Promoting safer use of injectable medicines

A template standard operating procedure for:
prescribing, preparing and administering injectable medicines in clinical areas

Introduction

The use of injectable medication has many healthcare benefits for patients. The complexities associated with the prescription, preparation and administration of injectable medicines means that there are greater potential risks for patients than for other routes of administration. Weak operating systems increase the potential risk of harm, and safe systems of work are needed to minimise these risks.

Step 1: Prescribing

1.1 All prescriptions for injectable medicines must specify the following:
   • patient’s name;
   • prescriber’s signature;
   • the approved medicine name;
   • the dose and frequency of administration;
   • the date and route of administration;
   • the allergy status of the patient.

1.2. Where relevant, the prescription, or a readily available local protocol, must specify the following:
   • brand name and formulation of the medicine;
   • concentration or total quantity of medicine in the final infusion container or syringe;
   • name and volume of diluent and/or infusion fluid;
   • rate and duration of administration;
   • stability information to determine the expiry date of the final product;
   • type of rate-control pump or device(s) to be used;
   • the age and weight of any patient under 16 years of age, where relevant;
   • date on which treatment should be reviewed;
   • arrangements for fluid balance or clinical monitoring should be made on an individual patient basis and according to local protocol and clinical need.

Step 2: Preparation

2.1 General
2.1.1 Read all prescription details carefully and confirm that they relate to the patient to be treated.
2.1.2 Ensure that the area in which the medicine is to be prepared is as clean, uncluttered and free from interruption and distraction as possible. Ideally, preparation should take place in an area dedicated to this process.
2.1.3 Assemble all materials and equipment: sharps bin for waste disposal, medicine ampoule(s)/vial(s), diluent, syringe(s), needle(s), alcohol wipes, disposable protective gloves, clean re-usable plastic tray.
Check the following:
• expiry dates;
• damage to containers, vials or packaging;
• that medicines were stored as recommended, e.g. in the refrigerator.

2.1.4 Beware of the risk of confusion between similar looking medicine packs, names and strengths. Read all labels carefully.

2.1.5 Check that:
• the formulation, dose, diluent, infusion fluid and rate of administration correspond to the prescription and product information;
• the patient has no known allergy to the medicine (see 1.1);
• you understand the method of preparation.

2.1.6 Calculate the volume of medicine solution needed to give the prescribed dose. Write the calculation down and obtain an independent check by another qualified healthcare professional.

2.1.7 Prepare the label for the prepared medicine (see standard 2.7).

2.1.8 Cleanse your hands according to local policy.

2.1.9 Put on a pair of disposable protective gloves.

2.1.10 Use a 70% alcohol wipe or spray to disinfect the surface of the plastic tray.

2.1.11 Assemble the syringe(s) and needle(s). Peel open wrappers carefully and arrange all ampoules/vials, syringes and needles neatly in the tray.

2.1.12 Use a ‘non-touch’ technique, i.e. avoid touching areas where bacterial contamination may be introduced, e.g. syringe-tips, needles, vial tops. Never put down a syringe attached to an unsheathed needle.

2.1.13 Prepare the injection by following the manufacturer’s product information or local guidelines, and the relevant guidance in standards 2.2 to 2.7.

2.2 **Withdrawing solution from an ampoule (glass or plastic) into a syringe**

2.2.1 Tap the ampoule gently to dislodge any medicine in the neck.

2.2.2 Snap open the neck of glass ampoules, using an ampoule snapper if required.

2.2.3 Attach a needle to a syringe and draw the required volume of solution into the syringe. Tilt the ampoule if necessary.

2.2.4 Invert the syringe and tap lightly to aggregate the air bubbles at the needle end. Expel the air carefully.

2.2.5 Remove the needle from the syringe and fit a new needle or sterile blind hub.

2.2.6 Label the syringe (see standard 2.7).

2.2.7 Keep the ampoule and any unused medicine until administration to the patient is complete to enable further checking procedures to be undertaken.

2.2.8 If the ampoule contains a suspension rather than solution, it should be gently swirled to mix the contents immediately before they are drawn into the syringe.

2.2.9 The neck of some plastic ampoules is designed to connect directly a syringe without use of a needle, after the top of the ampoule has been twisted off.

2.3 **Withdrawing a solution or suspension from a vial into a syringe**

2.3.1 Remove the tamper-evident seal from the vial and wipe the rubber septum with an alcohol wipe. Allow to dry for at least 30 seconds.

2.3.2 With the needle sheathed, draw into the syringe a volume of air equivalent to the required volume of solution to be drawn up.

2.3.3 Remove the needle cover and insert the needle into the vial through the rubber septum.

2.3.4 Invert the vial. Keep the needle in the solution and slowly depress the plunger to push air into the vial.

2.3.5 Release the plunger so that solution flows back into the syringe.
2.3.6 If a large volume of solution is to be withdrawn, use a push-pull technique. Repeatedly inject small volumes of air and draw up an equal volume of solution until the required total is reached. This ‘equilibrium method’ helps to minimise the build-up of pressure in the vial.

2.3.7 Alternatively, the rubber septum may be pierced with a second needle to let air into the vial as solution is withdrawn. The tip of the vent needle must always be kept above the solution to prevent leakage.

2.3.8 With the vial still attached, invert the syringe. With the needle and vial uppermost, tap the syringe lightly to aggregate the air bubbles at the needle end. Push the air back into the vial.

2.3.9 Fill the syringe with the required volume of solution then draw in a small volume of air. Withdraw the needle from the vial.

2.3.10 Expel excess air from the syringe. Remove the needle and exchange it for a new needle or a sterile blind hub.

2.3.11 The vial(s) and any unused medicine should be kept until administration to the patient is complete.

2.3.12 If the vial contains a suspension rather than solution, it should be gently swirled to mix the contents, immediately before they are drawn into the syringe.

2.4 Reconstituting powder in a vial and drawing the resulting solution or suspension into a syringe

2.4.1 Remove the tamper-evident seal from the vial and wipe the rubber septum with an alcohol wipe. Allow to dry for at least 30 seconds.

2.4.2 Use the procedure in 2.2 above to withdraw the required volume of diluent (e.g. water for injections or sodium chloride 0.9%) from ampoule(s) into the syringe.

2.4.3 Inject the diluent into the vial. Keeping the tip of the needle above the level of the solution in the vial, release the plunger. The syringe will fill with the air which has been displaced by the solution (if the contents of the vial were packed under a vacuum, solution will be drawn into the vial and no air will be displaced). If a large volume of diluent is to be added, use a push-pull technique (see above).

2.4.4 With the syringe and needle still in place, gently swirl the vial(s) to dissolve all the powder, unless otherwise indicated by the product information. This may take several minutes.

2.4.5 Follow the relevant steps in 2.3 above to withdraw the required volume of solution from the vial into the syringe.

2.4.6 Alternatively, the rubber septum may be pierced with a second needle to let air into the vial as solution is withdrawn. The tip of the vent needle must always be kept above the solution to prevent leakage.

2.4.7 If a purpose-designed reconstitution device is used, the manufacturer’s instructions should be read carefully and followed closely.

2.5 Adding a medicine to an infusion

2.5.1 Prepare the medicine in a syringe using one of the methods described in 2.2 to 2.4 above.

2.5.2 Check the outer wrapper of the infusion container is undamaged.

2.5.3 Remove the wrapper and check the infusion container itself in good light. It should be intact and free of cracks, punctures/leaks.

2.5.4 Check the infusion solution, which should be free of haziness, particles and discolouration.

2.5.5 Where necessary, remove the tamper-evident seal on the additive port according to the manufacturer’s instructions or wipe the rubber septum on the infusion container with an alcohol wipe and allow to dry for at least 30 seconds.
2.5.6 If the volume of medicine solution to be added is more than 10% of the initial contents of the infusion container (more than 50ml to a 500ml or 100ml to a 1litre infusion), an equivalent volume must first be removed with a syringe and needle.

2.5.7 Inject the medicine into the infusion container through the centre of the injection port, taking care to keep the tip of the needle away from the side of the infusion container. Withdraw the needle and invert the container at least five times to ensure thorough mixing before starting the infusion.

2.5.8 Do not add anything to any infusion container other than a burette when it is hanging on the infusion stand since this makes adequate mixing impossible.

2.5.9 Before adding a medicine to a hanging burette, administration must be stopped. After the addition has been made and before administration is re-started, the contents of the burette must be carefully swirled to ensure complete mixing of the contents.

2.5.10 Check the appearance of the final infusion for absence of particles, cloudiness or discolouration.

2.5.11 Label the infusion (see standard 2.7).

2.6 Diluting a medicine in a syringe for use in a pump or syringe-driver

2.6.1 Prepare the medicine in a syringe using one of the methods described above.

2.6.2 Draw the diluent into the syringe to be used for administration by the pump or syringe-driver. Draw in some air (slightly more than the volume of medicine needed) and remove the needle.

2.6.3 Stand the diluent syringe upright. Insert the needle of the syringe containing the medicine into the tip of the diluent (administration) syringe and add the medicine to it. Alternatively, a disposable sterile connector may be used to connect two syringes together directly.

2.6.4 Check the following:

- the total volume of injection solution in the syringe is as specified in the prescription and that the infusion can be delivered at the prescribed rate by the administration device chosen;
- the rate of administration is set correctly on the administration device and according to the manufacturer's instructions.

2.6.6 Fit a blind hub to the administration syringe and invert several times to mix the contents.

2.6.7 Remove the blind hub. Tap the syringe lightly to aggregate the air bubbles at the needle end. Expel the air and refit the blind hub.

2.6.8 Carefully check the syringe for cracks and leaks and then label it (see standard 2.7), especially noting the requirements specific to syringe drivers.

2.6.9 Check that the rate of administration is set correctly on the device before fitting the syringe, priming the administration set and starting the infusion device.

2.7 Labelling injection and infusion containers

2.7.1 All injections should be labelled immediately after preparation, except for syringes intended for immediate push (bolus) administration by the person who prepared them. Under no circumstances should an operator be in possession of more than one unlabelled syringe at any one time, nor must an unlabelled syringe be fitted to a syringe driver or similar device.

2.7.2 Labels used on injectable medicines prepared in clinical areas should contain the following information:

- name of the medicine;
- strength;
- route of administration;
- diluent and final volume;
- patient's name;
• expiry date and time;
• name of the practitioner preparing the medicine.

2.7.3 Place the final syringe or infusion and the empty ampoule(s)/vials(s) in a clean plastic tray with the prescription for taking to the patient for administration.

**Step 3: Administration of an injectable medicine**

**3.1 Before administering any injection**

3.1.1 Check all the following:
• patient’s name, hospital/NHS Number or date of birth or address;
• prescriber’s signature;
• the approved medicine name;
• the dose and frequency of administration;
• the date and route of administration;
• the allergy status of the patient.

3.1.2 Also check, where relevant:
• brand name and formulation of the medicine;
• concentration or total quantity of medicine in the final infusion container or syringe;
• name and volume of diluent and/or infusion fluid;
• rate and duration of administration;
• type of rate-control pump or device(s) to be used;
• the age and weight of any patient under 16 years of age, where relevant;
• date on which treatment should be reviewed.

3.1.3 Check that the medicine is due for administration at that time and has not already been given.

3.1.4 Assemble everything you need including any flushing solution(s) needed.

3.1.5 Explain and discuss the procedure with the patient.

3.1.6 Check any infusion already in progress. It should be free of haziness, particles and discolouration.

3.1.7 Check that an appropriate access device is in place. Flush it immediately before and after administration of a medicine, and between doses of different medicines administered consecutively, according to local policy. Also check the administration site for signs of leakage, infection or inflammation.

**3.2 Administration of injections – general**

3.2.1 Check infusions. They should be free of haziness, particles and discolouration.

3.2.2 Use aseptic (non-touch) technique at all times.

3.2.3 Attach administration sets to infusion containers carefully, on a flat surface and using the technique appropriate to the type of container.

3.2.4 Prime the access device according to local policy immediately before starting an infusion.

3.2.5 Before adding a medicine to a hanging burette, administration must be stopped. After the addition has been made and before re-commencement, the contents of the burette must be carefully swirled to ensure complete mixing.

**3.3 After administration**

3.3.1 After completion of an intermittent infusion, flush the access device according to local policy.

3.3.2 Ask the patient to report promptly any soreness at the injection site or discomfort of any sort.

3.3.3 Make a detailed record of administration.
• Discard the empty ampoules/vials from which the injection was prepared and any unused medicine. Ampoules or vials should never be used to prepare more than one injection unless specifically labelled by the manufacturer for ‘multi-dose’ use.

3.4.5 Re-check the administration site for signs of leakage, infection or inflammation and continue to monitor the patient, contents of the infusion container and the rate of infusion according to local policy.

3.4.6 Check that arrangements for monitoring fluid balance or clinical parameters have been made. Ensure that relevant documentation is made available for subsequent regular monitoring to take place.