Opioid antagonism in humans: a primer on optimal dose and timing for central mu-opioid receptor blockade

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Non-human animal studies outline precise mechanisms of central mu-opioid regulation of pain, stress, affiliation and reward processing. In humans, pharmacological blockade with non-selective opioid antagonists such as naloxone and naltrexone is typically used to assess involvement of the mu-opioid system in such processing. However, robust estimates of the opioid receptor blockade achieved by opioid antagonists are missing. Dose and timing schedules are highly variable and often based on single studies. Here, we provide a detailed analysis of central opioid receptor blockade after opioid antagonism based on existing positron emission tomography data. We also create models for estimating opioid receptor blockade with intravenous naloxone and oral naltrexone. We find that common doses of intravenous naloxone (0.10–0.15 mg/kg) and oral naltrexone (50 mg) are more than sufficient to produce full blockade of central MOR (>90% receptor occupancy) for the duration of a typical experimental session (~60 min), presumably due to initial super saturation of receptors. Simulations indicate that these doses also produce high KOR blockade (78–100%) and some DOR blockade (10% with naltrexone and 48–74% with naloxone). Lower doses (e.g., 0.01 mg/kg intravenous naloxone) are estimated to produce less DOR and KOR blockade while still achieving a high level of MOR blockade for ~30 min. The models and simulations form the basis of two novel web applications for detailed planning and evaluation of experiments with opioid antagonists. These tools and recommendations enable selection of appropriate antagonists, doses and assessment time points, and determination of the achieved receptor blockade in previous studies.

INTRODUCTION

A variety of psychological processes are thought to be modulated by the brain’s mu-opioid system, including reward [1], pain [2], stress [3], and social bonding [4, 5]. A popular method to study the mu-opioid system in humans is the pharmacological blockade of opioid receptors with antagonistic drugs. Opioid receptor antagonists bind opioid receptors, but in contrast to agonists they do not generally produce a response by the cell (although some, e.g., naloxone, may act as inverse agonists under certain conditions [6]). Opioid antagonists such as naloxone and naltrexone have a high affinity to mu-opioid receptors (MOR) and thereby prevent other ligands (including endogenous ones) from binding to this receptor type. Therefore, when antagonism with these drugs blocks a behavior, the behavior is assumed to be mu-opioid-dependent [7].

Most opioid antagonists available for human use are non-selective for opioid receptor subtypes and also bind to kappa-opioid receptors (KOR) with high affinity and to delta-opioid receptors (DOR) with low affinity (Table S1). To enable causal inferences about mu-opioid receptor functions based on pharmacological blockade, it is optimal to select an antagonist, a dose, and an assessment time point that results in complete blockade of MOR while causing minimal interference with other receptor types.

Antagonist doses used in basic human research to block the mu-opioid system are often based on plasma concentration, estimates from single positron emission tomography (PET) studies, or on conventions (e.g., 0.10–0.15 mg/kg intravenous naloxone and 50 mg oral naltrexone), and they vary considerably. For example, reported intravenous naloxone doses used in studies of endogenous mu-opioid function are as low as 0.006 mg/kg [8] and as high as 6 mg/kg [9]. Concurrent KOR and DOR blockade is seldom considered when selecting doses.

PET and dual-detector systems use radiolabeled ligands to quantify in vivo receptor binding in the human brain [10]. Because antagonist drugs prevent accumulation of the radiotracer in the brain, positron emission-based techniques can also be used to estimate receptor blockade with these drugs (Supplement) [11, 12]. Here we synthesize the available PET and dual-detector data and create models for estimating the amount and duration of central opioid receptor blockade with various doses of commonly used opioid antagonists. In line with previous interpretations of blockade estimates, we define full blockade as >90% receptor occupancy [13, 14]. This overview will help determine the achieved MOR blockade in previous studies and evaluate the possibility of DOR and KOR blockade or carry-over effects affecting the results or complicating inferences. It will also enable selection of the appropriate antagonist drugs, doses, assessment time points and intersession intervals for future studies.

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MATERIALS AND METHODS

We synthesized and further analyzed the available evidence from PET and dual-detector studies on opioid receptor blockade with naloxone and naltrexone. This project used data extracted from published articles and did therefore not require ethical approval. Studies containing blockade data were located using a semi-systematic approach, based on Web of Science searches and examination of references in relevant papers (Supplement). Model specification, fitting, and diagnostics were performed in R [15] using the packages minpack.lm [16], investr [17], nistsools [18], miceNls [19], opcr [20], and aomisc [21]. First, we used non-linear least squares regression to model the relationship between antagonist dose and MOR blockade. Using data on MOR blockade half-life, we adjusted blockade estimates from the dose-blockade model according to a specified time since administration of an opioid antagonist. Next, we used the linpk package [22] and data on MOR blockade half-life and time-to-peak MOR blockade to describe the time-blockade relationship. Finally, we simulated DOR and KOR blockade from MOR blockade using the antagonists’ average receptor affinities (Table S1).

The available data on central opioid receptor blockade with nalmefene and GSK1521498 is summarized in the Supplement (Table S1, Figures S2, S3).

Modelling the dose-blockade relationship

Following Mayberg and Frost [13] and Rabiner et al. [23], we fitted a log-logistic model to the available data to describe the relationship between antagonist dose and central blockade at the measurement time point \( t_{\text{measure}} \). We specifically used a four-parameter model (Eq. 1) with the lower limit constrained to 0, the upper limit constrained to 100, and the Hill slope constrained to 1. The parameter \( ED_{50} \) (i.e., effective dose 50) in this model indicates the estimated dose at which 50% of the receptors are blocked [23].

\[
\text{Blockade}(Dose)_{\text{measure}} = \frac{100 - 0}{1 + \left( \frac{ED_{50}}{Dose + ED_{50}} \right)}
\]

A correction factor was applied to the dose-blockade curve to account for the delay between antagonist administration and MOR blockade assessment in the included studies (Eq. 2). This enabled extrapolation of MOR blockade at the administration time point \( t_{\text{admin}} \) assuming no absorption phase. Because MOR blockade with naloxone and naltrexone is eliminated exponentially [24, 25], we used an exponential decay model for the correction (Supplement). The elimination rate constant \( k \) in this model was calculated from available estimates of MOR blockade half-life (Table S3–S7), and the time \( t \) was set to \( 0-t_{\text{measure}} \) to reflect the temporal delay.

\[
\text{Blockade}(Dose)_{\text{admin}} = \frac{Dose \times 100}{Dose + ED_{50}} \times \exp(-k \times (0-t_{\text{measure}}))
\]

Describing the time-blockade relationship

To describe blockade over time, we adapted the pkpro model for calculating the concentration-time profile of a drug that can account for absorption, infusion duration, and administration of multiple doses [23]. It uses \( V \) and \( Cl \) parameterization where \( V \) is the volume of distribution and \( Cl \) is the clearance. \( Cl \) can be expressed as \( k \times V \), with \( k \) being the elimination rate constant calculated from the blockade half-life (see above). By setting \( V \) to 1, \( Cl \) defaults to \( k \) and we can substitute the dose input in the pkpro function with the estimated MOR blockade at \( t_{\text{admin}} \) (Eq. 2), assuming no absorption phase. Absorption can instead be handled by the pkpro function by specifying an absorption rate constant \( k_a \) which can be calculated from the elimination rate constant \( k \) and the time-to-peak blockade \( t_{\text{peak}} \).

The time-to-peak blockade was estimated from time series data of \([11C] \) carfentanil activity in the absence and presence of an antagonist. To obtain the absorption rate constant \( k_a \) that results in the MOR blockade-time profile of a bolus dose peaking at \( t_{\text{peak}} \) we used the Lambert W function \( W(x) \) implemented in the praca package [26] (Supplement, Equation S10–S17).

The time-blockade profiles produced by the pkpro function treat blockade as a truncated measure of the concentration of the antagonist in the brain. While blockade has an upper limit of 100%, the central concentration may exceed the level necessary to produce full MOR blockade. We allowed model estimates to exceed 100% to reduce underestimation of the duration of full MOR blockade and facilitate detection of excessive concentration levels that contribute to high DOR and KOR blockade.

Simulating delta- and kappa-opioid receptor blockade

Due to limited availability of PET and dual-detector data, we simulated blockade of other opioid receptor subtypes. DOR and KOR blockade was simulated from the MOR blockade by multiplying \( ED_{50} \) for MOR blockade with the affinity of the antagonist drugs to MOR relative to DOR and KOR (see e.g., [27]). We specifically used the relative average affinity from studies of cloned human opioid receptors expressed on Chinese hamster ovary (CHO) cells (Table S1). Time profiles of DOR and KOR blockade assumed similar absorption and elimination rate as for blockade of MOR. Simulation-based estimates were compared to the available PET and dual-detector data on DOR and KOR blockade with naloxone and naltrexone.

RESULTS

Naloxone

Mu-opioid receptor blockade with intravenous naloxone. Two PET [12, 13] and two dual-detector studies [24, 28] have used \([11C] \) carfentanil to quantify MOR blockade with intravenous naloxone (Table 1). One PET study used a single dose of intravenous naloxone [12] while the other used four different doses [13]. The dual-detector studies used two [24] and eight [28] different doses. Due to the similarities in the protocols used across these studies, we considered the data suitable for quantitative synthesis (Supplement). Timing information was available from a dual-detector study with intravenous naloxone [24] and a PET study with intranasal naloxone [29].

Modeling the dose-blockade relationship. Blockade estimates in available PET and dual-detector studies (Table 1) are based on the mean signal recorded between 45–65 min after intravenous naloxone administration. Assuming a linear decrease in blockade within this 20-minute time window (Supplement, Figure S1), the reported blockade estimates approximately correspond to the blockade observed halfway (i.e., 10 min) through this section of the recording, i.e., at 55 min after intravenous naloxone administration \( t_{\text{measure}} \). Figure 1 displays the relationship between dose and MOR blockade (in black) for intravenous naloxone–55 min after administration. For the log-logistic model \( (RMSE = 9.44; \text{Pseudo-}R^2 = 0.92); \) Shapiro-Wilk test: \( W = 0.96, p = 0.62; \) Levene’s test: \( F_1, 14 = 0.06, p = 0.81 \), we obtained the parameter estimate \( ED_{50} (SE) = 0.0023 \text{mg/kg} (0.0004) \). Although dual-detector studies tended to report higher blockade estimates than PET studies, the difference in \( ED_{50} \) between the two methods was not statistically significant \( (\Delta ED_{50} = 0.0032 \text{mg/kg}, SE = 0.0016, \quad t_{4} = 2.05, \quad p = 0.06; \) Supplement, Equation S1, S2).

Blockade half-life estimates for naloxone obtained in PET [29] and dual-detector studies [24] are 100 and 120 min respectively. Using an average blockade half-life of 110 min \((SE = 10)\), we obtained the elimination rate constant \( k = 0.006 \) (Supplement, Equation S18) and the following adjusted formula for calculating MOR blockade as a function of dose at \( t_{\text{admin}} \) assuming no absorption phase:

\[
\text{MOR blockade}(Dose)_{\text{admin}} = \frac{Dose \times 100}{Dose + 0.0023} \times \exp(-0.006 \times (0 - 55))
\]

Describing the time-blockade relationship. Time series data from dual-detector studies with \([11C] \) carfentanil [11, 24, 25] show a maximum reduction in activity from the control condition (i.e., no naloxone) at 23–29 min after administration of intravenous naloxone. When naloxone was administered 5 min before \([11C] \) carfentanil.
carfentanil, maximum signal reduction occurred 29 min later with 2 mg naloxone [24], and 23 [11, 25], and 26 [24] minutes later with 1 mg/kg naloxone ($M = 24.7, SE = 1.2$; Supplement). This suggests that it takes ~25 min after $t_{\text{admin}}$ for naloxone to be distributed to the brain and occupy a maximum amount of central MOR after a single intravenous bolus of naloxone. With the elimination rate constant $k = 0.006$, and $t_{\text{max}} = 25$, we get an absorption rate constant of $k_a = 0.126$ (Equation S19). Together with the adjusted dose-dependent blockade estimate at $t_{\text{admin}}$, these timing-based parameters enable estimation of time-blockade profiles of various bolus doses of intravenous naloxone with the pkprofile function (Fig. 2).

### Table 1. Overview of positron emission studies used for modeling the dose-blockade relationship of intravenous naloxone and oral naltrexone.

| Antagonist Method           | N     | Measurement time point | MOR blockade |
|----------------------------|-------|------------------------|--------------|
| **Intravenous naloxone**   |       |                        |              |
| Frost et al. (1985) [12]   | 1     | 35–65 min              | 1 mg/kg 83%  |
| Frost and Mayberg (1990) [13] | PET  | 45–65 min              | 0.001 mg/kg 19% |
| Frost and Mayberg (1990) [13] | PET  | 45–65 min              | 0.01 mg/kg 65% |
| Frost and Mayberg (1990) [13] | PET  | 45–65 min              | 0.1 mg/kg 97%  |
| Frost and Mayberg (1990) [13] | PET  | 45–65 min              | 1.0 mg/kg 98%  |
| Kim et al. (1997) [24]     | 8     | 45–65 min              | 0.002 mg/kg 43% |
| Kim et al. (1997) [24]     | 8     | 45–65 min              | 0.03 mg/kg 81% |
| Villemagne et al. (1994) [28] | Dual-detector | 45–65 min | 0 mg/kg 0% |
| Oral naltrexone            |       |                        |              |
| Rabiner et al. (2011) [23] | PET  | < 8 h                  | 2 mg 27%     |
| Rabiner et al. (2011) [23] | PET  | < 8 h                  | 5 mg 43%     |
| Rabiner et al. (2011) [23] | PET  | < 8 h                  | 5 mg 46%     |
| Rabiner et al. (2011) [23] | PET  | < 8 h                  | 5 mg 58%     |
| Rabiner et al. (2011) [23] | PET  | < 8 h                  | 15 mg 61%    |
| Rabiner et al. (2011) [23] | PET  | < 8 h                  | 50 mg 92%    |
| Rabiner et al. (2011) [23] | PET  | < 8 h                  | 50 mg 98%    |

--- = not reported. * Information provided by J. James Frost.

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Fig. 1 The effect of intravenous (IV) naloxone dose on opioid receptor blockade ~55 min after administration. The bottom x-axis displays untransformed doses while the top x-axis shows the corresponding log$_{10}$-transformed doses. The dashed horizontal line indicates full (90%) receptor blockade. MOR blockade (solid black curve) is based on the data in Table 1 (black dots). DOR (dot-dashed red curve) and KOR blockade (long-dashed blue curve) were approximated from MOR blockade using the relative receptor affinities of naloxone (Table S1). Semitransparent bands indicate 95% confidence band (black band), or range based on highest and lowest reported affinity ratio (blue and red bands; Table S1). The estimated $ED_{50}$ for MOR, DOR and KOR blockade was 0.0023 ($SE = 0.0004$), 0.094 and 0.018 mg/kg, respectively.

Delta- and kappa-opioid receptor blockade with intravenous naloxone. Using relative average affinity values (Table S1), we estimated that $ED_{50}$ would be 41 times greater for DOR than MOR and 8 times greater for KOR than MOR. By multiplying $ED_{50}$ for MOR blockade with naloxone’s affinity to MOR relative to DOR and KOR, we obtain $ED_{50} = 0.094$ mg/kg for DOR blockade and $ED_{50} = 0.018$ mg/kg for KOR blockade. This results in the following equations for approximating DOR (Eq. 4) and KOR blockade (Eq. 5) at $t_{\text{admin}}$ assuming no absorption phase and $k = 0.006$:

$$\text{DOR blockade}(Dose)_{\text{admin}} = \frac{Dose \times 100}{Dose + 0.094} \times \exp(-0.006 \times (0 - 55))$$

(4)

$$\text{KOR blockade}(Dose)_{\text{admin}} = \frac{Dose \times 100}{Dose + 0.018} \times \exp(-0.006 \times (0 - 55))$$

(5)
Mu-opioid receptor blockade with oral naltrexone

Single dose: Several studies have used PET and dual-detection systems to investigate MOR blockade with single doses of oral naltrexone. Approximately 2 h after administration of 50 mg oral naltrexone, the $[^{11}C]$carfentanil signal in the brain matched the signal recorded 35–65 min after intravenous administration of 1 mg/kg naltrexone [25], suggesting almost complete blockade of mu-opioid receptors [13]. Consistent with this, Rabiner et al. [23] report that 50 mg oral naltrexone produced 95% mu-opioid receptor blockade within 8 h after administration. The same dose maintained >90% blockade at ~49 h after administration in the study by Lee et al. [25]. The observed blockade in this study decreased to 80% at ~73 h, 46% at ~121 h, and 30% at ~169 h after administration of naltrexone. Based on these data, Lee et al. [25] estimated the blockade half-life of naltrexone in the brain to be 72 h. Lower doses of oral naltrexone also produce substantial levels of blockade. Within 8 h of administration, 2, 5, and 15 mg blocked 27, 49, and 61% of the receptors, respectively [23].

Bednarczyk et al. [32] administered 12.5, 50, and 100 mg oral naltrexone and measured blockade after 3, 24, 72, and 144 h (see also [33]). The blockade estimates from this study are unfortunately unavailable. A study of obese subjects using the non-selective opioid $[^{11}C]$diprenorphine found that a single dose of 150 mg oral naltrexone produced 90% opioid receptor blockade 2 h after administration [34].

A dose-blockade curve ($RMSE = 6.96$; Pseudo-$R^2 = 0.92$; Shapiro-Wilk test: $W = 0.98, p = 0.93$; Levene’s test: $F_{1.9} = 0.08, p = 0.79$) based on data obtained within 8 h of administration was available from Rabiner et al. [23] (Table S1 and Fig. 3). This yielded an $ED_{50}$ of 5.59 mg ($SE = 0.80$) for naltrexone blockade with oral naltrexone and the following formula for converting dose to blockade:

$$\text{MOR blockade}(\text{Dose}) = \frac{\text{Dose} \times 100}{\text{Dose} + 5.59}$$

Insufficiently detailed timing information prevented generation of time-blockade profiles for this antagonist.

Repeated administration: The effect of repeated naltrexone administration on mu-opioid receptor availability has been investigated with PET in abstinent alcohol-dependent patients and in obese subjects. Following four days of treatment with 50 mg oral naltrexone, MOR blockade reached 95% in a sample of abstinent alcohol-dependent patients [14]. High levels of MOR blockade (75-97%) was observed in a similar patient sample after daily administration of 50 mg oral naltrexone for three days [35]. In an $[^{11}C]$diprenorphine study of obese subjects who had completed seven days of treatment with oral naltrexone, opioid receptor blockade was 70-80% in those who had received 16 mg/day and 90% in those who had received 32 and 48 mg/day [34]. Due to the non-selectiveness of the radiotracer and naltrexone’s preference for MOR (Table S1), it is possible that the estimated blockade in this study is an underestimation of MOR-specific blockade.

Delta- and kappa-opioid receptor blockade with oral naltrexone. The estimated affinity of naltrexone to MOR was 79 times greater...
Pharmacological blockade of a receptor system is a common method for probing the function of that receptor system in the human brain. Positron emission techniques yield data on the achieved level of blockade, but for studies of mu-opioid receptors, existing practices vary widely with regards to doses and assessment timing [43]. Here, we have synthesized the available PET and dual-detector data, and created models and web applications for calculating central opioid receptor blockade with the commonly used opioid antagonists naloxone and naltrexone. General recommendations for selecting optimal antagonist drugs, doses, and timings in basic human research based on our models and simulations are outlined in Table 2.

To simplify planning of future studies and evaluation of past studies with opioid antagonists, we have designed two web applications using the R package Shiny [44] that incorporate the models for intravenous naloxone (https://martintrostheim.shinyapps.io/planoxone/) and oral naltrexone (https://martintrostheim.shinyapps.io/plantrexone/) presented here. Key features of these applications include estimation of MOR, DOR and KOR blockade over time and across relevant dosing options in human clinical and experimental research, and estimation of total drug amount and cost for planned studies. The source code for both web applications is available on GitHub (https://github.com/martintrostheim/opioid-antagonist-planner).

Pharmacokinetic modeling of opioid antagonists typically focuses on plasma levels. However, for psychopharmacological studies it is important to understand the kinetics of the antagonist in the brain. Available PET and plasma data indicate that the central receptor blockade half-life of intravenous naloxone (110 min) and oral naltrexone (72 h) correspond relatively closely to the plasma half-life of these antagonists during the terminal phase (75 min for intravenous naloxone [45] and 96 h for oral naltrexone [46]), but not during the distribution phase. This suggests that plasma level would be a poor proxy for receptor blockade during the distribution phase and that modeling the elimination from the brain, as approximated here, is needed to inform psychopharmacological experiments in sufficient detail. These novel analytical tools can also aid interpretation of reported effects in the literature, since the presented models yield several insights into how previously used doses affect opioid receptors at the time of assessment. This is especially useful when interpreting the literature using naloxone, which has a relatively short half-life in the brain. Bolus doses of intravenous naloxone used in basic human research are often as large as or larger than 0.10–0.15 mg/kg (e.g., [47–49]). The initial bolus is sometimes supplemented with continuous infusion or an additional bolus (e.g., [49–51]), indicating that many authors may have underestimated the duration of the full blockade with these doses (~60 min). As our model shows, such supplements are only necessary if researchers want to assess the effect of full MOR blockade on outcomes measured more than an hour after the initial bolus (see Fig. 3). Yet, testing typically occurs within 15–60 min after the initial bolus. Lower doses may be sufficient to produce full MOR blockade, but for a shorter period of time. For example, our model estimates 0.01 mg/kg to maintain full MOR blockade for ~30 min. Combining a low bolus dose with continuous infusion can greatly extend the duration of the full MOR blockade for only a fraction of the cost of a high bolus dose (Table 2). An added benefit of using lower doses is that lower doses typically yield weaker and/or fewer side effects.

Studies using naltrexone to probe the endogenous mu-opioid system often administer 50 mg orally and begin outcome assessment ~60–120 min later (e.g., [52–55]). However, some studies use a higher dose of 100 mg (e.g., [56, 57]). The available PET and dual-detector data indicate that 50 mg is more than sufficient to produce full MOR blockade. This dose likely produces central concentration of naltrexone in excess of the dose required

**Fig. 3** The effect of oral (PO) naltrexone dose on opioid receptor blockade within 8 h of administration. The bottom x-axis displays untransformed doses while the top x-axis shows the corresponding log10-transformed doses. The dashed horizontal line indicates full (90%) receptor blockade. MOR blockade (solid black curve) is based on data (black dots) from Rabiner et al. [23] (Table 1), DOR (dotted red curve) and KOR blockade (long-dashed blue curve) were approximated from MOR blockade using the relative receptor affinities of naltrexone (Table S1). DOR (dot-dashed green curve) and KOR blockade (long-dashed blue curve) were estimated from MOR blockade within 8 h of administration. Blockade of DOR and KOR within 8 h of oral naltrexone administration can then be simulated with the following models:

\[
\text{DOR blockade} = \frac{\text{Dose} \times 100}{\text{Dose} + 441.83} \quad (7)
\]

\[
\text{KOR blockade} = \frac{\text{Dose} \times 100}{\text{Dose} + 11.19} \quad (8)
\]
to completely block mu-opioid receptors. Weerts et al. [14] observed high level of and low variability in MOR blockade with 50 mg oral naltrexone. However, this ceiling effect could also be a result of the repeated dosing schedule used in this study. Compared to acute doses, repeated dosing would likely cause naltrexone to accumulate, thereby increasing the blockade and extending its duration. This is consistent with the finding that daily administration of an oral naltrexone dose lower than 50 mg (i.e., 32 mg) can result in full opioid receptor blockade [34].

The earliest available MOR blockade estimate with oral naltrexone was collected ~2 h after administration and indicates that waiting 2 h is sufficient to reach full MOR blockade with 50 mg [25]. Considering that naltrexone plasma levels peak 1 h after oral administration [46], it is possible that the central blockade peaks in less than 2 h.

Exponential decay processes are considered to be complete after five to ten half-lives [58]. Assuming a blockade half-life of ~110 min with naloxone, we estimate that a washout period of 9 h should be sufficient to eliminate the MOR blockade. This allows researchers to arrange experimental sessions on consecutive days in within-subjects designs using naloxone. In contrast, naltrexone’s half-life is estimated to 72 h. After 1 week, which is a common intersession interval for within-subjects designs with oral naltrexone, a minimum intersession interval of 15 days would be necessary (i.e., five times the 72-hour blockade half-life [58]; see Supplement for further discussion).

Because the opioid antagonists currently marketed for human use are non-selective, there is a concern that DOR and KOR blockade could complicate inferences about MOR functions. The available data and the simulations presented here indicate that naloxone and naltrexone can produce considerable KOR and DOR blockade depending on the dose. Our results need further validation against human PET data as we used highly variable receptor affinity data from CHO cells to simulate human DOR and KOR blockade from MOR blockade. While using a lower naloxone dose could reduce DOR and KOR blockade, this comes at the cost of a shorter duration of full MOR blockade from a bolus injection (see Fig. 3). Applying a range of doses optimized for each receptor type can help disentangle the effects of MOR, DOR and KOR blockade on the outcome of interest. For researchers primarily interested in the mu-opioid system, more selective antagonists like GSK1521498 (Supplement) could be a viable alternative to naloxone and naltrexone provided that detailed timing information becomes available.

The models and recommendations presented here should be considered in light of certain limitations. Here, we have used 90% receptor occupancy as the threshold for full blockade [13, 14], but we cannot exclude that up to 10% of unblocked receptors may fulfil some endogenous functions.

While our models, recommendation, and web applications are based on data from multiple studies, it is important to note the limited availability of data on opioid receptor blockade with naloxone and naltrexone. Many of the studies tested a small sample of participants, and some blockade estimates are based on a dose applied to a single participant. Due to the variability in design and results between studies, researchers may want to adjust how different data sources are weighted when estimating blockade with our web applications. We have therefore enabled users to tweak all parameters of the models, including ED50, time-to-peak, half-life, and affinity ratios. Data from future PET studies can be used to validate and improve our models and recommendations (see Supplement for further discussion).

The majority of data informing this overview and models were collected from male participants. Men and women are pharmacokinetically and pharmacodynamically different [60], and these differences might affect the opioid receptor blockade.
produced by antagonist drugs. Many of the studies summarized here either present no analysis of gender effects [13, 24, 25, 28, 31, 32, 35, 37, 41], tested men only [23, 30, 36, 61], or tested a single participant [11, 12]. The few studies that report analyses of gender effects report no significant differences in receptor blockade between men and women [14, 27, 38, 40]. Note that these studies used large doses (e.g., 50–150 mg oral naltrexone) that produce full MOR and KOR blockade in most participants. However, Weerts et al. [14] report no significant relationship between gender and the low DOR blockade produced by daily dosing of 50 mg oral naltrexone. More data is needed determine the generalizability of our models to men and women separately.

The accuracy of opioid receptor blockade estimates with opioid antagonists depends in part on the radiotracer’s affinity to the various opioid receptor subtypes. $[^{11}C]$Carfentanil (MOR:DOR:KOR ratio = 1:137:1796) [62], $[^{11}C]$MeNTI (700:1:3250) [63], and $[^{11}C]$GR103545 (810:26800:1) [64] are all highly selective ligands, suggesting that blockade estimates based on the activity of these radiotracers are minimally influenced by their binding to other opioid receptor subtypes. $[^{11}C]$LY2795050 (36:213:1) [65] and $[^{11}C]$naltrexone) in these studies may have resulted in ceiling effects on MOR blockade and thereby prevented the detection of significant predictors. The inconsistent results from an already low number of studies using high and repeated doses highlight the need for more data to determine drug- and disease-related predictors of receptor blockade with opioid antagonists. Such data are key to evaluating the generalizability of our models and recommendations beyond generally healthy people.

The purpose of this primer is to help researchers select antagonist doses that can block endogenous ligands (as modeled by carfentanil) from binding to central opioid receptors. While the information synthesized here may also have implications for the choice of opioid antagonist doses in future clinical research, selection of doses for treatment purposes should primarily be based on knowledge about the relationship between opioid antagonist dose and clinical outcomes. Naloxone and naltrexone are competitive antagonists, meaning that highly potent opioids such as fentanyl, or high doses of opioids like heroin or oxycodone, may overcome the blockade produced by these antagonists. Larger doses or repeated administration of opioid antagonists may therefore be necessary in treatment settings to prevent opioid abuse and to reverse opioid overdose [69].

Pharmacological blockade of the endogenous MOR system with an antagonist such as naloxone and naltrexone is a commonly used method of investigating the role of this system in human psychological processes. While more data on the opioid receptor blockade produced by these antagonists are needed, we hope that this overview and the accompanying tools can aid researchers in evaluating past antagonist studies and in selecting appropriate drugs, doses, assessment time points, and intercession intervals for future studies.

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COMPETING INTERESTS
The authors declare no competing interests.

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