

PATIENT GROUP DIRECTION (PGD)

Administration of OXYBUPROCAINE 0.4% eye drops By Registered Practitioner working for University Hospitals of Derby and Burton

Documentation details

Reference no:	UHDB 114
Version no:	2.1
Valid from:	19/09/2024
Review date:	19/03/2027
Expiry date:	18/09/2027

Change history

Version number	Change details	Date
2.1	Addition of Orthoptist to PGD working group	20.05.24

Glossary

Abbreviation	Definition



1. PGD template development (PGD Working Group)

PGD Working Group Membership (minimum requirement of consultant, pharmacist and a registered professional who can work under a PGD (or manages the staff who do). If this is a review of existing PGD, <u>replace</u> previous names with the individuals involved for this version

Name	Designation	
VICKI MEREDITH	SENIOR SISTER OUTPATIENTS	
PARAMJIT KAUR	SENIOR SISTER OUTPATIENTS	
MR ANIL KUMAR	CONSULTANT OPHTHALMOLOGIST	
SUZANNE SMITH	PHARMACIST	
ANDREW CASTLE	ORTHOPTIST	

Where an antimicrobial is included, confirm the name, designation and date of the antimicrobial pharmacist who has reviewed this version

Name of antimicrobial pharmacist	Designation	Date Reviewed
n/a	n/a	n/a

2. Organisational authorisations

The PGD is not legally valid until it has had the relevant organisational authorisation.

University Hospitals of Derby & Burton NHS Foundation Trust authorises this PGD for use by the services or providers listed below:

Authorised for use by the following organisation and/or services All UHDB sites and in clinics operated by UHDB staff at peripheral sites

Limitations to authorisation

Organisational Authorisation (legal requirement).

Role	Name	Sign	Date
Medicines Safety Pharmacist	Ellie Gunner	Signed copy held by Pharmacy	19/09/2024
Pharmacist: Medicines Safety Officer, Chief Pharmacist or assigned deputies)			

Additional signatories (required as per legislation and locally agreed policy)			
Role	Name	Sign	Date
Pharmacist	SUZANNE SMITH	Signed copy held by Pharmacy	10/09/2024
Assistant Clinical Director (Ophthalmology)	LOLA LAWUYI	Signed copy held by Pharmacy	10/09/2024
Senior Sister Outpatients QHB	VICKI MEREDITH	Signed copy held by Pharmacy	16/09/2024
Senior Sister Outpatients RDH	PARAMJIT KAUR	Signed copy held by Pharmacy	18/09/2024

Local enquiries regarding the use of this PGD may be directed to UHDB.PGDgovernance@nhs.net

Section 7 provides a registered health professional authorisation sheet. Individual professionals must be authorised by name to work to this PGD.



3. Characteristics of staff

Qualifications and professional registration	 Qualified NMC Registered Nurse HCPC Registered Orthoptist Health Care Professionals (who can legally operate under PGDs) who have undergone additional training to administer drug outlined in this PGD and been assessed as competent.
Initial training	 Completion of all Essential-to-role training as outlined in the UHDB PGD policy. Individual has read and understood full content of this PGD and signed authorisation (section 7) The Registered Healthcare Professional will undertake training and will ensure he/she is competent in all aspects of this treatment.
Competency assessment	Staff operating under this PGD are encouraged to review their competency using the <u>NICE Competency Framework for health</u> <u>professionals using patient group directions</u> Individuals operating under this PGD are personally responsible for ensuring they remain up to date with the use of all medicines included in the PGD - if any training needs are identified these should be discussed with either the authorising manager (section 7) or the manager within the PGD working group (section 1) so that further training can be provided as required.
Ongoing training and competency	- Health care professionals must complete annual basic life support and anaphylaxis training to administer drugs under this PGD.
The decision to supply any medication rests with the individual registered health professional who must abide by the PGD and any associated organisation policies.	

4. Clinical condition or situation to which this PGD applies

Clinical condition or situation to which this PGD applies	Local anaesthesia for removal of foreign bodies, or for examination purposes (see inclusion criteria).
Criteria for inclusion Use BNF/BNFC/SPC. Take into account any clinical guidelines or policies that are available locally or nationally, e.g. BASHH/NICE/JCVI	 Patients identified as requiring a topical ocular anaesthetic. Schirmers Lacrimal syringing Pachymetry Tonometry Minor procedures Botulinum procedures Removal of foreign body/rust ring Eye irrigation Ultrasonography/ A scan biometry Intra-vitreal injections
Criteria for exclusion	 Consent not gained Previous local or systemic reactions to the medicine Known or suspected hypersensitivity to any of the ingredients Penetrating injuries Pregnant patients Breastfeeding patients Pre-term neonates
Cautions including any relevant action to be taken	• Ensure emergency drugs and equipment, including adrenaline, are available for the treatment of anaphylaxis and emergencies, according to local policy.
Action to be taken if the patient is excluded	 Record reasons for exclusion in patient notes Inform doctor Advise patient on alternative treatment
Action to be taken if the patient or carer declines treatment	 Document advice given Inform doctor Advise patient on alternative treatment
Arrangements for referral for medical advice	Inform doctor or suitably qualified specialist in ophthalmology. Discuss potential consequences/referral/records to be kept. The practitioner is expected to use their own clinical judgement and refer patients to OOHs GP/ A&E / Minor Injuries unit / Walk-in centre as they see fit. Provide appropriate details e.g. Eye casualty opening times

5. Description of treatment

Name, strength & formulation of drug	Oxybuprocaine hydrochloride 0.4% eye drops single dose (minims)
Legal category	Prescription Only Medicine (POM)

Route / method of administration	Topical into the eye Patients, especially children, should be asked to compress the lachrymal sac for a minute following application to reduce systemic absorption. The anaesthetised eye should be protected from dust and bacterial contamination.
Indicate any off-label use	None
(if relevant)	
Dose and frequency of administration	1-3 drops (depending on indication) Instil 1 – 3 drops at 90 second intervals:
Duration of treatment	1 drop should be sufficient to allow tonometry, A further drop after 90 seconds for e.g. fitting of contact lenses
	Three drops at 90 second intervals for removal of foreign body or incision for minor procedures.
	Corneal sensitivity is normal again after about one hour.
	Each single dose unit should be discarded after use.
Quantity to be supplied (leave blank if PGD is administration ONLY)	n/a administration only
Storage	 Stock must be securely stored according to UHDB medicines policy and in conditions in line with SPC as detailed below: Store below 25°C. Do not freeze. Protect from light. Each single dose unit should be discarded after use.
Drug interactions	 The following interactions have been identified and should be considered where it is known a patient is on the following medicines: None Known
	A detailed list of drug interactions is available in the SPC, which is available from the electronic Medicines Compendium website: www.medicines.org.uk
Identification & management of adverse reactions	 Transient stinging and blurring of vision may occur on instillation. Cornea may be damaged by prolonged application.
	Systemic absorption may be reduced by compressing the lacrimal sac at the medial canthus for a minute during and following the instillation of the drops.
	A detailed list of adverse reactions is available in the SPC, which is available from the electronic Medicines Compendium website: www.medicines.org.uk

University Hospitals of Derby and Burton NHS Foundation Trust

Management of and reporting procedure for adverse reactions to the Medicines and Healthcare professionals and patients/carers are encouraged to reporting scheme on: https://wilowcard.mirra.cov.uk Averse reactions Healthcare products Regulatory Agency (MHRA) using the Yellow Card reporting scheme on: https://wilowcard.mirra.cov.uk Record all adverse reactions (moderate harm or above as per NRLS definition) should be reported via trust incident management system (e.g. Datix) to ensure duty of candour and learning from harm during clinical use. Anaphylaxis/Emergency Treatment at Derby and main Queens site: Call the crash team via switchboard Anaphylaxis/Emergency treatment at all other sites: Call P39 Summon help Maintain airway CPR Follow trust resuscitation guidelines Written Information to be given to patient or carer Pollow trust resuscitation guidelines None routinely required for administration in department. May give copy of marketing authorisation holder's patient information leaflet (PLU) which can be obtained from www.medicines.org.uk if required Any stinging or blurring of vision after application should be transient. Do not druch or rub the eye after application The anaesthetic effect will subside after about an hour. Do not dwear contact lenses during the treatment and until normal vision is restored. Do not wear contact lenses during the treatment and until normal vision is restored. Do not wear contact lenses during the treatment and which may include: in the event of an adverse reaction. The individual/carer should be advised to s		
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 batch number and expiry date (if applicable e.g. injections and implants) 		
implants)		
		 advice given, including advice given if excluded or declines
treatment		treatment
details of any adverse drug reactions and actions taken		



 Confirm whether <u>supplied and/or administered</u> and that this was done via Patient Group Direction (PGD) Records should be signed and dated (or a password controlled e-records). All records should be clear, legible and contemporaneous.
If you are not recording in ePMA (or other electronic system which has ability to generate audit reports) then a record of all individuals receiving treatment under this PGD should also be in the clinical area for audit purposes as per UHDB PGD policy.

6. Key references

Key references	 Electronic Medicines Compendium <u>http://www.medicines.org.uk/</u> Electronic BNF <u>https://bnf.nice.org.uk/</u> NICE Medicines practice guideline "Patient Group Directions" <u>https://www.nice.org.uk/guidance/mpg2</u>
	https://medusa.wales.nhs.uk

7. Registered health professional authorisation sheet

PGD Name [version]: Ophthalmology-Oxybuprocaine 0.4% [V2.1] PGD ref: UHDB 114

Valid from: 19/09/2024 Expiry date: 18/09/2027

Before signing check that the document you have read is published on Koha or is an in-date hard-copy with all necessary authorisations signed in section 2. The Name/Version/Ref of the document you have read MUST match this authorisation form.

Registered health professional

By signing this patient group direction you are indicating that

a) You agree to and understand all content and commit to only work within this framework.

b) You have completed any core PGD e-Learning or training records on My Learning Passport or within your department.

c) You meet the staff characteristics and have completed any additional learning/competency outlined in Section 3 of this PGD.

Patient group directions do not remove inherent professional obligations or accountability.

It is the responsibility of each professional to practise only within the bounds of their own competence and professional code of conduct.

I confirm that I have read and understood the content of this Patient Group Direction and that I am willing and competent to work to it within my professional code of conduct.

Name	Designation	Signature	Date

Authorising manager / Assessor

I confirm that the registered health professionals named above have declared themselves suitably trained and competent to work under this PGD. I give authorisation on behalf of University Hospitals of Derby & Burton NHS Foundation Trust for the above named health care professionals who have signed the PGD to work under it.

Name	Designation	Signature	Date

Note to authorising manager

Score through unused rows in the list of registered health professionals to prevent additions post managerial authorisation.

This authorisation sheet must be retained by a manager in the clinical department where the PGD is in-use to serve as a record of those registered health professionals authorised to work under this PGD.