Hydrogen-Ion Potential of Antibiotics According to the Environment Factors Temperature and Luminosity

Renata Maria Coelho Crepaldi
Cíntia Monteiro
Maria Angélica Sorgini Peterlini
Mavilde da Luz Gonçalves Pedreira

The objective of this experimental study was to measure the pH of antibiotics administered by intravenous infusion - ceftiraxone sodium, vancomycin hydrochloride, metronidazole, penicillin G potassium and amikacin sulfate - after reconstitution with sterile water and dilution with NaCl 0.9% or dextrose 5% in water, according to temperature and luminosity of the environment. The results showed that variation in the drugs’ pH was less than 1.0 value and that some antibiotics remained acidic after dilution and maintained this chemical profile in all situations studied, suggesting that the studied environmental factors did not change the solutions’ acid base characteristic. Some pH values measured characterize risk for the development of chemical phlebitis and infiltration, and it is important for clinical practice to emphasize the profile of intravenous solutions of antibiotics, considering method of dilution, and time to infusion.

Descriptors: Pediatric Nursing; Infusions Intravenous; Hydrogen-Ion Concentration; Drug Stability; Safety.

1 Supported by CNPq Process #476295/2004-1 and #502382/2007-4.
2 RN. E-mail: recrepaldi@hotmail.com.
3 Nursing Undergraduate Student, Departamento de Enfermagem, Universidade Federal de São Paulo, SP, Brazil. Scholarship holder, Basic Scientific Program CNPq. E-mail: cintiamonteiroms@hotmail.com.
4 Ph.D. in Nursing, Adjunct Professor, Departamento de Enfermagem, Universidade Federal de São Paulo, SP, Brazil. E-mail: maria.angelica@unifesp.br.
5 Ph.D. in Nursing, Adjunct Professor, Departamento de Enfermagem, Universidade Federal de São Paulo, SP, Brazil. E-mail: mpedreira@unifesp.br.
Potencial hidrogeniônico de antimicrobianos, segundo os fatores ambientais temperatura e luminosidade

O objetivo deste estudo experimental foi medir o pH dos antibióticos de administração intravenosa ceftriaxona sódica, cloridrato de vancomicina, metronidazol, penicilina G potássica e sulfato de amicacina, após reconstituição com água destilada e diluição com NaCl 0,9%, ou suro glicosado 5%, considerando a influência da temperatura e luminosidade ambientais, assim como do tempo de exposição, no comportamento químico desses fármacos. Os resultados demonstraram variações que não ultrapassaram 1,0 valor de pH e que alguns antimicrobianos, eminentemente ácidos após a diluição, mantiveram esse comportamento em todas as situações estudadas, não sugerindo a influência de fatores ambientais no comportamento químico das soluções. Como alguns valores de pH encontrados podem contribuir para o desenvolvimento de flebite química e infiltração, é importante enfatizar para a prática clínica em saúde, a necessidade de conhecer as características das soluções de infusão intravenosa, considerando tipo de diluição e tempo de infusão.

Descritores: Enfermagem Pediátrica; Infusões Intravenosas; Concentração de Íons de Hidrogênio; Estabilidade de Medicamentos; Segurança.

Introduction

Intravenous (IV) medication administration is highly relevant in clinical practice for the treatment of patients with acute or chronic conditions and for the prophylaxis of some diseases. Among evidence of the large-scale use of this treatment in the pediatric population, a study carried out with children hospitalized at pediatric units of a Brazilian hospital found the daily administration of 4.4 doses\(^1\).

Antibiotics stand out among the most used drugs in children. Due to the lack of a pharmaceutical form for the pediatric age range, IV administration of these drugs requires dilution in small volume and prolonged infusion time, at speeds of up to tens of milliliters per hour, so as to avoid hypervolemia, heart failure and toxicity\(^2\). Inquiries arise, however, about maintaining these drugs’ stability in hospital units’ environmental conditions.
The stability of a drug is a critical element for appropriate medication administration, as both the appropriateness and safety of therapy can be affected in case of pharmacological instability\(^{(3)}\). A drug is considered stable in a solution if at least 90\% of its concentration does not suffer decomposition\(^{(4)}\). The term pharmacological instability is applied to continuing and irreversible chemical reactions that result in different substances or products of degradation, which can be both therapeutically inactive and capable of causing toxic effects. Among different degradation routes, drugs can suffer hydrolysis, oxidation and reduction\(^{(3)}\).

The stability of drugs in solutions can be affected by many factors, such as environmental light, temperature, and the solution’s final concentration. Many drugs’ degradation is catalyzed by pH extremes, with reactions being affected by the presence of hydrogen or hydroxide ions. Drugs’ reaction rates are generally lower in intermediary than in high or low pH values. This evidences the importance of maintaining a constant hydrogen-ion concentration in the solution. The consequence of an altered pH can be both decreased chemical stability or precipitation of one or more drugs. Moreover, pharmacological instability can lead to alterations in hydrogen-ion concentration\(^{(4)}\).

A solution’s pH represents the concentration level of this solution’s hydrogen ions. By convention, the pH measurement unit equals the negative logarithm of the hydrogen-ion concentration in a solution. The concentration of this ion in liquid water at 25\(^{\circ}\)C is 10\(^{-7}\)mol/L. As hydrogen ions are neutralized by hydroxide ions at this concentration, the neutral medium has a pH equal to seven, the acid medium, between zero and seven and the base medium, between seven and fourteen. Each unit decrease in pH represents a ten-time addition in the concentration of hydrogen ions\(^{(5-6)}\).

This research aims to provide tools for safe IV therapy practice in pediatrics, considering the lack of studies on the pH of antibiotics solution, submitted to environmental conditions verified in clinical practice. Thus, the goal is to assess the pH of five antibiotics for IV administration, considering the influence of external factors like environmental light and temperature during the time recommended for the IV infusion of these solutions.

**Purpose**

To measure the pH of five antibiotics after their reconstitution and dilution in NaCl 0.9\% and dextrose 5\% in water, considering the influence of temperature, environmental light and exposure time to these factors.

**Material and method**

This experimental study was carried out after approval by the internal ethical review board at the university the authors are affiliated with. Data were collected at a laboratory between April and October 2007. The study hypothesis was that environmental temperature and light, as well as, exposure time to these factors, provoke alterations in the pH behavior of solutions with antimicrobial agents for IV administration.

The sample comprised the pH of five antimicrobials used in pediatric hospital units of the university where the study was carried out\(^{(1)}\): ceftriaxone sodium, vancomycin hydrochloride, metrodinazole, penicillin G potassium and amikacin sulfate.

The following environmental conditions were considered: \(\alpha\) – environmental temperature of 22\(^{\circ}\)C and non-exposure to light (control); \(\beta\) – environmental temperature of 22\(^{\circ}\)C and exposure to light; \(\gamma\) – environmental temperature of 30\(^{\circ}\)C and non-exposure to light and \(\delta\) – environmental temperature of 30\(^{\circ}\)C and exposure to light.

For each situation, eight moments were chosen for pH measurement: immediately after reconstitution in sterile water or liquid presentation; 24h after reconstitution in refrigerated conditions; \(t_0\) – immediately after dilution in sodium chloride 0.9\% or dextrose 5\%; \(t_1\) – five minutes after dilution; \(t_2\) – 30 minutes after dilution; \(t_3\) – 60 minutes after dilution; \(t_4\) – 120 minutes after dilution and \(t_5\) – 24h after dilution.

The drugs’ reconstitution and dilution was performed according to the institution’s nursing protocol. After one hour of exposure to the environmental condition under analysis, the powder drugs were reconstituted with sterile water, resulting in concentrations of 100mg/mL for ceftriaxone sodium and vancomycin hydrochloride and 500 000U/mL for penicillin G potassium. After the reconstitution, a sample was taken from the vial to verify the pH. Another sample was used to dilute the antibiotics. The remainder was kept in the vial for refrigeration and pH was measured again after 24h of refrigeration.

As metrodinazole and amikacin sulfate are distributed in liquid form, pH values were obtained from the commercial solution at concentrations of 5mg/mL and 250mg/mL, respectively.

All drugs were diluted in NaCl 0.9\% and dextrose 5\% (SG 5\%), at different times. The final concentration...
of each drug was 5mg/mL for amikacin sulfate, metrodinazole and vancomycin hydrochloride, 20mg/mL for ceftriaxone sodium and 50 000U/mL for penicillin G potassium. The final volume of the solution was set at 20mL for all drugs. The only non-diluted drug was metrodinazole, as its pharmaceutical form has the ideal concentration for IV administration i.e., according to the manufacturer, 5mg/mL.

After the drugs had been diluted and exposed to environmental situations, a sample of the solution was removed from the syringe and stored in a glass Becker to verify pH values at intervals from t0 to t5.

A digital ExStik™ PH 100 meter manufactured by Extech® (USA) was used. This is a millivoltmeter linked to an electrode selective for hydrogen ions, with a scale that converts the voltage of the electrode into pH units. The accuracy of the meter covers an interval up to ±0.01 on the pH scale. Calibration was performed according to the manufacturer’s instructions.

Environmental light was obtained through white light fluorescent lamps. Two digital thermometers, model 7429, brand TFA® Dostmann (Reicholzhein, Baden-Württemberg, Germany) were used for thermal control of the laboratory environment.

Table 1 – Hydrogen-ion potential (pH), mean and standard deviation of antimicrobial agents’ pH immediately after reconstitution with sterile water and 24-hour storage in refrigerator, according to temperature and environmental light. São Paulo, 2007

<table>
<thead>
<tr>
<th>Antimicrobial</th>
<th>Condition</th>
<th>Immediately after reconstitution</th>
<th>24 hours</th>
<th>Mean</th>
<th>Standard deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ceftriaxone sodium</td>
<td>α</td>
<td>6.96</td>
<td>6.95</td>
<td>6.96</td>
<td>0.007</td>
</tr>
<tr>
<td></td>
<td>β</td>
<td>7.05</td>
<td>6.89</td>
<td>6.97</td>
<td>0.113</td>
</tr>
<tr>
<td></td>
<td>γ</td>
<td>6.48</td>
<td>6.41</td>
<td>6.45</td>
<td>0.049</td>
</tr>
<tr>
<td></td>
<td>θ</td>
<td>6.44</td>
<td>6.49</td>
<td>6.47</td>
<td>0.035</td>
</tr>
<tr>
<td>Vancomycin hydrochloride</td>
<td>α</td>
<td>2.91</td>
<td>2.93</td>
<td>2.92</td>
<td>0.014</td>
</tr>
<tr>
<td></td>
<td>β</td>
<td>2.92</td>
<td>2.95</td>
<td>2.94</td>
<td>0.021</td>
</tr>
<tr>
<td></td>
<td>γ</td>
<td>2.72</td>
<td>2.72</td>
<td>2.72</td>
<td>0.028</td>
</tr>
<tr>
<td></td>
<td>θ</td>
<td>2.96</td>
<td>2.92</td>
<td>2.94</td>
<td>0.028</td>
</tr>
<tr>
<td>Penicillin G potassium</td>
<td>α</td>
<td>6.76</td>
<td>6.63</td>
<td>6.70</td>
<td>0.092</td>
</tr>
<tr>
<td></td>
<td>β</td>
<td>6.76</td>
<td>6.51</td>
<td>6.64</td>
<td>0.177</td>
</tr>
<tr>
<td></td>
<td>γ</td>
<td>6.58</td>
<td>6.55</td>
<td>6.57</td>
<td>0.021</td>
</tr>
<tr>
<td></td>
<td>θ</td>
<td>6.68</td>
<td>6.66</td>
<td>6.67</td>
<td>0.014</td>
</tr>
</tbody>
</table>

A small variation in the pH of all drugs diluted in NaCl 0.9% was observed, with greater modifications after 24 hours than at other times. Increases in pH values were observed for all antibiotics, except for amikacin sulfate at 30ºC. Vancomycin hydrochloride and amikacin sulfate showed the most acid pH values: 3.10 and 4.94, respectively. The least acid values came from ceftriaxone sodium and crystalline penicillin, with 6.88 and 6.17, respectively (Figure 1).
Using dextrose 5% for dilution, modifications are observed in some antibiotics, mainly at 24 hours of exposure. At that moment, the pH values of ceftriaxone sodium and vancomycin hydrochloride increased in all conditions and that of amikacin sulfate in conditions without exposure to light. For penicillin G potassium, pH values went down in all situations. Like when diluted in NaCl 0.9%, vancomycin hydrochloride and amikacin sulfate showed the most acid pH values: 3.11 and 5.01, respectively. The highest values were found for ceftriaxone sodium and crystalline penicillin: 7.01 and 6.74, respectively (Picture 2).

It should be observed that pH values of penicillin G potassium increase after 24 hours when diluted in NaCl 0.9% in all situations. When diluted in SG5%, however, it becomes more acid after 24 hours. The pH value of vancomycin hydrochloride remained quite acid, even after dilution.

Small variations in pH values of metrodinazole occurred over time, with considerable pH after 24 hours of exposure in all environmental conditions (Picture 3).
Crepaldi RMC, Monteiro C, Peterlini MAS, Pedreira MLG.

Figure 2 – Hydrogen-ion potential (pH) of antimicrobial agents diluted in dextrose 5%, according to environmental condition and exposure time. São Paulo, 2007

Figure 3 – Hydrogen-ion potential (pH) of metronidazole, according to environmental condition and exposure time. São Paulo, 2007

The individual analysis of each drug diluted in physiological saline solution shows the mean and standard deviation of the antibiotics in the four situations as follows: ceftriaxone sodium (6.20±0.311), vancomycin hydrochloride (3.21±0.080), penicillin G potassium (5.87±0.185) and amikacin sulfate (5.01±0.030). After dilution in SG5%, the following values are observed: ceftriaxone sodium (6.56±0.212), vancomycin hydrochloride (3.33±0.087), penicillin G potassium (6.60±0.158) and amikacin sulfate (5.08±0.064). And, finally, for metrodinazole, the mean value was 5.50±0.350. Moreover, data demonstrate that, when diluted in SG5%, all antimicrobials showed higher mean pH values than what was found for the other thinner (Table 2).
It should be highlighted that, although analogue pH values were to be expected for the drugs in commercial form, variations in pH values were observed, even when the drugs came from the same manufacturer (Pictures 1 and 3).

Table 2 – Mean and standard deviation in pH of antimicrobial agents at six moments after dilution, according to environmental condition and type of dilution. São Paulo, 2007

<table>
<thead>
<tr>
<th>Antimicrobial</th>
<th>Condition</th>
<th>pH NaCl 0.9%</th>
<th>pH Dextrose 5%</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Mean</td>
<td>Standard deviation</td>
</tr>
<tr>
<td>Ceftriaxone sodium</td>
<td>α</td>
<td>6.32</td>
<td>0.276</td>
</tr>
<tr>
<td></td>
<td>β</td>
<td>6.13</td>
<td>0.351</td>
</tr>
<tr>
<td></td>
<td>γ</td>
<td>6.22</td>
<td>0.315</td>
</tr>
<tr>
<td></td>
<td>θ</td>
<td>6.13</td>
<td>0.302</td>
</tr>
<tr>
<td>Vancomycin hydrochloride</td>
<td>α</td>
<td>3.27</td>
<td>0.071</td>
</tr>
<tr>
<td></td>
<td>β</td>
<td>3.19</td>
<td>0.129</td>
</tr>
<tr>
<td></td>
<td>γ</td>
<td>3.18</td>
<td>0.047</td>
</tr>
<tr>
<td></td>
<td>θ</td>
<td>3.21</td>
<td>0.073</td>
</tr>
<tr>
<td>Penicillin G potassium</td>
<td>α</td>
<td>5.92</td>
<td>0.160</td>
</tr>
<tr>
<td></td>
<td>β</td>
<td>5.93</td>
<td>0.137</td>
</tr>
<tr>
<td></td>
<td>γ</td>
<td>5.83</td>
<td>0.233</td>
</tr>
<tr>
<td></td>
<td>θ</td>
<td>5.80</td>
<td>0.211</td>
</tr>
<tr>
<td>Amikacin sulfate</td>
<td>α</td>
<td>5.01</td>
<td>0.034</td>
</tr>
<tr>
<td></td>
<td>β</td>
<td>5.02</td>
<td>0.029</td>
</tr>
<tr>
<td></td>
<td>γ</td>
<td>5.01</td>
<td>0.040</td>
</tr>
<tr>
<td></td>
<td>θ</td>
<td>5.01</td>
<td>0.017</td>
</tr>
</tbody>
</table>

Discussion

Both phlebitis and infiltration are commonly identified complications in peripheral intravenous therapy. Phlebitis can be classified as chemical, mechanical or infectious. The cause of chemical phlebitis can derive, among other factors, from extreme pH values, leading to an inflammatory response in the inner layer of the vein that can lead to infiltration, edema, thrombosis and cell death(6).

A randomized clinical trial involving children hospitalized at a Brazilian institution showed that 55.3% of peripheral venous catheters were removed due to infiltration and 4.7% due to phlebitis(9).

The range of pH values that entails risks for IV therapy complications is below 5.5 and above 8.0(10). In this research, vancomycin hydrochloride and amikacin sulfate presented pH values below 5.5 after dilution. Metrodinazole also showed risky pH values, but only after 24 hours of exposure.

In nursing practice, professionals are often unaware of drugs’ pH, exposing patients to complications by administering more concentrated solutions in less time than recommended, due to the need for fluid restriction in some cases. Besides causing changes in the IV administration site, extreme pH values can catalyze many drugs’ degradation. This also derives from other factors, including temperature, environmental light and concentration of the solution. Drug instability can also lead to pH value alterations(3).

Light activates photodegradation reactions, such as oxidation and hydrolysis. Hence, the stronger a light source and/or the closer the photosensitive drug is placed to that source, the higher the degradation level will be(3). During IV administration of metrodinazole, protection against light is no source of concern in the hospital environment(11). In this research, metrodinazole showed low variation in the concentration of hydrogen ions when exposed to light.

An experimental research revealed that the pH of penicillin G potassium at 5000U/mL presented minimal variations after adding saline (mean 6.04) and dextrose (mean 5.60), solutions at 4, 25 and 37°C(12). When diluted in saline solution, the mean pH value of penicillin G potassium was close to literature values (5.87). When diluted in dextrose, however, the value was higher (6.60).

In general, increases in environmental temperature influence solutions’ degradation rate, causing pH
modifications and, consequently, pharmacological instability. Literature describes penicillin G potassium and cephalosporins as labile to heat\(^{(3)}\). This was observed for the penicillin, with a higher standard deviation in situations at 30°C.

Although ceftriaxone sodium showed low lability to high temperature, important increase in pH occurred after 24 hours of exposure in all situations and in both thinners. Another experiment evidenced that this antimicrobial at 100mg/mL in sterile water remained stable for 40 days at 4°C\(^{(13)}\). The analysis of pH variations in the same concentration of ceftriaxone sodium, after reconstitution and storage in a refrigerator for 24 hours, showed hardly considerable changes, revealing no reactions that would modify pH values.

Due to the lack of a pharmaceutical form for the pediatric age range, the reconstituted solutions are stored in refrigerated conditions and concentrated drugs are diluted in different volumes and solutions for IV infusion. Questions arise, however, about refrigeration time until the next administration and stability after dilution and IV administration in uncontrolled environmental situations\(^{(1)}\).

At some institutions, the reconstituted drugs are stored in refrigerators that sometimes do not maintain the recommended temperature between 4°C and 8°C\(^{(14)}\). These drugs are removed from the refrigerator different times, exposed to environmental temperature and returned to the refrigerator without knowledge about their stability. Besides ceftriaxone sodium, vancomycin hydrochloride and penicillin G potassium were also stored for 24 hours, with small variations in their pH values.

In this experiment, the antibiotics were diluted in recommended solutions, i.e. NaCl 0.9% and SG5%, with pH values of 5.5 and between 3.5 and 6.5, respectively\(^{(10)}\). Antimicrobials in general are unstable in alkaline (pH above 8) or acid (pH below 4) solutions\(^{(15)}\). Therefore, the used thinners do not represent risk for instability, but should be stored at temperatures below 25°C without freezing\(^{(11)}\).

Thus, the pH of each component in a mixture should be previously verified in order to avoid therapeutic harm and other complications\(^{(3)}\). When reconstituted, vancomycin hydrochloride showed pH values below 4, representing a risk factor for pharmacological instability for the solution stored until the next administration.

For practical purposes, it is important to plan IV therapy, to know drugs’ physical-chemical characteristics, such as the hydrogen potential, and to determine the dilution method and infusion time, guaranteeing safe IV administration without complications.

**Conclusion**

The environmental situations that simulate clinical practice provoked changes in the antimicrobials’ pH values of less than 1.0. Amikacin sulfate and vancomycin hydrochloride stand out as risky drugs for the development of local complications of intravenous therapy. In clinical practice, it is important to know the characteristics of IV infusion solutions, considering dilution type and infusion time. Therefore, it is fundamental for clinical health practice to continue this type of research, using other parameters to analyze the pharmacological stability of intravenous solutions, such as high performance liquid chromatography.

**Acknowledgements**

We thank the nurses Ariane Ferreira Machado and Lidiane Lopes Reis, doctoral and master’s students at the Nursing Graduate Program, Universidade Federal de São Paulo, for their collaboration.

**References**


