Carbamide peroxide gel stability under different temperature conditions: is manipulated formulation an option?

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Nowadays the use of gel containing carbamide peroxide (CP) prepared in Pharmacy is a normal practice in the population. However, the quality of this product is questionable concerning its stability. The aim of this study is was to synthesize and to analyze this drug alone or associated to Carbopol gel through analytical methodology compatible with the routine of the Pharmacies. The reaction between urea and hydrogen peroxide was carried out at different resting times: 24 hours (CP 24 powder) and 48 hours (CP48 powder) after the mixture. Both products were associated with Carbopol 940® gel 1.5% (G) generating G24 and G48 samples. The stability of powders (CP24 e CP48) and the formulations (G24 and G48) were evaluated as a function of time (15, 40 and 45 days) and thermal variation (refrigeration: 8 °C±1; thermal shock 32 ±1 /8 °C±1; stove: 32 ±1), using a standard titration method. As a result, only under refrigeration the CP24 and CP48 contents remained stable during the period of 45 days. An interesting finding was that G24 and G48 presented greater stability for at least 45-days under refrigeration and thermal shock conditions, and up to 30 days under stove conditions. The results for the G24 and G48 were slightly higher than those obtained for the control. Therefore, we were able to conclude that association with Carbopol 940® Gel 1.5 % provided greater CP stability and that manipulated formulations containing CP may be viable for use in a period of 45 days under refrigeration conditions. The titration proved to be an effective technique for the analysis of CP with or without Carbopol 940® gel 1.5%.

Uniterms: Carbamide peroxide/quantitative analysis. Carbamide peroxide/stability. Dental bleaching. Carbopol 940® gel 1.5%.

Atualmente, a utilização de gel contendo peróxido de carbamida manipulado em Farmácia é uma prática comum na população. No entanto, a qualidade deste produto é questionada, sobretudo no que se refere à estabilidade deste fármaco. O objetivo deste trabalho consiste na avaliação da viabilidade de sintetizar e analisar quantitativamente este fármaco associado ou não a um gel de Carbopol através de metodologia analítica compatível com a rotina das Farmácias. A reação entre a uréia e o peróxido de hidrogênio foi realizada em tempos diferentes de repouso após a mistura, 24 h para sintetizar o pó PC 24 e 48 h para o pó PC 48. Estes pós foram associados a um gel (G) de Carbopol 940® 1,5 %, originando as amostras G24 e G48. A estabilidade dos pós (PC 24 e PC 48) e das formulações (G24 e G48) foi avaliada em função do tempo (15, 40 e 45 dias) e da variação térmica (refrigeração: 8 °C±1; choque térmico: 32 °C±1/8 °C±1 e estufa: 32 °C±1), através da técnica de titulometria. Os resultados indicam que unicamente sob refrigeração o CP24 e o CP 48 mantiveram-se estáveis no período de 45 dias. O G24 e o G48 apresentaram estáveis por pelo menos 45 dias nas condições de refrigeração e choque térmico e por 30 dias na condição estufa. Os resultados obtidos para o G24 e G48 foram ligeiramente superiores aos obtidos para o controle. Além disso, é possível concluir que a associação do PC com o gel de Carbopol 940® 1,5 % promoveu um aumento na estabilidade do PC e que as preparações manipuladas contendo PC são viáveis para uso durante um período de 45 sob refrigeração. A titulometria mostrou-se uma técnica eficaz para a análise do PC associado ou não ao gel de Carbopol 940® 1,5 %.


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INTRODUCTION

Recently, dental bleaching procedures became more popular as a result of the increasing awareness of the importance of having healthy white teeth (Gordon, Chirrienssen, 2005). Dental darkening may be caused by extrinsic factors (e.g. smoking, food or beverages with pigments), or intrinsic factors either congenital (e.g. imperfect dentinogenesis) or acquired (e.g. use of tetracyclines, fluorine or trauma) (Goldstein, 2000). Nowadays, two bleaching techniques are commonly used. The home-bleaching technique implies on using a low concentration of CP gel dental strips for an average period of 2 to 3 weeks, and is becoming increasingly popular. On the other hand, the office technique, although older, was not frequently used until recently due to the difficulties associated with the hydrogen peroxide solution application and the need for absolute isolation (Papathanasiou et al., 2002; Wetter et al., 2009).

The carbamide peroxide (CP), also known as urea peroxide, hydrogen peroxide-urea or urea-peridrol, was initially used as an anti-inflammatory in the great wars to treat periodontal diseases (Gonçalvez et al., 2001; Gratão et al., 2003). Starting in 1960, it has been used in the form of antiseptic solutions (10 % CP) for the treatment of gingivitis in children and adults by the orthodontist William Klusmier, in which the tooth whitening occurred as a side effect of the anti-septic property. After this event, in 1989, Haywood and Heymann published the first article about tooth whitening with CP at concentrations ranging from 10 % to 20 % (Gonçalvez et al., 2001; Compoy et al., 2001; Goldstein, 2000).

The action mechanism of bleaching consists in the oxidation reaction of the CP bleaching agent, which in contact with the saliva or gingival tissue is broken down into hydrogen peroxide and urea. The urea produces ammonia and carbon dioxide, contributing to the maintenance of an alkaline pH, which potentiates the action of the bleaching agent (Sun, 2000). The hydrogen peroxide diffuses through the organic matrix of enamel and dentin, releasing free radicals that break O₂ carbon rings of high molecular weight from darker pigmentation, converting them into clearer smaller molecules (Compoy et al., 2001; Rodrigues et al., 2005; Goldberg et al., 2009). Nevertheless, for bleaching gel to act on enamel and dentin, it is necessary for it to diffuse through the dental tissues. It is known that the greater the surface tension of a liquid, the lower its ability to diffuse through a surface or to wet it. After the bleaching treatment changes have been observed in surface texture (Potocnik et al., 2000) mineral content (Attin et al., 2005), chemical composition (Justino, Demarco, 2004) and loss by tooth brushing abrasion (Wiegand, Attin, 2004). The addition of surfactants to hydrogen-peroxide-based bleaching gel resulted in a significantly more extensive bleaching (Canepelle, Torres, 2011).

The other application for CP is in the treatment for plaque, gingival health and caries, so antibacterial activity has been studied (Nunes et al., 2011). Plaque accumulation and resulting caries or periodontal disease is a frequent problem in patients with special-care needs. Lazarchik and Haywoode reviewed in 2010 the antibacterial properties of CP and the effects of CP on saliva, plaque, caries and gingival health. They addressed the challenges involved in the research needed regarding the use of tray-applied CP materials in special-care patients. The results showed that CP at 10 % may be a great promise for improving the oral health of many special-care patients, including elderly patients, patients with cancer and patients with dry mouth.

The technique described by Haywood and Heymann was improved when the CP was associated with the Carbopol® gel at 1.5 %. Carbopol is a thickening agent, of thixotropic nature, used in the process to stabilize the dissociation of the bleaching agent, releasing oxygen more slowly, increasing the time of action of the product formulation and adhesion to tooth structure (Compoy et al., 2001; Apple, Reus, 2003).

The bleaching agents, however, are chemically very unstable (Baratier et al., 2004). The commitment of their stability is related to factors such as temperature, light, moisture, pH, impurities and others (Ansel et al., 2000). Thus, stability tests to predict monitoring and to determine the validity and the ideal storage conditions are needed.

The products used for dental bleaching could be prepared by the industry and by the pharmacist. The manipulated products have questionable quality, because of the low stability of CP. Therefore, the studies about applicable analytical methodologies in pharmacy practice contribute to the quality certification of the products.

The aim of this work was to analyze the synthesis of CP, through the reaction of hydrogen peroxide and urea for 24 and 48 hours and subsequent percentage evaluation of CP in the synthesized product. Moreover, the work was to study the incorporation of active Carbopol 940® gel at 1.5 % and stability evaluation of both the CP and the CP gel as a function of time and temperature variation, with the aim to establish a protocol for monitoring its adaptive stability reality of pharmacies.

METODOLOGY

The CP was synthesized using urea (All Chemistry) and hydrogen peroxide 130 V (All Chemistry) in equal
proportions. Urea was crushed in porcelain mortar and then hydrogen peroxide 130 V was added. The mixture was divided into two portions, covered with plastic wrap and kept at rest for 24 h and 48 h respectively, in the dark, resulting in the samples CP24 and CP48. After this period, both portions were vacuum filtered and the solids were stored in desiccators for 24 hours (USP 27 2004; Zanardi et al., 2007).

For the Carbopol 940® gel at 1.5 % containing CP preparation, 16 % of CP24 was incorporated, taking into account its level, in 60 % Carbopol 940 gel at 1.5 %. It was also added 0.3 % of sodium fluoride as a desensitizing agent. The same procedure was repeated for CP48, resulting in samples G24 and G48, respectively. The controls used for this experiment consisted of CP in the dust from Merck (Dust Control) and the gel of CP 16 % of Whiteness Perfect® (FGM Control).

Samples CP24, CP48, G24 and G48 and controls were stored in bottles, milky plastic and syringes respectively sealed, and then exposed to three different temperature conditions for the experiment: a) Thermal Shock: 32 °C±1/8 °C±1 °C; b) Stove: 32 °C±1 °C and c) Refrigeration: 8±1 °C.

The CP determination in the samples and controls, stored under different conditions, was done by quantitative iodometric (titrimetric), taking as a basis the methodology described in the United States Pharmacopeia (USP 27, 2004). It was done in triplicate at predetermined dates, divided into 4 different times totaling 45 days. Then, 100.00 mg of CP were weighed, and this amount was transferred to an Erlenmeyer flask of 500 mL. For the gel, 625.00 mg were weighed on a CP 500 mL flask, which corresponds to 100.00 mg of CP. Later, distilled water, acetic acid PA, potassium iodide and sodium molybdate 10% (aqueous solution) were added. The solution was homogenized and then maintained under light for 10 min. The liberated iodine was titrated with aqueous 0.05 M sodium thiosulfate TS. Near the turning point, 3.00 mL of 2% starch indicator were added, changing the color from dark blue to colorless.

Considering that each mL of 0.05M sodium thiosulfate TS is equivalent to 4.704 mg of CP, the concentration percentage was calculated for the samples and controls. The average return and standard deviation of the results were calculated. The effects of different storage conditions (8 °C ± 1; 32 °C / 8° C ± 1; 32 °C ± 1), time (15, 30 and 45 days), formulation (G24, G48 and Control) and their interactions were assessed using factorial design analysis for three factors by ANOVA followed by Tukey post-test (α = 0.05). It was used SPSS (Statistical Package for the Social Sciences) 17.0 for Windows.

RESULTS AND DISCUSSION

Table I shows the results of the CP content in the samples and controls, obtained by titrimetric assay at different temperature conditions in which they were exposed for a period of 45 days.

The concentration of CP in the samples CP24, CP48 and control at time zero (after preparation), showed an average concentration of 98.31%, 99.01% and 97.85%, respectively (Table I); values were considered within pharmacopoeial limits of 96% to 102 (USP 27, 2004), with no significant statistically difference between them (Tukey test, p<0.05). Threshold values of concentration of CP gel incorporated into Carbopol 940® were not found in the literature.

<table>
<thead>
<tr>
<th>Samples</th>
<th>CP Content (%) ± S.D</th>
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<tbody>
<tr>
<td></td>
<td>8 °C±1</td>
</tr>
<tr>
<td></td>
<td>Days</td>
</tr>
<tr>
<td></td>
<td>15</td>
</tr>
<tr>
<td>CP24</td>
<td>98.31 ± 0.44</td>
</tr>
<tr>
<td>CP48</td>
<td>99.01 ± 0.88</td>
</tr>
<tr>
<td>Merck® Control</td>
<td>97.85 ± 0.49</td>
</tr>
<tr>
<td>(powder)</td>
<td>97.85</td>
</tr>
<tr>
<td>G24</td>
<td>100.39 ± 0.64</td>
</tr>
<tr>
<td>G48</td>
<td>100.16 ± 0.58</td>
</tr>
<tr>
<td>FGM Control</td>
<td>99.75 ± 0.55</td>
</tr>
<tr>
<td>(gel)</td>
<td>99.26</td>
</tr>
</tbody>
</table>
The results demonstrated in Table I showed that the samples CP24 and CP48 in the refrigeration condition (8 °C ± 1) had no statistical difference compared to their control at time zero, 15 and 30 days (tukey test, p< 0.05). However, in 45 days the CP48 and control showed a significant CP decrease of 3.7 % and 2.6 % respectively (Table I). This decrease is within the acceptable limit variation of 5 % in stability studies recommended in RE nº1.5 (Brazil, 2005).

Regarding the heat shock condition (32 °C ± 1 / 8 °C ± 1), the CP24 control remained stable during the first 15 days, differing from CP48. After 45 days of exposure, the CP24 and CP48 samples showed a decrease in the CP levels of 64 %. Only the control remained stable with a reduction of only 5.2 %.

In stove conditions (32 °C ± 1), CP24 and CP48 samples showed a substantial decrease in the CP level that started at 15 days of exposure, resetting the CP content over time. This may be related to the decomposition of CP in ammonia (urea), oxygen and water (hydrogen peroxide) at temperatures above 30 °C (Gratac et al., 2003; Basting, 2006; Leonard et al., 1994). The dust control in this condition showed a slight CP decrease after 30 days, but remained constant at the end of 45 days. This ambiguity of behavior for the CP24 and CP48 in comparison to controls, tested under different conditions can be attributed to different experimental conditions in the methods of obtaining these substances. The resolution RE nº01/05 on pharmaceutical active establishes that stability depends on temperature, humidity, light and other factors relating to the product itself (physical and chemical properties). In addition, one must consider other aspects that can interfere with stability of assets such as: presence or absence of excipients, the dosage form type, manufacturing process and material packaging (Brazil, 2005).

The samples containing the CP associated with the Carbopol 940® gel 1.5 % (G24 and G48), when subjected to cooling showed statistical difference in CP content, when compared to control.

With regard to exposure of the same samples to heat shock (32 °C ± 1 / 8 °C ± 1), it was found that all samples were stable during the first 15 days. After 30 days, the G24 had a decreased percentage of CP which was not observed for G48 and control. In 45 days the G24 was the formulation that showed the most reduced levels of CP; however, this reduction was not significant, indicating stability of these formulations at the end of 45 days. The control gel at the end of 45 days showed a content change, demonstrating the instability of this vehicle in the tested condition. The G48 formulation was stable for 45 days on the studied condition temperature.

The display results under stove (32 °C ± 1) showed that the increase in temperature led to a decreased CP percentage after 30 days for all samples; however, this reduction was more significant for the control FGM (8.40%). After 45 days, there was a reduction of 6.39 % for G24, 10.27 % for G48 and 10.46 % for the control. According to Aulton (2005) such behavior is expected due to the drastic conditions of storage. Increasing the temperature at 10 °C produces a 2-5 fold increase in the drugs degradation.

Figure 1 shows the statistical analysis of the gel containing CP in the different experimental conditions (time, storage conditions and formulation). Factor analysis applied significant difference p (probability values) <0.01 for the main factors by the ANOVA for three factors. The comparison of means by Tukey test (α = 0.05) has shown that the storage of CP gel at low temperatures (8 ± 1 °C) is more appropriate, and therefore recommended it as the condition for storage of the Carbopol 940®gel 1.5 % containing CP. It is possible to verify that the temperature exerts a strong influence on the CP decay associated to Carbopol 940®gel 1.5 %. Higher temperatures cause oscillation of the decay of this drug. This influence can be observed mainly for the CP powder. It is possible to show that the drug contents decrease as a function of time, since the observed differences are significant fortnightly. This observation is particularly useful for estimating an expiration date for the compounded product, through the results one can estimate a validity of at least 45 days for the product stored in the refrigerator.

A comparison of results between the G24, G48 and FGM control in different conditions, demonstrated that the G24 e G48 showed higher CP content than the control. This result is positive in demonstrating that manipulated formulations were similar to industrialized formulations, setting the feasibility and safety in the preparation of this formulation in masterful scale. Considering the results for the G24 and G48, one can realize that the time of synthesis of CP is not directly influenced by the quality of the final product, since there was no significant difference in CP content for these samples.

Considering these results, it was possible to show a greater reduction in CP level in powder than in Carbopol 940® gel 1.5 %. Studies that evaluate the stability of the drug in isolated solid state and in the presence of excipients are important during the stage of pre-formulation of a product. Studies of stability of the formulation as a whole prioritize the evaluation of the drugs behavior over time considering the drug, the mixture of excipients or used vehicles, as well as the interaction between them, in different conditions (Brazil, 2005; Mamede et al., 2006). The present results have highlighted the importance that the
dosage form provides to the system as a whole, protecting the active ingredient from degradation caused by the agents tested (Baratieri et al., 2004).

CONCLUSION

In relation to the CP synthesis, one can say that it was feasible to carry it through because it is an easy and affordable methodology. Based on the temperature conditions described and in accordance with the valuation method (titration), the results showed that the best form of storage for the CP dust is under refrigeration (8 °C ± 1), since the results indicate no significant decrease in CP levels when exposed to this condition. The association of CP in the Carbopol 940® gel 1.5 % provided greater CP stability, showing the importance of the dosage form in the protection of the active substance, having no significant difference between samples prepared with synthesized CP at different times.

This paper presented relevance with regard to its applicability in monitoring the CP content in powder and Carbopol 940 gel 1.5 % to be easily adapted into a pharmacy routine. Thus, it can be ensured that the manipulated formulations in the Carbopol 940 gel at 1.5 % containing CP are viable for use in a period of 45 days when stored under refrigeration.

REFERENCES


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