Abstract

Purpose: To evaluate the stability of an extemporaneously prepared 7% chloral hydrate syrup under different conditions of storage and dispensing.

Methods: Three batches of 7% chloral hydrate syrup were prepared. Each batch was stored in 50 light-resistant glass containers of 60 mL with child-resistant caps and in two bottles of 1000 mL to simulate two forms of dispensing, mono and multi-dose, respectively. Twenty five mono-dose bottles and a multi-dose bottle of each batch were stored under room conditions (20 ± 1°C) and the rest of the samples were stored in the fridge (5 ± 2°C). The physical, chemical and microbiological stability was evaluated for 180 days. Stability was defined as retention of at least 95% of the initial concentration of chloral hydrate, the absence of both visible particulate matter, or color and/or odor changes and the compliance with microbiological attributes of non-sterile pharmaceutical products.

Results: At least 98% of the initial chloral hydrate concentration remained throughout the 180-day study period. There were no detectable changes in color, odor, specific gravity and pH and no visible microbial growth. These results were not affected by storage, room or refrigeration conditions or by the frequent opening or closing of the multi-dose containers.

Conclusions: Extemporaneously compounded 7% chloral hydrate syrup was stable for at least 180 days when stored in mono or multi-dose light-resistant glass containers at room temperature and under refrigeration.

PALABRAS CLAVE
Jarabe de hidrato de cloral; Sedación; Compulación; Pediatría; Estabilidad

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Introduction

The chloral hydrate is a sedative drug widely used in pediatrics, especially to administer to children before diagnostic procedures such as computerized axial tomography scan and magnetic resonance imaging that requires patient immobility. It is also used in intensive care units, pediatric emergency departments and dental surgery to decrease stress and anxiety caused by technological procedures, light and sound levels and the presence of strangers. It is a cheap, effective and safe drug that can be easily handled and administered when performing nasofibroscopy. Chloral hydrate is quickly absorbed and metabolized into the liver, erythrocytes and kidneys to trichloroethanol (active metabolite) and to trichloroacetic acid (inactive metabolite). The binding to the proteins of the active metabolite is 70-80% and its half-life is from four to twelve hours. The rapidity with which it induces sleep (30 to 60 min) is attributed to the active principle, while the long-lasting action (4 to 8 hours) is due to its active metabolite. A sedative dose (50-100 mg/Kg) has minimal effects on breathing and blood pressure and the reflexes are slightly suppressed. The patient can, therefore, be fully awake. Chloral hydrate is generally well tolerated when a single dose is given. It has a low toxicity as long as the recommended dose is not exceeded and it is administered for a short period of time. Chloral hydrate as sleep inducer for electroencephalogram recordings in children of one to five years old of age is more effective than midazolam.

It achieves a significantly deeper level of sedation compared with other sedative agents and children remain calmer when undergoing echocardiography.

Chloral hydrate occurs as transparent colorless crystals, with a penetrating slightly acrid odor and a slightly bitter caustic taste. It is highly soluble in water. Because of its unpleasant taste and its irritant action on the gastric mucosa, oral administration of dilute solutions is advisable. From the chemical point of view chloral hydrate is a gemdiol, near infrared studies, in aqueous solution, showed the existence of a labile equilibrium between the gemdiol and a dimolecular 1:1 complex of the aldehyde and water (eq. 1). Besides, it has been proposed that, in slightly acid or neutral aqueous solutions, chloral hydrate decomposes by an oxidation-reduction process forming dichloroacetaldehyde, trichloroacetic acid and hydrochloric acid (eq. 2). Exposure to light and heat can speed up this degradation process.

On the other hand, it has been described that chloral hydrate decomposes in diluted aqueous solution of sodium hydroxide (eq. 3) by first-order kinetics catalyzed by hydroxyl ions.

At present, there is no commercially available formulation of chloral hydrate, thus it is considered an orphan drug. Because of this shortage, it is prepared as an extemporaneous formulation in hospital pharmacies. The 10% oral solution of chloral hydrate is codified by USP 30 Ed., whereas the 7% chloral hydrate syrup is codified by the Argentine Pharmacopoeia 6th Ed. It is known that, for its texture and palatability, properly flavored syrups effectively mask taste and they are greatly preferred by pediatric patients. In our Institution, the 7% chloral hydrate syrup is prepared and stored in 100, 250, 500 and 1000 mL of multi-dose light-resistant glass containers. The main production demand of the syrup has increased significantly from 6.4 L/year during 2009 and 2010 to 20 L/year during 2011, which forced us to prepare larger batches to be stored in the pharmacy until dispensed. This situation points out the need for stability studies which allow establishing adequate storage conditions and an expiration date based on stability studies. Notice that the frequent opening of the multi-dose bottles during the utilization period could increase the concentration of sugar and of chloral hydrate due to water evaporation. Besides, aqueous solution of 10% chloral hydrate produces a pH of 3.5-5.5 which can lead to hydrolysis (inversion) of sucrose, with the subsequent loss of the syrup consistency and the possibility of microorganism fermentation.

Few stability studies of liquid formulations of chloral hydrate have been carried out. However, both stability studies on 7% chloral hydrate syrup nor multidose containers considering the possible opening and closing of the containers were found, and they were not designed to assign an expiration date or to define optimal storage conditions. In a previous study it has been proposed that the decrease in the pH of the formulations of chloral hy-
drate can be used as an indicator of the degradation of the active principle.

In this context, the objective of this work is to evaluate the physical, chemical and microbiological stability of 7% chloral hydrate syrup both under room and refrigeration conditions in mono and multi-dose bottles.

**Materials and Methods**

Chloral hydrate \((C_3H_5Cl_2O_2, USP\) grade, Parafarm®, Bs. As., Arg.) was assayed by means of titration, according to FA 8 Ed\(^{16}\). The determinations were performed in quintuplicate which showed a 100.6 ± 0.4% concentration of \(C_3H_5Cl_2O_2\). Refined sucrose (Ledesma®, Salta, Arg.), Alcohol 96° (Porta®, Córdoba, Arg.), NaOH 1.0 N and 0.1 N solutions (Anedra®, Bs.As., Arg.) and distilled water were also used.

**Preparation of 7.0% chloral hydrate syrup**

Three batches of 5.0 L of 7% chloral hydrate syrup, flavored with bitter orange fluid extract 30% (Parafarm®, Bs. As., Arg.) were prepared in accordance with the procedure described in Appendix to observe reproducibility and robustness of the gathering and control methods. Each batch was stored in 50 light-resistant glass containers of 60 mL with child-resistant caps and in two bottles of 1000 mL, with the same characteristics, to simulate two dispensing forms, mono- and multi-dose, respectively.

**Stability evaluation**

Twenty five mono-dose bottles and a multi-dose bottle of each batch were stored under room conditions in an air-conditioned room and the rest of the samples were stored in the fridge. The temperature of both conditions was daily monitored using a digital thermo-hygrometer, registering values of 20 ± 2°C y 5 ± 2°C for room and refrigeration conditions, respectively. All the samples were labeled and stored for 180 days. For physical and chemical stability, one mono-dose container and an aliquot of 35 mL from the multi-dose containers of each chloral hydrate syrup batch, at the mentioned temperature conditions, were collected on days 0, 7, 15, 30, 45, 60, 75, 90 and 180.

- **Physical Stability Evaluation.** At each time point, physical stability was assessed by visual examination and by determination of specific gravity, according to USP <841> method 1 using a 25-mL calibrated pycnometer\(^{12}\). Physical stability was defined as the compliance of the specific gravity according to the USP specifications (± 1.30 g/mL)\(^{12}\) and the absence of either visible particulate matter, or color and/or odor changes.

- **Chemical Stability Evaluation.** Chemical stability was assessed following the concentrations of chloral hydrate and HCl in the samples. Chloral hydrate concentrations were determined by titrimetry with 1.0 N sodium hydroxide adapting USP30 Chloral Hydrate 10% Oral Solution assay to 7% syrup. In addition the time required to produce the reaction between chloral hydrate and sodium hydroxide was determined to be 10 min in the syrup instead the 2 min described for the 10% oral solution. Besides, the syrup samples were measured by weight instead of volume to avoid errors due to its high viscosity. Briefly, approximately 27.7 g of 7% chloral hydrate syrup were weighed in an appropriate glass-conical flask. Then, 25.0 mL of sodium hydroxide were added and mixed for 10 min. Finally, 4 drops of 1% phenolphthalein alcoholic solution were added. The excess of sodium hydroxide, regarded as \(A\), was immediately titrated with 1.0 N sulfuric acid (Anedra®, Bs.As., Arg.). For the second titration, 5.55 g of the syrup were weighed in a 100 mL conical flask and titrated with 0.1 N sodium hydroxide solution after adding 10 drops of 1% phenolphthalein alcoholic solution. This solution was regarded as \(B\). The weight in mg of chloral hydrate in the amount of 7% chloral hydrate syrup taken for the first titration was calculated by the formula:

\[
165.4 (A - 0.5 B) \quad \text{eq. 4}
\]

A coefficient variation of 0.47% for this method was obtained from 10 replicated titrations.

For pH determinations, the samples were thermostatted at 25°C in a water bath. The pH values were recorded with an ADWA AD 8000 pH-meter, with an Ag/AgCl-reference electrode, calibrated at the same temperature with commercial reference buffer solutions of pH 4.01 and 7.00 (HACH®, USA). The stability-indicating capability of the titration and pH determination methods was also evaluated in this work. An aliquot of 1.0 M HCl was added to three 100 mL samples of 7% chloral hydrate syrup to get HCl concentrations equivalent to 1 and 5% of chloral hydrate degradation, according to equation 2. The resulting pHs were 2.50 ± 0.09 and 2.05 ± 0.09, for HCl concentrations equivalent to 1 and 5% of chloral hydrate degradation, respectively. The chemical stability of the formulation was defined as not less than 95% of the initial drug concentration remaining in the samples and a pH value not less than 2.05. Results are expressed as the mean of three batch determinations for each storage condition.

- **Microbiological stability evaluation.** Microbiological assessment of chloral hydrate syrup, only on the
samples stored under the most unfavorable conditions (multi-dose containers at room temperature), was carried out on days 0, 30, 60, 90 and 180. The samples were subjected to microbiological evaluation in order to determine if they meet the microbiological attributes of non-sterile pharmaceutical products which is set as total aerobic microbial count below $10^2$ cfu/mL, total combined yeasts/molds count below 2 cfu/mL and absence of *Pseudomonas aeruginosa*, *Staphylococcus aureus*, *Escherichia coli* and *Salmonella spp*. Limit content of microorganisms was performed according to USP<1111> and ANMAT N° 7667 disposition18.

Results and discussion

Table 1 shows some relevant properties of the three batches of 7% chloral hydrate syrup. All the test samples were practically colorless and free of visible particles. Table 2 shows the chemical and physical stability descriptors after six months of assay. In addition, no particles were observed in the bulk solution or in the container closures. Besides, little or no chloral hydrate loss occurred in any of the samples at any storage temperature throughout the study and there was no significant change in pH since it remained within the range of 2.97 to 3.09 which suggests that there was no chemical degradation. Specific gravity after 6 months of storage under refrigeration and room temperature complies with USP specifications. Accordingly, no increase in concentration of chloral hydrate due to vehicle evaporation was observed in the multi-dose containers even after 6 months of storage, and in spite of having been opened 9 times for sampling during the stability experiments.

Microbiological tests were negative in the three batches which prove that the product complies with official specifications after 6 months of storage. All these aspects support the conclusion that, for at least 180 days, 7% chloral hydrate in a sucrose-based, orange flavored syrup, in light-resistant glass containers of 1000 and 60 mL can be stored for at least 180 days. The three batches under study at 20°C and 5°C showed excellent physical stability as well as practically no loss in chloral hydrate concentration. Levels of HCl as degradation product were well below the acceptable limits. Therefore, hospital pharmacies, can set an expiration date of 6 months under any of the conditions studied.

Conclusions

This study provides conclusive information about the period and the storage conditions in which chloral hydrate syrup can be utilized as a sedative in pediatrics. A 7% chloral hydrate in a sucrose-based, orange flavored syrup, in light-resistant glass containers of 1000 and 60 mL can be stored for at least 180 days. The three batches under study at 20°C and 5°C showed excellent physical stability as well as practically no loss in chloral hydrate concentration. Levels of HCl as degradation product were well below the acceptable limits. Therefore, hospital pharmacies, can set an expiration date of 6 months under any of the conditions studied.

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References

Table 2. Chemical and physical stability descriptor

<table>
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<th>Container type</th>
<th>Storage temperature (°C)</th>
<th>Parameters</th>
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<tbody>
<tr>
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<td></td>
<td></td>
<td>7</td>
</tr>
<tr>
<td>5</td>
<td>Chloral hydrate (% ± SD)a</td>
<td>99.7 ± 0.1</td>
<td>98.9 ± 0.2</td>
</tr>
<tr>
<td></td>
<td>pH ± SD</td>
<td>3.03 ± 0.04</td>
<td>3.04 ± 0.02</td>
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<tr>
<td></td>
<td>Specific gravity (mg/mL ± SD)</td>
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<td>1.325 ± 0.001</td>
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<tr>
<td>20</td>
<td>Chloral hydrate (% ± SD)a</td>
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<td>98.9 ± 0.4</td>
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<td>pH ± SD</td>
<td>3.02 ± 0.06</td>
<td>3.03 ± 0.02</td>
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<tr>
<td></td>
<td>Specific gravity (mg/mL ± SD)</td>
<td>1.325 ± 0.001</td>
<td>1.325 ± 0.001</td>
</tr>
<tr>
<td>5</td>
<td>Chloral hydrate (% ± SD)a</td>
<td>99.7 ± 0.6</td>
<td>99.1 ± 0.7</td>
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<td>pH ± SD</td>
<td>3.02 ± 0.05</td>
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<tr>
<td></td>
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<td>1.325 ± 0.001</td>
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</table>

*Mean values from the three batches. aExpressed as% of initial chloral hydrate concentration.

SD: standard deviation.

Appendix. Procedure for compounding 1,000 mL of 7.0% chloral hydrate syrup

1. Dissolve 70.0 g of chloral hydrate in 25.0 mL of distilled water under heating at 80-90°C, in a 1,000 mL glass conical-flask.
2. Add slowly 900 mL of simple syrup to this solution, under constant stirring to ensure complete mixing.
3. Mix 30 mL of bitter orange fluid extract with 58 mL of 96° alcohol and distilled water to complete 100 mL. Add 10 drops of this solution to the syrup and bring to a final volume of 1000 mL with simple syrup under vigorous mixing.
4. Transfer the solution to light-resistant glass containers of adequate volume.
5. Label the bottles with an expiration date of 180 days after preparation, and store either at room temperature or under refrigeration.
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