Antinociceptive local effect of the combination of dexketoprofen trometamol and chlorhexidine gluconate in a formalin test: an additive effect

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INTRODUCTION

Pain is a complex response, caused by different elements that evokes the physiological, sensory and emotional pain process (Szallasi, 2010). Generally, is the consequence of local injury that affects the anatomical, biochemical and sensory equilibrium. When the damage is exposed to bacteria contamination, a dual etiology may trigger different pathways of the local inflammation (Arnstein, 2013).

Painful conditions can be controlled by the local application of analgesics, such as non-steroidal anti-inflammatory drugs (NSAIDs), which inhibit cyclooxygenases (COXs) activity both central and peripherally (Derry et al., 2017). Dexketoprofen trometamol (DXT) is the water soluble S(+) enantiomer of the NSAID ketoprofen (Mauleón et al., 1996). Previous studies demonstrated the local effect of ketoprofen (Derry et al., 2015) and DXT (Isiordia-Espinoza et al., 2014; Sagir et al., 2013) and other studies showed analgesic efficacy of DXT in different post-operative conditions (Esparza-Villalpando et al., 2016; Rodríguez, Arbós, Amaro, 2008). The local administration of NSAIDs is desirable for elective interventions where the clinician can access directly to the lesion site. Theoretically, if the molecules are administered locally obtaining acceptable analgesia, then systemic intake can be reduced dramatically, thus decreasing the incidence of side-effects as well (Arnstein, 2013; Derry et al., 2017). In the surgical context, the procedure itself is susceptible to complications such as surgical site infection (SSI), especially in areas were microbial flora is abundant. This complication occurs in up to 30% of all surgical procedures (Bruce et al., 2001).

The use of skin antisepsics decrease the colonization of bacteria and therefore reduces the events of SSI (Lee
et al., 2013). Chlorhexidine gluconate (CHX) is a topic antiseptic with antibacterial and antifungal clinical efficacy in different conditions, it is effective against gram-positive and gram-negative bacteria, some viruses and presents good tolerability and safety profile for the patient (Edmiston et al., 2010).

Animal pain models, can be used to test antinociceptive effect of drug combinations by evaluating the previous standardize response of rodents. The formalin test is a pain model used to study inflammatory pain by injecting noxious chemical irritant (formalin) into the paw of rodents (Szallasi, 2010; Krzyzanowska, Avendaño, 2012), allowing to determine not only the individual response of analgesic molecules; but the effect of certain combinations (showing either possible antagonism, synergism or additive effect of analgesic molecules under defined conditions). A drug delivery system (DDS) that allows analgesic and antibacterial properties would be desirable for the control of post-operative pain, especially when the bacterial etiology is related to painful response (such as surgical procedures or post-surgical infections). Thus, the aim of the present study was to evaluate the antinociceptive effect of the combination of DXT and CHX in formalin pain model.

MATERIAL AND METHODS

Animals

Thirty six female Wistar rats (6 per group) of 6-7 weeks of age (200-250 g), were used in the experiment. The animals were obtained from the Animal Center of the Guanajuato University and were housed on a 12 h light-dark cycle at 24± 2 °C, with free access to food and water. On the day of the experiment, animals were acclimatized to the laboratory conditions for at least 2 hours before the test. All animals were used once and were sacrificed immediately after the test. The study was carried out according to the Official Mexican Norm NOM-062-ZOO-1999 (Technical specifications for the production, care, and use of laboratory animals) and the Guidelines on Ethical Standards for Investigation of Experimental Pain in Animals (Zimmermann, 1983). Animals were used once during the protocol and then sacrificed in a CO₂ chamber at the end of each experiment. DXT (Stein Labs®, San José, Costa Rica) was dissolved at the moment of the experiment in des-ionized water, obtaining a concentration of 12 mg/mL. CHX (Sigma Aldrich®, EUA), 20% water solution, was dissolved in des-ionized water in 3 different concentrations (0.02%, 0.3% and 2.5% respectively).

Measurement of nociceptive response

Antinociception was assessed by the formalin test. The rats were placed in transparent plastic cylindrical chambers with mirrors placed in a 45° angle to assure complete viewing of the injected paw. The rats were injected into the dorsal surface of the hind paw with 50 µL of dilute formalin (5%) alone or mixed with the appropriated concentrations of formulations using a 30-gauge needle. The rats were observed for nociceptive behavior immediately after formalin injection. All measurements were made by an independent blinded evaluator. Nociceptive behavior was quantified as the number of flinches of the injected paws in one minute (min) periods, every 5 min up to 60 min after injection. Time-courses of nociceptive response for each group were constructed as mean number of flinches in each time. For the two phases of formalin test, the areas under the curve (AUC) were calculated by trapezoidal method (Allison et al., 1995): the first phase included the measures of the minutes 0-15; and the second phase the minutes 15-60. Group-response curves for each phase were constructed using six animals for each group. The percentage of antinociception (%AN) was calculated as follow:

\[
%AN = \frac{AUC_{Formalin} - AUC_{Treatment group}}{AUC_{Formalin}} \times 100
\]

Experimental design

The animals were injected with a total solution of 50 µL per experiment, containing fixed volumes that kept the following concentrations: formalin 5%, DXT 12 mg/mL and CHX (with different concentrations as mentioned before). Proportions v/v are shown in the Table I. All solutions were prepared freshly the day of the test and were put together directly in the same syringe right before its application.

Statistical analysis

The antinociceptive responses are presented as Mean±Standard Error (S.E.M.), the AUC/%AN difference between experimental groups was assessed by ANOVA with planned contrast: a) G1 vs G2, G3, G4, G5, G6; contrast b) G2 vs G3, G4, G5, G6; contrast c) G3 vs G4, G5, G6; contrast d) G4 vs G5, G6 and contrast e) G5 vs G6 and the assumption of normality was assessed by Shapiro-Wilks test. The same contrasts were performed to both phase 1 and phase 2. All analysis were performed by statistical software R ver. 3.4.0.
RESULTS

The test for normality (Shapiro-Wilks) showed data distribution not different to normal (P > 0.05). The time course of the nociceptive response is shown on Figure 1. The formalin test showed two well defined phases, the phase 1 (1-11 minutes) and the phase 2 (21-61 minutes) representing tonic acute pain and inflammatory pain respectively.

The %AN and the p-values of different comparisons are shown on the Figures 2 and 3. Control group showed the lower value of %AN in both phases, the group DXT only (G2), and the other groups not showed statistical difference. Additionally, the group of CHX only (G3), show similar antinociceptive effect to DXT only (G2). The group of the combination between DXT+CHX 0.02% (G4) showed lower antinociceptive effect than groups G5 and G6, but only on the phase 1. The group of CHX only (G3) showed lower antinociceptive effect than the combination groups (G4, G5 and G6), but only on the phase 2.

DISCUSSION

This work confirmed the antinociceptive effect of DXT when is locally administered in the formalin test, using the 50% effective dose of DXT previously reported by Isiordia et al. (2014). The local use of NSAID’S can provide acceptable pain relief, commonly reported in acute conditions. Ketoprofen is one of the molecules with this effect (Derry et al., 2015), supporting the logical election of DXT for this investigation. Moreover, this work focuses on the antinociceptive effect of the combination between DXT and CHX in the formalin model of inflammatory pain. The concentrations used in this study were selected to:

<table>
<thead>
<tr>
<th>Groups</th>
<th>µL of Formalin</th>
<th>µL of DXT†</th>
<th>µL of CHX‡</th>
<th>% of CHX</th>
</tr>
</thead>
<tbody>
<tr>
<td>G1 (Control)</td>
<td>50</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>G2 (DXT only)</td>
<td>25</td>
<td>25</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>G3 (CHX 0.3% only)</td>
<td>25</td>
<td>-</td>
<td>25</td>
<td>0.3</td>
</tr>
<tr>
<td>G4 (DXT + CHX 0.02%)</td>
<td>25</td>
<td>12.5</td>
<td>12.5</td>
<td>0.02</td>
</tr>
<tr>
<td>G5 (DXT + CHX 0.3%)</td>
<td>25</td>
<td>12.5</td>
<td>12.5</td>
<td>0.3</td>
</tr>
<tr>
<td>G6 (DXT + CHX 2.5%)</td>
<td>25</td>
<td>12.5</td>
<td>12.5</td>
<td>2.5</td>
</tr>
</tbody>
</table>

† Dexketoprofen trometamol, ‡ Chlorhexidine

FIGURE 1 - Time course of the nociceptive behavioral response of the experimental groups (n=6 rats each group). Each symbol represents the mean with S.E.M., over the 60 min post-injection observation period.
based on previously reported minimal inhibitory concentration (MIC) for CHX (Edmiston et al., 2010; Hope, Wilson, 2004), since they are considered non-cytotoxic (Faria et al., 2007; Faria et al., 2009; Lee et al., 2010; Li et al., 2009). As mentioned before, beyond the classic employments of formalin test, this research tool also can discard possible antagonistic reactions between the components of new pharmacological combinations; in

FIGURE 2 - Antinociceptive effects (%AN) of control and experimental groups in the first phase of formalin test. Data are present in Mean±S.E.M. of n=6 rats for each group. The letters represent the planned contrast ANOVA: a) G1 vs G2, G3, G4, G5, G6; b) G2 vs G3, G4, G5, G6; c) G3 vs G4, G5, G6; d) G4 vs G5, G6 and e) G5 vs G6.

FIGURE 3 - Antinociceptive effects (%AN) of control and experimental groups in the second phase of formalin test. Data are present in Mean±S.E.M. of n=6 rats for each group. The letters represent the planned contrast ANOVA: a) G1 vs G2, G3, G4, G5, G6; b) G2 vs G3, G4, G5, G6; c) G3 vs G4, G5, G6; d) G4 vs G5, G6 and e) G5 vs G6.
our case, it was selected as the departure point to propose an analgesic/antiseptic DDS.

The behavioral response showed the classical two phases induced by the formalin test (Gonzalez et al., 2011; Miranda et al., 2011; Krzyzanowska, Avendaño, 2012). Phase 1 results from a direct stimulation of nociceptors and phase 2 involves a period of sensitization during the inflammatory process through peripheral mechanisms (Le Bars, Gozariu, Cadden, 2001). The expected result was that the combination between both did not showed a decreased antinociceptive effect, conversely, the combination showed an increased antinociceptive effect than the drug alone, but these differences were not significative.

To our knowledge, an antinociceptive effect related to local application of CHX has not yet been reported. Haraji and Rakhshan showed in a clinical trial an analgesic effect of the CHX gel in post-operative pain condition. As the authors hypothesized, the antiseptic feature of the CHX is possibly related to the reduction in painful inflammation mediators caused by bacteria (Haraji, Rakhshan, 2015). However, these results were inconclusive and more studies are needed to confirm these findings. Even when such conclusion may be indirectly related to the present results, they cannot be correlated since the nociceptive response of the formalin test is related to a noxious chemical stimulus and no to bacterial presence. In this context, the possible mechanism involved in the antinociceptive or analgesic response of the CHX should be analyze in deeper specialized models, including the analysis of specific pain pathways, expression of potential mediators or even the agonism/antagonism of certain receptors. Due to its chemical composition and ionic charge, CHX molecules may modulate the nervous conduction responsible of pain transmission, or even may contribute by blocking specific peripheral ionic channels. The bactericidal effect of CHX is a result of the binding of the CHX cationic molecules to negatively charged bacterial cell walls and extramicrobial complexes (McDonnell, Russell, 1999). A possible reason of the antinociceptive effect observed may involve the positively charged CHX interacting with the membranes of the peripheral nerve endings, thus creating the modulation of the action potential evoked by the formalin. However, any of these hypotheses must be meticulously analyzed in new experiments.

The surgical procedures involve wounds that will heal by primary or secondary intention. In the procedures involving “clean” cavities, the surgical infection prevalence is around 3% to 5%; but, when the procedures involve infected sites, dirty or necrotic tissue; the surgical infections increase up to 10% to 30% (White, 2009). The oral cavity is considered clean/contaminated and the risk for surgical infection is latent. The wound infection results from the dynamic interactions between the presence of pathogens and the susceptibility of the host. The pathways of the surgical infection are complex; however, the most common signs are erythema, pain, local temperature and swelling (Gardner, Frantz, Doebbeling, 2001).

The presence of pain is part of the local inflammatory reaction, and thus, it will be expected to decrease at the same rhythm that the etiology is controlled and the inflammation is auto-limited. To increase the quality of life of the patients, analgesic/antiseptic DDS will play a dual role, by modulating the normal painful response of inflammation while favoring the local septic ethology. When pain is poorly controlled, a pain-related infection would reduce the immune response to infection. Therefore, the control of pain is as important as the treatment of infection itself (White, 2009). The use of CHX could provide an extra analgesic effect in combination with DXT and at the same time covering the antimicrobial role to prevent SSI.

Since our results not only demonstrated that the presence of CHX didn’t affect the local antinociceptive effect of DXT (confirmed in our results), but the molecule itself showed to have antinociceptive capacity, and the presence of an additive effect between both molecules; a new favorable result must be concluded. Further efforts will be addressed not only to fully understand this hypothesis and the mechanisms involved, but to include new experiments analyzing decreasing combinatory doses in order to perform an isobolographic analysis (Tallarida, 2001), looking for possible pharmacological synergism.

CONFLICT OF INTEREST

All authors declare no competing interests.

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