# PATIENT GROUP DIRECTION (PGD)

**Supply of a combined oral hormonal contraceptive (COC)** **by Community Pharmacists and Pharmacy Technicians in England working in a pharmacy registered to provide the NHS Pharmacy Contraception Service**

**DRAFT v 0.2**

Version 3.0

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| **Change History** |
| **Version and Date** | **Change details** |
| Version 11 April 2023 | PGD approved |
| Version 1.127 April 2023 | Exclusion added relating to Zoely® only  |
| Version 2.01 December 2023 | Update to include initiation of oral contraception.andUpdated PGD development group members.Statement added in exclusion criteria regarding consideration of lactose/sucrose content in individual products. |
| Version 3.020 June 2025 | Included pharmacy technicians as an additional professional group.Updated Short Life Working Group members.Statement added in exclusion criteria regarding consideration of lactose/sucrose content in individual products.Clarified statement on management of people using 30mcgEE/LNG combination who have missed two pills in week one of cycle in dose and frequency section. Added statement on advice when used in combination with GLP-1 agonists. Clarified statement on Qlaira.Clarification of quantity to be supplied for ongoing suppliesAdded statement on depressed mood and depression in written information and further advice to be given to individual.Added statement on advice on desogestrel and risk of meningioma. Updated references. Update to amend Annex B |

This Patient Group Direction (PGD) must only be used by pharmacists and pharmacy technicians who have been named and authorised by their organisation to practise under it (See Appendix A). The most recent and in date final signed version of the PGD must be used.

**PGD DEVELOPMENT GROUP**

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| Date PGD template comes into effect:  | 20th June 2025 |
| Review date | September 2025 |
| Expiry date:  | 31st March 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 was approved by the Faculty of Sexual and Reproductive Healthcare (FSRH) in November 2022.

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| **Name** | **Designation** |
| Dr Cindy Farmer | Vice President, General Training 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) |
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| Katie Girling | British Pregnancy Advisory Service (BPAS) |
| Julia Hogan | CASH Nurse Consultant MSI Reproductive Choices |
| Kate Devonport | National Unplanned Pregnancy Advisory Service (NUPAS) |
| Chetna Parmar | Pharmacist adviser Umbrella  |
| Helen Donovan | Royal College of Nursing (RCN) |
| Carmel Lloyd | Royal College of Midwives (RCM) |
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| Kirsty Armstrong  | National Pharmacy Integration Lead, NHS England |
| Dipti Patel | Local authority pharmacist  |
| Emma Anderson | Centre for Pharmacy Postgraduate Education (CPPE) |
| Dr Sarah Pillai | Associate Specialist |
| Alison Crompton | Community pharmacist |
| Andrea Smith | Community pharmacist |
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| Tracy Rogers | Director, Medicines Use and Safety, Specialist Pharmacy Service  |
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| Jo Jenkins  | Lead Pharmacist PGDs and Medicine Mechanisms Specialist Pharmacy Service |
| Rosie Furner (Working Group Co-ordinator) | Specialist Pharmacist PGDs and Medicine Mechanisms Specialist Pharmacy Service |

**ORGANISATIONAL AUTHORISATIONS**

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| **Name**  | **Job title and organisation**  | **Signature** | **Date** |
| **Senior doctor** **Claire Fuller** | National Medical Director, NHS England  |  | 20/06/2025 |
| **Senior pharmacist****David Webb** | Chief Pharmaceutical Officer, NHS England  | Text  Description automatically generated | 20/06/2025 |
| **Person signing on behalf of** [authorising body](https://www.nice.org.uk/guidance/mpg2/chapter/recommendations#authorising-body)**David Webb** | Chief Pharmaceutical Officer, NHS England  | A close-up of a signature  AI-generated content may be incorrect. | 20/06/2025 |

1. Characteristics of staff

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| **Qualifications and professional registration** | GPhC registered pharmacist or pharmacy technician able to practise under Patient Group Directions (PGDs). |
| **Initial training** | The pharmacist or pharmacy technician authorised to operate under this PGD must have undertaken appropriate education and training and be competent to undertake clinical assessment of individuals ensuring safe provision of the medicines listed in accordance with the specification. To deliver this service, the pharmacist or pharmacy technician should have evidence of competence in the clinical skills and knowledge covered in the CPPE and/or the NHS England e-learning for healthcare (elfh) modules listed in section 5 of the [**NHS Pharmacy Contraception Service specification.**](https://www.england.nhs.uk/long-read/nhs-pharmacy-contraception-service/) The pharmacist or pharmacy technician has completed training and is up to date with service requirements for safeguarding children and vulnerable adults. |
| **Competency assessment** | * Pharmacists or pharmacy technicians operating under this PGD must have declared their competence and must be authorised by a manager within their organisation to provide the service (see [**Appendix A**](#AppendixA)).
* Pharmacists or pharmacy technicians operating under this PGD are encouraged to review their competency using appropriate competency framework tools, such as the[**NICE Competency framework: For health professionals using patient group directions.**](https://www.nice.org.uk/guidance/mpg2/resources)
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| **Ongoing training and competency** | * Pharmacists or pharmacy technicians 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 undertaken as required.
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| The decision to supply any medication rests with the individual pharmacist or pharmacy technician who must abide by the PGD and any associated organisational policies.  |

1. Clinical condition or situation to which this PGD applies

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| **Clinical condition or situation to which this PGD applies** | * This PGD applies to the [**NHS Pharmacy Contraception Service**](https://www.england.nhs.uk/long-read/nhs-pharmacy-contraception-service/) only:
	+ Initiation of oral contraception for contraceptive purposes
	+ Review and ongoing supply of oral contraception for contraceptive purposes where previously initiated in primary care or sexual health clinics (or equivalent).
 |
| **Criteria for inclusion** | * Individual (age from menarche up to and including 49 years of age) presenting for:
	+ Initiation of first-time oral contraception
	+ Initiation of oral contraception after a pill free break
	+ Initiation of a new (to the individual) oral contraceptive
	+ Ongoing supply of their current oral contraception.
* Able to confirm blood pressure and BMI, as per the specification, prior to supply.
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| **Criteria for exclusion** | * Individuals under 16 years of age and assessed as not competent using [**Fraser Guidelines**](https://learning.nspcc.org.uk/child-protection-system/gillick-competence-fraser-guidelines).
* Individuals 16 years of age and over and assessed as lacking capacity to consent.
* Established pregnancy. Note – risk of pregnancy with a negative pregnancy test is not an exclusion.
* Known hypersensitivity to the active ingredient or to any constituent of the product - see [**Summary of Product Characteristics**](https://www.medicines.org.uk/emc) (SPC).
* Some COC products contain lactose/sucrose. Individuals with rare hereditary problems of galactose intolerance, total lactase deficiency, fructose intolerance, glucose-galactose malabsorption or sucrase- isomaltase deficiency should not take these medicines. Where applicable, check product excipients before supplying.
* Less than 21 days after childbirth (for deliveries over 24 weeks gestation).
* Breastfeeding and less than six weeks post-partum.
* Not breastfeeding and less than 6 weeks post-partum with other risk factors for venous thromboembolism (VTE).
* Individuals aged 50 years and over.
* Significant or prolonged immobility.

**Cardiovascular disease*** Individuals aged 35 years and over who currently smoke or stopped smoking less than one year ago (this includes vaping and the use of e-cigarettes).
* Body Mass Index (BMI) equal to or greater than 35kg/m2.
* Blood pressure greater than 140/90mmHg or controlled hypertension.
* Multiple risk factors for cardiovascular disease (CVD) (such as smoking (including vaping/use of e-cigarettes), diabetes, hypertension, obesity, and dyslipidaemias).
* Current or past history of ischaemic heart disease, vascular disease, stroke, or transient ischaemic attack.
* Current or past history of venous thromboembolism.
* Complicated valvular or congenital heart disease, e.g., pulmonary hypertension, history of subacute bacterial endocarditis.
* First degree relative (a person's parent, sibling, or child) with venous thromboembolism which first occurred when they were under 45 years of age.
* Known thrombogenic mutations, e.g., factor V Leiden, prothrombin mutation, protein S, protein C and antithrombin deficiencies.
* Cardiomyopathy with impaired cardiac function.
* Atrial fibrillation.

**Neurological Conditions*** Current or past history of migraine with neurological symptoms including aura at any age.
* Migraine without aura, first attack occurred when on a method of contraception containing an estrogen.
* **Zoely® and desogestrel containing products** – individuals with a meningioma or a history of meningioma.

**Cancers*** Past or current history of breast cancer.
* Undiagnosed breast symptoms (for initiation only)
* Carrier of known gene mutations associated with breast cancer, e.g., BRCA1or 2.
* Malignant liver tumour (hepatocellular carcinoma).

**Gastro-intestinal Conditions*** Viral hepatitis, acute or flare (for initiation only).
* Benign liver tumour (hepatocellular adenoma).
* Severe decompensated cirrhosis.
* Gall bladder disease, currently symptomatic or medically managed.
* Any bariatric or other surgery resulting in malabsorption.
* Cholestasis (related to past combined hormonal contraceptive use).

**Other conditions*** Imminent planned major surgery (COC should be stopped at least 4 weeks prior to planned major surgery or expected period of limited mobility).
* Diabetes with end organ disease (retinopathy, nephropathy, neuropathy).
* Positive anti-phospholipid antibodies (with or without systemic lupus erythematosus).
* Organ transplant, with complications.
* Known severe renal impairment or acute renal failure.
* Acute porphyria.

**Medicines*** Individuals using enzyme-inducing drugs/herbal products or within 4 weeks of stopping them.
* Interacting medicines (other than enzyme inducers) including any medicines or herbal products purchased– see current British National Formulary (BNF) [**http://www.bnf.org**](http://www.bnf.org) or individual product SPC [**http://www.medicines.org.uk**](http://www.medicines.org.uk).
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| **Cautions including any relevant action to be taken** | * If the individual is less than 16 years of age, an assessment based on [**Fraser guidelines**](https://learning.nspcc.org.uk/child-protection-system/gillick-competence-fraser-guidelines#skip-to-content) must be made and documented.
* If the individual is less than 13 years of age, the pharmacist should speak to the local safeguarding lead and follow the local safeguarding policy.
* If there are reasons to believe an individual aged 16 or over lacks capacity, an assessment of capacity to consent should be conducted and recorded in their notes. Particular consideration should be given to any concern of sexual assault or sexual violence in vulnerable adults.
* Discuss with appropriate medical/independent non-medical prescriber any medical condition or medication of which the pharmacist is unsure or uncertain.
* Individuals taking lamotrigine should be advised that COC may interact with lamotrigine; this could result in reduced seizure control or lamotrigine toxicity.
* 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 COC is not contra-indicated it may be less effective and so these individuals should be advised to consider Long-Acting Reversible Contraception (LARC).
* Individuals should be advised that it is possible that medications that induce diarrhoea and/or vomiting (e.g. orlistat, laxatives, GLP-1 agonists) could reduce the effectiveness of COC.
* Individuals receiving GLP-1 agonists must use effective contraception. Note some GLP-1 agonists may reduce the effectiveness of oral contraception and additional barrier methods are recommended - refer to SmPC and [**FSRH advice**](https://www.fsrh.org/Public/Public/Documents/FSRH-statement-Glucagon-like-peptide-1-agonists-and-oral-contraception-Feb-2025.aspx) regarding GLP 1 agonists and contraception. Provide [**FSRH patient information leaflet (PIL).**](https://www.fsrh.org/Public/Public/Documents/FSRH-statement-Glucagon-like-peptide-1-agonists-and-oral-contraception-Feb-2025.aspx)
* **The option of LARC should be discussed with all individuals, in particular those with medical conditions for whom pregnancy presents an unacceptable risk and those on a pregnancy prevention plan. If this option is accepted, individuals should be signposted to where they can access LARC.**
* **If an individual is known to be taking a medication which is known to be harmful to pregnancy, a highly effective form of contraception is recommended. Highly effective methods include the LARC methods: intrauterine device, intrauterine system and implant. If a LARC method is unacceptable/unsuitable and a COC is chosen, then an additional barrier method of contraception is advised. See** [**FSRH advice**](https://www.fsrh.org/standards-and-guidance/documents/fsrh-ceu-statement-contraception-for-women-using-known/)**.**
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| **Action to be taken if the individual is excluded or declines treatment**  | * Explain the reasons for exclusion to the individual and document in the clinical record.
* Record reason for declining treatment in the clinical record.
* Where required, refer the individual to a suitable health service provider, if appropriate, and/or provide them with information about further options.
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1. Description of treatment

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| **Name, strength & formulation of drug** | * See [**Appendix B**](#AppendixB)**.**
* This PGD does not restrict which brands can be supplied – local formularies/restrictions should be referred to. Please refer to your local integrated care board (ICB) formulary for further information.
* See [**http://www.mhra.gov.uk/spc-pil/**](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 effects and interactions. COC containing ≤30micrograms ethinylestradiol in combination with levonorgestrel or norethisterone is a reasonable first-line choice of COC to minimise cardiovascular risk.
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| **Legal category** | POM |
| **Route of administration** | Oral |
| **Off label use** | Best practice advice is given by the FSRH and is used for guidance in this PGD and may vary from the SPC. This PGD includes inclusion criteria, exclusion criteria and dosage regimen which are outside the market authorisation for many of the available products, but which are included within FSRH guidance. Specifically:* the use of tailored COC regimen is outside the manufacturer’s licence but is supported by the FSRH. The regimen detailed within this PGD are permitted under this PGD.

Medicines should be stored according to the conditions detailed in the manufacturers’ guidance. However, in the event of an inadvertent or unavoidable deviation of these conditions the Responsible Pharmacist must be consulted. Where medicines have been assessed by a Responsible Pharmacist, 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 the Responsible Pharmacist.Where a medicine is recommended for off-label use, 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** | FSRH guidance states that COC can either be taken following a standard or tailored regimen. Individuals should be given information about both standard and tailored COC regimen to broaden contraceptive choice.**Monophasic COC products/regimen*** Monophasic COC can either be taken as a standard regimen or in a tailored regimen depending on the choice of the individual.
* The regimens which can be advised are detailed below:

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| **Type of regimen** | **Period of COC use** | **Hormone (pill) free interval** |
| **Standard use** |
| Standard use | 21 days (21 active pills) | 7 days |
| **Tailored use** |
| Shortened hormone-free interval | 21 days (21 active pills) | 4 days |
| Extended use (tri-cycling) | 9 weeks (3x21 active pills) | 4 or 7 days |
| Flexible extended use | Continuous use (≥21 days) of active pills until breakthrough bleeding occurs for 3–4 days | 4 days |
| Continuous use | Continuous use of active pills | None |

* For the monophasic regimen detailed above, a single tablet is to be taken at the same time each day, starting on day 1-5 of the menstrual cycle with no need for additional precautions.
* Individuals should have access to clear information (either written or digital) to support tailored COC use.

**Monophasic everyday, phasic and phasic everyday COC products/regimens** * For monophasic everyday, phasic, and phasic everyday regimens, a single tablet is to be taken at the same time each day, starting on day 1-5 of the menstrual cycle with no need for additional precautions.

The exceptions to this are: * + Qlaira®, which should be started on day 1, or if not, additional precautions should be used for 9 days after starting.
	+ Zoely®, which should be started on day 1, or if not, additional precautions should be used for 7 days after starting.
* Thereafter, follow manufacturer’s instructions for individual product use.

**For all COC products/regimens*** COC can be started at any time after day 5 of the menstrual cycle if it is reasonably certain that the individual is not pregnant. Additional precautions are then required for 7 days after starting (9 days for Qlaira®).
* When starting or restarting the COC as quick start after levonorgestrel emergency contraception, additional contraception is required for 7 days, and a pregnancy test should be performed 21 days after the last unprotected sexual intercourse.
* In line with FSRH guidance, individuals using hormonal contraception should delay restarting their regular hormonal contraception for 5 days following ulipristal acetate use. Avoidance of pregnancy risk (i.e. use of condoms or abstinence from intercourse) should be advised until fully effective. For COC this is 7 days after re-starting this method (9 days for Qlaira®).
* If, in a current user of 30mcg EE/LNG COC, two to seven pills are missed in the first week of pill taking, it may be appropriate to offer UPA-EC and restart COC on a quick starting basis. The individual should be referred to a prescriber in this specific circumstance. See [**FSRH guidance**](https://www.fsrh.org/Common/Uploaded%20files/documents/fsrh-ceu-statement-upa-coc-restart-november-2020.pdf)**.**
* For guidance on changing from one contraceptive method to another, and when to start after an abortion and postpartum, refer to the [**FSRH guidance**](https://www.fsrh.org/standards-and-guidance/documents/fsrh-clinical-guidance-quick-starting-contraception-april-2017/).
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| **Duration of treatment** | * For as long as the individual requires COC, meets the inclusion criteria, and has no contraindications to the use of COC.
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| **Quantity to be supplied**  | * Initiation - **Supply of up to three months** in appropriately labelled original packs.
* Ongoing Supply- **Supply of up to twelve months** in appropriately labelled original packs. **A minimum of six months** should be supplied unless there are clinical reasons not to. Such reasons should be documented in the individual’s clinical record.
* For all supplies, be aware that the regimen to be taken may not be reflected in the dosage information printed on the product packaging or within the supplied PIL – ensure full details of the regimen to be followed are supplied.
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| **Drug interactions** | All concurrent medications and herbal products, including those purchased, should be considered 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**](https://www.medicines.org.uk/emc), the BNF [**www.bnf.org**](http://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/**](https://www.fsrh.org/standards-and-guidance/documents/ceu-clinical-guidance-drug-interactions-with-hormonal/). |
| **Identification & management of adverse reactions** | A detailed list of adverse reactions is available in the individual product SPC, which is available from the electronic Medicines Compendium website: [**www.medicines.org.uk**](http://www.medicines.org.uk) and BNF [**www.bnf.org**](http://www.bnf.org) The following possible adverse effects are commonly reported with COC (but may not reflect all reported adverse effects):* Nausea
* Breast tenderness
* Headache and migraine
* Temporary disturbances of bleeding patterns
* Change in mood, including depression
* Fluid retention
* Change in libido
* Skin changes, including acne

Serious adverse effects - these are less common, but the risks should be discussed with the individual:* Venous thromboembolic events (VTE)
* Arterial thromboembolic events (ATE) (including ischaemic heart disease)
* Strokes (e.g., transient ischaemic attack, ischaemic stroke, haemorrhagic stroke)
* Hypertension
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| **Management of and reporting procedure for adverse reactions** | * Record all adverse drug reactions (ADRs) in the individual’s medical record.
* Pharmacists, pharmacy technicians and individuals/carers are encouraged to report suspected adverse reactions to the Medicines and Healthcare products Regulatory Agency (MHRA) using the Yellow Card reporting scheme: [**http://yellowcard.mhra.gov.uk**](http://yellowcard.mhra.gov.uk)
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| **Management of and reporting procedure for patient safety incidents** | * The pharmacy is required to report any patient safety incidents in line with the [**https://www.gov.uk/government/publications/clinical-governance-approved-particulars**](https://www.gov.uk/government/publications/clinical-governance-approved-particulars).
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| **Written information and further advice to be given to individual**  | * Provide patient information leaflet (PIL) with the original pack.
* Individuals should be informed about the superior effectiveness of LARC.
* Individuals should be provided with written information or a link to a trusted online resource to support safe, effective COC use.
* Explain mode of action, side effects, 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 noting that the risks of using COC could outweigh the benefits.
* **Serious symptoms:** the individual should stop taking the COC and seek medical help urgently if they experience calf swelling, heat or pain in the calf, shortness of breath, chest pain or haemoptysis. The individual should seek advice if they experience their first ever migraine or develops aura with existing migraine.
* Individuals should be advised that current use of COC is associated with a small increased risk of breast cancer which reduces with time after stopping COC.
* Individuals should be advised that current use of COC for more than 5 years is associated with a small increased risk of cervical cancer; risk which reduces over time after stopping COC and is no longer increased by about 10 years after stopping.
* Individuals should be advised that current use of COC is associated with an increased risk of VTE/ATE.
* Individuals using COC should be advised about reducing periods of immobility during travel.
* Individuals trekking to high altitudes (above 4500m or 14500 feet) for periods of more than 1 week may be advised to consider switching to a safer alternative contraceptive method.
* Individuals should be advised to stop COC and to switch to an alternative contraceptive method at least 4 weeks prior to planned major surgery or an expected period of limited mobility.
* Advise on action if vomiting or severe diarrhoea occurs and missed pill advice – see [**FSRH guidance**](https://www.fsrh.org/standards-and-guidance/documents/combined-hormonal-contraception/).
* Advise that non enzyme inducing antibiotics do not interact with COC and if these are prescribed, COC should be continued as normal, with no additional precautions required.
* Depressed mood and depression are well-known reported undesirable effects of hormonal contraceptive use. Depression can be serious and is a well-known risk factor for suicidal behaviour and suicide. Women should be advised to speak to the pharmacist or pharmacy technician where medication was initiated by the pharmacy, or their general practice in case of mood changes and depressive symptoms, appearing shortly after initiation of the treatment.
* Provide [**FSRH PIL**](https://www.fsrh.org/Public/Public/Documents/FSRH-statement-Glucagon-like-peptide-1-agonists-and-oral-contraception-Feb-2025.aspx) on GLP-1 agonists and contraception as appropriate (see cautions).
* Recommend the use of condoms and offer advice on safer sex practices and possible need for screening for sexually transmitted infections (STIs), where appropriate.
* Ensure the individual has contact details of any appropriate local services/sexual health services.
* Advise individual to seek advice from a pharmacist, doctor, or other prescriber before starting any new medications or herbal products, including those purchased.
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| **Advice / follow up treatment** | * The individual should be advised to seek medical advice in the event of an adverse reaction.
* The individual should seek further advice if they have any concerns.
* The individual should be advised on how to obtain future supplies.
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| **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 is 16 years of age or over and not competent, record action taken.
* Name of individual, address, date of birth.
* GP contact details where appropriate.
* Service provided – Initiation or ongoing supply of oral contraception
* Relevant past and present medical and sexual history, medication history (to include over the counter, herbal medications, supplements and recreational drug use), smoking status and family history.
* Results of biometrics and measurements e.g., BMI, blood pressure, height, and weight (if supplied by the individual)
* Any known allergies and nature of reaction.
* Name and registration number of pharmacist or pharmacy technician.
* Name of medication supplied.
* Date of supply.
* Dose amount.
* Quantity supplied.
* Advice given, including advice given and action taken 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 follow up and/or referral arrangements made.
* Any supply outside the terms of the product marketing authorisation.
* Recorded that supply is via PGD.

Records should be signed and dated (or be a password-controlled e-record) and securely kept for a defined period in line with the specification.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 the specification. |

1. Key references

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| **Key references** | * NHS Pharmacy Contraception Service Specification : [**https://www.england.nhs.uk/long-read/nhs-pharmacy-contraception-service/**](https://www.england.nhs.uk/long-read/nhs-pharmacy-contraception-service/)
* Electronic Medicines Compendium [**http://www.medicines.org.uk/**](http://www.medicines.org.uk/)
* Electronic BNF [**https://bnf.nice.org.uk/**](https://bnf.nice.org.uk/)
* NICE Medicines practice guideline “Patient Group Directions” [**https://www.nice.org.uk/guidance/mpg2**](https://www.nice.org.uk/guidance/mpg2)
* Fraser guidelines [**https://learning.nspcc.org.uk/child-protection-system/gillick-competence-fraser-guidelines#skip-to-content**](https://learning.nspcc.org.uk/child-protection-system/gillick-competence-fraser-guidelines#skip-to-content)
* FSRH Clinical Guideline: Combined Hormonal Contraception (January 2019, Amended October 2023) [**https://www.fsrh.org/standards-and-guidance/documents/combined-hormonal-contraception/**](https://www.fsrh.org/standards-and-guidance/documents/combined-hormonal-contraception/)
* FSRH CEU Guidance: Drug Interactions with Hormonal Contraception (May 2022) [**https://www.fsrh.org/standards-and-guidance/documents/ceu-clinical-guidance-drug-interactions-with-hormonal/**](https://www.fsrh.org/standards-and-guidance/documents/ceu-clinical-guidance-drug-interactions-with-hormonal/)
* FSRH UK Medical Eligibility Criteria for Contraceptive Use (UKMEC). (April 2016, Amended September 2019) [**https://www.fsrh.org/Public/Public/Standards-and-Guidance/uk-medical-eligibility-criteria-for-contraceptive-use-ukmec.aspx?hkey=e1816a9c-d7b1-4c64-8130-f6c013b1149a**](https://www.fsrh.org/Public/Public/Standards-and-Guidance/uk-medical-eligibility-criteria-for-contraceptive-use-ukmec.aspx?hkey=e1816a9c-d7b1-4c64-8130-f6c013b1149a)
* FSRH Clinical Guideline: Quick Starting Contraception (April 2017) [**https://www.fsrh.org/standards-and-guidance/documents/fsrh-clinical-guidance-quick-starting-contraception-april-2017/**](https://www.fsrh.org/standards-and-guidance/documents/fsrh-clinical-guidance-quick-starting-contraception-april-2017/)
* FSRH Clinical Guideline: Problematic Bleeding with Hormonal Contraception (July 2015) [**https://www.fsrh.org/standards-and-guidance/documents/ceuguidanceproblematicbleedinghormonalcontraception/**](https://www.fsrh.org/standards-and-guidance/documents/ceuguidanceproblematicbleedinghormonalcontraception/)
* FSRH CEU Statement: Response to Recent Publication Regarding Banh, et al. (November 2020)

[**The effects on ovarian activity of delaying versus immediately restarting combined oral contraception after missing three pills and taking ulipristal acetate 30 mg**](https://www.fsrh.org/Common/Uploaded%20files/documents/fsrh-ceu-statement-upa-coc-restart-november-2020.pdf)* FSRH statement for clinicians on Glucagon-like peptide-1 (GLP-1) agonists and oral contraception (January 2025)

[**https://www.fsrh.org/Public/Public/Documents/FSRH-statement-Glucagon-like-peptide-1-agonists-and-oral-contraception-Feb-2025.aspx**](https://www.fsrh.org/Public/Public/Documents/FSRH-statement-Glucagon-like-peptide-1-agonists-and-oral-contraception-Feb-2025.aspx)* FSRH Patient Information Leaflet on Glucagon-like peptide-1 (GLP-1) agonists and oral contraception (February 2025)

**https://www.fsrh.org/Public/Public/Documents/FSRH-statement-Glucagon-like-peptide-1-agonists-and-oral-contraception-Feb-2025.aspx** |

Appendix A - Registered pharmacist and pharmacy technician authorisation sheet

**PGD combined oral contraceptive pill (COC) Version 3.0**

**Valid from: 20th June 2025 Expiry: 31st March 2026**

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

**Registered pharmacist or pharmacy technician**

By signing this PGD, you are indicating that you agree to its contents and that you will work within it.

PGDs do not remove inherent professional obligations or accountability.

It is the responsibility of each pharmacist or pharmacy technician 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 PGD and that I am willing and competent to work to it within my professional code of conduct.** |
| **Name** | **Designation** | **Signature** | **Date** |
|  |  |  |  |
|  |  |  |  |
|  |  |  |  |
|  |  |  |  |
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|  |  |  |  |

**Authorising manager**

|  |
| --- |
| **I confirm that the registered pharmacists and pharmacy technicians named above have declared themselves suitably trained and competent to work under this PGD. I give authorisation on behalf of insert name of organisation for the above-named pharmacists and pharmacy technicians 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 pharmacists and pharmacy technicians to prevent additions post managerial authorisation.

This authorisation sheet should be retained to serve as a record of those registered pharmacists and pharmacy technicians authorised to work under this PGD.

Appendix B – Name, strength & formulation of drug

Any combined oral contraceptive preparation listed in the British National Formulary (BNF) can be supplied under this PGD. This includes, but may not be limited to the following:

**Monophasic**

| **VMP/AMP Name**  | **VMP/AMP Snomed Code**  | **VMPP/AMPP Snomed Code** | **Supplier Name** |
| --- | --- | --- | --- |
| **Ethinylestradiol 20microgram / Desogestrel 150microgram tablets** | **41876511000001101** | **1235811000001101** |   |
| Bimizza 150microgram/20microgram tablets | 29910811000001108 | 29911111000001107 | Morningside Healthcare Ltd |
| Gedarel 20microgram/150microgram tablets | 17346911000001108 | 17347011000001107 | Gedeon Richter (UK) Ltd |
| Mercilon 150microgram/20microgram tablets | 208311000001105 | 2619611000001105 | Organon Pharma (UK) Ltd |
| **Ethinylestradiol 20microgram / Gestodene 75microgram tablets** | **41876611000001102** | **3049011000001109** |   |
| Akizza 75microgram/20microgram tablets | 38335711000001105 | 38335911000001107 | Morningside Healthcare Ltd |
| Femodette tablets | 3049211000001104 | 3049411000001100 | Bayer Plc |
| Millinette 20microgram/75microgram tablets | 17353311000001100 | 17353411000001107 | Gedeon Richter (UK) Ltd |
| **Ethinylestradiol 30microgram / Desogestrel 150microgram tablets** | **41876811000001103** | **1015611000001103** |   |
| Cimizt 30microgram/150microgram tablets | 21730911000001106 | 21731311000001100 | Morningside Healthcare Ltd |
| Gedarel 30microgram/150microgram tablets | 17348811000001102 | 17349111000001102 | Gedeon Richter (UK) Ltd |
| Marvelon tablets | 524211000001108 | 2620511000001105 | Organon Pharma (UK) Ltd |
| **Ethinylestradiol 30microgram / Drospirenone 3mg tablets** | **41876911000001108** | **1056411000001101** |   |
| Dretine 0.03mg/3mg tablets | 27979911000001107 | 27980011000001108 | Theramex HQ UK Ltd |
| Ellanite 0.03mg/3mg tablets | 34181511000001103 | 34181611000001104 | Kent Pharma (UK) Ltd |
| Lucette 0.03mg/3mg tablets | 23649211000001107 | 23649311000001104 | Gedeon Richter (UK) Ltd |
| Yacella 0.03mg/3mg tablets | 29911411000001102 | 29911511000001103 | Morningside Healthcare Ltd |
| Yasmin tablets | 439011000001108 | 2646411000001106 | Bayer Plc |
| **Ethinylestradiol 30microgram / Gestodene 75microgram tablets** | **41877011000001107** | **3048411000001108** |   |
| Akizza 75microgram/30microgram tablets | 38340211000001108 | 38340311000001100 | Morningside Healthcare Ltd |
| Femodene tablets | 3048811000001105 | 3048911000001100 | Bayer Plc |
| Katya 30/75 tablets | 11753211000001109 | 11753311000001101 | Kent Pharma (UK) Ltd |
| Millinette 30microgram/75microgram tablets | 17351511000001107 | 17351811000001105 | Gedeon Richter (UK) Ltd |
| **Ethinylestradiol 30microgram / Levonorgestrel 150microgram tablets** | **41877111000001108** | **961711000001104** |   |
| Ambelina 150microgram/30microgram tablets | 38738111000001100 | 38738411000001105 | Crescent Pharma Ltd |
| Elevin 150microgram/30microgram tablets | 18358111000001100 | 18358211000001106 | Genesis Pharmaceuticals Ltd |
| Levest 150/30 tablets | 16614111000001103 | 16614411000001108 | Morningside Healthcare Ltd |
| Maexeni 150microgram/30microgram tablets | 24677511000001100 | 24677611000001101 | Lupin Healthcare (UK) Ltd |
| Microgynon 30 tablets | 42111000001107 | 3050411000001108 | Bayer Plc |
| Ovranette 150microgram/30microgram tablets | 492611000001103 | 1974811000001100 | Pfizer Ltd |
| Rigevidon tablets | 17346711000001106 | 17346811000001103 | Gedeon Richter (UK) Ltd |
| **Ethinylestradiol 35microgram / Norethisterone 1mg tablets** | **41877311000001105** | **1082311000001104** |   |
| Norimin 1mg/35microgram tablets | 403611000001106 | 2078211000001102 | Pfizer Ltd |
| **Ethinylestradiol 35microgram / Norethisterone 500microgram tablets** | **41877411000001103** | **1253611000001103** |   |
| Brevinor 500microgram/35microgram tablets | 312411000001108 | 2077911000001105 | Pfizer Ltd |
| **Ethinylestradiol 35microgram / Norgestimate 250microgram tablets** | **4187761100001100** | **1319111000001107** |   |
| Cilique 250microgram/35microgram tablets | 31364011000001106 | 31364211000001101 | Gedeon Richter (UK) Ltd |
| Lizinna 250microgram/35microgram tablets | 22562211000001105 | 22562311000001102 | Morningside Healthcare Ltd |

**Monophasic everyday**

| **VMP/AMP Name** | **VMP/AMP Snomed Code** | **VMPP/AMPP Snomed Code** | **Supplier Name** |
| --- | --- | --- | --- |
| **Drospirenone 3mg / Estetrol 14.2mg tablets** | **41095211000001102** | **41092211000001107** |   |
| Drovelis 3mg/14.2mg tablets | 41092311000001104 | 41092411000001106 | Gedeon Richter (UK) Ltd |
| **Ethinylestradiol 20microgram / Drospirenone 3mg tablets** | **21711311000001107** | **21707111000001101** |  |
| Eloine 0.02mg/3mg tablets | 30195711000001107 | 30196011000001101 | Bayer Plc |
| Microgynon 30 ED tablets | 3052511000001108 | 3052611000001107 | Bayer Plc |
| **Estradiol 1.5mg / Nomegestrol 2.5mg tablets** | **22403311000001100** | **22311311000001109** |  |
| Zoely 2.5mg/1.5mg tablets | 22311511000001103 | 22311711000001108 | Theramex HQ UK Ltd |

**Phasic**

| **VMP/AMP Name** | **VMP/AMP Snomed Code** | **VMPP/AMPP Snomed Code** | **Supplier Name** |
| --- | --- | --- | --- |
| Logynon tablets | 3213311000001106 | 3213811000001102 | Bayer Plc |
| TriRegol tablets | 17444111000001106 | 17444211000001100 | Gedeon Richter (UK) Ltd |
| Synphase tablets | 4432011000001108 | 4433611000001106 | Pfizer Ltd |

**Phasic everyday**

| **VMP/AMP Name** | **VMP/AMP Snomed Code** | **VMPP/AMPP Snomed Code** | **Supplier Name** |
| --- | --- | --- | --- |
| Logynon ED tablets | 3215011000001109 | 3215811000001103 | Bayer Plc |
| Qlaira tablets | 15470011000001100 | 15470111000001104 | Bayer Plc |