



This Patient Group Direction (PGD) must only be used by registered Community Pharmacists who have been named and authorised by their organisation to practice under it. The PGD must only be used in conjunction with a local authority commissioned service specification for Emergency Contraception. The most recent and in date final signed version of the PGD should be used.

Patient Group Direction

for the supply and/or administration of

Ulipristal acetate 30mg tablet

by registered Community Pharmacists for

Emergency Hormonal Contraception (EHC)

in Cheshire and Merseyside

Version number: 2.0

Effective From: June 1st 2016 Expires: May 31st 2019

Change history

Version number	Change details	Date
1	Original version developed by Onyia, Mullin, Stubbs, Knight, Carrol, Geoghegan, Cartwright & Major –introduced in April 2014, expires March 31 st 2016	April 2014
2	Completely reviewed and updated (February 2016) Takes into account NICE MPG 2 guidance & revised GMC prescribing guidance	March 2016

PGD approval/ development

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PGD authorisation

Na	ime	Job title and organisation	Signature	Date
Senior Pharmacist & Lead Author	John. P Hampson GPhC No= 2025614	Public Health Specialist, Cheshire West and Chester Council	Joh P Aanfr.	22/04/2016
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Person signing on behalf of authorising body ¹	Fiona Johnstone	Director of Public Health Wirral Borough Council	Had	9/5/2016

¹ Clinical governance or safety lead of the Local Authority , usually the Director of Public Health or Chief Executive

Community Pharmacist agreement to practise under the Ulipristal Acetate 30mg tablets Patient Group Direction for Community Pharmacists

I have read and understood the Patient Group Direction and agree to supply and/or administer this medicine only in accordance with this PGD

Name	GPhC Number	Signature	Date

Authorised to practice by Superintendent (or person acting on behalf of Superintendent)*

Full Name (print)	
GPhC number	
Signature	
Date	

Agrees to maintain a current list of the names of individuals who may implement this PGD and to keep this with a pharmacy master copy of the PGD.

"Person acting on behalf of Superintendent" is usually the pharmacist Area or Branch manager.

^{*} Has responsibility to ensure that only fully competent, qualified and trained professionals implement this PGD.

Training and competency of registered Community Pharmacists

	Requirements of registered community pharmacists working under the PGD
Qualifications and professional registration	Community Pharmacists currently registered with the General Pharmaceutical Council (GPhC), who are working in a pharmacy contracted to NHS England (Mersey) or NHS England (Cheshire, Warrington and Wirral).
Initial training	As a minimum requirement, this must be at the same level, covering the same learning objectives and competencies as the Centre for Pharmacy Postgraduate Education (CPPE) e-learning programme for emergency contraception and safeguarding.
Competency assessment	The pharmacist must satisfy the requirements of Self-declaration of Competence for Community Pharmacy for Emergency Contraception. The Pharmacist should be able to demonstrate the competencies specified in NICE's Competency Framework for Health Professionals using Patient Group Directions. http://www.nice.org.uk/mpc/goodpracticeguidance/GPG2.jsp
Ongoing training and competency	The pharmacist must maintain a regular self-assessment declaration of competency every two years or sooner if appropriate. In addition to the statutory requirement for Continuing Professional Development (CPD), each pharmacist is expected to maintain an up to date awareness of developments in emergency contraception.

Clinical condition		
Clinical condition	Emergency contraception between 72 – 120 hours of unprotected sexual	
or situation	intercourse (UPSI) or suspected failure of a contraceptive method (eg	
to which this PGD	barrier method or missed pills).	
applies		
Inclusion criteria	A woman of child bearing age AND presenting within 72 to 120 hours of	
	UPSI.	
	 Can also include women presenting between 72 – 120 hours. 	
	o with failure of barrier or normal contraceptive method (see Appendix	
	B).	
	OR with severe diarrhoea and/or vomiting which may have reduced	
	oral contraceptive efficacy.	
	Patient has received ulipristal acetate emergency contraception but has verified within three hours of taking it (provided they are still within 120).	
	vomited within three hours of taking it (provided they are still within 120	
	hours of UPSI).	
	Special notes on age	
	Less than 18 years: A risk assessment should be undertaken to determine	
	whether the child is at risk of harm. If you have a concern, the matter should	
	be discussed with the local safeguarding lead.	
	Less than 16 years: Must be competent as assessed under the Fraser	
	Guidelines on consent to medical treatment.	
	Less than 13 years: The matter must be discussed with the local	
	safeguarding lead.	
	The pharmacist must be aware of their local safeguarding contact	
	numbers.	
Exclusion criteria	Woman unable to attend in person.	
	Hypersensitivity to the active substance or any of the excipients (e.g.	
	lactose, povidone K30, croscarmellose and magnesium stearate) or	
	patient has previously experienced any severe clinical problems with	
	hormonal contraception.	
	Women with hereditary problems of galactose intolerance, Lapp lactase	
	deficiency or glucose – galactose malabsorption problems.	
	Up to 72 hours since UPSI. Advise woman that levonorgestrel is	
	available - refer to levonorgestrel PGD.	
	Confirmed pregnancy.	
	Previous use of ulipristal acetate within this menstrual cycle. NB a second	
	supply for patients who vomit within 3 hours of taking ulipristal acetate	
	(provided the repeat dose is still within 120 hours of UPSI) is allowed	
	under this PGD. Following termination of pregnancy , consider the	
	 date of termination as the last menstrual period. Previous use of levonorgestrel containing emergency hormonal 	
	Previous use of levonorgestrei containing emergency normonal contraception within this menstrual cycle/	
	Any earlier episodes of UPSI which took place more than 120 hours ago	
	and within this menstrual cycle.	
	and within this menstrual cycle.	
	and within this menstrual cycle.	
	and within this menstrual cycle.	

• Currently taking any medicine which induces hepatic enzymes

Carbamazepine, Eslicarbazepine, Oxcarbazepine

Phenobarbital, Phenytoin, Primidone

Rufinamide, Topiramate

Rifampicin, Rifabutin

Ritonavir (long term), Efavirenz, Nevirapine

Fosphenytoin

St John's Wort (hypericum perforatum)

Bosentan

Aprepitant

For enzyme inducers, this exclusion also covers women who have stopped the medicines listed that induce hepatic enzymes within the last 28 days.

 The following drugs have also been reported to interfere with progestogen containing contraceptives:-

Aprepitant

Bosentan

Crizotinib

Dabrafenib

Efavirenz

Fosaprepitant

Vemurafenib

The manufacturers recommend alternative forms of contraception. Because of the sparsity of information on the nature of these interactions and the potential for teratogenicity in some cases, these patients should be referred for specialist management.

- Uncontrolled severe asthma (where asthma is not controlled despite oral corticosteroid treatment).
- Severe hepatic impairment.
- Post partum patients (within 21 days) are not considered at risk of pregnancy and so are excluded from treatment.

Cautions (including any relevant action to be taken)

- An IUD is the most effective means of post coital contraception and this option must be discussed with the woman. In instances where an IUD is acceptable to the woman, continue to supply ulipristal acetate in case the IUD fitting is not done or proves unsuitable.
- Severe intestinal malabsorption syndromes e.g. Crohn's disease
 The FRSH advise that oral contraception may be less reliable in women
 with malabsorption due to severe small bowel disease or resection.
 Women with these conditions should be encouraged to consider an IUD
 as the preferred method of emergency contraception.
- Breast feeding For women who are breast feeding inform them that
 breast feeding is not recommended for 7 days after taking ulipristal
 acetate. The manufacturers advise that women who are breast feeding
 should feed their baby immediately before taking the tablet, then pump
 and discard the milk for 7 days after taking the ulipristal acetate. Breast
 feeding can be resumed after 7 days. If the woman is unable or unwilling
 to comply with this advice she is excluded from treatment with ulipristal
 acetate under this PGD consider supply under levonorgestrel PGD or

Arrangements for referral for medical advice	refer to GP or Community Sexual and Reproductive Health Clinic. The following drugs should be given 1.5 hours before or 1.5 hours after ulipristal acetate:- Dabigatran, digoxin, fexofenadine. Know the referral pathway into local sexual and reproductive health services or how to contact the local lead doctor for sexual and reproductive health for medical advice.
Action to be taken if patient excluded	 Discuss reasons for exclusions. Refer immediately to Community Sexual and Reproductive Health Clinic or GP if appropriate. An intrauterine device (IUD) may be fitted up to 5 days after unprotected intercourse or up to 5 days after likely ovulation. Provide an emergency supply of condoms. Consider supply and administration of levonorgestrel (refer to levonorgestrel PGD). Visitors from countries outside the EU are entitled to access this free service on the NHS.
Action to be taken if patient declines treatment	 Discuss reasons patient declines treatment. Consider the supply and administration of levonorgestrel. Refer immediately to Community Sexual and Reproductive Health Clinic or GP if appropriate.

Details of the medicine

Name, form and strength of medicine	Ulipristal acetate 30mg tablet	
Legal category	POM	
Indicate any off-label/ unlicensed use	Not applicable	
Route/method of Administration	Oral It is recommended that the woman takes the tablet while still in the pharmacy. If the tablet is not taken in the pharmacy, advise the woman to take the tablet as soon as possible.	
Dose and frequency	ONE 30mg tablet to be taken as soon as possible after the 72hrs but no later than 120 hours after UPSI. If woman vomits within 3 hours of taking the dose, then a second pack may be issued if the woman is able to take the repeated dose within 120 hours following UPSI. Administration while the patient is present should be encouraged and supported, although this is voluntary.	
Quantity to be administered and/or supplied	One pack containing one Ulipristal acetate 30mg tablet	
Maximum or minimum treatment period	See dose and frequency section above	
Adverse effects	Most common side effects may include; Headache Nausea Abdominal pain Vomiting Dysmenorrhea Dizziness (shouldn't drive or operate machinery if affected) Any serious adverse effects must be reported to the MHRA via the yellow card scheme.	

Records to be kept

- Where possible*, a consultation proforma for the "Supply and Administration of EHC" should be fully completed and signed for all consultations, irrespective of whether a supply is made. The following details should be recorded:
 - Valid informed consent has been given
 - o Patient's name, address (optional) and date of birth
 - o Name of GP
 - o Dose given
 - o Date of supply
 - A record of the counselling about encouragement to consider an IUD
 - o Advice given
 - Advice given if patient excluded or declines treatment
 - Details of any adverse reactions and actions taken
 - Signature, GPhC number and name of pharmacist who administered or supplied the medication
 - o Document if the dose is administered on the premises
 - The supply must be entered in the Patient Medication Record (PMR)
 - o All records should be clear, legible and contemporaneous.

This can be recorded via a paper or electronic version (or both)

- A "Fraser Ruling Assessment of Competency" form must be completed for all women under 16 years of age.
- * The Human Medicines Regulations 2012² confirms that in the case of supply of oral contraception, the requirements for recording information are relaxed. It is therefore reasonable for pharmacists to exert their professional judgement when supplying a woman with EHC who does not wish to provide any information.

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² http://www.legislation.gov.uk/uksi/2012/1916/regulation/253/made, paras 1,2,3 & 4

Patient information

Written information to be given to patient or carer	Give copy of the patient information leaflet and discuss as required e.g. failure rate (approx. 2 women out of 100 will become pregnant despite taking EHC. An IUD has negligible failure rate). Supply woman with appropriate leaflets and information about local Sexual and Reproductive Health services.
Follow-up advice to be given to patient or carer	 Woman should be advised to have a pregnancy test after 3 weeks to check for failure of EHC. Ulipristal acetate is not intended for use during pregnancy and should not be taken by any woman suspected or known to be pregnant. Limited human data regarding pregnancy exposure to ulipristal acetate do not suggest any safety concern. However, if a woman does become pregnant, she must inform her doctor. Advise patient that she could still become pregnant. If next period is delayed by more than 7 days or is abnormal in any way (light, heavy or painful), woman should seek medical advice. Emphasise that the tablet is for emergency use only and is not as effective as a regular method of contraception. She must continue to use another method for the remainder of the cycle. Also suggest that the woman makes a medical appointment to obtain regular contraception where appropriate. Seek medical advice if there is any lower abdominal pain because this could signify an ectopic pregnancy. Advise that the patient may be at risk of sharing sexually transmitted infections (STIs) and the need for condom use. Patients may be asymptomatic. Further advice, screening and treatment can be accessed from Community Sexual and Reproductive Health Services or their GP. If the patient wishes to resume hormonal contraception, they should do so AFTER 5 days. Patient should be advised to abstain from sex or use a condom during these 5 days because no other hormonal contraception can be used during this period. When restarting oral contraception after this "gap" (i.e. on day 6), additional barrier method must be used for the requisite number of days as outlined in appendix B. Breastfeeding is not recommended for 7 days after taking ulipristal acetate. During this time it is recommended to express and discard the breast milk in order to stimulate lactation. Advise that if vomiting occurs within 3 hours of taking ulipristal acetate to immediately return to the ph

APPENDICES

Appendix A: Key References

Faculty of Sexual & Reproductive Healthcare. *Clinical guidance: Emergency contraception.* London, Clinical Effectiveness Unit, 2012

http://www.fsrh.org/pdfs/CEUguidanceEmergencyContraception11.pdf

Faculty of Sexual & Reproductive Healthcare. *EMA Review of emergency contraception and weight Update.* London, Clinical Effectiveness Unit, 2014

http://www.fsrh.org/pdfs/FSRH_ECDecisionGuide.pdf

Electronic Medicines Compendium. *Summary of Product Characteristics: ellaOne 30mg tablet* . Paris, eMC (Laboratoire HRA Pharma), January 2015

https://www.medicines.org.uk/emc/medicine/22280 accessed 12th January 2016.

British National Formulary. *BNF: 70.* London, Pharmaceutical Press, 2015 Also electronic version www.bnf.org accessed 9th March 2016

General Medical Council. Good practice in prescribing and managing medicines and devices.

London: GMC, 2013 (updated 2014)

http://www.gmc-uk.org/Prescribing_guidance.pdf_59055247.pdf

Dickson N. The GMC's stance on Avastin. BMJ 2015; 350:h204

National Institute for Health and Care Excellence. *Medicines Practice Guideline 2:*

Patient Group Directions. London: NICE, 2013

https://www.nice.org.uk/guidance/mpg2/resources/patient-group-directions-1779401941189

National Institute for Health and Care Excellence. *Public health guideline 51: Contraceptive services for under 25s.* London: NICE , 2014

 $\underline{https://www.nice.org.uk/guidance/ph51/resources/contraceptive-services-for-under-25s-1996413367237}$

National Institute for Health and Care Excellence. *NICE local government briefing 17: Contraceptive services*. London: NICE, 2014

https://www.nice.org.uk/guidance/lgb17/resources/contraceptive-services-60521151465925

Cheng L, Che Y, Gulmezoglu AM. Interventions for emergency contraception (Review). *Cochrane Database of Systematic Reviews* 2012

Levonorgestrel and ulipristal remain suitable emergency contraceptives for all women regardless of bodyweight. *Letter, Medicines and Healthcare Products Regulatory Agency.* London: MHRA, 2014

Faculty of Sexual & Reproductive Healthcare. *Clinical Effectiveness Unit Statement: Missed pill recommendations*. London, Clinical Effectiveness Unit, 2011 http://www.fsrh.org/pdfs/CEUStatementMissedPills.pdf

Appendix B: Common reasons for usual method failure

- Misplaced / dislodged diaphragm / incorrect insertion / torn / removed too early.
- Condom breakage /leakage /ejaculation on/near external genitalia.
- IUD: If complete or partial expulsion is identified or mid cycle, removal of the IUD is deemed necessary and EHC should be considered. Also indicated if the device has expired i.e. more than 3 (Jaydess), 5 or 10 years since insertion, depending on type.
- · Miscalculation of fertility awareness method.
- Reduced contraceptive protection because of e.g. severe diarrhoea and vomiting which may have reduced oral contraceptive efficacy.
- Missed or late contraceptive pill COC or Progestogen only contraceptive (POP) (further notes available in BNF chapter 7.3.1).
- Combined vaginal ring (CVR) (NuvaRing ®)
 - o Left in for more than 4 weeks
 - o New ring hasn't been inserted following the 7 day break
 - Ring has been expelled from the vagina either spontaneously or during intercourse and not replaced within 3 hours
 - Expelled more than once per cycle
- Allowed more than 14 weeks (98 days) to elapse since the last medroxyprogesterone acetate (Depo Provera or Sayana Press) contraceptive injection.
- Using contraceptive patch which becomes partially or fully dislodged or missed the change day (refer to current SPC for further advice).
- Barrier method failure in women on the COC/POP/Progestogen implants or CVR who are also taking liver enzyme inducing drugs (and for 28 days after stopping the liver enzyme inducing drugs).
- Progestogen implant: Indicated if the implant has expired (more than 3 years since insertion).
 Also if there has been a failed barrier method or UPSI during, or in the 28 days following, the use of liver enzyme inducing drugs.

N.B. For "missed pill" failures – please see tables overleaf

Appendix B continued - Common reasons for usual method failure

"Missed pill" failures which DO warrant EHC ULIPRISTAL ACETATE ADMINISTRATION

wiissed pili Tallures	s which DO warrant EHC ULIPRISTAL ACETATE ADMINISTRATION
Combined pills 21 active tablets	Week 1 – If 2 or more pills are missed in the first week of pill taking and client has had UPSI either in the pill free week or in the first 7 days of the packet give EHC. Following this the COC should be continued with additional barrier contraception until pills have been taken on 7 consecutive days.
	Week 2 and 3 – If 2 or more pills are missed in the middle or last week of pill taking the woman should be advised to continue taking her pills and use condoms for 7 days in case of further missed pills. If there are not 7 pills left in the pack following the missed pills she should continue with the next pack without her usual 7 day break. If the pill free interval is avoided in this way she does not need emergency contraception.
	If there is uncertainty over which pills have been missed, EC should be given, along with advice to continue taking the pill and use a barrier method until a further 7 consecutive pills have been taken.
	EHC is indicated if there has been a failed barrier method or UPSI during or within the 7 days after a vomiting or severe diarrhoeal illness.
	EHC is indicated if there has been a failed barrier method or UPSI during, or in the 28 days following, the use of liver enzyme inducing drugs.
Progestogen-only pill (POP)	Indicated if one or more POPs have been missed or taken more than 3 hours late and UPSI has occurred in the 2 days following this. The POP should be continued with additional barrier contraception until pills have been taken correctly on two consecutive days. Exception is a desogestrel POP which can be taken up to 12 hours late without loss of contraceptive cover.
	EHC is indicated if there has been a failed barrier method or UPSI during or within the 2 days after a vomiting or severe diarrhoeal illness.
	Indicated if there has been a failed barrier method or UPSI during, or in the 28 days following, the use of liver enzyme inducing drugs.

N.B the above recommendations regarding additional barrier methods are in addition to the 5 day "gap" mentioned in the "follow-up advice to be given to patient or carer"

Potential contraceptive failures which do NOT warrant EHC use

Combined pills 21 active tablets	No indication if only one COC pill has been missed from the first 7 pills in a pack, as long as the last 7 pills in the previous pack were taken without omissions. Additional barrier contraception for 7 days is recommended, in case of further omissions.
	No indication if pills have been missed from the middle 7 pills in the pack as long as the first 7 pills were taken correctly. The COC should be continued to the end of the pack. Additional barrier contraception for 7 days is recommended, in case of further omissions.
	No indication if pills have been missed from the last 7 pills in the pack as long as the next pack is started without a pill-free interval. Additional barrier contraception for 7 days is recommended, in case of further omissions
Depo-provera	No indication if Depo Provera is given up to 2 weeks late (up to 14 weeks from the previous injection). The injectable can be given and no additional barrier method is required.

Appendix C: Advice to Young People Under 16

In considering the provision of advice or treatment on contraception, doctors and other professional staff need to take special care not to undermine parental responsibility and family stability. The doctor or other professional should therefore always seek to persuade the young person to tell the parents or guardian (or other person in loco parentis), or to let her inform them, that contraceptive advice is being sought and the nature of any advice or treatment that is given. It should be most unusual for a doctor or other professional to provide advice or treatment in relation to contraception to a young person under 16 without parental knowledge or consent.

Exceptionally, there will be cases where it is not possible to persuade the young person either to inform the parents or to allow the doctor or other professional to do so. This may be, for example, where family relationships have broken down. In such cases, a doctor or other professional would be justified in giving advice and treatment without parental knowledge or consent, provided they followed the Fraser Guidelines.

FRASER GUIDELINES

In law young people under 16 years are entitled to confidentiality in the same way as over 16 year olds. In 1985 Lord Fraser established the current legal position that a doctor or other professional can give contraceptive advice or treatment to a person under 16 without parental consent providing they are satisfied that:

- The young person will understand the risks and benefits of the treatment offered and the advice given.
- The young person cannot be persuaded to tell his or her parents or allow a health professional to inform them that he or she is seeking contraception advice.
- The young person is likely to begin or continue having intercourse with or without contraceptive treatment.
- Unless he or she receives contraceptive advice the young person's physical or mental health is likely to suffer.
- It is in the young person's best interests to give them contraceptive advice or treatment.

Reference Gillick v West Norfolk & Wisbech Area Health Authority (1984) AC 1121 ALL ER

 Where there are concerns about children and young people's welfare appropriate actions should be taken to address those concerns, working to agreed local policies and procedures. Refer to Safeguarding Children Flow Chart for Referral.

MEDICOLEGAL ASPECTS

Medical legal aspects regarding supply to under 16 year olds

1. It's illegal for them to be having sex

Answer: It is illegal for a man to have sexual intercourse with a girl under age 16 years. The girl is not committing any offence. The historical background to this Act was the need to have some structure to prevent child prostitution.

2. You are aiding and abetting an illegal act

Answer: Taking action after an event to minimise its ill consequences cannot be interpreted as aiding and abetting-any more than the investigation and treatment of sexually transmitted infection would be.

The medical Defence Union opinion is that aiding and abetting would only be involved if a person actually were present at the time of the sexual intercourse and was encouraging it.

3. The Age of Consent is 16 years

Answer: In English Law the validity of consent depends upon the capacity of the person to understand. The House of Lords considered the specific case of consent to contraceptive treatment in a ruling (Gillick v West Norfolk and Wisbech Area Health Authority and the Department of Health and Social Security, delivered October 1989). Attached is the advice which was issued after this by the Department of Health in the Handbook of Contraceptive Practice 1990 edition, pages 92 and 93.

Note that the exceptional nature of providing emergency contraception under protocol to young persons under 16 is confirmed by the actual numbers seen and considered under the protocol, compared to the numbers of older women.

Note also the young person is fully entitled to confidentiality. The guidance in paragraph 2 is that a doctor or other professional should always seek to persuade the young person to tell, or to permit to inform. No information should be given without the young person's consent and consent to disclosure given under pressure or undue persuasion would not be valid. The pharmacists training package includes a role play of the type of discussion which is valid and appropriate.

4. It shouldn't be allowed for the very young, it will just encourage them

Answer: Note that there is a lower age limit for sale of alcohol and for sale of cigarettes, but no lower age limit for the sale of condoms. Any deterrent effect in differential use is not immediately obvious!!

Rosemary Kirkman

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Contraceptive advice and treatment for young people under 16

(HN(81)5/LASS(81)2 has now been replaced by the following text which forms the Appendix to HC(86)1/HC(FP)(86)1/LAC(86)3 "Family Planning Services for Young People" issued in March 1986 – this also applies to England and Wales only).

- 1. The following guidance draws the attention of health authorities and others concerned to the considerations doctors and other professionals need to have in mind when providing contraceptive advice and treatment to young people under 16, and to the circumstances in which such advice and treatment can be given without parental knowledge or consent. The guidance results from a review of that in Section G of the Memorandum of Guidance on the Family Planning Service, as specified in the Appendix to HN(81)5 and LASSL(81)2, in the light of the House of Lords, decision in the case of Gillick v West Norfolk and Wisbech Area Health Authority and the Department of Health and Social Security delivered last October.
- 2. In considering the provision of advice or treatment on contraception doctors and other professional staff need to take special care not to undermine parental responsibility and family stability. The doctor or other professional should therefore always seek to persuade the young person to tell the parents or guardian (or other person in loco parentis)*, or to let them inform them, that contraceptive advice is being sought and the nature of any advice or treatment that is given. It should be most unusual for a doctor or other professional to provide advice or treatment in relation to contraception to a young person under 16 without parental knowledge or consent.
- 3. Exceptionally, there will be cases where it is not possible to persuade the young person either to inform the parents or to allow the doctor or other professional to do so. This may be, for example, where family relationships have broken down. In such cases, a doctor or other professional would be justified in giving advice and treatment without parental knowledge or consent, provided they were satisfied:

- that the young person could understand their advice and had sufficient maturity to understand what was involved in terms of the moral, social and emotional implications;
- that they could neither persuade the young person to inform the parents, nor to allow them to inform them, that contraceptive advice was being sought;
- that the young person would be very likely to begin, or continue having, sexual intercourse with or without contraceptive treatment;
- that without contraceptive advice or treatment, the young person's physical or mental health, or both would be likely to suffer;
- that the young person's best interests require them to give contraceptive advice, treatment or both without parental consent.
- 4. Decisions about whether to prescribe contraception in such cases are for a doctors clinical judgement, if a doctor who is not the young person's general practitioner has formed the view, after due consideration of the points made above, that it is in the best interest of the young person to prescribe contraception without parental knowledge or consent, it may be advisable and helpful for them, with the young person's agreement, to discuss the matter in confidence with her own general practitioner before making his decision.
- **5.** In organising contraceptive services for young people, health authorities may find it helpful to make separate, less formal arrangements that those for older age groups. The staff should be experienced in dealing with young people and their problems.

*Where the parental rights and duties in respect of a young person are vested in the local authority (by virtue of a care order or a parental rights resolution under Section 3 of the Child Care Act 1980) the authority must be treated as the young person's parents for the purposes of giving consent to medical treatment in respect of a young person under 16. Where the authority does not have parental rights, the natural parent's rights are not affected. Where a young person has been committed to the care of a local authority under wardship proceedings, the consent of the High Court must be obtained by the local authority. Where a local authority shares the parental rights and duties with another person, the consent of the local authority is sufficient unless the other person indicates an objection.

EMERGENCY CONTRACEPTION For Pharmacy Use with PGD Only

