High Risk Medication Alert – Intravenous Potassium Chloride

The former Australian Council for Safety and Quality in Health Care developed a national alert system for high risk medicines. The aim of this national communication strategy was to:

- warn health leaders and professionals about serious known medication-related hazards or risks
- provide tools to effectively ensure action to reduce these risks
- set out responsibilities for system change.

Concentrated intravenous potassium chloride is the subject for the attached alert.

Inappropriate injection of undiluted intravenous potassium chloride has been implicated in fatalities in Australia and overseas.

States, territories and hospitals have already been proactive through development of high risk medicine policies, procedures and guidelines that include implementation of risk management strategies for intravenous potassium. This includes removal the concentrated presentation of intravenous potassium chloride from general patient care areas.

This alert was based on international and Australian work to make a range of recommendations. Health service organisations should consider these along with other more contemporary strategies to ensure safe and appropriate practices are in place for the management of high risk medicines such as intravenous potassium chloride.

The Australian Commission on Safety and Quality in Health Care (ACSQHC) gratefully acknowledges the contribution of many groups who worked together to contribute to the content of this alert.

The ACSQHC can be contacted through its website at www.safetyandquality.gov.au or by email mail@safetyandquality.gov.au.

Please note that the asterisked (*) references within the attached alert are no longer available via web links on the Commission’s website. Additional references and more current information, that

- consider intravenous administration of concentrated potassium chloride as a never event,
- recommend compliance with access and storage restrictions,
- include the need for similar robust safeguards for ALL concentrated electrolyte solutions, and
- promote the use of standardised premix solutions to support safe administration of potassium chloride

are provided on the next page.

“The way to prevent tragic deaths from accidental intravenous injection of concentrated potassium chloride is excruciatingly simple – organisations must take it off floor stock of all units. It is one of the best examples I know of ‘forcing function’ – a procedure that makes a certain type of error impossible.” Lucien L. Leape, M.D. Harvard School of Public Health
1. NHS Specialist Pharmacy Service (SPS). Archived patient (medication) safety alerts from the NPSA and SPS resources to support their implementation. www.sps.nhs.uk/articles/patient-medication-safety-alerts-from-the-npsa-and-sps-resources-to-support-their-implementation/


MEDICATION ALERT!
From the Medication Safety Taskforce of the Australian Council for Safety and Quality in Health Care

The purpose of this alert is to provide frontline health professionals and administrators with information on high risk medications that have the potential to cause serious or catastrophic harm to patients. The intention is to raise awareness of the potential harm and provide a strategy for local level response.

Alert 1, October 2003

Intravenous POTASSIUM CHLORIDE can be fatal if given inappropriately

For the attention of Chief Executive Officers and Directors of Nursing, Pharmacy, and Medical Services; Doctors, Nurses and Pharmacists
For implementation immediately

Wrong ampoule (Australia)
A patient indicated that the cannula site in her hand was becoming painful. An ampoule of normal saline was selected from the medication cupboard in order to flush the cannula site. The patient quickly became distressed and stopped breathing within a few minutes. The ampoule that was thought to be normal saline was actually potassium chloride. The patient could not be resuscitated.1

Preparation error (Australia)
Two ampoules each containing 10 millimoles of potassium chloride were added directly to a running large-volume parenteral fluid without mixing. The patient received a bolus dose of potassium chloride and had a cardiac arrest.2

Overseas Experience
The risks associated with intravenous potassium chloride are well known. It has been identified as the drug most commonly implicated in fatal incidents in acute care facilities. This alert is based on similar recommendations from the UK3, USA4 & Canada5.

Tools and Tips
Tools to action this alert can be found on the Council website at www.safetyandquality.org

Critical incidents have been associated with the preparation and administration of intravenous (IV) potassium chloride indicating that patients are at risk. Ampoules of potassium chloride must be diluted before use.

Three types of error have been identified routinely5

- Wrong ampoule
  Potassium chloride ampoules are mistaken for ampoules of similar appearance, such as sodium chloride 0.9% (normal saline) when reconstituting a drug for injection. Consequently, the patient is administered an accidental overdose of potassium.

- Cognitive mix-up
  The intent is to select frusemide (a diuretic), but a potassium chloride ampoule is selected by mistake and administered. This type of cognitive error is thought to arise due to the frequent use of potassium chloride in patients who are taking frusemide; conditioning staff to the familiar pairing of the two drugs.

- Preparation error
  An intravenous infusion of potassium chloride is prepared incorrectly.

Errors have a single common cause
Incidents have a common root cause—potassium chloride ampoules are available as medication stock in wards and other patient care areas.

Recommendations
1. REMOVE AMPOULES OF POTASSIUM CHLORIDE FROM WARD STOCK AND REPLACE WITH PREMIXED SOLUTIONS.
Due to the risk associated with intravenous potassium chloride, ampoules of potassium chloride SHOULD NOT be kept as a stock item in wards.

2. In critical areas where high concentrations and doses of potassium chloride are necessary, do a risk assessment to determine whether it is appropriate to keep the ampoules as a stock item and develop a protocol for safe preparation and use.

3. Assess the storage of potassium chloride ampoules and premixed solutions to ensure they are stored separately and are readily identifiable from preparations with similar packaging.

The recommendations also apply to ampoules of potassium phosphate or other concentrated potassium salts.
Successful implementation of the actions below requires the commitment of personnel from all clinical areas.

Many acute care facilities have already implemented safety controls for IV potassium chloride in their institution—it is recommended that all facilities evaluate their current safety controls for IV potassium chloride against the actions recommended below.

**CHIEF EXECUTIVE OFFICERS**

1. Form and resource a multidisciplinary team to action the recommendations in this alert, and review and evaluate progress (see review and evaluation below). Team members would include representatives from the Drug and Therapeutics Committee, the Risk Management Department or Quality Department, and patient care teams.

2. The team should be given a mandate to reduce the error potential of potassium chloride and define an implementation strategy (including timelines). The team should provide regular updates to the CEO and/or the appropriate hospital committee outlining progress toward preventing incidents with intravenous potassium chloride.

**DRUG AND THERAPEUTICS COMMITTEES**

3. Develop clear therapeutic guidelines for the use of potassium chloride. Sample guidelines are available on the S&Q Council website. Guidelines should include the following points:

   3.1 Oral, instead of IV potassium chloride should be used for the treatment of hypokalemia whenever clinically feasible.

   3.2 Prescribing of all IV potassium chloride should be in millimoles (mmol).

   3.3 Prescribing and use of standardised premixed solutions containing potassium chloride should be encouraged.

   3.4 Provide a clear definition of the maximum concentration of potassium chloride allowable in an IV solution.

   3.5 Specify the maximum hourly rate and daily limits of potassium chloride that a patient may receive (by central or peripheral lines); and recommended infusion rate, infusion pump requirements, and patient monitoring.

4. Once the guidelines describing safe administration of potassium chloride are approved, ensure that they are readily available and accessible in all wards. Review regularly. Consider developing summary charts of key messages for ready reference; see the S&Q Council website for examples.

5. Review the concentrations of potassium chloride ampoules and premixed solutions available hospital-wide. Consider rationalising the range of concentrations (eg only stock the ‘10 mmol in 10 mL’ ampoules).

**DIRECTORS OF MEDICAL SERVICES, PHARMACY AND NURSING**

Where commercially prepared premixed potassium chloride infusions are available, these products should be procured and introduced, and IV potassium chloride ampoules withdrawn from use. Where this is not feasible, safe on-site preparation and administration protocols should be developed.

6. Undertake a specific multidisciplinary review (by doctors, nurses, and pharmacists) in each ward, department, and clinic with the following aims.

   6.1 Identify if potassium chloride ampoules are available. Identify any barriers to the removal of the ampoules. If no barriers exist, remove all potassium chloride ampoules from the area and replace with premixed solutions. In critical areas where potassium chloride ampoules are to be retained, a risk management policy should be developed and staff education on strategies to minimise risk should be undertaken.

   6.2 Ensure that appropriate concentrations of premixed IV solutions are available in adequate quantities in wards.

   6.3 Ensure prescribing practices are standardised to match the available premixed solutions.

**PHARMACISTS**

7. Evaluate practices for storing IV potassium chloride preparations in the pharmacy and on wards to reduce the likelihood of substitution errors.

8. Assess the range of premixed potassium chloride solutions available and ensure adequate supply for each area.

9. **Where facilities and staff are available**, have the pharmacy aseptic dispensing service prepare premixed potassium chloride products that are not available commercially. Otherwise, follow the protocol for safe on-site preparation.
NURSES

10. Prescriptions with directions such as “KCl 20 mmol IV now” or “give KCl 10 mmol IV bolus” should be considered incomplete and unacceptable. Orders without instructions for dilution and infusion rate should not be accepted. The word “bolus” should never be used for IV potassium chloride solution orders.

11. Consider instituting a double-check policy for administration of IV potassium chloride—have two practitioners check the correct product, dose, dilution, labelling, route and rate before administration, as per the safe on-site preparation protocol.

12. Consider adding auxiliary fluorescent warning labels to IV potassium chloride preparations.

13. Question any nonstandard order for an IV solution with potassium chloride.

14. **Where facilities and staff are available**, advocate having the pharmacy prepare any nonstandard solutions that are deemed necessary but are unavailable in a premixed form.

15. When the above options are not available, keep potassium chloride ampoules on the ward in a medicine cupboard (preferably locked) and store separately from other ampoules with similar appearance.

DOCTORS

16. Standardise prescribing of IV potassium chloride—prescribe in millimoles rather than ‘milligrams per litre’ or ‘percent’.

17. Ensure orders for IV potassium chloride have rate, route, dilution and administration instructions fully specified.

18. Prescribe premixed (standard concentration) potassium chloride infusions where possible.

TRAINING

19. Include the issue of potassium chloride injury and preventive system safeguards as an item for discussion during orientation programs for nurses, doctors, and pharmacists, and as part of continuing education training.

REVIEW AND EVALUATION AT FACILITY LEVEL

Resources must be made available to evaluate progress at an appropriate time, eg after 6 months. For example:

- Are premixed solutions being used? Audit the distribution of potassium chloride ampoules & premixed solutions pre and post system change.
- Are doctors prescribing, and nurses administering premixed solutions? If not, why not? Communicate with staff.
- Are ‘near miss’ incidents relating to IV potassium chloride reported and assessed? Communicate with staff.
- Are ampoules or premixed solutions being transferred between clinical areas? Assess protocols.
- Which areas have retained potassium chloride ampoules, and why? Assess safety controls in these areas.
- To what extent are non-standard IV potassium solutions (ie solutions not available as commercially prepared premixes) being used? How and where are they prepared? Assess the range of products available.
- Have regular meetings and monitor progress. Survey staff regarding knowledge of policies and guidelines.
- Comment on this alert system, your experience in implementation and share your knowledge and tools via the feedback form on the S&Q Council website.

FURTHER INFORMATION

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References

1. Case report supplied courtesy of the Monash University National Centre for Coronial Information (MUNCCI)
2. Case report – personal communication P. Thornton

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