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RESEARCH PAPER

Anandamide transport inhibition by ARN272 attenuates nausea-induced behaviour in rats, and vomiting in shrews (Suncus murinus)

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BACKGROUND AND PURPOSE
To understand how anandamide transport inhibition impacts the regulation of nausea and vomiting and the receptor level mechanism of action involved. In light of recent characterization of an anandamide transporter, fatty acid amide hydrolase-1-like anandamide transporter, to provide behavioural support for anandamide cellular reuptake as a facilitated transport process.

EXPERIMENTAL APPROACH
The systemic administration of the anandamide transport inhibitor ARN272 ([4-(5-(4-hydroxy-phenyl)-3,4-diaza-bicyclo[4.4.0]deca-1(6),2,4,7,9-pentaen-2-ylamino)-phenyl)-phenylamino-methanone]) was used to evaluate the prevention of LiCl-induced nausea-induced behaviour (conditioned gaping) in rats, and LiCl-induced emesis in shrews (Suncus murinus). The mechanism of how prolonging anandamide availability acts to regulate nausea in rats was explored by the antagonism of cannabinoid 1 (CB1) receptors with the systemic co-administration of SR141716.

KEY RESULTS
The systemic administration of ARN272 produced a dose-dependent suppression of nausea-induced conditioned gaping in rats, and produced a dose-dependent reduction of vomiting in shrews. The systemic co-administration of SR141716 with ARN272 (at 3.0 mg·kg⁻¹) in rats produced a complete reversal of ARN272-suppressed gaping at 1.0 mg·kg⁻¹. SR141716 alone did not differ from the vehicle solution.

CONCLUSIONS AND IMPLICATIONS
These results suggest that anandamide transport inhibition by the compound ARN272 tonically activates CB1 receptors and as such produces a type of indirect agonism to regulate toxin-induced nausea and vomiting. The results also provide behavioural evidence in support of a facilitated transport mechanism used in the cellular reuptake of anandamide.

Abbreviations
ARN272, [4-(5-(4-hydroxy-phenyl)-3,4-diaza-bicyclo[4.4.0]deca-1(6),2,4,7,9-pentaen-2-ylamino)-phenyl)-phenylamino-methanone]; CB1, cannabinoid 1; CTA, conditioned taste avoidance; FAAH, fatty acid amide hydrolase; FLAT, FAAH-1-like anandamide transporter; TR, taste reactivity
Introduction

The Cannabis sativa plant has been known for centuries to exert therapeutic effects in the treatment of nausea and vomiting. More recently, cannabinoid agonists such as Δ^2-tetrahydrocannabinol have been found to be as effective as anti-emetic dopamine antagonists in human clinical trials (Carey et al., 1983; Tramer et al., 2001). The anti-emetic properties of cannabinoid agonists have been found to extend beyond humans to other emetic species, attenuating vomiting in ferrets (Simoneau et al., 2001; Van Sickle et al., 2001), cats (McCarthy and Borison, 1981) and the house musk shrew Suncus murinus (Kwiatkowska et al., 2004; Parker et al., 2004). Despite this long history, knowledge of how the endogenous cannabinoid system mediates nausea and vomiting is still incomplete.

Comparable to the effects of plant-derived cannabinoids, there is a body of evidence among animal models implicating the endocannabinoid anandamide, as important in the regulation of nausea and vomiting. The administration of exogenous anandamide has been found to have anti-emetic properties in the least shrew (Darmani, 2002) and in ferrets (Van Sickle et al., 2005). Deactivation of anandamide occurs through intracellular hydrolysis and is known to be mediated by the enzyme fatty acid amide hydrolase, FAAH (Desarnaud et al., 1995; Cravatt et al., 1996; McKinney and Cravatt, 2005). As such, prolonging the activity of endogenous anandamide through the inhibition of its degradation has also been found to reduce vomiting, specifically in the house musk shrew Parker et al., 2009) and in the ferret (Sharkey et al., 2007). FAAH inhibition has also been shown to attenuate nausea-induced responding, interfering with conditioned gaping reactions in rats (Cross-Mellor et al., 2007). These findings suggest that anandamide acts within the endocannabinoid system to regulate both nausea and vomiting.

Anandamide is known to be an endogenous agonist at cannabinoid 1 (CB1) receptors (Devane et al., 1992). The mechanism by which anandamide exerts its anti-emetic and anti-nausea effects is thought to be CB1 receptor mediated. The anti-emetic effects of exogenous anandamide administration have been found to be reversed by the CB1 antagonist, AM251 (Van Sickle et al., 2005), and the suppression of nausea by the FAAH inhibitor URB597 was reversed by the CB1 antagonists AM251 and SR141716 in rat (Cross-Mellor et al., 2007) and the house musk shrew respectively (Parker et al., 2009).

The specific mechanisms behind how anandamide signalling is terminated are still unfolding. How anandamide re-enters the post-synaptic cell appears to conform to a twofold process, cellular reuptake and subsequently cellular degradation. Previously, cellular reuptake has been hypothesized as occurring through either passive membrane diffusion driven by FAAH metabolism (Glaser et al., 2003), or by some previously unknown selective carrier system (Ligresti et al., 2004; Hillard et al., 2007). In support of the latter, Fu et al. (2012) recently reported the molecular identity of a facilitated anandamide transport mechanism, FAAH-1-like anandamide transporter (FLAT). As an isoform of the FAAH molecule, FLAT was found to bind anandamide selectively, but not other structurally similar molecules such as 2-AG, and without enacting any catalytic activity (Fu et al., 2012). Concurrently, Fu et al. (2012) identified an antagonist of anandamide transport, ARN272 ([4-(5-(4-hydroxy-phenyl)-3,4-diaza-bicyclo[4.4.0]deca-1-(6,2,4,7,9-pentaen-2-ylamino)-phenyl)-phenylamino-methanone]), which produced analgesic effects in rodent models of nociceptive and inflammatory pain in a CB1 dependent manner. The present study evaluated the potential of ARN272 to also reduce nausea and vomiting in animal models.

While emetic species are used to explore the regulation of vomiting, the subjective experience of nausea requires more consideration in animal models. Conditioned taste avoidance (CTA) is a measure that has often been used to evaluate the nauseating potential of drugs in rats (the extent to which a taste previously paired with an emetic agent is avoided through the amount a rat drinks in a consumption test). However, problematic evidence for CTA as a model of nausea in rats has been found. Anti-emetic drugs do not generally interfere with the establishment of CTA, (for review, see Parker et al., 2008). Also, rats not only avoid tastes paired with nauseating drugs, but they also avoid tastes paired with drugs they choose to self-administer (Berger, 1972; Reicher and Holman, 1977). Therefore, the CTA test cannot be considered a selective measure of nausea in rat. Physiological regulation of nausea can be studied, however, in the rat using their distinctive pattern of disgust reactions, most prominently conditioned gaping reactions (Parker et al., 2011). Despite the rat’s inability to vomit, the detection mechanism of nausea is still present, with similar orofacial musculature being activated by the gaping reaction as the orofacial vomiting reaction in emetic species (Travers and Norgren, 1986). Conditioned gaping reactions occur both when intra-orally infused with a bitter-tasting solution of quinine, as well as when exposed to cues (taste or context) previously paired with a drug, which produces vomiting in emetic species (Parker et al., 2008). Moreover, and unlike CTA, only drugs with emetic properties produce conditioned gaping reactions when paired with a flavour or contextual stimulus, and anti-emetic drugs consistently prevent the establishment of nausea-induced conditioned gaping in rats (Limebeer and Parker, 2000; 2003). Therefore, conditioned gaping can be used as a selective measure of nausea in rat.

The experiments reported here investigate the impact of anandamide transport inhibition on the endocannabinoid systems regulation of nausea and vomiting. Experiment 1 evaluated the potential of systemic administration of ARN272 to attenuate LiCl-induced conditioned gaping in rats and the potential of the CB1 receptor antagonist/inverse agonist SR141617 to reverse the ARN272-suppressed conditioned gaping response. The extent to which rats avoided a taste paired with the nausea inducing agent LiCl was also assessed using a CTA measure, to re-assert the selectivity of conditioned gaping in the measurement of nausea. Experiment 2 evaluated the potential of systemic administration of ARN272 to regulate LiCl-induced vomiting in the house musk shrew (S. murinus). Here, we provide behavioural support for anandamide reuptake occurring through a facilitated transport mechanism from the ARN272-suppressed gaping in rats and attenuated vomiting in shrews. As such, it is reasoned that prolonging the synaptic availability of endogenous anandamide augmented its action to tonically
activate CB₁ receptors and extend the anti-emetic and nausea attenuating properties of anandamide.

Methods

Animals

All animal care and experimental procedures complied with the recommendations of the Canadian Council on Animal Care and were approved by the Animal Care Committee of the University of Guelph. All studies involving animals are reported in accordance with the ARRIVE guidelines for reporting experiments involving animals (Kilkenny et al., 2010; McGrath et al., 2010). A total of 58 naïve male Sprague-Dawley rats (Charles River Lab, St. Constantr, QC, Canada) were used and equally distributed among the groups, with subjects ranging from 98 to 814 days of age. The sexes did not significantly differ in vomiting frequency in any analysis; therefore, males and females were pooled in all reported analyses. The shrews were bred at 1.0 mg·mL⁻¹ (1.0 mg·kg⁻¹ dose) at a volume of 1 mL·kg⁻¹. The dose of 1.0 mg·kg⁻¹ SR141716 was chosen based on prior effectiveness in reversing the breakpoint and reinstatement of nicotine self-administration (Forget et al., 2009), while also being found to not to potentiate the effects of emetic agents unlike doses of SR141716 at 2.5 mg·kg⁻¹ or higher (Parker et al., 2003).

The LiCl drug treatment 0.15 M (Sigma-Aldrich, St. Louis, MO, USA) used in all experiments was prepared in sterile water and administered at volumes of 20 mL·kg⁻¹ (127 mg·kg⁻¹) in rats in experiment 1 (Limebeer and Parker, 2000), and 60 mL·kg⁻¹ (390 mg·kg⁻¹) in shrews in experiment 2 (Rock et al., 2011).

Apparatus

The taste reactivity (TR) chamber consisted of a clear Plexiglas box (29 × 29 × 10 cm) resting on a glass surface. Two 60 W lights suspended from the apparatus illuminated the chamber. A mirror mounted at a 45° angle below the glass surface facilitated viewing of the ventral surface of the rat, specifically any orofacial responses. Each rat, prior to being placed in the chamber, was connected to an infusion pump (KDS100; KD Scientific Inc., Holliston, MA, USA) via a section of PE 90 tubing attached to their intra-oral cannula, which ran through a hole in the lid of the TR chamber. All orofacial and somatic responses were recorded during the session via a video camera (Sony DCR-HC28 Handy Cam, New York, NY, USA) connected directly to a desktop PC using Roxio VideoWave Premiere Suite 8 video capture program (Corel Corporation, Ottawa, ON, Canada).

Vomiting in shrews was measured in a clear Plexiglas chamber (22.5 × 26 × 20 cm) illuminated by a 60 W light suspended from the chamber’s floor. A mirror was mounted at a 45° angle beneath the chamber floor, which allowed for clear viewing of the ventral surface of the shrew, and an observer counted the number of vomiting episodes.

Procedure

Experiment 1: potential of ARN272 to attenuate LiCl-induced conditioned gaping, and reversal of ARN272-suppressed gaping by SR141716. All rats were surgically implanted with intra-oral cannula under isoflurane anaesthesia as described by Limebeer et al. (2010). Following recovery from surgery (3 days), rats received a single adaptation trial to habituate them to the chamber and the infusion procedure. During the adaptation trial, rats were placed individually in the TR chamber and received a 2 min intra-oral infusion of water (reverse osmosis water infused at 1 mL min⁻¹). On the following day, rats received the first of two conditioning trials (separated by 72 h). On each conditioning trial, rats received a pretreatment injection of ARN272 or VEH 120 min prior to the conditioning trials. During conditioning trials, rats were intra-orally infused with a saccharin solution (0.1%) for 2 min (1 mL min⁻¹) and orofacial and somatic reactions were recorded on video. Immediately following the saccharin infusion, the rats were injected with LiCl (0.15 M) or saline, and then returned to their home cage. Two additional groups were added (after ARN272 at 3.0 mg·kg⁻¹ attenuated gaping)
where a pretreatment of ARN272 at 3.0 mg·kg$^{-1}$ or VEH was given 120 min prior, and with SR141716 30 min prior, to each conditioning trial. The groups were VEH-Saline (VEH-SAL), $n = 9$; VEH-LiCl, $n = 8$; 0.1 mg·kg$^{-1}$ ARN272-LiCl, $n = 9$; 1.0 mg·kg$^{-1}$ ARN272-LiCl, $n = 8$; 3.0 mg·kg$^{-1}$ ARN272-LiCl, $n = 8$; 1.0 mg·kg$^{-1}$ SR-3.0 mg·kg$^{-1}$ ARN272, $n = 8$; 1.0 mg·kg$^{-1}$ SR-VEH, $n = 8$.

Seventy-two hours following the second conditioning trial, the rats received a drug-free TR test. During the TR test, rats were re-exposed to a 2 min intra-oral infusion of saccharin solution and their oro-facial and somatic responses again recorded. All video recordings were later scored by a rater blind to the experimental conditions using ‘The Observer’ (Noldus Information Technology Inc., Leesburg, VA, USA).

Following the TR test, the rats were returned to their home cages and at 16:00 h, their water bottles were removed to begin a water deprivation regime in preparation for the CTA test. At 08:00 h the following morning, the rats received a one-bottle test in which a graduated tube of 0.1% saccharin solution was placed on the home cage, and the amount consumed was recorded at 30 and 120 min intervals. A one-bottle test was used as there is evidence to suggest it is more sensitive in detecting between group differences in the strength of taste avoidance than a two-bottle test where both water and saccharin are made available, (Batsell and Best, 1993).

Results

**Experiment 1: systemic ARN272 suppressed LiCl-induced conditioned gaping in rats, and was reversed by the CB$_1$ receptor antagonist SR141716**

**Gaping measure.** The systemic administration of ARN272 produced a dose-dependent suppression in nausea-induced conditioned gaping in rats, effects that were reversed by pretreatment with the CB$_1$ receptor antagonist SR141716. Figure 1 presents the mean number of gapes on the drug-free test day by drug pretreatment group. The one-way ANOVA revealed a significant effect of drug pretreatment, $F(6, 51) = 10.83$, $P < 0.001$; subsequent post hoc Bonferroni tests revealed that ARN272 3.0 significantly attenuated gaping as compared with all groups other than VEH-SAL ($P < 0.01$), which also differed from all other groups ($P < 0.01$).

**CTA measure.** All pretreatment groups demonstrated greater taste avoidance than the VEH-SAL group at both time intervals (30, 120 min) in that less saccharin was consumed. There were no saccharin consumption differences specifically between the pretreatment conditions that received LiCl, at any of the time intervals. Figure 2 presents the mean cumu-

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**Figure 1**

Mean (+SEM) number of gapes by rats on drug-free test day, in experiment 1, by each of the groups. VEH-SAL ($n = 9$), VEH-LiCl ($n = 8$), ARN272 0.1 mg·kg$^{-1}$ ($n = 9$), ARN272 1.0 mg·kg$^{-1}$ ($n = 8$), ARN272 3.0 mg·kg$^{-1}$ ($n = 8$), ARN272 3.0 mg·kg$^{-1}$ + SR 1.0 mg·kg$^{-1}$ ($n = 8$), VEH + SR 1.0 mg·kg$^{-1}$ ($n = 8$). ***$P < 0.001$ indicates that group ARN272 3.0 gaped less than VEH, ARN272 0.1, SR 1.0 and ARN272 3.0 + SR 1.0, and that group VEH-SAL gaped less than all other groups. The number of rats that gaped in each group is indicated above each bar.
The one-way ANOVAs revealed significant differences in the mean cumulative amount of saccharin consumed at 30 min, $F(6, 51) = 34.66, P < 0.001$, and at 120 min, $F(6, 51) = 27.66, P < 0.001$. Subsequent post hoc Bonferroni tests revealed the VEH-SAL group as drinking significantly more saccharin than all other groups ($P < 0.001$) at each time period.

Experiment 2: systemic ARN272 reduced LiCl-induced vomiting in shrews

**Vomiting measure.** The systemic administration of ARN272 produced a dose-dependent reduction of vomiting in shrews. Figure 3 presents the mean number of LiCl-induced vomiting episodes for shrews pretreated with VEH, ARN272 9.0 and ARN272 18.0. The one-way ANOVA revealed a significant effect of drug pretreatment, $F(2, 18) = 3.75, P < 0.05$, with planned comparisons revealing that ARN272 18.0 significantly attenuated vomiting as compared with VEH ($P < 0.05$).

Discussion

Inhibition of anandamide transport by pretreatment with ARN272 reduced both LiCl-induced conditioned gaping in rats and LiCl-induced vomiting in the house musk shrew. The present study’s findings were consistent with existing evidence that increased anandamide availability through FAAH inhibition also attenuates nausea-induced gaping in rats (Cross-Mellor et al., 2007). Both the attenuation of nausea by FAAH inhibition (Cross-Mellor et al., 2007) and the attenuation of nausea by ARN272 were reversed by pretreatment with a CB1 antagonist. As well, consistent with the reported anti-emetic effects of increasing anandamide availability through FAAH inhibition in shrews (Parker et al., 2009) and ferrets (Sharkey et al., 2007), anandamide transport inhibition by ARN272 reduced vomiting in the house musk shrew. Although the mechanism for this reversal was not specifically evaluated here, the suppression of vomiting by FAAH inhibition was reversed by the CB1 antagonist, SR141716 (Parker et al., 2009).

The increased doses required to attenuate vomiting as compared with those required to suppress conditioned gaping are likely due to the extremely high basal metabolism of the shrew. The common European white-toothed shrew *Crocidura russula* (from the same subfamily as the house musk shrew, *Crocidurinae*) has been found to have a metabolic quotient higher than that required by its small body weight (Vogel, 1976). In comparison to the rat, Durrer (1982) found the common European white-toothed shrew to exhibit an increased number of cells per cm$^2$ of liver tissue, an increase in mitochondrial volume and a relative larger liver volume. Such differences in the cytoarchitecture of liver hepatocytes between shrew and rat likely account for the higher dose requirements in shrew due to greater clearance by increased liver metabolism.
The use of endocannabinoid transport inhibitors have potential not only as a means to further elucidate the role and function of the endocannabinoid system, but also as therapeutic agents. The indirect agonism produced by anandamide transport inhibition seemingly augments the body’s own regulatory processes. Indirectly enhancing anandamide action in tissue where synthesis, release and degradation is already occurring, could provide a safer and more selective action than direct agonists (Di Marzo, 2008).

The present experiments suggest that anandamide transport inhibition tonically activates CB1 receptors to regulate nausea and vomiting, and provides in vivo support for a facilitated transport mechanism used in the cellular reuptake of anandamide.

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**Conflict of interest**

None.

**References**


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Fu et al. (2012) previously reported greater anandamide accumulation in cells expressing FLAT as compared with controls, and an elevation in extracellular anandamide in cells, which overexpress FLAT, suggesting a bidirectional mechanism of anandamide translocation. Endocannabinoids have a short duration of action (Di Marzo, 2008) and a highly localized form of neural communication (Wilson and Nicoll, 2001), being produced on demand as required. As such, it is logical that the termination of endocannabinoid signalling would need to be as efficient a process as the one required for its activation. The suggested involvement of FLAT in both the release and the reuptake of anandamide may impact how FLAT inhibition would alter the synaptic availability of anandamide in regions responsible for the control of nausea and vomiting. Should FLAT inhibition favour reuptake, synaptic availability would increase. Conversely, if export is favoured, synaptic availability of anandamide would decrease. If FLAT inhibition occurs without favour bidirectionally, it would be expected to prolong the presence of any available anandamide within the synapse. It is hypothesized then that FLAT inhibition acted to regulate nausea in rats and vomiting in shrews by prolonging the synaptic availability of endogenously produced anandamide. This extended agonist action of anandamide at CB1 receptors was evidenced by the reversal of ARN-suppressed gaping by a CB1 receptor antagonist. Here, the behavioural evidence suggests that the efficiency of anandamide synaptic removal occurs via a facilitated transport system, through FLAT. Further investigation of the mechanism by which ARN272 acts to regulate nausea and vomiting is however warranted.

The challenges associated with the many available compounds that pharmacologically target anandamide transport are their diverse off-target effects, most notably at higher concentrations, FAAH inhibition (Hillard et al., 2007). The inhibition of anandamide transport by ARN272 does appear to be selective in that the compound produced only a weak and incomplete inhibition of FAAH in vitro, and had little to no inhibitory effect on other endocannabinoid metabolizing enzymes such as monoacylglycerol lipase, (Fu et al., 2012). There are further benefits to using indirect agonism by a selective transport inhibitor as compared with FAAH inhibition to understand the role of anandamide within the endocannabinoid system. The catalytic activity of the FAAH enzyme has been found to impact other N-acyl ethanlamidases as well as anandamide; such as oleoyl ethanolamide and palmitoylethanolamide (Bracey et al., 2002; Kathuria et al., 2003), bioactive molecules that act on non-cannabinoid receptors. As such, the use of FAAH inhibition to isolate anandamide-mediated effects remains problematic.

Existing evidence to suggest that anti-nausea treatments do not interfere with CTA learning in rats (Rabin and Hunt, 1983; Rudd et al., 1998; Limebeer and Parker, 2000; Limebeer et al., 2012) was supported in that pretreatment with ARN272 did not attenuate CTA responding. Only conditioned disgust, the gaping reaction specifically produced by the nausea induced by LiCl, was blocked by pretreatment with ARN272. As such, the present findings suggest that ARN272 is not interfering with learning per se, instead, it interfered with LiCl-induced nausea selectively necessary for the production of gaping reactions, but not CTA (see Parker et al., 2009).


