GUIDELINES FOR
Extended Interval Gentamicin/Tobramycin Dosing/Monitoring in ADULT patients
Legacy Health System 2002

1. These guidelines are NOT to be used in:
   - Patients with severe renal disease (CrCl<30ml/min) or requiring dialysis. Use traditional kinetics to calculate an appropriate dose.
   - Patients with gram positive infections where aminoglycoside is used for synergy
   - Enterococcal Endocarditis (multiple doses/day more effective)
   - Neonates or pediatric patients
   - Pre-op surgical prophylaxis
   - Patients with a history of allergy to aminoglycosides
   - Patients with a history or signs of a hearing loss or vestibular dysfunction
   - Deteriorating renal function

2. These guidelines should be used with caution in:
   - Patients with chronic ascites or serious liver disease
   - Pregnancy (no data on fetal pharmocokinetics and toxicity)
   - Patients with rapid clearance of drug (e.g. cystic fibrosis, burns>20% of BSA)
   - Patients who have received multiple courses of aminoglycosides in the past

3. Dosing gentamicin and tobramycin
   Utilize aminoglycoside dosing calculator on pharmacy website:
   - Fill in required information (pt name, sex, location, weight, height, age, SrCr, Vd (normal or large Vd which may include burn, postpartum, leukemia, trauma, cystic fibrosis, and edema) and time of infusion (60min for QD dosing, 30min with traditional dosing)
   - Select An appropriate peak and trough and fill in those blanks.
     Goal peak: >10 x MIC for maximize efficacy. Examples of target maximum serum concentrations (Cmax) based on infection: moderate(Cmax14-16mcg/ml), severe (Cmax 16-20mcg/ml). Also consider susceptability of infecting organism. Note these target levels are much higher than with conventional dosing (goal peak 4-10 depending on site of infection., goal trough <2. See Global RX website for specific recommendations)
     Goal trough: "0"mcg/ml at 24 hours. A drug-free interval of 3 - 5 hours minimizes toxicity and permits the reversal of the adaptive post-exposure resistance. (goal trough < 0.5mcg/ml for at least 4 hours).
     Goal 12 hour level around 3mcg/ml

   - Select a "realistic dose" and interval based on the information provided by the calculator. Adjust the dose to achieve the goal levels as above
   - Once completed, print out the calculator information and attach it to the pharmacy monitoring sheet or use as part of your chart note.

If a dosing interval longer than 48 hours would be necessary to achieve a drug free interval of 3-5 hours, then an alternative antimicrobial agent should be considered. Also, in patients with high clearance (e.g., young adults, cystic fibrosis, burns, etc) dosing intervals shorter than 24 hrs may be more appropriate to prevent treatment failure caused by prolonged (>4hr) drug free periods.

IF pharmacy website is down, a reasonable dose can be determined by:
   - Calculate IBW: IBW (females) = 45.5 + (2.3 x inches > 60)
     IBW (males) = 50 + (2.3 x inches > 60)
   - Calculate adjusted weight if obese (actual weight > 20% over IBW)
     Adj BW = IBW + 0.4(actual BW - IBW)
   - Calculate Creatinine Clearance (ml/min) using IBW = 140 - age (IBW) x 0.85 if female
     (72) (serum Cr)
   - Determine if normal Vd or Large Vd (patients with large Vd may include burn, postpartum, leukemia, trauma, cystic fibrosis, and edema)
   - Calculate Dose based on actual body weight OR adjusted weight if obese using table below:

<table>
<thead>
<tr>
<th>Estimated CrCl</th>
<th>Normal Vd</th>
<th>Large Vd</th>
<th>Dosing Interval</th>
</tr>
</thead>
<tbody>
<tr>
<td>&gt;80ml/min</td>
<td>5mg/kg</td>
<td>7mg/kg</td>
<td>q24h</td>
</tr>
<tr>
<td>60-79ml/min</td>
<td>4mg/kg</td>
<td>5.5mg/kg</td>
<td>q24h</td>
</tr>
<tr>
<td>50-59ml/min</td>
<td>3.5mg/kg</td>
<td>5mg/kg</td>
<td>q24h</td>
</tr>
<tr>
<td>30-49ml/min</td>
<td>2.5mg/kg</td>
<td>3.5mg/kg</td>
<td>q24h</td>
</tr>
<tr>
<td>&lt;30ml/min</td>
<td>calculate dose with traditional kinetics</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

***When dosing/monitoring, always consider site/type of infection and patient’s other disease states/risk factors

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4. Aminoglycoside levels:
   a. When to order levels:
      - Aminoglycoside serum concentration MUST be obtained with first dose if:
        - Creatinine Clearance < 50ml/min
        - Intensive Care Unit patient (except one time doses)
        - Severe infection
        - Patient on concomitant nephrotoxic drugs (e.g. amphotericin, cyclosporine, vancomycin)
        - Patient at higher risk for acute renal failure (received contrast media in the prior 72 hours, age>65yo, diabetic, underlying renal dysfunction, received multiple courses of aminoglycoside in the past)
      - Must obtain aminoglycoside level later in therapy if:
        - Patient maintained on therapy ≥ 5 days
        - Renal function changes (SrCr increases by 0.5mg/dl above baseline) Consider DC aminoglycoside in this patient
        - After any dose change that was based on a previous high or low level
        - Plan to send home for continued therapy (obtain prior to DC)
      - Repeat level at least every 7 days for patients on long term therapy
   b. When to draw level:
      - Draw one level 12 hours after start of infusion
      - Draw two levels if expect patient to exhibit unusual kinetics (fast clearance, large Vd) e.g.at 6 and 13 hours in order to perform kinetic calculations
   c. Interpreting levels
      - Specifically timed drug levels (e.g. 12hr) should be plotted on the dosing nomogram graph (goal at 12hours (≤ 3mg/L). If level draw right at 12hr, use table below. Levels draw at other times can be interpreted based on the nomogram graph. This table/graph requires the interval be lengthened for high levels. Alternatively, the dose could be decreased to allow for a drug free interval.

<table>
<thead>
<tr>
<th>Level Draw</th>
<th>Dosing Interval</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 3</td>
<td>q24h</td>
</tr>
<tr>
<td>3 - 5</td>
<td>q36h</td>
</tr>
<tr>
<td>5 - 7</td>
<td>q48h</td>
</tr>
<tr>
<td>&gt; 7</td>
<td>DC protocol and switch to traditional dosing</td>
</tr>
</tbody>
</table>

- Any level drawn ≤4 hours prior to the next dose should be near zero (<<1 mg/L) to allow for a drug free period. If level near zero, continue same dose and interval. If level >1mg/L, extend dosing interval.

5. Routine Monitoring Guidelines:
   a. Is the patient getting better (WBC decreasing, temp decreasing, etc)?
   b. Serum Creatinine and BUN. Obtain at baseline and at least every 3 days
   c. CBC w/diff
   d. Temperature curve
   e. Daily in/out record
   f. Culture results
   g. Addition of potentially nephrotoxic agents or contrast media
   h. Signs of vestibular/auditory dysfunction if on therapy >5 days (dizziness, nausea, tinnitus, sense of fullness in the ears, hearing loss). Notify physician to consider DC of the aminoglycoside should these symptoms occur.
   i. What is the projected course of therapy (long term)?

6. Required documentation/communication:
   a. Initiation and completion of pharmacokinetic monitoring form
b. Daily patient chart note

c. Daily clindoc entry in echart

d. Communication between LIS/inpatient hospital if patient being transferred to their service. FAX copy of monitoring form with pertinent chart notes and patient information to facilitate adequate transition (fax# 225-6130). In addition, call LIS RPh if complicated patient or specific information needs to be communicated (225-6100)

**Background Information**

1. The rational for the pulse dosing of aminoglycosides is rooted in the following observations:

   a. Aminoglycosides exhibit a significant **post-antibiotic effect** (PAE) against aerobic GNB both in vitro and in vivo. The PAE refers to the continued suppression of bacterial growth despite the decline of the antimicrobial concentration to zero. The duration of this effect (2 - 8 hrs) depends on several factors, chief among them is the height of the preceding aminoglycoside peak. The PAE phenomenon suggests that the aminoglycoside serum level may be allowed to fall below the MIC of the pathogen without compromising antimicrobial efficacy. Animal studies suggest that the PAE duration may be shortened by neutropenia. In addition, in vitro studies suggest that the aminoglycoside PAE is extended by the addition of a β-lactam antibiotic.

   b. The **bactericidal action of aminoglycosides is concentration dependent**, i.e., the higher the peak/MIC ratio the higher the kill rate). The multiple daily dosing regimen usually results in relatively low peak/MIC ratios (<5), but when the same total daily dose is given as a single bolus (infused over 60 minutes), much higher ratios are obtained (>10). Ideally, the peak-to-MIC ratio should be >10. (MIC <2 for most organisms)

   c. **Aminoglycoside uptake into renal tubule cells and the inner ear appears to be saturated at relatively low serum levels**, suggesting that higher peaks do not necessarily result in a greater risk of toxicity. Also, serum troughs that are at or near zero may promote tissue drug disposition, shorten tissue exposure, and promote recovery. In addition to the well known risk factors (age, volume depletion, liver disease, co-administration of certain drugs, etc.), duration of exposure to the aminoglycoside appears to be a more important determinant of toxicity than the serum aminoglycoside level. Although definitive evidence is still lacking, animal and human studies strongly suggest that pulse dosing is less nephrotoxic. Further, when the data of 17 clinical studies are considered together, the conclusion that the pulse dosing of aminoglycosides is safer is inevitable.

   d. **Adaptive Resistance of Aminoglycosides.** In vitro studies have also demonstrated that some microorganisms can develop an adaptive resistance to AG when these drugs are present for sustained intervals. The mechanism of resistance is thought to be due to down regulation by allowing several hours for the organism to grow in a drug-free environment.

   e. **The post-antibiotic effect (PAE)** is generally considered a significant advantage of the pulse dosing regimens. However, the data is more equivocal for the single daily dosing regimen. The PAE phenomenon suggests that the aminoglycoside serum level may be allowed to fall below the MIC of the pathogen without compromising antimicrobial efficacy. Animal studies suggest that the PAE duration may be shortened by neutropenia. In addition, in vitro studies suggest that the aminoglycoside PAE is extended by the addition of a β-lactam antibiotic.

   f. **Aminoglycosides exhibit a significant post-antibiotic effect (PAE) against aerobic GNB both in vitro and in vivo.** The PAE refers to the continued suppression of bacterial growth despite the decline of the antimicrobial concentration to zero. The duration of this effect (2-8 hrs) depends on several factors, chief among them is the height of the preceding aminoglycoside peak. The PAE phenomenon suggests that the aminoglycoside serum level may be allowed to fall below the MIC of the pathogen without compromising antimicrobial efficacy. Animal studies suggest that the PAE duration may be shortened by neutropenia. In addition, in vitro studies suggest that the aminoglycoside PAE is extended by the addition of a β-lactam antibiotic.

**References**


5. Barza Ioanidis JPA, Cappelleri et al Single or multiple daily doses of aminoglycosides - a meta-analysis BMJ 1996; 312:338-345


