, in the $k_{\text{off}}$ values measured for compound 5 and LUF7565, with LUF7565 displaying the slowest $k_{\text{off}}$ of the measured unlabelled compounds using both probes. This indicates that for both these probes, a wide range of kinetic parameters can be measured. As seen in Fig. 7, the profile of the competition association curves with each of the three unlabelled compounds was comparable, and this is reflected in the similarities in the kinetic parameters obtained. To date, LUF7565 is one of the longest residence time unlabelled ligands reported at A3AR, and it may be that XAC-S-ser-S-tyr-X-BY630 will be a useful tool to find even longer RT compounds at this receptor.

Although a similar trend was seen for the $k_{\text{off}}$ values determined using [3H]PSB-11 and AV039, 10-fold higher $k_{\text{off}}$ values were determined for PSB-11 and compound 5 when using AV039, leading to compounds with apparently shorter RT. There are many differences in setup of the two assays which could lead to these differences and need to be kept in mind when comparing the data sets. One of the main differences is that the radioligand binding assay uses membrane preparations, whereas the BRET assays are performed on intact cells. The intact cell environment preserves all of the intracellular proteins that a GPCR can interact with, and these are known to be able to stabilise different conformations of the receptor which in turn may affect the binding kinetics of the ligands [44, 45]. As these interacting proteins include G proteins and arrestins, preservation of the intact cellular environment allows downstream signalling and internalisation to still occur. It is unlikely that internalisation plays a role in this study since XAC-S-ser-S-tyr-X-BY630 and AV039 have previously been shown to be competitive antagonists, and in confocal imaging studies, there was no indication that these ligands caused internalisation of the A3AR [33, 34]. Although it appears that these fluorescent ligands do not cause internalisation of the receptor, it is possible that the antagonist bound form of the receptor may interact with different adaptor proteins and this may underpin some of the differences observed. Differences in binding kinetics have also been observed previously when comparing measurements from membrane preparations and whole cells at other GPCRs including the dopamine D2L receptor [46] and histamine H1 receptor [29]. For the D2L receptor, it was suggested that differences in dissociation rates are a reflection of how the ligand interacts with the cell membrane [46]. Apart from receptor environment, the two assay setups were performed at different temperatures—radioligand binding at 10 °C and BRET at 37 °C—and it can be assumed from basic thermodynamic principles that both association and dissociation rates will increase with temperature. Due to the rapid kinetics of [3H]PSB-11, the competition association assay is challenging to perform at higher temperature [30], and since the plate reader used for the live cell assay lacks the capacity to be actively cooled, we are unable at the present time to directly test this. An increase in association and dissociation rate with temperature has been observed for the histamine H1 receptor [47] and the prolactin receptor [48].

In summary, this study has demonstrated that a NanoBRET-based assay can be applied to measure ligand-binding kinetics at the A3AR in intact living cells at physiological temperatures. The data shown here do, however, indicate that care needs to be taken when selecting a probe with the appropriate kinetics for the study of unlabelled ligands with different kinetic profiles. Thus, probes with very slow kinetics may be problematic for the determination of the kinetic parameters of unlabelled ligands with rapid kinetic properties. As the range of fluorescent ligands for GPCRs continues to expand [49], their use in NanoBRET-based assays should provide the required diversity of kinetic properties to evaluate the kinetic profiles of the broadest range of unlabelled ligands in a physiologically relevant setting.

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Compliance with ethical standards

Conflicts of interest Monica Bouzo-Lorenzo declares that she has no conflict of interest. Leigh A. Stoddart declares that she has no conflict of interest. Lizi Xia declares that she has no conflict of interest. Adriaan P. Ilzerman declares that he has no conflict of interest. Laura H. Heitman declares that she has no conflict of interest. Stephen J Briddon declares that he has no conflict of interest. Stephen J Hill declares that he has no conflict of interest.

Ethical approval This article does not contain any studies with human participants or animals performed by any of the authors.
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