Long-Term Visceral Hypersensitivity Following Induction of Chemical Colitis is Primarily Reduced via Kappa Opiate Pathways - A Comparative Study of Different Opioidergic Subtypes

D Engel¹,², M Stutz¹,², JC Miller¹,², B Flogerzi¹,², L Tovar¹,², Scheurer U¹,² and Gschossmann JM*¹,²

¹Department of Visceral Surgery and Medicine, Inselspital, Switzerland
²Department of Clinical Research, Inselspital, Switzerland

*Corresponding author: Juergen MGschossmann, Klinikum Forchheim, Department of Internal Medicine, Krankenhausstrasse 10, D-91301 Forchheim, Germany

Introduction

Irritable bowel syndrome (IBS) is a gastrointestinal syndrome characterized by chronic abdominal pain and altered bowel habits in the absence of any organic cause. Although IBS represents 25 to 50 percent of all referrals to gastroenterologists [1] and is the second most frequent cause for work absenteeism after common cold [2]. It remains a chronic condition without any generally accepted and successful therapy for all IBS patients. Clinical studies have traditionally brought attention to colonic motility as potential cause for the symptoms associated with IBS. One study described 3cpm (contractions per minute) in the unstimulated colon in patients with IBS compared to 6cpm in controls [3]. Other studies have described selective hypersensitization of visceral afferent nerves in the gut in IBS patients based upon the observation that many patients with IBS have lower tolerance for rectal balloon distension than controls [4,5]. Interestingly, epidemiological studies have revealed in IBS patients an increased prevalence of persistent...
alterations in visceral nociception following acute gastroenteritis. The extent to which these observations might contribute to the pathogenesis of IBS in the general population remains unclear, as only a relatively limited number of patients have been studied.

Since IBS generally presents as a complex of symptoms, treatment is based on the predominant symptom and subtype (constipation vs. diarrhea vs. mixed vs. unclassified). Lifestyle and dietary modification are of cornerstone importance and considered first line treatment. Further, in patients with IBS and abdominal pain, antispasmodics and antidepressants, anxiolytics and antibiotics are used [6,7]. Limited evidence suggests that antineuropathic, selective serotonin reuptake inhibitors, probiotics and herbal treatments may be useful in IBS patients [8]. However, some of the serotonin receptor antagonists/agonists used for causal therapy in IBS have been removed from the market, due to severe side effects such as ischemic colitis and serious cardiovascular complications. Hence at the time being, research is still in progress for a specific causal therapy with as little unwanted side effects as possible. Nociceptive mechanisms of potential interest in this context are opioidergic pathways, the glutamate receptor axis (e.g. NMDA receptor antagonists) as well as a combination of these two pathways. In the study of Burton and Gehbhart the pain threshold to CRD in rats with or without acute inflammation of the colon was elevated after administration of the κ-opiate receptor agonist (ORA) U50.488 [9]. Others found that the analgesic effect of fentanyl can be potentiated by a co-application of ketamine [10].

Endogenous opioids are distributed throughout the whole organism and produce a diversity of biologic effects such as analgesia, motoric or stress responses. Opioids bind to three different receptor subtypes, the κ-, μ- and δ-opioid receptors (OR). κ-OR are mainly found in the tunica muscularis as well as in the myenteric plexus (Auerbach) where they have an impact on analgesia and motility. μ-OR of the gastrointestinal tract are primarily located in the tunica mucosa, submucosa, as well as in the submucosal plexus (Meissner). Finally, the δ-OR are found in the submucosal and myenteric plexus and the peripheral nerve fibers [11]. Given the hypomotile effect of opioids, their use in treatment of IBS may be accompanied by problematic unwanted effects such as constipation because of diminished intestinal passage and secretion [12,13]. Further gastrointestinal side effects may include emesis and nausea, but also reduced motility of the gallbladder and elevated tonus of the sphincter oddi. Thus, it would be desirable to identify an opioidergic substance with maximal analgesic potency and minimal side effects. This holds especially true because nausea, vomiting and constipation often represent symptoms of IBS patients. In our study, we focused on two main objectives. First, to reproduce the known long-term effect of a chemically induced colitis on visceral hypersensitivity to tonic distension in Lewis rats in an animal model with phasic colorectal distensions as stimulus in order to simulate the hyperalgesic status of IBS patients following acute gastrointestinal infection. Secondly, to study the impact of different opioid receptor agonists and the NMDA antagonist ketamine on long-term visceral hypersensitivity following induction of TNB colitis. We aimed to compare the clinically used μ-ORA’s fentanyl and morphine, the δ-ORA SNC80 as well as the NMDA antagonist ketamine with the κ-ORA U50.488.

Material and Methods

Animals

As laboratory animals male Lewis rats (Harlan, Horst, Netherlands) weighing between 200 and 320g are used. They housed individually in temperature-controlled rooms with a 12h dark–light cycle. Access to water and food was free. Before induction of the TNB colitis as well as before implantation of the electrodes, the animals were fasted for 24 hours. The state department of veterinary services approved the study.

Drugs

Every drug is given in a 1ml solution. When a combination of two drugs were used, each drug is given in a 0.5ml solution to reach the targeted volume of 1ml. The dose of each drug was chosen based upon the literature. For the κ-ORA 1S,2S-U50.488 (18mg/kg BW i.p. [14], Sigma-Aldrich Chemie, Buchs (SG), Switzerland) H₂O dist. was used as solvent. The μ-ORA fentanyl (0.05mg/kg BW i.p. [15], central pharmacy of the university hospital of Berne, Switzerland) and morphine (3mg/kg BW i.p. [15,16], central pharmacy of the university hospital of Berne, Switzerland) was used as solvent. For the δ-ORA SNC80 (10mg/kg BW i.p. [18], Sigma-Aldrich Chemie, Buchs (SG), Switzerland) DMSO (Sigma-Aldrich Chemie, Buchs (SG), Switzerland) was used as solvent.

Induction of Colitis

To induce a local colorectal inflammation, 0.4ml of 100% 2,4,6-trinitro-benzene-sulfonic (TNB) acid (Sigma-Aldrich Chemie, Buchs (SG), Switzerland) and 0.4ml of 50% ethanol (test group) or an equal volume of 0.9% saline (control group) was injected via a nelathon catheter (outer diameter 3mm; Medicoplast, Illingen, Germany) under combined anaesthesia with ketamine hydrochloride (1ml/kg BW i.p., Ketamin 10%®, Sanofi-Ceva, Dusseldorf, Germany) and xylazin hydrochloride (0.05ml/animal i.p., Xylazin 2%®, Sanofi-Ceva, Dusseldorf, Germany). The catheter was inserted 8cm into the colon. This model has been frequently used and is well characterized [19].

Implantation of EMG Electrodes

To quantify the visceromotor response (VMR) to phasic colorectal distension (CRD), the activity of the abdominal wall...
musculature was recorded by electromyography (EMG). For this purpose, we implanted small-sized electrodes into the lateral abdominal wall of the animals following median laparotomy under combined anesthesia with ketamin hydro-chloride (1ml/kg BW i.p., Ketamin 10%®, Sanofi-Ceva, Duesseldorf, Germany) and xylazin hydrochloride (0,05ml/animal i.p., Xylazin 2%®, Sanofi-Ceva, Duesseldorf, Germany). As bipolar electrodes, teflon-insulated silver wires (Cooner Wire Co., Chatsworth, CA/USA) were used, passing through the abdominal wall musculature. After surgical intervention, animals could recover for one week. After this week, animals were put together in groups of five with food and water ad libidum.

**Timetable**

First forty male Lewis rats were divided into two groups of 20 animals, a TNB colitis (test) and a saline group (control). Five weeks after induction of a TNB colitis or colorectal injection of saline, the EMG electrodes were implanted. After a recovery period of one-week, phasic CRD experiments were started. The VMR to CRD was studied with eight test substances or their combinations (saline, U50.488, fentanyl, morphine, SNC80, ketamine, ketamine/fentanyl or ketamine/morphine). In the process of the testing, the animals were given a recovery period of three days after each series of stimulation (Figure 1). The order of the injected substances was identical in all animals and every substance was used in each animal. Therefore, the sequence of administration was not randomised.

![Figure 1: Timetable.](image)

**Measurement of the Visceromotor Response**

The CRDs were performed with pressure-controlled air inflation of a 4,5cm flexible balloon, which was connected to a computer-controlled barostat (Distender Series IIITM barostat, G&J Electronics, Willowdale, Ontario/Canada). The VMR to the phasic CRDs was recorded and measured with an EMG device (ISOLAB/IDAA, Intestinal Data Acquisition and Analysis, Fa. Standard Instruments, Karlsruhe, Deutschland). The VMR to CRDs in the rat has been extensively characterized by Ness [20]. Under slight anesthesia with isoflurane, the lubricant-coated balloon (sonography lubricant, central pharmacy of the university hospital of Berne, Switzerland) was inserted into the colorectum, positioned about 1cm proximal of the anus and fixed to the tail with adhesive
Afterwards, the animals were placed in a Boreman cage and the EMG electrodes connected with the EMG recording system. The study protocol consisted of phasic CRDs (every minute 40 mmHg for a period of 30 sec). Before the application of a test substance, we performed three CRD’s as basic measurement (measurement A). The application of the test substance was followed by 15 CRD’s. The first three and the last three CRD’s were denoted measurement B and C, respectively. The first 3 CRDs and the subsequent generated 15 CRDs were 11 min apart. The EMG was recorded during the whole trial, starting 1 min before the first CRD and ending 1 min after the last CRD (Figure 2). The drugs were administered intraperitoneal, either 1 min (Ketamine) after measurement A or 1 min before measurement B (saline, 1S,2S-U50.488, £Fentanyl, morphine and SNC80).

Statistical Analyses

To evaluate the effect of a chemically induced colitis on visceral nociception, the difference in EMG activity (in µV) over time, resulting in an area under the curve (µV*time), between period C and B in the test group (TNB) versus the control group (saline) was compared. In this first part of the experimental protocol, a possible adaptation in the course of time was analysed. Therefore, only the first and the last three distensions were studied. To quantify the impact of the different ORA’s and ketamine, the differences in EMG activity between the measurement C and A were compared. Since ketamine is hypothesized to act only on activated pain pathways, the last three measurements of the repetitive distensions were evaluated. This protocol was consistently applied to all the substances to compare the results. Therefore, a negative value represents a decrease of VMR and thus a decrease in visceral sensory function. A mean value for the three consecutive distensions was calculated. Due to the non-normal distribution, the Wilcoxon rank sum test as well as the Wilcoxon signed rank test for statistical analysis were used. All values are given as the median of the area under the curve. A p-value of ≤ 0.05 was considered significant.

Results

Influence of TNB colitis on VMR

After i.p. administration of saline, an increase of the VMR to CRD was recorded during the following 15 repetitive CRD’s in the colitis group whereas the control group showed a decrease resulting in a significant difference between the test results (differences of the measurement C and B) of the treatment and the control group (49 vs. -161; p = 0.006) (Figure 3).

Influence of different ORA’s on VMR in TNB Animals

- **κ-ORA:** Compared to saline-treated animals, VMR to CRD significantly decreased from measurement A to C after application of U50.488 in TNB animals (183 vs. -135; p=0.003).

- **µ-ORA:** After application of both fentanyl and morphine (18 vs. 183; p=0.012 for fentanyl, 4 vs. 183; p=0.002 for morphine) a significant reduction of VMR in TNB-treated animals was recorded compared to saline-treated rats.

- **δ-ORA:** Compared to saline, the application of the δ-ORA SNC80 did not have any statistically significant effect on the VMR (183 vs. -105; p 0.067) (Figure 4).

Influence of Ketamine on VMR in colitis Animals

The comparison of VMR to CRD during measurement A and C did not show any significant reduction after application of ketamine compared to saline (85 vs. 183; p=0.376) (Figure 4).

Influence of Ketamine Combined with µ-ORA on VMR in colitis Animals

The combination of ketamine with fentanyl caused a significantly greater decrease of the VMR (0 vs. 183; p=0.027) compared to saline. Similar findings were seen for the combination of ketamine with morphine compared to saline (-31 vs. 183; p=0.002). The comparison of the effect of ketamine combined with fentanyl to the κ-ORA U50.488 shows a slightly stronger effect of U50.488 (0 vs.-135; p=0.094). The same counts for the combination of ketamine with morphine compared to U50.488 (-31 vs. -135; p=0.068) (Figure 4).
Figure 4: Differences of the measurements C versus A of the VMR in μV*time. Only the measurements of the TNB colitis animals are displayed. The boxplot shows the median, the 25th and 75th percentile, as the minimum and maximum. *= p-value ≤ 0.05, **= p-value ≤ 0.005, ***= p-value ≤ 0.0005

Discussion

Since alterations in visceral sensory function are considered a major etiological factor in the pathogenesis of IBS, opioidergic neurotransmitter pathways represent an attractive target to treat IBS. Unfortunately, unwanted side effects of several opioids prevent their widespread use in this context. Given the fact that these side effects depend on the type of ORA, the effect of different ORA’s on visceral nociception deserves particular attention to identify a suitable ORA with maximal impact on visceral sensory function and minimal side effects. The focus of our present study was the characterization of different opiate subtypes for pain modulation following induction of chemical colitis in a rat model. Therefore, the comparison of TNB versus control animals served as confirmation of prior results of tonic CRD and proof of concept for the present set-up of phasic CRD as suitable animal model for IBS. Based upon these data, the purpose of this study was not to determine a specific and IBS-limited effect of opiate subtypes in IBS (in contrast to control animals). On the contrary, the study aimed at defining an opiate subtype with maximal analgesic effect and, at the same time, minimal gastro-intestinal side effects as possible therapeutic tool for the treatment of IBS. In our study with male Lewis rats, a chemically induced colitis with TNB caused persisting visceral hyper-sensitivity thus simulating visceral hypersensitivity seen in IBS patients following acute gastrointestinal infection. Both the κ-ORA U50.488 and the μ-ORA’s fentanyl and morphine reduced VMR to CRD with U50.488 displaying the most analgesic potential. No further analgesic effect could be observed by combining ORA’s with the NMDA antagonist ketamine.

Influence of TNB Colitis on VMR to CRD

In our series of phasic CRD tests, it was possible to reproduce the already known influence of TNB colitis on the VMR to tonic distensions. Contrary to our earlier experiments, in the present study we modified our experimental set-up and used phasic distensions as stimulus. In the control animals, we observed a reduction of the EMG response during the 15 phasic CRD’s after an i.p. application of saline. This finding can be explained by adaptation to the distension stimulus in healthy animals. This mechanism appears to fail in TNB-pretreated animals. In these latter animals, VMR to CRD raises significantly during the series of distensions. Similarly, to our results, among others, Zhou et al. described an increase in sensitivity and hyperalgesia following the induction of a TNB colitis in the rat model [21]. This was tested by using mechanic and thermal stimulation of the hind-paws. Zhou et al. concluded that this somatic hypersensitivity reflects tonic impulse input from the inflamed colon along with central sensitization.

Influence of Different ORA’s on VMR in Colitis Animals

κ-ORA: Both in TNB and control animals, the application of the κ-ORA U50.488 led to a significant reduction of the VMR to the CRD stimulus with a greater potency in the TNB-pretreated colon. The analgesic effect of κ-ORA on visceral nociception has been controversially discussed in the literature [22,23]. Danzebrink et al. postulated that only the systemic but not the intrathecal application of the κ-ORA led to an antinociceptive effect in visceral pain [24]. Other reports suggested that subcutaneous, but not central administration of κ-ORAs inhibited the VMR to CRDs [25].
In their model, Larsson et al. used mice and distensions were performed with increasing as well as repeated steady distensions lasting 10s and with 5 min intervals. U-69,593 was used as κ-ORA. Kitamura et al. showed that intravenous administration of U50,488 produced strong visceral antinociception in the uninflamed colon [26]. This effect was completely antagonized by naloxone. It has been shown that κ-, but not µ- and δ-ORAs, given intra-arterial attenuated the response to the noxious stimulus. For this purpose, Su et al. performed recordings of mechanosensitive pelvic nerve afferent fibres of the decentralized S1 dorsal root in anesthetized rats. This antinociceptive activity of κ-ORAs such as U50,488 during inflammation is partly mediated by NO [27]. The peripheral injection of U50,488H was enhanced by the injection of the NO-releasing agent FK409 and reversed by the administration of the NO scavenger PTIO [28]. Another group demonstrated that the peripheral antinociception induced by the κ-ORA bremazocine was mediated by L-arginine/NO/cGMP activation [29]. The long-term persistence of the analgesic effect after induction of chemical colitis could be explained by the theory that the corresponding receptors are either not localized or restricted to the site of inflammation or they are not affected by the colitis. Other groups suggest a peripheral upregulation of κ-opioid receptors in or following colonic inflammation [30-32].

µ-ORA Fentanyl: Following an intraperitoneal application of fentanyl, we observed a decreased pain inhibitory effect on visceral nociception compared to morphine. We are not able to explain why fentanyl leads to a lesser reduction in pain than morphine in our set-up. Theoretically, the use of both ORA at different concentrations could have changed this finding.

µ-ORA Morphine: In our study, the administration of morphine was associated with a significant pain reduction to CRD in TNB-treated animals. Galantin et al. discovered that the µ-opioid morphine is highly active as analgesic against peritoneal visceral pain states in rats [33]. Similar effects have been described earlier, where systemic injections of morphine dose-dependently inhibited the VMR to CRD in rats with uninflamed or inflamed colon [22].

δ-ORA SNC80: In our study, application of the δ-ORA SNC80 reduced the EMG response to the visceral stimulus. Although this effect did not reach statistical significance in our study due to the largely dispersed data, it supports earlier findings by Sengupta et al. who described in a behavioural study that the δ-ORA SNC80 attenuated the responses of pelvic nerve afferent fibers in an uninflamed as well as in an inflamed colon but with significantly greater potency in the inflamed colon [27]. The greater potency of κ-ORAs in the TNB-inflamed condition suggests a peripheral upregulation of κ-OR's in colonic inflammation. The existence of distinct subtypes of δ-opioid receptors, δ1 and δ2, has been proposed by several research groups [32,34]. This hypothesis has been supported by the introduction of specific agonists and antagonists for both receptor subtypes. Bilsky et al. [35] reported that antinociception in response to intracerebroventricular application of SNC80 was fully antagonized by the DALCE/Cys-Delt combination but only partially blocked when these δ1 and δ2 receptor antagonists were separately given [36]. Therefore, these authors postulated that SNC80 is an unspecific δ-agonist. Pacheco et al. described the peripheral occurrence of δ1 and δ2-receptors and their antinociceptive function as result of the activation of the L-arginine/NO/cGMP pathway [37].

Influence of Ketamine on VMR in Colitis Animals

In our study, the intraperitoneal application of ketamine as non-competitive antagonist of the NMDA receptor [38,39] did not lead to any statistically significant decrease of the VMR to CRD. One possible explanation could be that the VMR in our pain model was not mediated by NMDA receptors at all. It can also be argued that the applied pressure for CRD in our experimental set-up did not translate into a pain response high enough to activate the glutamate pathways. This might indicate that the application of ketamine is only effective after the onset of pain. This excludes a prophylactic use of the NMDA antagonist. Alternatively, the missing effect of ketamine in areas of chronic inflammation might be the result of an up-regulation of spinal NMDA receptors. As described by Kolhekar [40], spinal NMDA receptors are directly involved in local mechanisms that lead to enhanced spinal transfer of visceral nociceptive information and thus may participate in mechanisms leading to central hyperexcitability and visceral hyperalgesia. Zhou et al. also suggest a role for colonic myenteric plexus NMDA receptors in the development of neuronal plasticity and visceral hypersensitivity in the colon [21]. Up-regulation of NMDA receptor subunits may reflect part of the basis for chronic visceral hypersensitivity in conditions such as post-infectious irritable bowel disease. Thus, in our study, the used dose of ketamine could have been below the necessary level to show a significant effect.

Influence of Ketamine Combined with µ-ORA on VMR in Colitis Animals

Nadeson et al. showed in their work the possibility to potentiate the impact of intrathecal fentanyl by additional administration of ketamine and to reduce the amount of fentanyl used by a co-application of ketamine [10]. In our study, we were not able to increase the antinociceptive effect of fentanyl by adding ketamine when both drugs were given by a non-spinal way. This combination has also been studied by several other research groups with mainly contradictory results [41-43]. The reason for these confounding results remains unclear. In our experiments, we were not able to further reduce the VMR to CRD by co-application of ketamine. This discordance of findings might be a result of the different models. Hirota et al. postulated a competitive interaction of ketamine with the µ- and κ-OR as functional µ- and κ-OR-antagonist that causes a non-opioidergic analgesic mechanism [44].
Limitations

Variability of the data can at least partially be explained by the difference in behaviour of the animals even before the beginning of the measurements. Some were calm, while others were already agitated resulting in higher baseline electromyographic activity and EMG measurements. Some animals did not survive the entire protocol. Thus, the total number of animals in each test series differed between 11 and 15. Another potential weak point of the present study is the lack of dose-response curves for the different substances which were tested. The respective concentrations for each test substance were determined by literature search. The aim of a follow-up study must be to confirm the conclusions of our study by including dose-response curves. In this context, both the absolute and relative expression of opiate receptor subtypes in the colon should be quantified in colitis and control animals. Finally, all substances were given in the same order in every animal, therefore no randomisation was performed. Thus, an adaption of the test animals to the CRD stimulus can not be excluded.

The irritative bowel syndrome is a very common disease. However, even in the year 2019, it cannot be treated satisfyingly. Post-infectious hyperalgesia may play a crucial role in the development of IBS at least in a subpopulation of IBS patients. The pathways of how to counteract this altered post-infectious visceral nociception have not been fully characterised yet. In earlier studies, we were able to demonstrate in Lewis rats that a chemically induced inflammation of the colon mucosa causes long-lasting visceral hyperalgesia in response to tonic visceral stimuli [45-47]. In our present series of tests, we were able to confirm these findings in an experimental set-up with phasic distensions. We also demonstrated that, at the applied doses, the κ-opioid receptor agonist U50.488 has the most potent analgesic effect on viscero-sensory function to CRD in Lewis rats following induction of a chemical colitis compared to the μ-ORA fentanyl and morphine, the δ-ORA SN80 and the NMDA antagonist ketamine. Furthermore, we could show that the μ-ORA morphine also possesses a positive effect on the visceral hyperalgesia in the TNB-colitis rats. In our study, the application of ketamine alone or a combination of ketamine and the ORA's missed to reduce or further reduce the VMR compared to the application of opioids alone. To conclude, our experiments demonstrate a strong analgesic effect of the selective κ-ORA U50.488 on long-term nociception in response to colorectal stimulation in a chemically induced colitis model. This is particularly noteworthy given the fact that this opiate receptor agonist unfolds its analgesic effect without the unwanted side effects of other opiate subtypes.

Acknowledgements

The statistical analysis was performed in close collaboration with the Institute for Mathematical Statistics of the University of Berne, Switzerland. This Study was supported by a grant-in-aid of the University of Berne and by unrestricted educational grants of the Novartis Stiftung für medizinisch-biologische Forschung, AstraZeneca, Nycomed and Solvay (all grants of JMG).
17. Suarez-Roca H, Jose Antonio Silva, Jose Luis Arcaya, Luis Quintero, William Maixner, et al. (2006) Role of mu-opioid and NMDA receptor activations in the maintenance and development of repeated swim-stress-induced thermal hyperalgesia. Behav Brain Res 167(2): 205-211.

18. Gallantine EL, TF Meert (2005) A comparison of the antinociceptive and adverse effects of the mu-opioid agonist morphine and the delta-opioid agonist SNC80. Basic Clin Pharmacol Toxicol 97(1): 39-51.

19. Hoffmann P, J Zeeh, J Lakshmanan, V Wu, F Procaccino, et al. (1997) Increased expression of transforming growth factor alpha precursors in acute experimental colitis in rats. Gut 41(2): 195-202.

20. Ness TJ, GF Gebhart (1988) Colorectal distension as a noxious visceral stimulus: physiologic and pharmacologic characterization of pseudoaffective reflexes in the rat. Brain Res 450(1-2): 153-169.

21. Zhou Q, Donald D Price, Robert M Caudle, G Nicholas Verne (2008) Visceral and somatic hypersensitivity in a subset of rats following TNBS-induced colitis. Pain 134(1-2): 9-15.

22. Sengupta JN (1999) Effects of kappa opioids in the inflamed rat colon. Pain 79(2-3): 175-185.

23. Junien JL, P Riviere (1995) Review article: the hypersensitive gut-peripheral kappa agonists as a new pharmacological approach. Aliment Pharmacol Ther 9(2): 117-126.

24. Danzebrink RM, SA Green, GF Gebhart (1995) Spinal mu and delta, but not kappa, opioid-receptor agonists stimulate responses to noxious colorectal distension in the rat. Pain 63(1): 39-47.

25. Larsson M (2003) A model for chronic quantitative studies of colorectal sensitivity using balloon distension in conscious mice: effects of opioid receptor agonists. Neurogastroenterol Motil 15(4): 371-381.

26. Kitamura TM Ogawa, Y Yamada (2009) The kappa-opioid agonist (−)-bremazocine elicits peripheral antinociception by activation of the l-arginine/nitric oxide/cyclic GMP pathway. Life Sci 78(1): 54-60.

27. Hustveit, OA Mauser, I Oye (1995) Interaction of the chiral forms of ketamine with phencyclidine, sigma and muscarinic receptors. Pharmacol Toxicol 77(6): 355-359.

28. Mody I, JF MacDonald (1995) NMDA receptor-dependent excitotoxicity: the role of intracellular Ca2+ release. Trends Pharmacol Sci 16(10): 356-359.

29. Kolhekar R, GF Gebhart (1996) Modulation of spinal visceral nociceptive transmission by NMDA receptor activation in the rat. J Neurophysiol 75(6): 2344-2353.

30. Carstensen M, AM Moller (2010) Adding ketamine to morphine for intravenous patient-controlled analgesia for acute postoperative pain: a qualitative review of randomized trials. Br J Anaesth 104(4): 401-406.

31. Malec D, M Mandryk, S Fidecka (2008) Interaction of memantine and ketamine in morphine- and pentazocine-induced antinociception in mice. PharmacolRep 60(2): 149-155.

32. Sarton E (2001) The involvement of the mu-opioid receptor in ketamine-induced respiratory depression and antinociception. Anesth Analg 93(6): 1495-1500.

33. Hirota K (1999) Stereoselective interaction of ketamine with recombinant mu, kappa, and delta opioid receptors expressed in Chinese hamster ovary cells. Anesthesiology 90(1): 174-182.

34. Adam R, Tobias Liebregts, Juergen M Gschossmann, Constanze Krippner, Franziska Scholl, et al. (2006) Severity of mucosal inflammation as a predictor for alterations of visceral sensory function in a rat model. Pain 123(1-2): 179-186.

35. Gschossmann JM (2002) Effect of transient chemically induced colitis on the visceromotor response to mechanical colorectal distension. Eur J Gastroenterol Hepatol 14(10): 1067-1072.

36. Gschossmann JM, T Liebregts, I Buenger, M Ruwe, G Gerken, G Holtmann, et al. (2004) Long-term effects of transient chemically induced colitis on the visceromotor response to mechanical colorectal distension. Dig Dis Sci 49(1): 96-101.