

Patient Group Direction For Insertion Of The Progestogen-Only Intra-Uterine Device (LNG-IUD) In NHS Grampian, Orkney And Shetland

Lead Author:

Adapted from FSRH/SPS PGD Insertion of the Progestogen-Only Intra-Uterine Device (LNG-IUD) Version 2.0 – Date published August 2023

Approver:

NoS PGD Group

Authorisation:

NHS Grampian

Signature:

Signature:

NoS Identifier:

NoS/PGD/LNG_IUD/1408

Review Date:

February 2026

Expiry Date: July 2026

Date Approved by NoS: 23rd September 2023

NHS Grampian, Orkney, Shetland, have authorised this Patient Group Direction to help individuals by providing them with more convenient access to an efficient and clearly defined service within the NHS Boards. This Patient Group Direction cannot be used until Appendix 1 and 2 are completed.

Uncontrolled when printed

Version 2

Revision History for NoS:

Nos PGD that has been adapted and/or superseded		PGD supersedes NHSG/PGD/IUS/Levonorgestrel/MGPG1063, Version 2.1 (NoS Version)	
Date of change	Summary of Changes		Section heading
August 2023	Reference	e to NoS Appendix 1 and 2	Authorisation
June 2023	Removed SPS advised training and added TURAS Initial NoS PGD training link added		Initial Training
June 2023	Added in statement about capacity under the age of 13 and the legislation statement added		Criteria for inclusion
June 2023	Additional facilities and supplies added as per NoS Sexual health Template added.		Additional facilities and supplies
June 2023	Information added about recording on HEPMA Records added		Records

FSRH/SPS Revision

Change History	у
Version and	Change details
Date	
Version 1.0	New template
August 2020	
Version 1.1	Addition of Jaydess [®] ▼ Levonorgestrel 13.5 mg intrauterine system as
November	a black triangle product.
2020	Acute porphyria added as exclusion.
Version 1.2	Levosert® license revised to usage period from 5 to 6 years for when
March 2021	indication is for contraception.
	Dose and frequency of administration section amended to read:
	Levonorgestrel 52mg Intrauterine System (Levosert®) - effective
	for up to 6 years or until contraception no longer required if
	individual is over the age of 45 years of age at time of insertion.
Version 1.3	Benilexa One Handed® 52mg levonorgestrel-releasing intrauterine
September	system added to Name, strength and formulation of drug and Dose
2022	and frequency of administration sections.
	eLFH PGD e learning added to training section
Version 2.0	Updated template. Amendments to exclusion, cautions, dose and
April 2023	frequency of administration and adverse effects sections to align with
	updated FSRH IUC guidance. Minor formatting/wording changes to
	align with other SPS PGD reproductive health templates.

Review date: The review date for a PGD needs to be decided on a case-by-case basis in the interest of safety. The expiry date should not be more than 3 years, unless a change in national policy or update is required.

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Authorisation

This specimen Patient Group Direction (PGD) template has been produced by FSRH/SPS and adapted by North of Scotland PGD Group (NoS) to assist NHS Boards. NHS Boards should ensure that the final PGD is considered and approved in line with local clinical governance arrangements for PGDs.

The qualified health professionals who may administer LNG-IUD under this PGD can only do so as named individuals. It is the responsibility of each professional to practice within the bounds of their own competence and in accordance with their own Code of Professional Conduct and to ensure familiarity with the manufacturer's product information/Summary of Product Characteristics (SmPC) for all vaccines administered in accordance with this PGD.

NHS Board governance arrangements will indicate how records of staff authorised to operate this PGD will be maintained. Under PGD legislation there can be no delegation. Administration of the LNG-IUD has to be by the same practitioner who has assessed the patient under the PGD.

All authorised staff are required to read the PGD and sign the Agreement to Administer Medicines Under PGD (Appendix 1).

A Certificate of Authorisation (<u>Appendix 2</u>) signed by the authorising professional/manager should be supplied. This should be held in the individual health professional's records, or as agreed within the individual Health Board.

This PGD has been produced for NoS by:					
Doctor	Dr Linda Sandilands	Signature	hhhih	Date Signed	18/09/2023
Pharmacist	Alison Smith	Signature	(Denotin)	Date Signed	22/09/2023
Nurse	Julia Penn	Signature	Julia Penn	Date Signed	21/09/2023

Approved for use within NoS by:

NoS Group Chair	Signature	Date Signed
Lesley Coyle	The second second	22/09/2023

Authorised and executively signed for use within NoS by:

NHS Grampian Chief Executive	Signature	Date Signed	
Professor Caroline Hiscox P.P. Adam Coldwells, Deputy Chief Executive	Almhus	23/09/2023	

PGD Development Group

Date PGD template comes into effect: August 2023

Review date February 2026 Expiry date: July 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 February 2023.

Name	Designation
Dr Cindy Farmer	Vice President, General Training FSRH
Michelle Jenkins	Advanced Nurse Practitioner, Clinical Standards Committee 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)
Sim Sesane	CASH Nurse Consultant MSI Reproductive Choices
Kate Devonport	National Unplanned Pregnancy Association (NUPAS)
Chetna Parmar	Pharmacist adviser Umbrella
Heather Randle	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	Consultant
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

1. Characteristics of Staff

Qualifications and professional registration	Current contract of employment within a Local Authority or NHS commissioned service or an NHS Trust/organisation.	
registration	Registered healthcare professional listed in the legislation as able to practice under Patient Group Directions.	
	Registered nurses and midwives as recognised by the Nursing and Midwifery Council (NMC).	
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.	
	Recommended 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. Individuals working under this PGD should hold an in date FSRH Letter of Competence Intrauterine Techniques (LoC IUT) which has been achieved or recertified within the last 5 years.	
	Have undertaken NoS PGD module training on <u>TURAS</u> Learn.	
	PGD users should have read thoroughly and be familiar with the <u>FSRH IUC guidance</u> .	
	The healthcare professional has completed locally required training (including updates) in safeguarding children and vulnerable adults or level 2 safeguarding or the equivalent.	
	The healthcare professional must ensure that they have an up to date certificate for Basic Life Support (BLS) and anaphylaxis as required by the employing Trust/organisation	
Competency assessment	Individuals operating under this PGD must be assessed as competent (see Appendix 1 and 2) or complete a self-declaration of competence for LNG-IUD contraception insertion.	
Ongoing training and competency	Individuals operating under this PGD are personally responsible for ensuring 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. CONTRACTOR CONTRACTOR	
	 FSRH LoC IUT must be recertified every 5 years. Organisational PGD and/or medication training as required by employing Trust/organisation. 	
	Competent to assess the Individuals capacity to understand the nature and purpose of the medicine administration in order to give or refuse consent.	

 Have undertaken NHS e-anaphylaxis training or equivalent (including annual updates) which covers all aspects of the identification and management of anaphylaxis
 Maintain their skills, knowledge and their own professional level of competence in this area according to their Code of Professional Conduct
 Have knowledge and familiarity of the following;
 <u>SmPC</u> for the medicine(s) to be administered in accordance with this PGD.

The decision to administer any medication rests with the individual registered health professional who must abide by the PGD and any associated organisational policies.

Clinical condition or situation to which this PGD applies

2. Clinical condition or situation to which this PGD applies				
Clinical condition or	Contraception			
situation to which				
this PGD applies				
Criteria for inclusion	 Individual (age from menarche to 55 years) presenting for contraception. Informed consent given. An individual under 16 years of age may give consent for the administration of LNG-IUD, provided they understand fully the benefits and risks involved. The individual should be encouraged to involve a parent/guardian, if possible, in this decision. Where there is no parental involvement and the individual indicates that they wish to accept the administration, administer should proceed, if the healthcare professional deems the individual to have the legal capacity to consent. The Age of Legal Capacity (S) Act 1991, s2 (4) states that 'a person under the age of 16 years shall have legal capacity to consent on their own behalf to any surgical, medical or dental procedure or treatment where, in the opinion of a qualified medical practitioner attending them, they are capable of understanding the nature and possible 			
Critoria for exclusion	consequences of the procedure or treatment.			
Criteria for exclusion	 Informed consent not given. Individuals under 16 years of age and assessed as not competent using Fraser Guidelines. Individuals 16 years of age and over and assessed as lacking capacity to consent. If under 13 years of age this PGD cannot be used and the healthcare professional should speak to the local Child Protection lead if any concerns and follow the local child protection policy, otherwise refer to GP or paediatrics. Known hypersensitivity to the active ingredient or to any constituent of the product - see Summary of Product Characteristics. Risk of pregnancy Any reported unprotected sexual intercourse (UPSI) within the last 3 weeks. 			

- Over 48 hours and less than 4 weeks postpartum (note the LNG-IUD can be fitted immediately post-partum, post termination of pregnancy, ectopic pregnancy or miscarriage)
- Postpartum sepsis
- Post-abortion sepsis
- Gestational trophoblastic disease with decreasing or. persistently elevated β-hCG levels or malignancy

Refer to the FSRH CEU clinical guideline Intrauterine Contraception and clinical guidance 'switching' for specific guidance about starting and switching IUC:

Insertion of new device (no current IUC in situ)

- Any reported unprotected sexual intercourse (UPSI) since day 5 of a natural cycle, AND within the last 3 weeks.
- If any UPSI >3 weeks ago where menstruation has not since occurred - negative pregnancy test required prior to insertion.

Changing to a new device (current IUC insitu and in

Any reported unprotected sexual intercourse (UPSI) within the last 7 days

Changing to a new device (current IUC insitu but out of date)

- Any reported unprotected sexual intercourse (UPSI) within the last 3 weeks
- If UPSI >3 weeks ago- negative pregnancy test required prior to insertion

Cardiovascular Disease

- Development of ischaemic heart disease, transient ischaemic attack or stroke whilst using the LNG-IUD.
- For individuals with pre-existing arrhythmia, Eisenmenger physiology, single ventricle (or Fontan) circulation, long QT syndrome or impaired ventricular function, a vasovagal reaction could pose a serious risk of a significant cardiac event and therefore IUC procedures should be undertaken in a hospital setting.

Cancers

- Current or past history of breast cancer.
- Malignant liver tumour (hepatocellular carcinoma).
- Cervical cancer (awaiting treatment)
- Endometrial cancer
- Cervical cancer (resulting in radical trachelectomy)

Gastro-intestinal conditions

- Severe decompensated cirrhosis.
- Benign liver tumour (hepatocellular adenoma).

Infections

- Current or recurrent pelvic inflammatory disease (PID)
- Known chlamydial infection either symptomatic or asymptomatic
- Known gonorrhoea infections either symptomatic or asymptomatic
- Current purulent cervicitis or vaginitis
- Known pelvic tuberculosis
- HIV infection with CD4 <200cells/mm³

Anatomical abnormalities

Distorted uterine cavity; congenital or acquired abnormality distorting the uterine cavity, including fibroids, incompatible with LNG-IUD insertion.

Other Conditions

- Unexplained vaginal bleeding suspicious of a serious medical condition, present before commencing the method
- Organ transplant with complications
- Acute porphyria
- Previous endometrial ablation

Cautions including any relevant action to be taken

- 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.
- Individuals taking anticoagulants or antiplatelets refer to **FSRH CEU Statement Management of women taking** anticoagulants or antiplatelet medications who request intrauterine contraception or subdermal implants
- Liaison with an individual's MDT or clinical specialist may be required with certain conditions (e.g. inherited bleeding disorders, cardiac disease, taking anticoagulants, Ehlers-Danlos syndromes (EDS), Postural tachycardia syndrome (PoTS).
- Individuals at risk of an adrenal crisis will usually need to increase their steroid dose prior to, and for 24 hours after, IUC insertion and should ideally have their IUC procedure scheduled for early morning.
- If an individual with PoTS has a history of postural syncope, advice should be sought from their cardiologist, as it may be recommended that insertion should be undertaken in a hospital setting.
- Individuals with cardiac arrhythmias (other than long QT) discuss with relevant clinician.
- Discuss with appropriate medical/independent non-medical prescriber any medical condition or medication of which the healthcare professional is unsure or uncertain.

- Explain the reasons for exclusion to the individual and document in the consultation record.
- Record reason for decline in the consultation record.
- Where required refer the individual to a suitable health service provider if appropriate and/or provide them with information about further options.

3. Description of treatment		
Name, strength & formulation of drug	Levonorgestrel 13.5 mg intrauterine system (Jaydess®▼) Levonorgestrel 19.5mg intrauterine system (Kyleena®) Levonorgestrel 52mg intrauterine System (Levosert®) Levonorgestrel 52mg intrauterine system (Mirena®) Levonorgestrel 52mg intrauterine system (Benilexa One Handed®) Note: • This PGD does not restrict which brands can be supplied – local formularies/restrictions should be referred to and the above list edited to reflect local formularies. • See http://www.mhra.gov.uk/spc-pil/ or http://www.medicines.org.uk for further information and further brand information including full details of adverse	
Legal category	effects and interactions.	
Black triangle	Jaydess® ▼ Levonorgestrel 13.5 mg intrauterine system is a black triangle product. This information was accurate at the time of writing. See product SPCs at www.medicines.org.uk for indication of current black triangle status.	
Route of administration	Intra-uterine Insert using aseptic or no-touch technique as per FSRH guidance on intrauterine contraception	
Off label use	Best practice advice is given by the FSRH and is used for guidance in this PGD and may vary from the Summary of Product Characteristics (SPC). This PGD specifically includes inclusion criteria and dosage regimens which are outside the market authorisation for many of the available products but which are included within FSRH guidance: When used for contraception only, any 52mg LNG-IUD maybe retained until contraception no longer required in individuals over 45 years of age at time of insertion Mirena® – effective for up to 6 years Initial insertion after day 7 of the menstrual cycle if it is reasonably certain that the individual is not pregnant Postpartum insertion between 4-6 weeks	

Medicines should be stored according to the conditions detailed in the Storage section below. However, in the event of an inadvertent or unavoidable deviation of these conditions the local pharmacy or Medicines Management team must be consulted. Where medicines 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 medicines for use lies with pharmacy/Medicines Management.

Where a medicine is recommended off-label consider, as part of the consent process, informing the individual/parent/carer that the medicine is being offered in accordance with national guidance but that this is outside the product licence.

Dose and frequency of administration

- One LNG-IUD to be inserted (after removal of previous LNG-IUD if required).
- Insert on day 1-5 of the menstrual cycle with no need for additional protection
- The LNG-IUD can be inserted at any time after day 5 of the menstrual cycle if it is reasonably certain that the individual is not pregnant. Additional contraception is then required for 7 days after insertion of the LNG-IUD.
- For guidance on changing from one contraceptive method to another, and when to start after an abortion and postpartum, refer to the Faculty of Sexual and Reproductive Healthcare (FSRH) guidelines.

Frequency of LNG-IUD insertion:

- Levonorgestrel 13.5mg intrauterine delivery system (Jaydess®) - effective for up to 3 years
- Levonorgestrel 19.5mg intrauterine delivery system (Kyleena®) - effective for up to 5 years.
- Levonorgestrel 52mg intrauterine delivery system (Levosert®) - effective for up to 6 years or until contraception no longer required if individual is over the age of 45 years of age at time of insertion.
- Levonorgestrel 52mg intrauterine delivery system (Mirena®) - effective for up to 6 years or until contraception no longer required if individual is over the age of 45 years of age at time of insertion.
- Levonorgestrel 52mg intrauterine delivery system (Benilexa One Handed®) - effective for up to 6 years
- or until contraception no longer required if individual is over the age of 45 years of age at time of insertion.

Duration of treatment

For as long as individual requires contraception and has no contraindications to its use.

Quantity to be supplied	Single LNG-IUD is to be inserted per episode of care.
Storage	Medicines must be stored securely according to national guidelines.
Drug interactions	All concomitant medications should be checked for interactions. A detailed list of drug interactions is available in the individual product SPC, which is available from the electronic Medicines Compendium website www.medicines.org.uk the BNF www.bnf.org and FSRH CEU Guidance: Drug Interactions with Hormonal Contraception https://www.fsrh.org/standards-and-guidance/documents/ceu-clinical-guidance-drug-interactions-with-hormonal/ Refer to a prescriber if any concern of a clinically significant drug
Identification & management of adverse reactions	interaction. A detailed list of adverse reactions is available in the SPC, which is available from the electronic Medicines Compendium website: www.medicines.org.uk and BNF www.bnf.org The LNG-IUD is generally well tolerated. The following possible adverse effects are commonly reported with LNG-IUD (but may not reflect all reported adverse effects): Headache Disturbance of bleeding patterns Changes in mood Weight change Loss of libido Breast tenderness Acne Insertion complications may include infection, expulsion, or perforation. Individuals should be advised on the signs that
Additional facilities and supplies	 perforation. Individuals should be advised on the signs that these may have occurred and the action to take if they become concerned. Access to working telephone Suitable storage facilities An acceptable level of privacy to respect individual's right to confidentiality and safety Access to medical support (this may be via the telephone Clean and tidy work areas, including access to hand washing facilities or alcohol hand gel. A copy of this current PGD in print or electronically.) Suitable waste disposal facilities Immediate access to in-date anaphylaxis kit (IM adrenaline 1:1000) and emergency drugs including atropine and oxygen according to local protocol.

Management of and	Healthcare professionals and patients/carers are					
reporting procedure for adverse reactions	encouraged to report suspected adverse reactions to the Medicines and Healthcare products Regulatory Agency					
	(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 via organisation incident policy. 					
	Note certain LNG-IUDs have additional Risk Minimisation					
	materials (RMMs) to support safe use – organisations should					
	ensure any RMMs supplied for the product/s used within their					
	organisation are considered. See product profile at					
Written information	 www.medicines.org.uk for further information Provide patient information leaflet (PIL) provided with the 					
and further advice to	original pack.					
be given to individual	 Explain mode of action, side effects, risks and benefits of the medicine 					
	Advise about the risks of the medication including failure					
	rates and serious side effects and the actions to be taken.					
	Advise about the possible symptoms of serious sequelae e.g. infection, actoric programs and perfection an					
	infection, ectopic pregnancy, expulsion and perforation and when to seek clinical advice					
	Teach individual how to check threads and to seek clinical					
	advice if threads not felt					
	Advise when replacement of the LNG-IUD will be due.					
	Offer condoms and advice on safer sex practices and					
	possible need for screening for sexually transmitted infections (STIs)					
	Ensure the individual has contact details of local					
	service/sexual health services.					
Advice / follow up	The individual should be advised to seek medical advice in					
treatment	the event of an adverse reaction.					
December	Individual to seek further advice if they have any concerns					
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. o If individual over 16 years of age and not competent,					
	 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 and family history.					
	Any known allergiesDetails of insertion procedure to include:					
	 Details of insertion procedure to include: Name of registered health professional 					
	O Harrie of registered fleditif professional					

- Date of insertion
- Name/brand of LNG-IUD inserted
- Batch number and expiry date of administered
- Bimanual examination and speculum findings
- Uterine sounding
- Use of no touch technique
- Name of assistant/their role
- Analgesia or local anaesthetic used
- Problems encountered during insertion
- Advice given, including advice given if excluded or declines treatment
- Individual has been advised on the date/s for next appointment as required.
- 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 administration outside the terms of the product marketing authorisation and additional advice given relating to this and advice given (e.g. additional contraception for 7
- Recorded that administration is 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 line with local policy.

An electronic/HEPMA record of the screening and subsequent administration, or not of the medicine(s) specified in this PGD should be made in accordance with individual Health Board electronic/HEPMA recording processes.

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.

Key references

Key references (accessed January 2023)

- **Electronic Medicines Compendium** http://www.medicines.org.uk/
- Electronic BNF https://bnf.nice.org.uk/
- NICE Medicines practice guideline "Patient Group Directions" https://www.nice.org.uk/guidance/mpg2
- FSRH Clinical Guideline: Intrauterine contraception (March https://www.fsrh.org/documents/ceuguidanceintrauterinecont
 - raception/ Faculty of Sexual and Reproductive Health Drug Interactions

- with Hormonal Contraception May 2022 https://www.fsrh.org/documents/ceu-clinical-guidance-druginteractions-with-hormonal/
- Faculty of Sexual and Reproductive Healthcare (2016) UK Medical Eligibility Criteria for Contraceptive Use. https://www.fsrh.org/documents/ukmec-2016/
- Faculty of Sexual and Reproductive Healthcare (2016 Clinical Guideline: Quick Starting Contraception (April 2017) https://www.fsrh.org/standards-and-guidance/current-clinicalguidance/quick-starting-contraception/
- Faculty of Sexual and Reproductive Healthcare (2019) Service standards for record keeping https://www.fsrh.org/standards-andquidance/documents/fsrh-service-standards-for-recordkeeping-july-2019/
- FSRH CEU Resource: New one-handed, reloadable 52mg levonorgestrel-releasing intrauterine system https://www.fsrh.org/news/fsrh-ceu-resource-new-onehanded-reloadable-52mg-levonorgestrel/ (2021)



Appendix 1 - Healthcare Professional Agreement to Administer Medicine(s) Under Patient Group Direction

:		(Insert name)
Working within:		e.g. Area, Practice
Agree to administer the medic Direction:	cine(s) contained within the fo	ollowing Patient Group
	ion For Insertion Of The (LNG-IUD) In NHS Gran Shetland, Version 2	
administer the medicine(s) un	riate training to my profession nder the above direction. I ag r out with the recommendation	ree not to act beyond my
Signed:		
Print Name:		
Date:		
Profession:		
Professional Registration number/PIN:		



Appendix 2 - Healthcare Professionals Authorisation to Administer **Medicine(s) Under Patient Group Direction**

The Lead manager/Professional of each clinical area is responsible for maintaining records of all clinical areas where this PGD is in use, and to whom it has been disseminated.

The Senior Nurse/Professional who approves a healthcare professional to administer the medicine(s) under this PGD is responsible for ensuring that they are competent, qualified and trained to do so, and for maintaining an up-to-date record of such approved persons.

The Healthcare Professional that is approved to administer the medicine(s) under this PGD is responsible for ensuring that they understand and are qualified, trained and competent to undertake the duties required. The approved person is also responsible for ensuring that administration is carried out within the terms of the direction, and according to their individual code of professional practice and conduct.

Patient Group Direction For Insertion Of The Progestogen-Only Intra-Uterine Device (LNG-IUD) In NHS Grampian, Orkney And Shetland, Version 2

Local clinical area(s) where the listed healthcare professionals will operate under this PGD:

Name of Healthcare Professional	Signature	Date	Name of Manager	Signature	Date

Patient Group Direction For Insertion Of The Progestogen-Only Intra-Uterine Device (LNG-IUD) In NHS Grampian, Orkney And Shetland, Orkney Version 2

Name of Healthcare Professional	Signature	Date	Name of Manager	Signature	Date