Faculty of Family Planning and Reproductive Health Care
Clinical Effectiveness Unit

A unit funded by the FFPRHC and supported by the University of Aberdeen and the Scottish Programme for Clinical Effectiveness in Reproductive Health (SPCERH) to provide guidance on evidence-based practice

FFPRHC Guidance (July 2005)
The use of contraception outside the terms of the product licence

This Guidance provides information for clinicians and women considering the use of contraception outside the terms of the product licence. A key to the grades of recommendations, based on levels of evidence, is given at the end of this document.

Details of the methods used by the Clinical Effectiveness Unit (CEU) in developing this Guidance and evidence tables summarising the research basis of the recommendations are available on the Faculty website (www.ffprhc.org.uk).

Abbreviations (in alphabetical order) used include: CEU, Clinical Effectiveness Unit; COC, combined oral contraception/contraceptive; DMPA, depot medroxyprogesterone acetate; ENG, etonogestrel; IUD, copper-bearing intrauterine contraceptive device; LNG-IUS, levonorgestrel-releasing intrauterine system; NET-EN, norethisterone enantate; PGD, Patient Group Direction; PIL, Patient Information Leaflet; POC, progestogen-only contraception/contraceptive; POEC, progestogen-only emergency contraception; POP, progestogen-only pill; RCT, randomised controlled trial; SPC, Summary of Product Characteristics; UPSI, unprotected sexual intercourse; WHO, World Health Organization; WHOMEC, WHO Medical Eligibility Criteria for Contraceptive Use; WHOSPR, WHO Selected Practice Recommendations for Contraceptive Use.

Background
Summaries of Product Characteristics (SPCs) provide information for clinicians on the licensed indications for contraceptives. The SPCs for most contraceptive methods are readily accessible (http://www.medicines.org.uk). SPCs provide information on how contraceptives should be administered including advice on the dose, when to start, and what to do when pills are missed or injections are late. However, SPCs often do not reflect current evidence and may be unnecessarily restrictive.

Evidence-based recommendations to guide clinicians and women on how to use contraceptive methods safely and effectively have been published by the World Health Organization (WHO). In addition, the Faculty of Family Planning and Reproductive Health Care (FFPRHC) adapted the WHO Selected Practice Recommendations for Contraceptive Use (WHOSPR) (1st edition), using formal consensus methods, for UK use. These Guidance documents provide evidence-based recommendations which at times are at odds with the terms of the product licences for contraceptives (e.g., advice on starting hormone contraception). Nevertheless, the recommendations are made on the basis of systematic review of currently available evidence.

It is unlikely that, in the near future, SPCs will be altered to reflect current evidence. Confusion has resulted from conflicting information for clinicians in various sources such as SPCs, the British National Formulary, and WHO publications. Also, manufacturers’ Patient Information Leaflets (PILs) provide women with information that conflicts with alternative evidence-based sources of patient information such as fpa (Family Planning Association) leaflets.

This evidence-based Guidance provides recommendations on prescribing outside the terms of product licences and provides support for clinicians and women using contraceptive methods outside their licensed indications. Readers are also referred to the FFPRHC Clinical Standards Committee document entitled Medicines Management in Contraception, Sexual and Reproductive Health Care Settings (currently in preparation).

How should clinicians prescribe contraceptives for uses other than those stated in the product licence?

1 Contraceptive, sexual and reproductive health care services should ensure that their organisational policy on use of medicines outside the terms of the product licence is incorporated into clinical practice (Grade C).

2 When prescribing contraceptives outside the terms of the product licence doctors should: take an appropriate clinical history; assess a woman’s priorities and preferences; discuss the evidence supporting use outside licence; document all this information clearly in the case records; and advise the woman of the benefits of informing her general practitioner (GP) (Grade C).

3 Nurse prescribers cannot prescribe medicines outside the terms of the product licence (Grade C).

4 Women should be informed when contraceptives are used outside the terms of the product licence and should be given appropriate and complementary written information in addition to the manufacturers’ Patient Information Leaflets (PILs) (Grade C).

5 Patient Group Directions (PGDs) can be developed to allow nurses and other health care professionals to supply and administer contraceptives. This may include use outside the terms of the product licence provided such use is justified by current best practice. The PDG must clearly describe the status of the product and when it is being used outside licence, and should include the reasons why such use is necessary (Grade C).
6 Pharmacists can dispense medicines outside the terms of the product licence if: the medicine is prescribed by a medical practitioner; in the best interests of the patient; and reasonable steps have been taken to ensure that the prescribing clinician knew that the medicine was to be used outside the terms of the product licence (Grade C).

Definitive guidance on medicine management, The Safe and Secure Handling of Medicines: A Team Approach, was published in 2005, and updated the 1998 Duthie report. This document highlights the need for policies on the safe use and handling of medicines throughout the National Health Service (NHS). Medicine management is an important area of clinical governance. Contraception, sexual and reproductive health care settings should ensure that there are written organisational policies for the use of medicines outside the terms of the product licence. This Guidance may assist in the writing of such policies.

Users have a right to information about all treatment options available to them. When providing information, clinicians should discuss individual needs and priorities that may influence choice. Users should be informed if medicines are being used outside the terms of the product licence and be told about the evidence, or current opinion, to support unlicensed use. To avoid conflicting information, women should be given appropriate and complementary written information that reflects current evidence (such as leaflets from the FPA) in addition to the manufacturers’ PILs.

Medicines that have no licence for use in the UK can be administered, but is usually within the context of a clinical trial and regulated by the Medicines and Healthcare products Regulatory Agency (MHRA). The regulations surrounding the use of unlicensed medicines are outside the scope of this Guidance.

Medicines, including contraceptives, which have a licensed indication in the UK, can be prescribed for the licensed indication but with supplementary instructions for aspects of use that fall outside the product licence. For example, the combined pill can be started up to and including Day 5 of the menstrual cycle without the need for additional contraceptive protection.

With the exception of the levonorgestrel-releasing intrauterine system (LNG-IUS) which is licensed for the management of idiopathic menorrhagia and protection from endometrial hyperplasia with oestrogen replacement therapy, no contraceptive method has been licensed in the UK other than for contraception. Nevertheless, contraceptive methods are commonly prescribed in the management of medical conditions: for example, combined oral contraception (COC) for menorrhagia. There appears to be no distinction in the regulations between prescribing a contraceptive for contraception but with instructions different from those in the product licence and prescribing a contraceptive solely for non-contraceptive benefits.

There are no SPCs for copper-bearing intrauterine contraceptive devices (IUDs) since these are regulated as medical devices rather than medicinal products. The medical devices regulations do not require SPCs. However, under the regulations, medical devices must be ‘CE (Kite) marked’ and therefore copper IUDs (which are CE marked) can be used in the UK without having a SPC. Whilst they do not have ‘a licence’, such devices are required to go through a similar regulatory process, which includes assessment by one of the European Union Medicines Regulatory bodies, such as the MHRA. All IUDs have physician and user instructions.

A nurse prescriber cannot prescribe medications for use outside the terms of the product licence. When prescribing a licensed medication for use outside the terms of the product licence, a clinician must:

- Be satisfied that the use of the medication outside licence meets the patient’s needs better than an appropriately licensed medication.
- Be satisfied that there is sufficient evidence and/or experience of using the medicine to demonstrate its safety and efficacy.
- Take responsibility for prescribing the medication and oversee the patient’s care, monitoring, and any follow-up, or arrange for another doctor to do so.
- Make a clear, accurate and legible record of all medicines prescribed and, when not following common practice, the reasons for prescribing the medicine.

When prescribing any medication a clinician should document in the case record details of information given to the user and the type and dose of medication prescribed. If the prescribing clinician is not the user’s own GP and the user has been seen and treated without a referral, the importance and benefits of keeping the GP informed should be highlighted, and the GP should be informed unless the patient specifically objects.

Patient Group Directions (PGDs) are written instructions for the supply and administration of a medicine (or medicines) where the patient need not be individually identified before presenting for treatment. A PGD can be developed locally by doctors, pharmacists and other health professionals and must meet certain legal criteria. Each PGD must be signed by a doctor or dentist, as appropriate, and a pharmacist, and approved by an appropriate body, usually a primary care or NHS Trust. PGDs can be developed to allow nurses and other health care professionals to supply and administer medicines.

A PGD can include a flexible dose range so that the health care professional can select the most appropriate dose for the patient. A document from the Scottish Executive Health Department on PGDs states that the use of any medicine should be consistent with the SPCs and any relevant guidance. However, medicines used outside the terms of the product licence can be included in PGDs provided such use is exceptional, justified by current best practice, and that such a PGD clearly describes the status of the product. Each PGD should clearly state when the product is being used outside the terms of the product licence and the documentation should include the reasons why this use is necessary.

The Royal Pharmaceutical Society (RPS) provides advice for pharmacists on the supply of medicines outside the terms of their product licence. The RPS states that pharmacists can prepare and dispense medicines outside the terms of the product licence in response to a prescription from a medical practitioner. The pharmacist must ensure that the supply is made in the best interests of the patient, and reasonable steps should be taken to ensure that the prescribing clinician knew that the medicine was to be used outside the terms of the product licence and what the consequences of that might be.

When are contraceptives prescribed outside the terms of the product licence?

Starting contraception outside product licence

Concerns have been raised about starting contraception on any day of the cycle other than Day 1 due to risk of ovulation and subsequent pregnancy. There are wide variations in the timing of ovulation and in menstrual cycle length in normally menstruating women. Studies in small numbers of women have investigated the effects of sex hormones on ovulation, cervical mucus and endometrial receptivity. There is wide inter-individual variation in follicular growth, which can occur even during...
Women can be advised that hormonal contraception (combined hormonal methods, POPs, implants and injectables) can be started at any other time in the cycle if it is reasonably certain they are not pregnant, but additional contraception is required for 7 days (or 2 days for POPs) (Grade C).

9 Women can be advised that the LNG-IUS can be inserted at any time in a cycle if it is reasonably certain that they are not pregnant. Additional contraceptive protection is required for 7 days if inserted after Day 7 (Grade C).

In general, a pragmatic and flexible approach to contraceptive provision should be taken based on current best evidence. Ideally, women should be encouraged to start hormonal contraception [combined hormonal pill and patch: progestogen-only pills (POPs), injectables or implants] on the first day of their menstrual cycle. However, if necessary, hormonal contraception can be started up to and including Day 5 of the menstrual cycle. In this situation, additional contraceptive protection is not required (although some women may still wish to use condoms). If a woman is >5 days after the start of menstruation, hormonal contraception may also be started rather than awaiting her next menstruation if it is reasonably certain she is not pregnant. A clinician can be reasonably sure that a woman is not pregnant if she has no signs or symptoms of pregnancy and meets any of the following criteria:  
- Has not had intercourse since the start of the last normal menses.
- Has been correctly and consistently using a reliable method of contraception.
- Is within 7 days after the start of normal menses.
- Is within 4 weeks postpartum.
- Is within 7 days post-abortion or miscarriage.
- Is fully or nearly fully breastfeeding, amenorrhoeic and <6 months postpartum.

A pregnancy test, if available, adds weight to the diagnosis (but only if >3 weeks since last intercourse).

Starting some hormonal contraceptives (progesterone-only implants and injectables) up to and including Day 5 of the cycle falls within the terms of the product licence. However, for other methods (POPs and combined hormonal methods) this is outside the terms of the product licence. Details of individual contraceptive methods are summarised in this section.

### Combined hormonal contraception

The SPCs for COCs suggest they should be started on Day 1 of the menstrual cycle. If the Day 1 start is missed, some SPCs advise delaying the start until Day 5, and in these circumstances condoms are advised for 7 days. Some SPCs advise condoms for 7 days if starting COCs on any day other than Day 1.

The SPC for the transdermal combined contraceptive patch suggests starting on Day 1. If starting after this time, condoms are advised for 7 days.

Most evidence on risk of ovulation with combined hormonal contraceptives relates to COC. Nevertheless, WHO Medical Eligibility Criteria for Contraceptive Use (WHOMECC) extrapolates all information regarding COCs to the transdermal combined contraceptive patch. COCs act primarily by reducing gonadotrophins, thus inhibiting ovulation. Seven consecutive COCs are required to inhibit ovulation and the remaining COCs maintain anovulation. The COC also has effects on cervical mucus and the endometrium, which contribute to
contraceptive efficacy. Studies in primates have shown that the COC can effectively inhibit ovulation even when started up to and including Day 5.
### Table 2 Progestogen-only contraception: licensed and unlicensed use

<table>
<thead>
<tr>
<th>Starting time</th>
<th>Licensed use as in the Summary of Product Characteristics (SPC)</th>
<th>Evidence-based use outside the terms of the product licence</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Progestogen-only pills (POPs)</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>First use</td>
<td>Start on Day 1; additional method required for 7 or 14 days.</td>
<td>Ideally start on Day 1 of the menstrual cycle. Can start to Day 5 without additional method. Start after Day 5 if it is reasonably certain* the woman is not pregnant; additional method for 2 days.</td>
</tr>
<tr>
<td>Following abortion or miscarriage</td>
<td>Start immediately following first-trimester abortion or miscarriage. (No advice given for second trimester.)</td>
<td>Start on day of surgical or second part of medical abortion (induced or spontaneous &lt;24 weeks); no additional method required. If started &gt;7 days after abortion an additional method required for 2 days.</td>
</tr>
<tr>
<td>Postpartum not breastfeeding</td>
<td>Start on Day 21 postpartum; no additional method required.</td>
<td>Contraception is not required before Day 21 postpartum. Ideally start on Day 21 postpartum for immediate contraceptive protection. Can be started before Day 21 if requested. If started after Day 21, additional method required for 2 days.</td>
</tr>
<tr>
<td>Breastfeeding</td>
<td>No advice given. Micronor® advice suggests waiting until infant weaned.</td>
<td>Start on or before Day 21 postpartum for immediate contraceptive protection. If started after Day 21, additional method required for 2 days.</td>
</tr>
<tr>
<td>Missed pills</td>
<td>&gt;27 hours since last pill (i.e. &gt;2 hours late). Additional method required for 7 or 14 days.</td>
<td>&gt;27 hours since last pill (i.e. &gt;3 hours late). The missed pill should be taken as soon as remembered; continue daily pill taking; additional method required for 2 days. EC is indicated if UPSI has occurred in the 2 days since missing pills.</td>
</tr>
<tr>
<td></td>
<td>Desogestrel pill is missed if &gt;36 hours since last pill (i.e. 12 hours late). Additional method required for 7 days.</td>
<td>Desogestrel-only pill is ‘missed’ if &gt;36 hours since last pill (i.e. 12 hours late). Missed pill should be taken as soon as remembered; continue daily pill taking; an additional method required for 2 days. EC is indicated if UPSI has occurred in the 2 days since missing POs.</td>
</tr>
<tr>
<td><strong>Progestogen-only implants</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>First use</td>
<td>Initiate between Days 1 and 5. (No mention of additional methods.)</td>
<td>Can start up to and including Day 5 without additional method. Start after Day 5 if it is reasonably certain* the woman is not pregnant; additional method for 7 days.</td>
</tr>
<tr>
<td>Following abortion or miscarriage</td>
<td>Start immediately following a first-trimester abortion. Following a second-trimester abortion, start between Days 21 and 28.</td>
<td>Start on day of surgical or second part of a medical abortion (induced or spontaneous &lt;24 weeks); no additional method required. If started &gt;7 days after abortion an additional method required for 2 days.</td>
</tr>
<tr>
<td>Postpartum not breastfeeding</td>
<td>Start between Days 21 and 28; no need for additional method.</td>
<td>Start between Days 21 and 28 postpartum for immediate contraceptive protection. If started after Day 28, additional method required for 7 days. Can be started before Day 21 but problematic bleeding can occur.</td>
</tr>
<tr>
<td>Breastfeeding</td>
<td>Start between Days 21 and 28; no need for additional method.</td>
<td>Start between Days 21 and 28 postpartum for immediate contraceptive protection. If started after Day 28, additional method required for 7 days.</td>
</tr>
<tr>
<td><strong>Progestogen-only injectables</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>First use</td>
<td>Start in the first 5 days of menstrual cycle without additional method.</td>
<td>Ideally start on Day 1 of the cycle. Can start up to and including Day 5 without additional method. Start after Day 5 if it is reasonably certain* the woman is not pregnant; additional method for 7 days.</td>
</tr>
<tr>
<td>Following abortion or miscarriage</td>
<td>For DMPA no advice given. NET-EN can be used immediately after abortion, no advice is given for miscarriage.</td>
<td>Start on day of surgical or second part of a medical abortion (induced or spontaneous &lt;24 weeks) no additional method required. If started &gt;7 days after abortion an additional method required for 7 days.</td>
</tr>
<tr>
<td>Postpartum not breastfeeding</td>
<td>Give DMPA within 5 days postpartum. Prolonged heavy bleeding can occur. NET-EN can be used immediately after birth.</td>
<td>Give DMPA within 5 days postpartum. Prolonged heavy bleeding can occur. NET-EN can be used immediately after birth.</td>
</tr>
<tr>
<td>Breastfeeding</td>
<td>DMPA should be delayed until at least 6 weeks postpartum. NET-EN can be used immediately after birth, although it is advised that breastfeeding should be withheld from neonates with severe or persistent jaundice requiring medical treatment.</td>
<td>Ideally start DMPA from 6 weeks postpartum. If other methods are unacceptable, then injection may be given &lt;6 weeks postpartum but ideally not before Day 21.</td>
</tr>
<tr>
<td>Repeat and late injections</td>
<td>DMPA should be given every 12 weeks. As long as injections are given no later than 5 days after due date no additional method is required. If the interval from the last injection is greater than 89 days (12 weeks + 5 days) then pregnancy should be excluded before the next injection is given and additional method used for 14 days.</td>
<td>Repeat DMPA should be given every 12 weeks and repeat NET-EN every 8 weeks. DMPA can be given up to 2 weeks early (10 weeks) or 2 weeks late (10 weeks) without the need for additional method. If DMPA and NET-EN are &gt;2 weeks late, contraceptive protection may be lost and protocols should be developed locally to guide women and clinicians in deciding on need for EC and timing of next injection.</td>
</tr>
</tbody>
</table>

*Reasonably certain the woman is not pregnant, has not had intercourse since the last normal menses; has been correctly and consistently using a reliable method of contraception; is within 7 days after normal menses; is within 4 weeks postpartum; is within 7 days post-abortion or miscarriage; is fully or nearly fully breastfeeding; amenorrhoeic and >6 months postpartum. Pregnancy testing adds weight to the diagnosis. DMPA, depot medroxyprogesterone acetate; EC, emergency contraception; NET-EN, norethisterone enantate; POP, progestogen-only pill; UPSI, unprotected sexual intercourse.
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penetrability were significantly reduced 12 hours after administration. Within 2 days of initiating POPs, cervical mucous changes and sperm penetration into the upper reproductive tract.

The SPCs for POPs containing levonorgestrel, norethisterone and etynodiol diacetate suggest starting on Day 1.37-42 Some SPCs suggest condoms for 7 days if starting on Day 1. Others suggest condoms for 7 days if starting after Day 1.43

The desogestrel-only pill works primarily by inhibiting ovulation.44 In addition, as for other POPs, there is thickening of cervical mucus and endometrial changes.45 Recent randomised controlled trials (RCTs) comparing the desogestrel-only pill to a levonorgestrel-only pill have shown that ovulation is inhibited in 97% and 60% of cycles, respectively.46 The SPC for the desogestrel-only POP suggests that it has to be started on Day 1 and that additional contraceptive protection is required for 7 days if starting after Day 1.47

Women should be encouraged to start all POPs on the first day of the menstrual cycle. However, as a result of the rapid effect on cervical mucus, it is unlikely that a Day 5 POP start in women with normal menstrual cycles would compromise contraceptive efficacy. The CEU supports recommendations from WHO3-4 that, if necessary, POPs can be started up to and including Day 5 without the need for additional contraceptive protection (unlicensed). In addition, POPs can be started after this time if it is reasonably certain that the woman is not pregnant. In this situation the CEU advise that additional contraceptive protection is required for 2 days (unlicensed) (Table 2).3

Manufacturers’ PILs do not support this advice. The CEU suggests that fpa leaflets, which do reflect this current advice, are given to women in addition to the PIL.

The progestogen-only implant

Serum etonogestrel (ENG) concentrations rapidly rise within 8 hours of insertion of the etonogestrel-only implant (Implanon®). Serum concentrations of ENG at 8 hours post-insertion (265.9 pg/ml) are higher than concentrations identified after 1 year of use (196 pg/ml) when ovulation is notably rare.48 Ovulation is likely to be inhibited quickly following insertion. The SPC advises initiation between Days 1 and 5 of the cycle.39

The WHO/SPR supports initiation of the progestogen-only implant up to and including Day 7 without the need for additional contraceptive protection.5 Nevertheless, the CEU supports the FFPHRHC UK version of this document which suggests, as for other hormonal contraceptive methods, that when necessary progestogen-only implants can be inserted up to and including Day 5 (licensed) without the need for additional contraceptive protection.5 The progestogen-only implant can be inserted at other times in the cycle if it is reasonably certain that the woman is not pregnant but additional contraceptive protection is advised for 7 days following insertion (unlicensed) (Table 2).3

The levonorgestrel-releasing intrauterine system

The LNG-IUS is licensed for contraception, the management of idiopathic menorrhagia and protection from endometrial hyperplasia with oestrogen replacement therapy.54 The main contraceptive effect of the LNG-IUS is due to local effects on the endometrium from 20 μg levonorgestrel released every 24 hours.53 Ovulatory cycles with follicular rupture usually occur. Functional and histological changes are evident within 1 month of insertion.56

The SPC suggests insertion of the LNG-IUS within 7 days of the onset of menstruation.54 No mention is made of additional contraceptive protection if inserted during this time. The CEU supports the WHO/SPR recommendations that the LNG-IUS can be inserted at any time in the first 7 days after the start of menstruation and not only during menstruation (licensed).5 If inserted up to and including Day 7 no additional contraceptive protection is required. If necessary the LNG-IUS can be inserted at any other time in the cycle, at the woman’s convenience, if it is reasonably certain that she is not pregnant (unlicensed).2 In this situation additional contraceptive protection is required for 7 days (Table 3).

Starting non-hormonal contraception

10 Women can be advised that an IUD can be inserted at any time in the menstrual cycle if it is reasonably certain she is not pregnant (Grade C).
Table 3  Intracutaneous contraception: licensed and unlicensed use

<table>
<thead>
<tr>
<th>Starting time</th>
<th>Licensed use as in the Summary of Product Characteristics (SPC) or physician information</th>
<th>Evidence-based use outside the terms of the product licence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Levonorgestrel-releasing intrauterine system (LNG-IUS)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>First use</td>
<td>Start in the first 7 days of the menstrual cycle no additional method required.</td>
<td>Ideally insert in the first 7 days of the menstrual cycle; no additional contraceptive protection required. Can be started at any time in the cycle if reasonably certain* the woman is not pregnant; an additional method is required for 7 days.</td>
</tr>
<tr>
<td>Following abortion or miscarriage</td>
<td>Can start immediately following abortion. No mention of miscarriage or second-trimester abortion.</td>
<td>Insert at the time of surgical or second part of a medical abortion (induced or spontaneous &lt;24 weeks); no additional method required. Otherwise delay insertion until 4 weeks post-abortion.</td>
</tr>
<tr>
<td>Postpartum not breastfeeding</td>
<td>Should delay insertion until 6 weeks postpartum. No mention of additional method.</td>
<td>Insert within first 48 hours or delay insertion until 4 weeks postpartum regardless of mode of delivery. Additional method advised for 7 days when inserted from 4 weeks postpartum.</td>
</tr>
<tr>
<td>Breastfeeding</td>
<td>Should delay insertion until 6 weeks postpartum. No mention of additional method.</td>
<td>Can be inserted from 4 weeks postpartum.</td>
</tr>
<tr>
<td>Copper intrauterine device (IUD)*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>First use</td>
<td>Can start an IUD in the first 7 days of the menstrual cycle with no additional contraception required.</td>
<td>Can be inserted at any time in the cycle if it is reasonably certain* she is not pregnant; no additional contraceptive protection is required.</td>
</tr>
<tr>
<td>Following termination of pregnancy or miscarriage</td>
<td>Insert immediately following first-trimester TOP. No mention of miscarriage or second-trimester TOP.</td>
<td>Insert at the time of surgical abortion or on the day of the second part of a medical abortion (induced or spontaneous &lt;24 weeks); no additional contraception required. Otherwise delay insertion until 4 weeks post-abortion; no additional contraceptive protection required.</td>
</tr>
<tr>
<td>Postpartum not breastfeeding</td>
<td>Delay insertion of an IUD until 6 weeks postpartum. No additional contraception required.</td>
<td>Start immediately postpartum and within first 48 hours or delay insertion until 4 weeks postpartum regardless of mode of delivery. Additional contraception is advised for 7 days when inserted from 4 weeks postpartum.</td>
</tr>
<tr>
<td>Breastfeeding</td>
<td>No mention of use in breastfeeding women</td>
<td>Start immediately postpartum and within first 48 hours or delay insertion until 4 weeks postpartum regardless of mode of delivery.</td>
</tr>
<tr>
<td>Emergency contraception</td>
<td>Advice unclear</td>
<td>Can be inserted up to 5 days after the first episode of UPSI. Can be inserted up to 5 days after the earliest expected date of ovulation (up to Day 19 in a regular 28-day cycle) regardless of the number of episodes of UPSI.</td>
</tr>
</tbody>
</table>

*Reasonably certain the woman is not pregnant: has not had intercourse since the last normal menses; has been correctly and consistently using a reliable method of contraception; is within 7 days after normal menses; is within 4 weeks postpartum for non-lactating women; is within 7 days post-abortion or miscarriage; is fully or nearly fully breastfeeding, amenorrhoeic and <6 months postpartum. Pregnancy testing adds weight to the diagnosis. Copper IUDs do not have to go through the same processes as other contraceptive methods to be licensed for use in the UK. If they have a CE Kitemark they can be used.

SPCs exist for LNG-IUS but are limited for copper IUDs. Physician and user information accompanies these devices. It is not entirely clear if use of IUD contraception required. Otherwise delay insertion until 4 weeks post-abortion; no additional contraceptive protection required.

**The copper-bearing intrauterine contraceptive device**

The copper IUD works primarily by inhibition of fertilisation due to the toxic effects of copper on ovum and sperm.57–59 A systematic review indicated that both pre-fertilisation (toxic effects) and post-fertilisation effects (preventing implantation) contribute to efficacy.55 As a result of this toxic effect, the copper IUD can work immediately after insertion. If fertilisation has already occurred, an IUD can be inserted to prevent implantation, its secondary mode of action.

Product information for the Nova-T® 380 (as an example) suggests insertion during or shortly after menstruation, and if pregnancy is excluded it may be inserted any time in the cycle.60 The CEU supports the WHOSPR and FFPRHC UK version which suggests that an IUD can be inserted at any time in the cycle if it is reasonably certain that the woman is not pregnant (Table 3).2,3

A copper IUD can be used as emergency contraception (EC) and inserted up to 5 days after the first episode of unprotected sexual intercourse (UPSI).59 An IUD can also be inserted as EC up to 5 days after the expected date of ovulation in a woman with a regular menstrual cycle (i.e. up to and including Day 19 in a regular 28-day cycle) (Table 3).2,51

### Starting contraceptives in special circumstances

Advice on starting hormonal contraception in special circumstances [such as following spontaneous or induced abortion, postpartum, when breastfeeding or following use of progestogen-only emergency contraception (POEC)] is covered in this section. Some advice was published by the CEU in previous Guidance and is summarised in Tables 1–3.62

#### Following miscarriage or abortion (<24 weeks’ gestation)

11 Women can be advised that all contraceptive methods can be started immediately following miscarriage or induced abortion occurring at <24 weeks’ gestation (Grade C).

Return of fertility is rapid following first-trimester abortion and can occur as soon as 2 weeks following abortion.63 SPCs do not give advice on starting COCs following...
second-trimester abortion. The SPC for the transdermal combined contraceptive patch suggests it can be started immediately following abortion at <20 weeks' gestation to provide immediate contraceptive protection. The SPC for the progestogen-only implants suggests that an implant can be inserted immediately following first-trimester abortion but should be delayed until between Days 21 and 28 following second-trimester abortion. The SPC for NET-EN suggests it can be started immediately following abortion but does not specify gestation.

Advice varies for POPs for women following abortion. Some POPs can be started immediately following first-trimester abortion but condoms are advised for 14 days. The desogestrel-only pill can be started immediately following first-trimester abortion with no need for condoms and can be started following second-trimester abortion “before menstruation”. Some POPs can be started the day after abortion with immediate contraceptive protection. The SPC for DMPA suggests it can be administered immediately following abortion although heavy or prolonged bleeding may occur. SPCs do not state whether this refers to first- and/or second-trimester abortion. The SPC for NET-EN suggests it can be started immediately following abortion but does not specify gestation.

There is little evidence or biological plausibility to suggest that use of progestogen-only methods increases the risk of venous thromboembolism (VTE). There appears to be no reason to delay insertion following second-trimester abortion. The SPC for the LNG-IUS suggests it can be inserted immediately after first-trimester surgical abortion. No mention is made of medical termination or abortion at other gestations. The information for the Nova-T 380 supports insertion immediately after first-trimester abortion. Insertion of an IUD immediately following induced abortion has advantages in that the woman is known not to be pregnant, is highly motivated and is already attending a health care setting. Systematic reviews (which included devices not currently in use) found that IUD insertion at the time of surgical abortion is safe and practical. Expulsion rates were higher following insertion at the time of a second-trimester abortion.

The WHO/SPR supports use of COC, POP, progestogen-only injectables and implants immediately after abortion without the need for additional contraceptive protection (although gestation is not specified). The WHO/SPR also supports insertion of an IUD or the LNG-IUS immediately after first-trimester abortion and generally immediately following second-trimester abortion.

The CEU suggests that the advice is the same for induced or spontaneous abortion at <24 weeks' gestation. By ‘immediately following abortion’ the CEU refers to the day of the surgical abortion procedure or the day of the second part of a medical abortion procedure. The CEU suggests that combined hormonal methods (pills and patch) and progestogen-only methods (pills, injectables and implants) should be started immediately or within 7 days following abortion (induced or spontaneous) at gestations <24 weeks to provide immediate contraceptive protection. If started after Day 7, additional contraceptive protection is advised for 7 days (or 2 days for POP). The CEU advises that an IUD or the LNG-IUS can be inserted immediately following abortion at <24 weeks' gestation.

### Postpartum (not breastfeeding)

12 Contraception is not required before Day 21 postpartum. Ideally, hormonal contraception should be started on Day 21 to provide immediate contraceptive protection. However, progestogen-only methods can be started before Day 21 if requested (Grade C).

13 All intrauterine methods can be inserted from 4 weeks postpartum; they can also be inserted within 48 hours of delivery (Grade C).

The earliest date of ovulation postpartum is believed to be Day 28; with contraceptive methods initiated on Day 21 providing immediate contraceptive protection. A small, prospective, comparative study included 22 postpartum non-breastfeeding women who had daily urine assays performed. All women menstruated within 12 weeks postpartum and first ovulation occurred on average 45 days postpartum. Another small longitudinal study investigated 30 postpartum women with assessment of salivary progesterone and found that the delivery to menstruation interval was 57 (± 7) days and that ovulation occurred on average at Day 23 (± 7) days.

The SPCs suggest starting COC from Day 21. The SPC for the transdermal contraceptive patch, however, suggests it should not be started until at least 4 weeks postpartum. The SPC's suggest some POPs can be started 7 days after a vaginal delivery if ambulant, with immediate contraceptive protection. Other SPCs suggest starting on Day 21 to provide immediate protection. If starting after Day 21, condoms are advised for 7 days. The SPC for the progestogen-only implant advises insertion between Days 21 and 28 postpartum. If starting COC from Day 21, SPCs suggest starting 7 days after a vaginal delivery if ambulant, with immediate contraceptive protection.

14 Breastfeeding women can be advised that: combined hormonal contraception can be used from 6 weeks to 6 months postpartum if no other method is acceptable; POPs can be started before Day 21 postpartum but if started after Day 21 additional contraceptive protection is required for 2 days; progestogen-only implants can be inserted before Day 21 postpartum but bleeding may be a problem; progestogen-only injectables can be given <6 weeks postpartum but should ideally be delayed until Day 21 postpartum (Good Practice Point).
‘Full breastfeeding’ can be ‘exclusive’ (when no other liquids or solids are given) or ‘almost exclusive’ (when milk, water or juice are given infrequently).62,69 Women who are fully breastfeeding, amenorrhoeic and ≤6 months postpartum can rely on the lactational amenorrhoea method to provide contraception with over 98% efficacy.62,69 Additional contraception is required when breastfeeding frequency decreases (particularly night feeds), menstruation recurs or when >6 months postpartum.62 ‘Partial’ or ‘token’ breastfeeding can vary from ‘high’ (where the vast majority of feeds are breastfeeds) to ‘minimal’ (where there are occasional irregular breastfeeds); this has less or little impact on fertility and additional contraception is advised.62

SPCs suggest combined hormonal methods should be avoided when breastfeeding. SPCs provide limited information on starting POPs in women who are breastfeeding. All highlight minimal effects on breast milk. For breastfeeding women, the use of POP before 6 weeks postpartum may be outside product licence.

The SPC suggests that the progesterone-only implant can be used by breastfeeding women.49 However, there is no recommendation on when to initiate this method. The SPC for DMPA suggests that it can be administered from 6 weeks postpartum if breastfeeding.58

The CEU has published Guidance on the use of contraceptives by breastfeeding women.62 Combined hormonal contraception should be avoided in the first 6 weeks postpartum due to effects on breast milk and infant growth. Combined hormonal contraception can be used without restriction from 6 months. Although use of combined hormonal contraception is not recommended between 6 weeks and 6 months, if feeding is established and other methods are unacceptable it can be considered (unlicensed).62

Studies have not supported a detrimental effect on breast milk or infant growth with POP use62 and therefore breastfeeding women can start POP up to Day 21 postpartum (outside licence for some POPs) without the need for additional contraception protection. If started after Day 21, additional contraceptive protection is required for 2 days (unlicensed).62

The CEU suggests delaying initiation of the progesterone-only implant until Day 21 postpartum if breastfeeding. However, if a woman wishes to have the progesterone-only implant inserted before Day 21 there are no data to suggest that this adversely affects breast milk or infant growth. Women should be advised that bleeding may occur.62

If breastfeeding women choose the progesterone-only injectable, this can be given before 6 weeks postpartum. However, initiation should ideally be delayed until Day 21 postpartum (unlicensed).62

Starting contraception following use of POEC

15 After POEC use, clinicians and women should discuss and consider individually the option of initiating a regular method of contraception prior to the onset of the next menstruation (Good Practice Point).

The SPC for POEC suggests that its use does not contraindicate the continuation of a regular method of contraception.70 The CEU has recommended previously that POEC does not provide effective contraceptive cover for the remainder of the cycle and effective contraception or abstinence must be advised.61 The WHOSPR suggests that contraceptive methods can be started at any time in the cycle if it is reasonably certain that the woman is not pregnant.2 It would be reasonable to consider initiating a method of contraception following POEC use if abstinence is unlikely or further condom failure or UPSI is a risk. Decisions on starting a method following POEC use should be considered on an individual basis. Following missed pills, women should be advised to resume hormonal contraception at their usual time as long as this is within 12 hours of taking POEC.61 It may be appropriate to initiate hormonal contraception, which can be discontinued should POEC fail and a pregnancy occur. Reassuringly, no hormonal method has been shown to be associated with an increased risk of fetal abnormality. Long-term options such as progesterone-only injectables and implants may be best delayed until the next menses as the associated amenorrhoea may delay the diagnosis of pregnancy. Nevertheless, there may be individual situations where they may be started and the woman followed up for a pregnancy test.

When hormonal contraception is late or missed

16 Women can be advised that if one or two 30–35 µg ethinylestradiol (EE)-containing COCs are missed (or one 20 µg EE-containing COC): the last missed pill should be taken as soon as remembered; pills should be continued daily at the usual time; additional contraceptive protection is not required; and EC is not indicated (Grade C).

17 Women can be advised that if three 30–35 µg EE-containing COCs are missed (or two 20 µg EE-containing COCs) the last missed pill should be taken as soon as remembered; an emergency pill taking continued. Additional contraceptive protection, such as condoms, is advised for 7 days. EC is only indicated if pills are missed in Week 1 and UPSI occurred in the pill-free week or in Week 1. If pills are missed in Week 3, the pill-free interval should be omitted (Grade C).

SPCs suggest that a COC is ‘late’ if >36 hours have elapsed since the last COC was taken (i.e. >12 hours behind schedule). However, ‘missed pill rules’ were updated in the WHOSPR2 and have been endorsed by the FFPHRHC71 and fpa (summarised in Table 1 and Figure 1). A ‘missed pill’ is a pill that a woman completely omits to take (i.e. 48 hours have elapsed since the last COC was taken).

COCs reduce gonadotrophins, thus inhibiting ovulation.18 Seven consecutive pills are regularly missed in the pill-free interval without losing contraceptive protection. Follicular activity is evident in the pill-free week or in Week 1. If pills are missed in Week 3, the pill-free interval should be omitted (Grade C).

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COCs (Figure 1).

Combined contraceptive patch

Evidence suggests that a combined contraceptive patch can be used for up to 9 days without reducing contraceptive efficacy. If a patch is used for up to an additional 48 hours, no additional contraceptive protection is required if a new patch is then applied. If, however, this is extended beyond 48 hours, additional contraceptive protection is required for 7 days when the new patch is applied. When the patch-free interval is extended, SPCs suggest the woman may not be protected from pregnancy. A new patch should be applied and additional contraceptive protection used for 7 days. In addition, if UPSI occurred during the extended patch-free interval, risk of fertilisation should be considered. These recommendations are within the terms of the product licence.

Progestogen-only pill

18 Women taking POPs containing levonorgestrel, norethisterone or etynodiol diacetate can be advised that if >27 hours have elapsed since taking the last pill (i.e. >3 hours late): the late pill should be taken as soon as remembered; the next pill should be taken at the usual time; additional contraceptive protection such as condoms is advised for 2 days; and if UPSI has occurred in the 2 days since late pill, EC is indicated (Grade C).

19 Women taking desogestrel-only pills can be advised that if >36 hours have elapsed since taking the last pill (i.e. >12 hours late): the late pill should be taken as soon as remembered; the next pill should be taken at the usual time; additional contraceptive protection such as condoms is advised for 2 days; and if UPSI has occurred in the 2 days since late pills, EC is indicated (Grade C).

SPCs for POPs containing levonorgestrel, norethisterone or etynodiol diacetate suggest if >3 hours late (i.e. 27 hours since taking last POP) additional contraceptive protection is required for 7 days. The SPC for the desogestrel-only pill suggests that a desogestrel-only pill is late when >36 hours have elapsed since the last pill was taken (i.e. 12 hours late). But, similarly, additional contraceptive protection is advised for 7 days. The desogestrel-only pill should be treated as other POPs when giving women advice on additional contraceptive protection or need to consider EC. However, this advice does not need to be followed until 12 hours late (i.e. >36 hours have elapsed since the last pill was taken).

Studies have suggested that after stopping desogestrel-only pills the minimum time to first ovulation is 7 days and on average is 17.2 days. An RCT showed that desogestrel-only pills inhibit ovulation in up to 97% of cycles. Therefore, there are women in whom ovulation can occur and cervical mucus thickening is providing contraceptive efficacy. POPs thicken cervical mucus and this effect is restored within 48 hours of taking a POP, thus additional contraceptive protection is required for only 2 days following missed or late POPs. EC is indicated if UPSI occurred during these 2 days before cervical mucus effect has peaked (Table 2).

Late progestogen-only injectables

20 Should a woman present late for her next contraceptive injection (i.e. >12 weeks to <14 weeks for DMPA, and >8 weeks to <10 weeks for NET-EN) the risk of pregnancy is extremely small. Additional contraceptive protection need not be recommended (Grade C).

The SPC for NET-EN suggests contraceptive efficacy is provided for 8 weeks. The SPC for DMPA advises that DMPA should be given every 12 weeks. As long as injections are given no later than 5 days after the due date, no additional contraceptive protection is recommended. If the interval from the last injection is greater than 89 days (i.e. 12 weeks and 5 days) pregnancy should be excluded before the next injection is given and additional contraceptive protection used for 14 days.

A small study in three women observed serum medroxyprogesterone concentrations and ovarian activity (by serum progesterone and LH) following a single DMPA injection. Concentrations of medroxyprogesterone gradually declined and remained relatively constant at 1 ng/ml for 2–3 months, declining thereafter to 0.2 ng/ml by the sixth month. Concentrations were undetectable at 7–9 months. Serum oestradiol remained at early to mid-follicular levels for 4–6 months after the injection. Oestradiol was raised to ovariatory concentrations only when serum medroxyprogesterone concentrations had declined to below 0.25–0.5 ng/ml. In some women, MPA can be identified in serum for as long as 9 months after injection. This would suggest that the DMPA can be given late without the need for additional contraceptive protection as the risk of ovulation is low.

In addition, there is abundant evidence of a delay in return to fertility after discontinuation of DMPA. A total of 796 Thai women stopped using DMPA and 125 had an IUD removed. All were followed up for 2 years. For women discontinuing an IUD the median delay to conception was 4.5 months. For women discontinuing DMPA the median delay was 5.5 months plus the estimated duration of the effect of the last injection, therefore a delay to conception of approximately 9 months can occur in some women.
The CEU recommends that the progesterone-only injectable DMPA should be given every 12 weeks and NET-EN every 8 weeks. An injection is ‘late’ when >12 weeks (for DMPA) or >10 weeks (for NET-EN) have elapsed since the last injection. However, despite late injections, effective contraception is still provided for up to 14 weeks (for DMPA) and to 10 weeks (for NET-EN). After this time, contraceptive protection may be reduced. The CEU supports WHOSPR recommendations that progesterone-only injectables can be given up to 2 weeks late (i.e. an interval of 14 weeks for DMPA and of 10 weeks for NET-EN) without the need for additional contraceptive protection.2 Beyond 2 weeks late, protocols for women who attend for progesterone-only injectables should be developed locally and take account of factors such as timing of any UPSI, duration of injectable use, and the presence or absence of amenorrhoea.

When the dose of oral contraceptives is increased

21 Women taking liver enzyme-inducing drugs who wish to use COC should choose a regimen containing at least 50 µg EE daily. Additional contraceptive protection, such as condoms, should be used until 4 weeks after the liver enzyme-inducing drug has been stopped (Grade C).

22 Women using POPs containing levonorgestrel, norethisterone or etynodiol diacetate who weigh >70 kg are advised to take two pills together every day (Good Practice Point).

Liver enzyme-inducing drugs

It is established UK practice to start a COC with a regimen containing at least 50 µg EE daily for women taking liver enzyme-inducing drugs.84–86 The most commonly used COC containing 50 µg EE (Ovran®, Wyeth Laboratories) was discontinued in 2002. An alternative preparation containing 50 µg mestranol (Norinyl-1®, Pharmacia) is available. Two studies have provided conflicting evidence on the bioequivalence of 50 µg EE and mestranol.87,88 Although inter-individual variation may occur, 50 µg EE and mestranol have been shown to be comparable.58 Liver enzyme-inducers increase the metabolism of oestrogen and progesterone99 and therefore an increased dose is required. The CEU suggests a regimen of two low-dose COCs (e.g. Loestrin 20®, plus Loestrin 30®; Marvelon® plus Mericon®, or two Microgynon 30®), providing a total daily dose of 50–60 µg EE but no trials have compared bioavailability to that of a single tablet. This use is unlicensed. Shortening the pill-free interval has also been advised.

Guidance from the CEU summarised potential drug interaction with liver enzyme-inducers and methods of hormonal contraception.96 Methods such as POPs and progesterone-only implants are not advised for women using liver enzyme-inducing drugs, while progesterone-only injectables, which are less affected.97

Weight over 70 kg

Some health professionals in the UK advise women weighing >70 kg using POPs containing levonorgestrel, norethisterone or etynodiol diacetate to take two pills together every day. Direct evidence to support this practice was not identified. Studies show that obese women who use levonorgestrel implants or vaginal rings91,92 have higher failure rates than non-obese women. For women using the etonogestrel-only implant, anovulation occurs in 100% of women. The levonorgestrel-only implant is less reliable in ovulation inhibition, especially in the first years of use. Data may be extrapolated to the effect on weight and POPs containing levonorgestrel, norethisterone or etynodiol diacetate (which are also less reliable at ovulation inhibition). Data from women recruited to an RCT and a non-comparative study were investigated.93 Cumulative pregnancy rates differed significantly with body weight at 5 and 7 years of use. Women weighing ≥80 kg experienced relatively higher failure rates. In a multinational study, 1198 women were randomised to Norplant® or Norplant II®.94 Pregnancies only occurred in Year 5 of use and both pregnancies in women weighing >70 kg. Thus women weighing >70 kg were more likely to have a method failure than women weighing <70 kg.94,95 The desogestrel-only pill reliably inhibits ovulation in 97% of cycles and does not appear to be influenced by weight.

No evidence was identified that taking two POPs per day is harmful and, until further direct evidence is available, the CEU continues to recommend that women weighing >70 kg using POPs containing levonorgestrel, norethisterone or etynodiol diacetate should be advised to take two pills together every day (unlicensed). Women using the desogestrel-only pill are advised to take only one pill per day regardless of weight.46

When the hormone-free interval is delayed, shortened or omitted

23 Women may be given advice regarding ‘tricycling’ combined hormonal contraception to avoid withdrawal bleeds, extending the active hormone-taking days to delay menses, or shortening the pill-free interval if using liver enzyme-inducing drugs (Good Practice Point).

Tricycling

Women can be advised to ‘tricycle’ packets of COCs for a variety of reasons: to prevent withdrawal bleeds; to reduce menstrual bleeding problems or premenstrual symptoms; and to avoid withdrawal headaches. The regimen is well tolerated and acceptable. Two RCTs have investigated bleeding patterns associated with tricycling COCs.96 Women taking COCs continuously reported fewer bleeding days requiring sanitary protection, more amenorrhoea, less menstrual pain, and less bloating than women using a 21-day COC regimen.97 The rate of spotting was the same in the two groups and both groups reported high satisfaction with bleeding patterns.96–98 A parallel, randomised, multicentre, open-label trial compared two 30 µg COCs containing 150 µg levonorgestrel.99 Women received an extended regimen (84 days of active COCs and 7 placebo days) or 13 cycles of the conventional 28-day regimen. Unscheduled bleeding occurred in women using the extended regimen but this settled with time. Pregnancy rates and side effects were similar in the two groups. Both regimens were well tolerated.

A small, non-randomised, prospective study reviewed the experiences of 50 women who used their standard COC for 6 weeks followed by a 7-day pill-free interval.100 Limited information was obtained due to inherent bias and confounding factors. Nevertheless, many of the women were using COC for menstrual related symptoms, which were reduced by extending the number of active weeks of pill taking. There is no evidence that breast cancer or cardiovascular disease is increased with this type of dosing regime.
The SPCs for COCs suggest they should be taken for 3 weeks followed by a pill-free week (or a placebo pill-taking week). SPCs suggest that the combined hormonal contraceptive patch may be used for up to 6 consecutive weeks before a patch-free week.\textsuperscript{33} In previous Guidance, the CEU supported tricycling COC packs to avoid withdrawal bleeds.\textsuperscript{6} The CEU suggests that patch use may be extended to 12 weeks as for COCs.

\textbf{Shortening the hormone-free interval}

An alternative regimen to tricycling has been investigated which involves shortening the hormone-free interval by increasing the number of active pills in a packet to 24.\textsuperscript{101} This has been advocated to reduce the risk of COC failures for women using liver enzyme-inducing drugs and may be useful for women who have had true pill failures. There is some evidence that reducing the hormone-free interval to 5 days may reduce the risk of ovarian follicular activity.\textsuperscript{102}

\textbf{Delaying menses}

A population study found that norethisterone is widely used by women in the UK to delay menstruation\textsuperscript{103} with peak prescribing (1100 items/month/million population) in July. Smaller peaks in prescribing were evident in December and January. Women like to be able to control and delay menses. Combined hormonal contraception can be used to achieve this either by regularly tricycling the COC or patch or, on occasions, increasing the number of pills or continuing to reapply a new patch each week to delay menses (unlicensed).

\textbf{Extended use of intrauterine methods}

\textbf{24 After counselling about declining fertility, contraceptive efficacy and risks associated with IUD insertion, women who have an IUD with \textgreater{}300 mm\textsuperscript{2} of copper inserted at age \textgreater{}40 years can be advised to retain the device until the menopause (Grade C).}

\textbf{25 Women can be advised that if the LNG-IUS is inserted at \textgreater{}45 years of age (and not being used in combination with oestrogen replacement therapy) it may continue to be used to provide contraception for 7 years (Grade C).}

It is generally accepted that non-hormonal contraception can be stopped 1 year after the last menstrual period (LMP) in a woman \textless{}50 years of age.\textsuperscript{104} If the LMP occurs in a woman \textless{}50 years of age, however, contraception should be continued until 2 years of amenorrhoea.\textsuperscript{29,105} It is accepted practice that a copper IUD with \textgreater{}300 mm\textsuperscript{2} copper which is inserted at \textgreater{}40 years can be retained until the menopause.\textsuperscript{59,104} This falls outside the manufacturers' recommended duration of use. Women should be counselled about declining fertility with age, the risks associated with insertion (infection, perforation, expulsion) and potential loss of contraceptive efficacy if used beyond the recommended life of the device and be given the choice to continue. There are data from RCTs of LNG-IUS contraceptive efficacy with 7 years of continuous use\textsuperscript{106,107} At present, the CEU advises that women aged \textgreater{}45 years at the time of LNG-IUS insertion can be counselled regarding ongoing contraceptive efficacy and the risks of removal and replacement and may choose to continue the LNG-IUS for 7 years although this is outside the terms of the product licence.\textsuperscript{104}

\textbf{When is POEC used outside the terms of the product licence?}

POEC is licensed for use as EC after UPSI or potential contraceptive failure. The current licence for POEC is for a 1.5 mg dose of levonorgestrel to be taken as a single dose as soon as possible and within 72 hours of UPSI or potential contraceptive failure.\textsuperscript{70,108} There are no RCTs to identify the optimal dose of POEC to prevent pregnancy. In addition, there are limited data on how POEC works if ovulation has already occurred. If taken prior to ovulation, POEC can inhibit ovulation for 5–7 days, by which time any sperm in the upper reproductive tract have lost their ability to fertilise an ovum.

\textbf{Use of POEC beyond 72 hours}

\textbf{26 POEC may be considered for use between 73 and 120 hours after UPSI, but women should be informed of the limited evidence of efficacy and offered the alternative option of an IUD (Good Practice Point).}

Previous CEU Guidance did not support the use of POEC beyond 72 hours.\textsuperscript{61} A large RCT by the WHO\textsuperscript{109} provided evidence that POEC continues to reduce the expected pregnancy rate if taken between 73 and 120 hours after UPSI. Numbers recruited to this arm of the WHO trial were small (214 women between 73 and 120 hours). Between 60% and 63% of expected pregnancies were prevented with the divided or the single-dose regimen taken between 73 and 120 hours after UPSI. Although numbers were small and results did not reach statistical significance, these data add weight to the theory that POEC does not suddenly stop working at 72 hours. No data were identified on use of POEC beyond 120 hours.

A randomised, double-blind, placebo-controlled trial\textsuperscript{110} showed that POEC (1.5 mg levonorgestrel as a divided regimen or 0.75 mg levonorgestrel as a single dose) administered in the follicular phase (pre-ovulation) interferes with the ovulatory process. The efficacy of POEC in inhibiting ovulation depends on the timing of POEC in relation to the pre-ovulatory LH surge. When POEC is given before the LH surge, when follicles are \textless{}15 mm in size, follicular rupture was prevented or ovulatory dysfunction (absent LH peak or LH peak after follicular rupture) was apparent in the 5 days following administration in between 79% and 86% of women.\textsuperscript{110} The larger the follicle at the time of POEC administration the less effective is prevention of follicular rupture. Therefore POEC may still be effective in preventing ovulation if given \textgreater{}72 hours after UPSI, particularly if given prior to the LH surge.

\textbf{Use of POEC more than once in a cycle}

\textbf{27 Women can be advised that POEC can be used more than once in a cycle if clinically indicated (Good Practice Point).}

The SPC states that repeated administration of POEC within a menstrual cycle is not advisable because of the possibility of disturbance of the cycle.\textsuperscript{70} Previous CEU Guidance supported use more than once in a cycle if clinically indicated.\textsuperscript{61} It is likely that if POEC is used again prior to the LH surge it will continue to be effective. A randomised, double-blind, placebo-controlled trial showed that when leading follicles did not rupture in the 5 days after taking POEC, ovulation occurred in up to 34.5% of...
women after this time.\textsuperscript{110} This suggests that further UPSI may be an indication for repeat POEC use.

Use of POEC will not induce abortion if the woman is already pregnant.

**Increased dose of POEC for women using liver enzyme-inducers**

28 Women using liver enzyme-inducing drugs should be advised to increase the dose of POEC and take 2.25 mg as a single dose as soon as possible and within 72 hours of UPSI (Grade C).

It is established practice in the UK to increase the dose of POEC by 50\% for women using liver enzyme-inducing drugs.\textsuperscript{6,61} This regimen is unlicensed and no studies have confirmed that this dose increase is required. Women should be informed of the lack of data to support the efficacy of POEC when also using liver enzyme-inducing drugs and should be offered an IUD (which is unaffected by concomitant medication). If POEC is chosen, the dose should be increased to 2.25 mg. At present, despite lack of evidence, the CEU continues to recommend this increase in dose.

Levonelle One-Step\textsuperscript{6} is available for pharmacy supply (a single tablet of 1.5 mg levonorgestrel) and should be available for NHS supply in Autumn 2005. There is no recommendation in SPCs regarding use for women using liver enzyme-inducing drugs but it is likely that this single tablet (1.5 mg) will be taken as soon as possible and within 72 hours of UPSI sex or potential contraceptive failure and the same dose repeated 12 hours later (100\% increase in dose) for women using liver enzyme-inducing drugs.

**Advance provision of POEC**

29 Advance provision of POEC and instructions on use can be offered to those women attending family planning and sexual health services to increase early use when required (Grade A).

Previous CEU Guidance supports advance provision of POEC for women attending family planning and sexual health services.\textsuperscript{61} Advance provision of POEC refers to provision in advance of need. The SPC provides information on indications for use but does not state that is has to be provided only at the time of unprotected intercourse. It is likely therefore that providing an advance supply of POEC is not outside the terms of the product licence.

RCTs have shown that in selected women advance provision is safe and effective and increases the early uptake of EC and may reduce the rate of unintended pregnancies without increasing the number of women having UPSI.\textsuperscript{111,112} A population-based study did not show that providing sexually active women with EC had any impact on abortion rates.\textsuperscript{113} Nevertheless, advance provision enabled most women who used the supply to do so within 24 hours of UPSI. A recently published RCT that investigated the effect of direct access to EC from pharmacies and advance provision on use of the regimen, pregnancies, sexually transmitted infections, contraceptive use and sexual behaviour\textsuperscript{114} found no difference in pregnancy rates between the groups. Women with supplies at home were more likely to use POEC if required without compromising ongoing contraceptive use or sexual behaviour.\textsuperscript{114} Although the CEU does not advocate advance provision of POEC to all women, clinicians may consider advance provision if appropriate (e.g. relying on barrier methods or travelling abroad).

**What evidence is available to support the use of contraceptive agents for non-contraceptive unlicensed indications?**

Women may use contraception primarily for pregnancy prevention but also gain from non-contraceptive health benefits.\textsuperscript{6} Women who are not at risk of pregnancy may wish to continue their contraceptive method. Women may be not at risk of pregnancy because they are relying on another method of contraception (e.g. sterilisation) but wish to use a contraceptive agent for non-contraceptive benefits. Medical eligibility criteria have been published by the WHO\textsuperscript{115} and refer to the risk–benefit ratio of contraception for women with certain medical conditions. If the medical eligibility criteria should be revised if women are using contraception to manage other conditions, since alternative treatment options may also have risks. Contraceptive agents are commonly used in the management of conditions for which there is little published evidence of their effectiveness (e.g. polycystic ovarian syndrome or premenstrual syndrome). Some contraceptives, notably the COC, have been shown to reduce the risk of ovarian\textsuperscript{116–120} and endometrial cancer.\textsuperscript{118,121–124} However, there is insufficient evidence to support the use of COC solely to prevent these cancers. The use of DMPA in women with sickle cell disease has been shown to reduce disease recurrence\textsuperscript{125} and is a suitable contraceptive should one be required. There is insufficient evidence to support use routinely to prevent disease exacerbation.

**Combined oral contraception**

**Management of menorrhagia**

30 Women may be advised that menstrual blood loss may be reduced with COC use (Grade C).

A Cochrane Review concluded that there is insufficient evidence to confirm that COC reduces menstrual blood loss.\textsuperscript{126} The one small RCT included showed a 43\% reduction in measured menstrual blood loss over two cycles with COC use.\textsuperscript{127} Despite the Cochrane Review, in clinical practice, women almost universally describe less bleeding with COC use. Guidance from the Royal College of Obstetricians and Gynaecologists (RCOG) supports the use of COC in reducing menstrual bleeding.\textsuperscript{128} In addition, data from small prospective studies confirmed a reduction in menstrual blood loss and dysmenorrhoea in women using oral contraception.\textsuperscript{129} For women with low ferritin at recruitment, concentrations increased with oral contraceptive use. The Oxford Family Planning Association contraceptive study showed that hospital referral for excessive periods, painful periods, irregular periods and other menstrual disorders was only one-half to three-quarters as common in women currently using oral contraceptives or stopping them within the previous 12 months than in non-users.\textsuperscript{130}

**Management of dysmenorrhoea and endometriosis**

31 Women may be advised that menstrual pain may be reduced with COC use (Grade C).

A Cochrane Review concluded that there was insufficient evidence to determine if COC use can reduce primary dysmenorrhoea.\textsuperscript{131} A subsequent small, randomised, double-blind, placebo-controlled trial showed a significant reduction in menstrual cramps with oral contraceptive use.\textsuperscript{132} A non-systematic review suggested COC was more effective than placebo in relief of menstrual pain but there was marked heterogeneity between the studies included.\textsuperscript{131}
A Cochrane Review identified one RCT which suggested COC use was less effective than gonadotrophin-releasing hormone (GnRH) agonists in the treatment of menopausal pain associated with endometriosis, but was as effective in reducing dyspareunia and non-menstrual pain.133

Management of menstrual bleeding abnormalities associated with progestogen-only implants or injectables

32 Women using progestogen-only implants or injectables who have menstrual abnormalities may consider the short-term use of a COC or NSAID after gynaecological problems and infection have been excluded (Grade C).

The UK version of WHOSPR recommends use of COC for women with problematic bleeding while using progestogen-only implants or DMPA.3 However, supporting evidence is limited. The WHOSPR suggested that treatment of light or heavy bleeding in women using progestogen-only injectables with oestrogen or non-steroidal anti-inflammatory drugs (NSAIDs) was likely to be of short term or no benefit. EE was moderately effective in decreasing the number of days of bleeding.134 The WHOSPR, however, supports the use of NSAIDs (ibuprofen or mefenamic acid), EE or COC in the management of bleeding associated with progestogen-only implants.7 Most data, however, refer to bleeding associated with the levonorgestrel-only implant (Norplant®).

Management of acne vulgaris

33 Women can be informed that COCs improve acne vulgaris (Grade A).

A Cochrane Review highlighted that COCs can improve acne vulgaris.135 Small randomised trials have shown significant reductions in acne lesions with COCs containing desogestrel,136 levonorgestrel137-140 and norgestimate.141 Dianette® (Schering Healthcare Ltd) contains 35 μg EE with 2 mg cyproterone acetate (a progestogen with anti-androgenic properties). Dianette is not licensed as a contraceptive, but for treatment of acne or hirsutism. A case-control study used data from the General Practice Research Database and, after adjustment for body mass index, smoking and androgenic disorders, showed a four-fold increase in the risk of VTE with Dianette compared to a COC containing levonorgestrel (odds ratio 3.9; 95% CI 1.1–13.4).142 Duration of use did not affect this risk. A combined, nested, cohort analysis and case-control study support this level of risk but no randomised trials have been performed, thus confounding and bias cannot be excluded.143 The Committee on the Safety of Medicines advised: “Dianette is not indicated solely as a contraceptive; it is a treatment option for women with severe acne, which has not responded to oral antibiotics, or for moderately severe hirsutism; it should be withdrawn 3–4 months after the treated condition has resolved”.144 If the hyperandrogenism, however, does not resolve and after counselling women may choose to continue or recommence Dianette.

Management of hypo-oestrogenism

34 Combined hormonal contraception may be useful in preserving bone mineral density (BMD) for women with a premature menopause but has no beneficial effect on BMD in women with anorexia nervosa (Grade B).

Studies have suggested that for women with hypo-oestrogenism, combined hormonal contraception may be clinically beneficial. The COC appears as effective as hormone replacement therapy (HRT) in preserving or increasing BMD in perimenopausal women.142 For women with premature menopause, use of COC may be more acceptable than using a traditional HRT. For women with hypogonadotrophic or normogonadotrophic amenorrhoea there is evidence that COC use maintains or improves BMD.146 For women with anorexia nervosa, however, use of COC has no beneficial effect on BMD.147,148

Progestogen-only pills and implants

Management of menstrual bleeding

35 POPs and implants are ineffective in the management of menstrual bleeding problems and should not be used for this purpose (Good Practice Point).

There are no recommended uses for POPs or progestogen-only implants other than for contraception. Nevertheless, these methods are used by some women to manage bleeding problems. Women using POPs can have a variety of bleeding patterns while using DMPA for treatment of menorrhagia (Grade B).

Progestogen-only injectables

Management of menorrhagia

36 Progestogen-only injectables can induce amenorrhoea and may be considered by women with menorrhagia (Grade B).

Menorrhagia is reported by almost half (46%) of DMPA users at 3 months.149 This increases with time to 53.3% at 6 months and 58.5% at 12 months. Research from the Special Programme of Research and Training in Human Reproduction of the WHO showed that women using DMPA tolerated far greater menstrual disturbances than women using other methods (combined pills, progestogen-only pills and vaginal rings).150 Women using DMPA were more likely to have been counselled regarding bleeding disturbances than women using other methods, which may explain less discontinuation with DMPA. Although there are no RCTs investigating DMPA in the management of menorrhagia, it may be helpful for some women.

Levonorgestrel-releasing intrauterine system

Duration of use in management of menorrhagia

37 Women can be advised that if the LNG-IUS is used to treat menorrhagia (and when not relying on it for contraception or in combination with oestrogen replacement therapy) then it may be continued beyond the usual 5 years of licensed use, if bleeding patterns remain acceptable (Good Practice Point).
The LNG-IUS is now licensed for use in idiopathic menorrhagia (5 years) and as protection from endometrial hyperplasia during oestrogen replacement therapy (4 years).104

References

121 Cancer and Steroid Hormone Study of the Centers for Disease Control and the National Institute of Child Health and Human Development. Combination oral contraceptive use and the risk of endometrial cancer. JAMA 1987; 257: 796–800.


137 Worret A, Wazradzuk JP, Andreas JO, Binder N. Acne resolution rates: results of a single-blind, randomized, controlled, parallel phase three trial with EE/CMO (Belara) and EE/LENG (Microgynon). Dermatology 2001; 203: 38–44.


This Guidance was developed by the Clinical Effectiveness Unit (CEU) of the Faculty of Family Planning and Reproductive Health Care (FFPRHC): Dr Gillian Penney (Director), Dr Susan Brechin (Co-ordinator); Ms Lisa Allerton (Research Assistant) in consultation with a multidisciplinary expert group of health care professionals involved in family planning and reproductive health care. The expert group comprised: Dr Caroline Boorer (Associate Specialist, Contraception and Sexual Health, Northumberland Health Care); Ms Lorraine Forster (Practice Development Nurse, The Sandyford Initiative, Glasgow); Dr Alyson Elliman (Lead Associate Specialist, Croydon PCT/FFPRHC Clinical Standards Committee Member, Professor Anna Glaser, Consultant in Sexual Health/Contraception, Northumbria Partnership NHS Trust, Consultant in Sexual and Reproductive Health, Centre for Contraception and Sexual Health, Nottingham); Dr Louise Massey (Locum Consultant in Public Health, Wyre Forest PCT, Brook House, Kidderminster/Trainee Member of the CEU/FFPRHC Education Committee Member, Dr Noel Mack (General Practitioner, Kenway, Aberdeenshire); Ms Elaine Ross (Community Services Pharmacist, Westholm, Woodend Hospital, Aberdeen); Dr Rachel Westwick (Career Grade Trainee, Margaret Pyke Centre, London/FFPRHC Education Committee Member). Written feedback was provided by: Ms Toni Belfield (Director of Information, FPA, London).

Evidence tables are available on the FFPRHC website. These summarise relevant published evidence on use of contraception outside product licence, which was identified and appraised in the development of this Guidance. The clinical recommendations within this Guidance are based on evidence whenever possible.

**Grades of Recommendations**

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<th>Evidence based on randomised controlled trials (RCTs)</th>
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<td>A</td>
<td>Evidence based on other robust or observational studies</td>
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<tr>
<td>B</td>
<td>Evidence is limited but the advice relies on expert opinion and has the endorsement of respected authorities</td>
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<tr>
<td>C</td>
<td>Good Practice Point where no evidence exists but where best practice is based on the clinical experience of the expert group</td>
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Electronic searches were performed for: MEDLINE (CD Ovid version) (1996–2005); EMBASE (1996–2005); PubMed (1996–2005); The Cochrane Library (to February 2005) and the US National Guideline Clearing House. The searches were performed using relevant medical subject headings (MeSH), terms and text words. The Cochrane Library was searched for systematic reviews, meta-analyses and controlled trials relevant to use of contraceptive methods for contraception and other conditions outside the terms of the product licence. Previously existing guidelines from the FFPRHC, the Royal College of Obstetricians and Gynaecologists (RCOG), the Department of Health, the British Medical Association (BMA), the Royal College of Nursing (RCN), the Royal College of General Practitioners (RCGP) and reference lists of identified publications were also searched. Similar search strategies have been used in the development of other national guidelines. Selected key publications were appraised according to standard methodology checklists before conclusions were considered as evidence. Evidence was graded as above, using a scheme similar to that adopted by the RCOG and other guideline development organisations.