

Name of product	Cyclophosphamide injection. Aseptically prepared from licensed sterile starting ma
Concentration	Concentration 20mg/ml in pre-filled syringes
Diluent	Sodium Chloride 0.9% w/v
Volume	Dose specific volumes in accordance with the national dose banding tables
Final container	Polypropylene or polycarbonate sterile luer lock syringe, graduated in millilitres an sealed with sterile syringe cap/hub. Syringe and syringe cap/hub integrity to be va The validation should conform to the Protocols for the integrity testing of syringes published by the NHS Pharmaceutical QA Committee (2nd Edition, April 2013). Air bubbles within the final product must be minimised so the end user can use wi need to remove air.
Starting materials	Licensed cyclophosphamide powder for reconstitution for injection. Licensed Sodium Chloride Injection 0.9% w/v CE marked sterile syringes and syringe caps/hubs
Labelling	Labelling must be compliant with the principles of labelling for safety and the BP G specification on unlicensed medicines. Tall Man lettering must be used for the drug name.
Label sample	An example label is provided below stating the minimum requirements only (the lab format is not restrictive and suppliers can use their preferred layout):- CYCLOphosphamide xxmg in xxml Sodium Chloride 0.9% w/v For Intravenous Injection Store in a Refrigerator at 2-8°C Protect From Light Expiry: dd/mm/yyy BN: XXXXXXX Keep out of the reach and sight of children Manufacturer's details MS XXXXXX <i>Caution Cytotoxic: Handle with care</i>
Batch Number	All products will have a unique batch identification number
	All materials and manufacturing processes will be latex free or clearly labelled if no
Latex status of - components - manufacturing process	