Gentamicin Dosing and Monitoring Challenges in End-Stage Renal Disease

Florczykowski B* and Storer A*
1 LSU Health- Shreveport, 1725 Claiborne Ave, Shreveport, LA 71130, USA
2 ULM College of Pharmacy, 1725 Claiborne Ave, Shreveport, LA 71130, USA

Abstract

The second leading cause of mortality in end-stage renal disease patients on intermittent hemodialysis is infection. Gentamicin is frequently used in this patient population due to the convenience of dosing only on hemodialysis days. Dosing recommendations vary among the most commonly used drug information databases. Differences are found in recommendations for both dosing and monitoring. Conflicting information found in drug information databases and primary literature can cause confusion among clinicians regarding the best way to dose and monitor gentamicin in ESRD patients on hemodialysis. While current practice utilizes post-hemodialysis gentamicin dosing, there is primary literature that introduces the concept of pre-dialysis dosing. Pre-dialysis gentamicin dosing may result in favorable pharmacokinetic profile.

Keywords: End-stage renal disease; Hemodialysis; Gentamicin

Introduction

The second leading cause of mortality in end-stage renal disease (ESRD) patients on intermittent hemodialysis is infection, preceded only by cardiovascular causes. The presence of indwelling catheters and repeated exposure to foreign materials during hemodialysis increase the risk of infection. Gentamicin is frequently used to treat infections in ESRD patients because of the convenient dosing profile in this patient population. The prolonged half-life in patients with ESRD allows for less frequent dosing, yet adequate therapeutic serum levels for extended time intervals. This pharmacokinetic profile allows for administration of intravenous gentamicin on scheduled outpatient hemodialysis days negating the need for an inpatient admission to administer IV antibiotics.

Dosing

Although gentamicin treatment is common in ESRD patients, determining the correct dose can be a challenge. Commonly used drug information databases have conflicting dosing recommendations for patients on hemodialysis (Table 1). Of the dosing references evaluated, Lexi-Comp recommends the most aggressive dosing strategies, including a loading dose as well as a higher maintenance dose. Micromedex has multiple dosing recommendations. One mimics the recommendation from the package insert and the other is a ½ reduction of the dose used for a patient with normal renal function. The most comprehensive set of recommendations is found in Facts and Comparisons. It provides three different dosing strategies which are a culmination of what is found in each of the other databases, but proposes a ⅔ reduction as opposed to the ½ suggested by Micromedex. Of note, the ⅔ reduction in Micromedex and ½ reduction in Facts and Comparisons reference different editions of the same book [1-3]. The Micromedex recommendation comes from a publication from 1994, whereas the Facts and Comparisons recommendation comes from a publication from 2007.

To further complicate gentamicin dosing, there is also conflicting data regarding which weight is used to determine the initial dose (Table 2). Some sources suggest using actual body weight while others suggest using ideal body weight. All of the references agree on using an estimated lean body weight formula to arrive at a dose for obese patients. However, only one database recommends using a dosing weight formula for underweight patients.

Timing of Dose

Since aminoglycosides are dialyzable, current practice is to infuse gentamicin doses at the completion of hemodialysis (Table 1). However, many hemodialysis units administer the infusion during the last 30 minutes of hemodialysis. With this practice, some of the gentamicin infused will be cleared during hemodialysis preventing the patient from achieving optimal peak levels and altering intended pharmacokinetics. Utilizing the higher end of the dosing range may be warranted in patients who receive gentamicin while still on the dialysis machine.

A novel method, pre-dialysis gentamicin dosing [4-18] has been described as being similar to once-daily dosing in patients with normal renal function [12]. This dosing strategy allows for higher gentamicin doses and serum concentrations. As with all aminoglycosides, higher drug concentrations lead to a greater rate and extent of bactericidal activity [5,6]. Because gentamicin clearance during dialysis approaches that of normal kidney function [8], a large portion of the pre-dialysis dose gets cleared and patients are not exposed to increased serum levels for prolonged durations. Although these studies show promising results, the pharmacokinetic profiles were obtained from computer extrapolations of small patient populations and have not been established in current practice. Larger prospective studies may be necessary prior to routinely adopting this dosing strategy for hemodialysis patients [19].

Monitoring

It is widely accepted that serum levels should be monitored closely due to the narrow therapeutic window of gentamicin. However, serum monitoring suggestions differ among drug information databases (Table 3). Ideal peak levels range between 4-6 mcg/mL for a urinary
A peak level should be drawn 30 minutes after completing the gentamicin infusion to allow for drug distribution. Peaks drawn prior to drug distribution will result in falsely elevated serum levels. Patients should remain in the hemodialysis unit for 30 minutes after the gentamicin infusion finishes to obtain an accurate peak level. Although 8-hour HD may reduce serum concentrations by approximately 50%, it is necessary to correctly monitor peak gentamicin levels and maximize efficacy.

<table>
<thead>
<tr>
<th>Database</th>
<th>Weight Used for Dosing in Obese</th>
<th>Weight Used for Dosing in Underweight Patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Package Insert</td>
<td>ABW × 1.13 x IBW</td>
<td>No recommendation</td>
</tr>
<tr>
<td>Lexi-Comp</td>
<td>IBW x DWo</td>
<td>No recommendation</td>
</tr>
<tr>
<td>Micromedex</td>
<td>ABW x DWo</td>
<td>DWu</td>
</tr>
<tr>
<td>Facts and Comparisons</td>
<td>ABW x DWo</td>
<td>No recommendation</td>
</tr>
</tbody>
</table>

\(^{1}\)Obese = >20% of IBW
ABW = Actual Body Weight, IBW = Ideal Body Weight, DWo = dosing weight for obese patients, DWu = dosing weight for underweight patients, 
DWo = IBW + 0.4(ABW-IBW), DWu = 1.13 x ABW

A peak level should be drawn 30 minutes after completing the gentamicin infusion to allow for drug distribution. Peaks drawn prior to drug distribution will result in falsely elevated serum levels. Patients should remain in the hemodialysis unit for 30 minutes after the gentamicin infusion finishes to obtain an accurate peak level. Although 8-hour HD may reduce serum concentrations by approximately 50%, it is necessary to correctly monitor peak gentamicin levels and maximize efficacy.

Table 1: Dosing recommendations.

<table>
<thead>
<tr>
<th>Database</th>
<th>Target Peak (mg/mL)</th>
<th>Target Trough (mg/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Package Insert</td>
<td>&lt;12</td>
<td>&lt;2</td>
</tr>
<tr>
<td>Lexi-Comp</td>
<td>4-6: Urinary Tract Infection 6-8: Serious Infection 8-10: Life Threatening Infection</td>
<td>0.5-1: Serious Infection 1-2: Life Threatening Infection</td>
</tr>
<tr>
<td>Micromedex</td>
<td>4-12</td>
<td>&lt;2</td>
</tr>
<tr>
<td>Facts and Comparisons</td>
<td>&lt;12</td>
<td>&lt;2</td>
</tr>
</tbody>
</table>

Table 2: Weight based dosing.

Table 3: Monitoring.

Table 4: Re-dosing.

Table 5: Database references.

A serum trough level should be drawn within 30 minutes of the next gentamicin dose. Because gentamicin is readily cleared by hemodialysis, patients will not reach true serum trough levels until dialysis is completed. Waiting for these results would be a significant time burden on hemodialysis units. For this reason, pre-dialysis levels are widely relied upon instead of true trough levels. Some drug information sources suggest re-dosing gentamicin when pre-dialysis levels fall within the range of 3-5 mcg/mL (Table 4).

Clearance

The amount of gentamicin cleared during hemodialysis is highly variable and multifactorial. Longer durations of hemodialysis, “high flux” dialyzer membranes, and increased flow rates on dialysis will all result in greater gentamicin clearance. Most references agree that approximately 50% of gentamicin is removed during an 8-hour hemodialysis session (Table 1). Typical hemodialysis sessions are 4 hours. Therefore, less than 50% of the drug may be removed. In addition to hemodialysis clearance, patients with residual renal function will continue to clear gentamicin between dialysis sessions.

Summary

As discussed above, gentamicin drug information found within drug databases is inconsistent which leads to challenges with gentamicin therapy in ESRD patients on hemodialysis. The abundance of research and primary literature available for gentamicin use in ESRD patients contributes to variety in dosing recommendations. Each database uses different references to arrive at their recommendations (Table 5), none of which have been shown to be more effective than another. In summary, there are many recommendations by reliable drug information sources for dosing gentamicin in ESRD patients on hemodialysis. Regardless of the dosing strategy implemented, correctly monitoring serum levels is essential to achieving maximum efficacy.
References


