

This Patient Group Direction (PGD) must only be used by registered healthcare professionals who have been named and authorised by their organisation to practice under it. The most recent and in date final signed version of the PGD should be used.

PATIENT GROUP DIRECTION (PGD)

Supply and/or administration of ulipristal acetate 30mg tablet for emergency contraception

in Rotherham, Doncaster, and South Humber (RDaSH) NHS Foundation Trust

Version Number 2.1

Change History		
Version and Date	Change details	
Version 1.0 March 2020	New template	
Version 2.0 March 2023	Updated template (no clinical changes to expired V1)	
Version 2.1 October 2023	Reworded exclusion and caution sections to reflect change in guidance re combined oral contraceptive, in line with updated FSRH guidance. Updated references.	

PGD DEVELOPMENT GROUP

Date PGD template comes into effect:	1 st March 2023
Review date	September 2025
Expiry date:	28 th February 2026

This PGD template has been peer reviewed by the Reproductive Health PGDs Short Life Working Group in accordance with their Terms of Reference. It has been approved by the Faculty for Sexual and Reproductive Health (FSRH) in October 2022.

This section MUST REMAIN when a PGD is adopted by an organisation.

Name	Designation
Dr Cindy Farmer	Chair General Training Committee
	Faculty of Sexual and Reproductive Healthcare (FSRH)
Michelle Jenkins	Advanced Nurse Practitioner, Clinical Standards Committee
	Faculty of Sexual and Reproductive Healthcare (FSRH)
Vicky Garner	Deputy Chief Midwife British Pregnancy Advisory Service (BPAS)
Gail Rowley	Quality Matron British Pregnancy Advisory Service (BPAS)
Katie Girling	British Pregnancy Advisory Service (BPAS)
Julia Hogan	CASH Nurse Consultant MSI Reproductive Choices
Kate Devonport	National Unplanned Pregnancy Association (NUPAS)
Chetna Parmar	Pharmacist adviser Umbrella
Helen Donovan	Royal College of Nursing (RCN)
Carmel Lloyd	Royal College of Midwives (RCM)
Clare Livingstone	Royal College of Midwives (RCM)
Kirsty Armstrong	National Pharmacy Integration Lead, NHS England
Dipti Patel	Local authority pharmacist
Emma Anderson	Centre for Postgraduate Pharmacy Education (CPPE)
Dr Kathy French	Specialist Nurse
Dr Sarah Pillai	Associate Specialist
Alison Crompton	Community pharmacist
Andrea Smith	Community pharmacist
Lisa Knight	Community Health Services pharmacist
Bola Sotubo	NHS North East London ICB pharmacist
Tracy Rogers	Director, Medicines Use and Safety, Specialist Pharmacy Service
Sandra Wolper	Associate Director Specialist Pharmacy Service
Jo Jenkins (Working Group Co-ordinator)	Lead Pharmacist PGDs and Medicine Mechanisms Specialist Pharmacy Service

ORGANISATIONAL AUTHORISATIONS

Name	Job title and organisation	Signature	Date
Dr Graeme Tosh	Medical Director	#	24/10/23
Stephen Davies	Chief Pharmacist	Danies	32/16/23
Sheila Lloyd	Director of Nursing and AHPs	Silajd	06/10/23
Anil Rajpal	Senior Pharmacist	Eng Rayed	7/11/23
Tina Proctor	Nurse Consultant Senior Representative of professional group using the PGD	Thirds	7/11/23

Committee Approval:

NAME	DATE
Medicines Management Committee	20.10.2023

Date of Unplementation 08.11.2023

1. Characteristics of staff

Qualifications and professional registration	Current contract of employment within the Local Authority or NHS commissioned service or the NHS Trust/organisation.
	Registered healthcare professional listed in the legislation as able to practice under Patient Group Directions.
Initial training	The registered healthcare professional authorised to operate under this PGD must have undertaken appropriate education and training and successfully completed the competencies to undertake clinical assessment of patients ensuring safe provision of the medicines listed in accordance with local policy.
	Suggested requirement for training would be successful completion of a relevant contraception module/course accredited or endorsed by the FSRH, CPPE or a university or as advised in the RCN training directory.
	Individual has undertaken appropriate training for working under PGDs for the supply and administration of medicines. Recommended training - eLfH PGD elearning programme
	The healthcare professional has completed locally required training (including updates) in safeguarding children and vulnerable adults or level 2 safeguarding or the equivalent.
Competency assessment	 Individuals operating under this PGD must be assessed as competent (see Appendix A) or complete a self-declaration of competence for emergency contraception. Staff operating under this PGD are encouraged to review their competency using the NICE Competency Framework for health professionals using patient group directions
Ongoing training and competency	 Individuals operating under this PGD are personally responsible for ensuring that they remain up to date with the use of all medicines and guidance included in the PGD - if any training needs are identified these should be addressed and further training provided as required. Organisational PGD and/or medication training as required by employing Trust/organisation.
	ation rests with the individual registered health professional any associated organisational policies.

2. Clinical condition or situation to which this PGD applies

Clinical condition or situation	To reduce the risk of pregnancy after unprotected sexual
to which this PGD applies	intercourse (UPSI) or regular non-hormonal contraception
	has been compromised or used incorrectly.
Criteria for inclusion	Any individual presenting for emergency contraception
	(EC) between 0 and 120 hours following UPSI or when
	regular non-hormonal contraception has been
	compromised or used incorrectly.
	No contraindications to the medication.
	Informed consent given.
Criteria for exclusion	Informed consent not given.
	Individuals under 16 years old and assessed as lacking
	capacity to consent using the Fraser Guidelines.
	Individuals 16 years of age and over and assessed as
	lacking capacity to consent.
	This episode of UPSI occurred more than 120 hours ago.
	N.B. A dose may be given if there have been previous
	untreated or treated episodes of UPSI within the current
	cycle if the most recent episode of UPSI is within 120
	hours.
	Known pregnancy (N.B. a previous episode of UPSI in
	this cycle is not an exclusion. Consider pregnancy test if
	more than three weeks after UPSI and no normal
	menstrual period).
	Less than 21 days after childbirth.
	Less than 5 days after miscarriage, abortion, ectopic
	pregnancy or uterine evacuation for gestational
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	trophoblastic disease (GTD).
	Known hypersensitivity to the active ingredient or to any
	component of the product - see <u>Summary of Product</u>
	Characteristics
	Use of levonorgestrel (LNG-EC) or any other progestogen in the provious 7 days (i.e. harmonal contracentian).
	in the previous 7 days (i.e. hormonal contraception
	including combined oral contraception, hormone
	replacement therapy (or use for other gynaecological
	indications).
	Concurrent use of antacids, proton-pump inhibitors or H ₂ -
	receptor antagonists including any non-prescription (i.e.
	over the counter) products being taken
	Severe asthma controlled by oral glucocorticoids.
	Individuals using enzyme-inducing drugs/herbal products
	or within 4 weeks of stopping.
	Acute porphyria
Cautions including any	All individuals should be informed that insertion of a
relevant action to be taken	copper intrauterine device (Cu-IUD) within five days of
	UPSI or within five days from earliest estimated ovulation
	is the most effective method of emergency contraception.
	If a Cu-IUD is appropriate and acceptable supply oral EC
	and refer to the appropriate health service provider.
	Ulipristal acetate (UPA-EC) is ineffective if taken after
	ovulation.
	If individual vomits within three hours from ingestion, a
	repeat dose may be given.
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	 Body Mass Index (BMI) >26kg/m2 or weight >70kg – individuals should be advised that though oral EC methods may be safely used, a high BMI may reduce the effectiveness. A Cu-IUD should be recommended as the most effective method of EC. Consideration should be given to the current disease status of those with severe malabsorption syndromes, such as acute/active inflammatory bowel disease or Crohn's disease. Although the use of UPA-EC is not contra-indicated it may be less effective and so these individuals should be advised that insertion of Cu-IUD would be the most effective emergency contraception for them and referred accordingly if agreed. Breast feeding – advise to express and discard breast milk for 7 days after UPA-EC dose. The effectiveness of UPA-EC can be reduced by progestogen taken in the following 5 days and individuals must be advised not to take progestogen containing drugs, including combined oral contraception, for 5 days after UPA-EC. UPA EC is generally not recommended in a missed pill situation. See section 'Written information and further advice to be given to individual'. If the individual is less than 16 years of age an assessment based on Fraser guidelines must be made and documented. If the individual is less than 13 years of age the healthcare professional should speak to local safeguarding lead and follow the local safeguarding policy. If the individual has not yet reached menarche consider onward referral for further assessment or investigation.
Action to be taken if the individual is excluded or declines treatment	 document in the consultation record. Record reason for decline in the consultation record. Offer suitable alternative emergency contraception or refer the individual as soon as possible to a suitable health service provider if appropriate and/or provide them
individual is excluded or	 after UPA-EC. UPA EC is generally not recommended a missed pill situation. See section 'Written information and further advice to be given to individual'. If the individual is less than 16 years of age an assessment based on Fraser guidelines must be made and documented. If the individual is less than 13 years of age the healthcare professional should speak to local safeguarding lead and follow the local safeguarding policy. If the individual has not yet reached menarche consideration onward referral for further assessment or investigation. Explain the reasons for exclusion to the individual and document in the consultation record. Record reason for decline in the consultation record. Offer suitable alternative emergency contraception or refer the individual as soon as possible to a suitable.

3. Description of treatment

N	Ulipristal acetate 30mg tablet
Name, strength & formulation of drug	Oliphistal acetate 30Hg tablet
Legal category	P
Route of administration	Oral
Off label use	Best practice advice given by Faculty of Sexual and Reproductive Healthcare (FSRH) is used for guidance in this PGD and may vary from the Summary of Product Characteristics (SPC). This PGD includes off-label use in the following conditions: Lapp-lactase deficiency Hereditary problems of galactose intolerance Glucose-galactose malabsorption Severe hepatic impairment Medicines should be stored according to the conditions detailed in the Storage section in this table. However, in the event of an inadvertent or unavoidable deviation of these conditions the local pharmacy or Medicines Management team must be consulted. Where drugs have been assessed by pharmacy/Medicines Management in accordance with national or specific product recommendations as appropriate for continued use this would constitute off-label administration under this PGD. The responsibility for the decision to release the affected drugs for use lies with pharmacy/Medicines Management. Where a drug is recommended off-label consider, as part of the consent process, informing the individual/parent/carer that the drug is being offered in accordance with national
Dose and frequency of	 guidance but that this is outside the product licence. One tablet (30mg) as a single dose taken as soon as
administration	possible up to 120 hours after UPSI.
Duration of treatment	 A single dose is permitted under this PGD. If vomiting occurs within 3 hours of UPA-EC being taken a repeat dose can be supplied under this PGD. Repeated doses, as separate episodes of care, can be given within the same cycle. Please note: If within 7 days of previous LNG-EC offer LNG-EC again (not UPA-EC) If within 5 days of UPA-EC then offer UPA-EC again (not LNG-EC) Appropriately labelled pack of one tablet.
Quantity to be supplied	
Storage	Medicines must be stored securely according to national guidelines and in accordance with the product SPC.
Drug interactions	A detailed list of drug interactions is available in the SPC, which is available from the electronic Medicines Compendium website: www.medicines.org.uk or the BNF www.bnf.org Refer also to FSRH guidance on drug interactions with
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hormonal contraception A detailed list of adverse reactions is available in the SPC. Identification & management which is available from the electronic Medicines Compendium of adverse reactions website: www.medicines.org.uk and BNF www.bnf.org The following side effects are common with UPA-EC (but may not reflect all reported side effects): Nausea or vomiting Abdominal pain or discomfort Headache Dizziness Muscle pain (myalgia) Dysmenorrhea Pelvic pain Breast tenderness Mood changes Fatigue The FSRH advises that disruption to the menstrual cycle is possible following emergency contraception. Healthcare professionals and patients/carers are Management of and reporting encouraged to report suspected adverse reactions to the procedure for adverse Medicines and Healthcare products Regulatory Agency reactions (MHRA) using the Yellow Card reporting scheme on: http://yellowcard.mhra.gov.uk Record all adverse drug reactions (ADRs) in the patient's medical record. Report any adverse reactions via organisation incident All methods of emergency contraception should be Written information and discussed. All individuals should be informed that fitting further advice to be given to a Cu-IUD within five days of UPSI or within five days individual from the earliest estimated ovulation is the most effective method of emergency contraception. Ensure that a patient information leaflet (PIL) is provided within the original pack. If vomiting occurs within three hours of taking the dose, the individual should return for another dose. Explain that menstrual disturbances can occur after the use of emergency hormonal contraception. Provide advice on ongoing contraceptive methods, including how these can be accessed. Repeated episodes of UPSI within one menstrual cycle the dose may be repeated more than once in the same menstrual cycle should the need occur. In line with FSRH guidance individuals using hormonal contraception should delay restarting their regular hormonal contraception for 5 days following UPA-EC use. Avoidance of pregnancy risk (i.e. use of condoms or abstain from intercourse) should be advised until fully effective. Advise a pregnancy test three weeks after treatment especially if the expected period is delayed by more than seven days or abnormal (e.g. shorter or lighter than usual), or if using hormonal contraception which may

Advice / follow up treatment	 affect bleeding pattern. Promote the use of condoms to protect against sexually transmitted infections (STIs) and advise on the possible need for screening for STIs. There is no evidence of harm if someone becomes pregnant in a cycle when they had used emergency hormonal contraception. Advise to consult a pharmacist, nurse or doctor before taking any new medicines including those purchased. The individual should be advised to seek medical advice
Advice / follow up treatment	 in the event of an adverse reaction. The individual should attend an appropriate health service provider if their period is delayed, absent or abnormal or if they are otherwise concerned. Pregnancy test as required (see advice to individual above). Individuals advised how to access on-going contraception and STI screening as required.
Records	 Record: The consent of the individual and If individual is under 13 years of age record action taken If individual is under 16 years of age document capacity using Fraser guidelines. If not competent record action taken. If individual over 16 years of age and not competent, record action taken Name of individual, address, date of birth GP contact details where appropriate Relevant past and present medical history, including medication history. Examination finding where relevant e.g. weight Any known medication allergies Name of registered health professional operating under the PGD Name of medication supplied Date of supply Dose supplied Quantity supplied including batch number and expiry date in line with local procedures. Advice given, including advice given if excluded or declines treatment Details of any adverse drug reactions and actions taken Advice given about the medication including side effects, benefits, and when and what to do if any concerns Any referral arrangements made Any supply outside the terms of the product marketing authorisation Recorded that administered/supplied via Patient Group Direction (PGD) Records should be signed and dated (or a password controlled e-records) and securely kept for a defined period in
Potoronae Number: 574	controlled e-records) and securely kept for a defined period in

line with local policy.
All records should be clear, legible and contemporaneous.
A record of all individuals receiving treatment under this PGD should also be kept for audit purposes in accordance with local policy.

4. Key references

Key references (accessed	Electronic Medicines Compendium http://www.medicines.org.uk/
September 2022 and July	Electronic BNF https://bnf.nice.org.uk/
2023)	NICE Medicines practice guideline "Patient Group Directions"
	https://www.nice.org.uk/guidance/mpg2
	Faculty of Sexual and Reproductive Health Clinical Guidance:
	Emergency Contraception - March 2017 (Amended July 2023)
	https://www.fsrh.org/standards-and-guidance/current-clinical-
	guidance/emergency-contraception/
	Faculty of Sexual and Reproductive Health Drug Interactions with
	Hormonal Contraception – May 2022
	https://www.fsrh.org/documents/ceu-clinical-guidance-drug-
	interactions-with-hormonal/
	Royal Pharmaceutical Society Safe and Secure Handling of
	Medicines December 2018
	https://www.rpharms.com/recognition/setting-professional-
	standards/safe-and-secure-handling-of-medicines

Appendix A – Example registered health professional authorisation sheet PGD Name/Version Valid from: Expiry:

Before signing this PGD, check that the document has had the necessary authorisations. Without these, this PGD is not lawfully valid.

Registered health professional

By signing this patient group direction you are indicating that you agree to its contents and that you will work within it.

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

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 RDASH 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 should be retained to serve as a record of those registered health professionals authorised to work under this PGD.

This information will be retained by the Chief Pharmacist of RDaSH NHS Foundation Trust.