



Policy Category	BC - Best Care		
Best Care Goals	<input checked="" type="checkbox"/> Safe	<input type="checkbox"/> Personal	<input type="checkbox"/> Connected <input type="checkbox"/> Effective
Applies To	National		
Version	Approval Authority	Effective from	Review by date
1	National Medical Director, WA	5/10/2021	5/10/2024

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1 Rationale

The purpose of this Clinical Protocol is to provide a guiding framework for the administration of aminoglycosides within Hospital at the Home services by Medical Practitioners and clinical staff.

2 Scope

The Clinical Protocol applies to Nationally for HATH clients treated with aminoglycoside medications.

3 Acceptance to HATH Criteria and Pathway

<p>RED Unacceptable for community admission to HATH Refer to ED/ Inpatient management. (May become suitable for HATH after ED or inpatient stabilisation)</p>	<ul style="list-style-type: none"> • Clients with previous vestibular or auditory toxicity related to aminoglycoside therapy. • Serious hypersensitivity reaction to an aminoglycoside. • Pregnancy. • Myasthenia Gravis
<p>ORANGE Requires discussion with Medical Governance prior to acceptance.</p>	<ul style="list-style-type: none"> • Chronically impaired kidney function (adults with a creatinine clearance less than 40 mL/minute and children with an estimated glomerular filtration rate less than 50 mL/minute/1.73 m²) • Rapidly deteriorating kidney function unrelated to sepsis • Request for treatment > 48 hours requires discussion with an Infectious Diseases or Respiratory Physician • Frail and elderly (eg 80 years or older)
<p>GREEN Accepted for HATH</p>	<ul style="list-style-type: none"> • Short- term empirical therapy of less than 48 hrs for serious gram negative infections (eg complex UTI) pending the outcome of investigations to instigate alternative treatment • Over 13 years, suitable for adult dosing and not under the care of a Paediatrician. • Client’s medical condition has been assessed as stable, has a clear diagnosis and prognosis and is at a low risk of rapid deterioration.

4 Pathology Work Up

- Serum creatinine levels to be confirmed and creatinine clearance calculated using Cockcroft–Gault Equation prior to commencement of aminoglycoside therapy and monitored regularly.
- Liver function tests

- Any relevant M C and S results to be obtained during admission

5 General Management

- Collect medical history including potential contraindications or risks such as
 - pre-existing auditory problems,
 - pre-existing vestibular problems (ie dizziness, vertigo or tinnitus),
 - neuromuscular disorders (myasthenia gravis, Parkinsonism),
 - decompensated liver disease,
 - severe cholestasis (serum bilirubin >90 micromol/L),
 - chronic renal failure or deteriorating renal function and
 - recent and current medication history.
- Aminoglycosides should **not generally** be used in patients with:
 - pre-existing significant auditory impairment
 - a pre-existing vestibular condition
 - a first-degree relative with aminoglycoside-induced auditory toxicity, since some people have a rare inherited genetic predisposition
- Note increased risk for clients
 - >70 years
 - those with recent history of aminoglycoside therapy
 - concurrent use of nephrotoxic medications such as NSAIDs, ACE inhibitors, diuretics etc.
- A **single** dose of an aminoglycoside **can** be used in patients:
 - with chronically impaired kidney function (adults with a creatinine clearance less than 40 mL/minute and children with an estimated glomerular filtration rate less than 50 mL/minute/1.73 m²) [NB2]
 - with rapidly deteriorating kidney function unrelated to sepsis
 - who are frail and elderly (eg 80 years or older)
- Ensure pathology results from referral source and results reported to medical governance doctor.

- Collaborate with medical governance doctor regarding any abnormal results.
- Complete nursing assessment, including weight and calculation of ideal body weight and Creatinine Clearance (as per Cockcroft-Gault formula below)
- Initiate intravenous access and commence intravenous therapy as prescribed.
- Educate client regarding medication, side effects.
- Access further investigations as requested by medical governance doctor.
- Liaise with medical governance doctor regarding ongoing management or referral to Infectious Diseases Physician (IDP) if clinically indicated prior to 48hours elapsing post commencement of treatment.

6. Medical Management / Treatment Plan

6.1 Background

Advantages of aminoglycosides

- Aminoglycosides are bactericidal and associated with rapid control of Gram-negative infections.
- Most community- and healthcare-associated Gram-negative pathogens are susceptible to aminoglycosides
- Aminoglycosides have a 'postantibiotic effect' that allows for effective once-daily therapy with reduced rates of toxicity.
- When combined with cell-wall-active drugs (eg beta lactams, glycopeptides), aminoglycosides are synergistic for enterococcal and streptococcal infections.
- Aminoglycosides rarely cause hypersensitivity reactions.
- Aminoglycosides are rarely associated with *Clostridium difficile* infection.

Disadvantages of aminoglycosides

- Aminoglycosides cause nephrotoxicity, usually associated with prolonged treatment courses (longer than 5 to 7 days) and pre-existing kidney impairment. Nephrotoxicity is generally reversible.
- Aminoglycosides cause vestibular and, less commonly, auditory toxicity, mostly associated with prolonged treatment courses. Vestibular and auditory toxicity are generally irreversible.

Aminoglycosides are therefore primarily used for short-term empirical therapy of serious Gram-negative infections, pending the outcome of investigations.

In a limited number of circumstances, they also have a role in directed therapy. These include:

- treatment of pathogens confirmed, or suspected to be, resistant to antibiotics more appropriate for longer term use
- initial combination therapy for *Pseudomonas aeruginosa* infections, until susceptibility results are available
- combination therapy for [brucellosis](#), [nontuberculous mycobacterial infections](#), and [tuberculosis](#)
- combination therapy for synergistic therapy of [streptococcal](#), [enterococcal](#) and [Bartonella](#) endocarditis.

6.2 Dosing

Gentamicin should be administered as once daily dosing (exceptions: endocarditis, ascites, and major burns).

Initial Dose - Once Daily Dosing

Calculate Ideal Body Weight

$$\text{Males: IBW (kg) = [Height (cm) - 152] x 0.9 + 50}$$

$$\text{Females: IBW (kg) = [Height (cm) - 152] x 0.9 + 45.5}$$

Calculate Creatinine Clearance from serum creatinine:

$$\text{Adult males: CrCl (mL/min) = } \frac{(140 - \text{age}) \times \text{weight (kg)}}{0.814 \times \text{serum creatinine (micromol/L)}}$$

Adult females: Multiply the above formula by 0.85

Step One – Initial Dose

13 – 60 years	5mg/kg/day (up to 480mg)
>60 years	4mg/kg/day (up to 400mg)

Step Two – Subsequent Dosing after initial dose is guided by creatinine clearance

Creatinine Clearance (mL/min)	Dosing Interval and maximum number of doses
>60	24 hours (at 0,24 and 48 hours)
40-60	36 hours (at 0 and 36 hours)
<40	Give initial dose once then seek advice from IDP or Microbiologist

6.3 Further doses beyond 48 hrs

No further doses should be administered beyond 48 hours unless there is a clear indication and/or there is no alternative from a safer antibiotic class.

Treatment beyond 48 hours must be discussed with Medical Governor who must be an Infectious Diseases Physician or a Respiratory Physician.

7 Monitoring

- Routine monitoring of aminoglycoside plasma levels is not required if the clinical plan is to cease therapy within 48 hours.
- Monitoring of plasma aminoglycoside levels is recommended if clients are to receive prolonged therapy ie
 - longer than 48hrs OR
 - for patients with altered pharmacokinetics such as
 - [Patients treated with renal replacement therapy.](#)
 - Patients with cystic fibrosis.
 - Patients with ascites.
 - Obese patients
 - Patients treated with chemotherapy that causes kidney dysfunction (eg cisplatin).

In this situation dosing should be based on a 24-hour Area under the Curve (AUC) based computerised method.

Please consult a Clinical Pharmacist if aminoglycoside monitoring is required.

- Nomograms for plasma aminoglycoside concentration monitoring that appeared in older versions of the “Antibiotic Therapeutic Guidelines” are no longer recommended. Likewise, trough concentrations are not recommended for monitoring as they underestimate exposure to aminoglycosides and potential for toxicity.

8 Medical Governance

- The client must have access to medical governance support for 24 hours per day, 7 days per week.
- Primary medical governance can be by referring medical specialists, credentialed referring GPs or by Silver Chain medical staff.
- Care delivery is planned and provided in consultation with the client, medical officer/specialist holding medical governance and nursing staff. Where the primary medical governor is unavailable a Silver Chain medical officer will provide the medical governance.
- In the instance when a client’s condition deteriorates the Silver Chain medical officer or nursing staff can confer with a hospital specialist colleague or an emergency department medical officer.
- The ED is not the primary escalation point unless it is an emergency

When governance is retained by a Silver Chain medical officer the client will have a medical review within 24 hours of admission and the medical officer will determine when the scheduled follow up and discharge will occur. A summary of the episode of care is sent to the referrer or the client’s GP at discharge.

9 Discharge Planning

- Ensure the client has an appointment arranged with own General Practitioner (GP) prior to discharge to ensure continuity of care.
- Ensure discharge summary has highlighted the key clinical concerns/risks you wish to hand over
- Fax client discharge summary to GP.



10. Guiding and Supporting Documents

Silver Chain Group documents that directly relate to and inform this Clinical Protocol are available with this document in the Policy Document Management System (PDMS).

Other documents that directly relate to and inform this Clinical Protocol are as follows:

- Principles of Aminoglycoside Use; eTG complete [Internet]. Published April 2019. Amended March 2020. © Therapeutic Guidelines Ltd (eTG March 2021 edition)

11. Document Details

Document Owner	Executive Medical Director, East Coast
Document Type	CP – Clinical Protocol
Consumer Participation	<input type="checkbox"/> Yes <input checked="" type="checkbox"/> Not Applicable
Functional Area	Acute
Risk Rating	Moderate
Periodic Review	36 months

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