

Investigation and Management of Post-Coital Bleeding UHL Gynaecology Guideline

Contents

1. Introduction and Who Guideline applies to	1
Table 1. Aetiology of PCB by anatomical location.	2
Table 2. Risk of cervical cancer in women presenting with post coital bleeding [7]	2
PCB WITH NORMAL EXAMINATION AND INVESTIGATIONS BY GP	4
PCB AND ABNORMAL EXAMINATION AND INVESTIGATIONS.....	5
3. Education and Training	6
4. Monitoring Compliance	6
5. Supporting References.....	6
6. Key Words	7
Appendix 1 - Post coital bleeding Clinic History sheet.....	8

1. Introduction and Who Guideline applies to

Post-coital bleeding (PCB) refers to spotting or bleeding that occurs after intercourse and is not related to menstruation. The point prevalence ranges from 0.7 to 9.0% with one report indicating that the annual cumulative incidence is 6% among menstruating women ^[1]. In women under the age of 35 years, the most important cause to exclude is chlamydial infection whilst in women over the age of 35 years it is cervical cancer. Post-coital bleeding is commonly caused by cervical or endometrial polyps, cervicitis as a result of chlamydia or gonorrhoea and vaginitis associated with trichomoniasis or candidiasis. Endometritis in the presence of an intrauterine contraceptive device (IUCD) can occasionally cause post-coital bleeding. PCB is rarely caused by cervical intraepithelial neoplasia and cervical cancer ^[2]. Reassuringly, 60% of naturally menstruating women with post-coital bleeding will have spontaneous resolution of symptoms within six months and half of these women will maintain resolution for two years ^[1].

This guideline supports medical and nursing staff working within gynaecology services at UHL but is also a useful point of reference for the community based Leicestershire GP and sexual health practices.

Table 1. Aetiology of PCB by anatomical location.

Anatomical location	Cause of PCB
Vaginal	Vaginitis Vaginal atrophy Vaginal cancer Vaginal endometriosis
Cervical	Ectropion Polyp Infection Cervical Intraepithelial Neoplasia Cervical cancer Cervical endometriosis Vascular malformation
Uterine	Endometrial polyp Endometritis Endometrial cancer
Other	Trauma Foreign Body

2. Guideline Standards and Procedures

The primary role of a PCB clinic is to exclude pre-cancerous or cancerous changes in the cervix. Post-coital bleeding is the presenting complaint in 11% of women with cervical cancer ^[5], however, PCB is a common symptom and the majority of women with post-coital bleeding will not have pre-invasive or invasive cervical disease. Reassuringly, 97.7% of women presenting with post-coital bleeding whose last cervical screening was negative or inadequate were found to have no significant pathology identified at colposcopy however, around 2% of women were found to have pre-invasive or invasive cancer justifying assessment by a gynaecologist in secondary care for women presenting with PCB ^[8]. Table 2 outlines the risk of cervical cancer in women presenting with post-coital bleeding stratified by age at presentation.

3,152 women are diagnosed with cervical cancer in the UK each year with the highest incidence rates found in women aged 30-34 ^[6]. Risk factors for cervical cancer include failure to participate in cervical screening, HPV infection, smoking, prolonged COCP use and immunosuppression ^[5]. A family history of cervical cancer is associated with an increase in risk of cervical cancer although no specific genetic mutations have been identified suggesting shared risk factors may be primarily responsible for any familial association ^[5]. Incidence rates of cervical cancer are expected to fall following the introduction of the HPV vaccination programme ^[5].

Age (years)	Risk of cervical cancer
20-24	1: 44,000
25-34	1: 5,600
35-44	1: 2,800
45-54	1: 2,400

Table 2. Risk of cervical cancer in women presenting with post coital bleeding ^[7]

Scope of PCB Clinics

There are no established guidelines from the American College of Obstetricians and Gynaecologists or the Royal College of Obstetricians and Gynaecologists or evidence from randomized clinical trials to base recommendations on diagnosis and treatment of post-coital bleeding. The National Health Service cervical screening programme recommends women with post coital bleeding should be referred for examination by a gynaecologist experienced in the management of cervical disease (for example a cancer lead gynaecologist) once common causes (infection and contraception) have been excluded in primary care ^[3]. Gynaecologists may refer such individuals for symptomatic colposcopic examination outside the cervical screening programme if cancer is suspected ^[3]. NICE recommendations for suspected cervical cancer referrals do not include symptoms but recommend referral if the appearance of the cervix is suspicious of cancer ^[4].

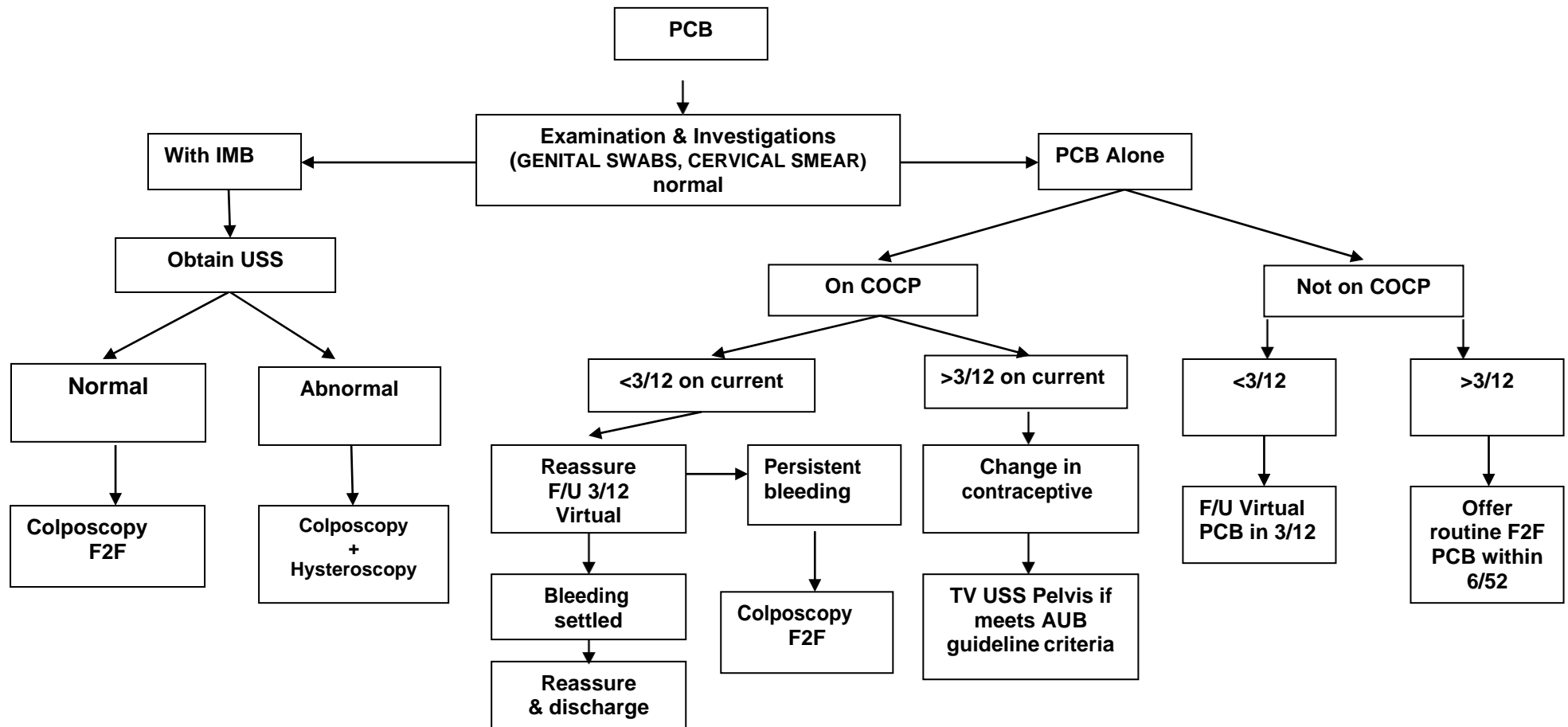
In UHL we have developed a PRISM pathway for PCB referrals to streamline their care. This includes exclusion of common and obvious causes of PCB (i.e. infections, polyps and contraceptive use) which enables these women to be seen by clinicians experienced in cervical diseases who are Colposcopy Accredited. Women are assessed within 6 weeks of referral. All women referred via the PRISM pathway must have an in-date cervical smear, a recent infection screen to exclude chlamydia, gonorrhoea, trichomonas and candida infections and a speculum examination must have been performed in primary care. Pregnancy should be considered and excluded as appropriate.

The majority of these women have concomitant symptoms of heavy menstrual bleeding and intermenstrual bleeding; it is beyond the scope of the PCB service to deal with these symptoms. Those with treatable conditions such as polyps, endometritis, cervicitis and ectropion will be offered appropriate treatments within these clinics. Women with no underlying cause for their post-coital bleeding will be referred back to the GP with advice for further referral to Benign Gynaecology if symptoms of HMB and IMB are persistent.

Name

Post.....

PCB WITH NORMAL EXAMINATION AND INVESTIGATIONS BY GP



PCB Post coital bleed

AUB Abnormal uterine bleeding

COCP Combined oral contraceptive pill

IMB Intermenstrual bleeding

TV Trans vaginal

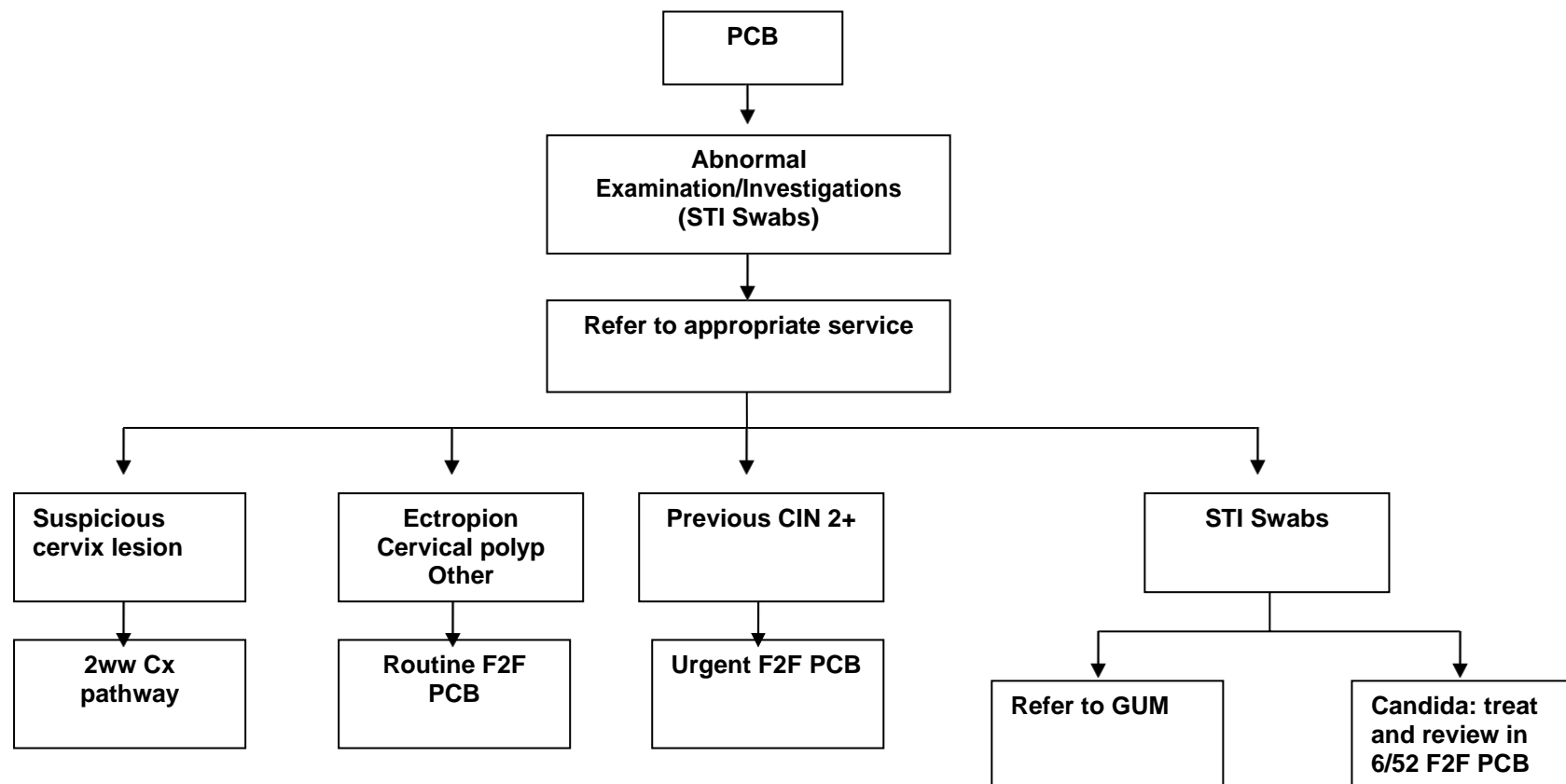
F2F Face to face

F/U Follow-up

USS Ultrasound scan

Clinic for F2F Booking. Colp and Hysteroscopy: H.Ball, V Shesha PMB, A.Banerji PMB

PCB AND ABNORMAL EXAMINATION AND INVESTIGATIONS



Colposcopy only: L.Spence, I.Silina, H.Ball Colp Clinics.

3. Education and Training

e-learning for Sexual and Reproductive Healthcare (e-SRH) supports healthcare professionals in acquiring the relevant knowledge needed for delivering sexual and reproductive healthcare. You will need an e-LfH account. [You can register on their website here](#) to access the [eSRH](#).

4. Monitoring Compliance

What will be measured to monitor compliance	How will compliance be monitored	Monitoring Lead	Frequency	Reporting arrangements
Number of referrals discharged after virtual consultation	Service evaluation Audit	Dr.V.Shesha	6 monthly	Audit Meeting presentation
Number needed to be seen F2F	Service evaluation Audit	Dr.V.Shesha	6 monthly	Audit Meeting presentation
Number diagnosed with Precancer or cancer.	Service evaluation Audit	Dr.V.Shesha	Yearly	Audit Meeting presentation

5. Supporting References

1. Trends in urology, Gynaecology & Sexual Health Volume 13, Issue 4, Article first published online: 25 JUL 2008.
2. NHSCSP Publication No. 20. (2020) Colposcopy and Programme Management: Guidelines for the NHS Cervical Screening Programme. NHSCSP.
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7. Michelle A.L. Godfrey, Manolis Nikolopoulos, Natalia Povolotskaya, Rashna Chenoy, Rekha Wuntakal, Post-coital bleeding: What is the incidence of significant gynaecological pathology in women referred for colposcopy? *Sexual & Reproductive Healthcare*, Volume 22, 2019, 100462, ISSN 1877-5756, <https://doi.org/10.1016/j.srhc.2019.100462>.
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9. Morgan S, Datta S. Intermenstrual and post-coital bleeding. *Obstetrics, Gynaecology & Reproductive Medicine*. 2017;27(12):379-84.
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6. Key Words

Candidiasis, Cervical cancer, Chlamydia, Colposcopy, Ectropion, Gonorrhoea, Hysteroscopy, Polyps, Spotting, Trichomoniasis

The Trust recognises the diversity of the local community it serves. Our aim therefore is to provide a safe environment free from discrimination and treat all individuals fairly with dignity and appropriately according to their needs.

As part of its development, this policy and its impact on equality have been reviewed and no detriment was identified.

EDI Statement

We are fully committed to being an inclusive employer and oppose all forms of unlawful or unfair discrimination, bullying, harassment and victimisation.

It is our legal and moral duty to provide equity in employment and service delivery to all and to prevent and act upon any forms of discrimination to all people of protected characteristic: Age, Disability (physical, mental and long-term health conditions), Sex, Gender reassignment, Marriage and Civil Partnership, Sexual orientation, Pregnancy and Maternity, Race (including nationality, ethnicity and colour), Religion or Belief, and beyond.

We are also committed to the principles in respect of social deprivation and health inequalities.

Our aim is to create an environment where all staff are able to contribute, develop and progress based on their ability, competence and performance. We recognise that some staff may require specific initiatives and/or assistance to progress and develop within the organisation.

We are also committed to delivering services that ensure our patients are cared for, comfortable and as far as possible meet their individual needs.

Development and approval record for this document			
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Approved by:	UHL Gynaecology Governance Committee		Date Approved: March 2022
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October 2021	1		New document
January 2025	2	V Shesha	Removed section' rationale behind PCB virtual clinics' as virtual clinics no longer occurring

Appendix 1 - Post coital bleeding Clinic History sheet

Clinician:

Date

Height

cm

Weight

kg

BMI

Age

Addressograph

Telephone

☐

Face to Face

☐

Referred by: GP

☐

Gynae

☐

Other

☐

GP opinion about the cervix: Suspicious ☐ Polyp ☐ Other ☐

Presenting Complaint(s):

☐ Post Coital bleeding

☐ Intermenstrual Bleeding

☐ Heavy Periods – Regular/Irregular

☐ Unscheduled PVB on HRT

☐ Peri-menopausal bleeding

☐ Postmenopausal PVB

Parity + {__vaginal;__CS}

LMP

Last Smear (Date) _____

Result ____

Previous Colposcopy/LLETZ

Histology

Duration:

Previous episodes:

Contraception:

History to exclude STI: (previous STI, partners, trauma etc)

PGHx (specify if)

PMHx

DHx

PSHx

Allergies

SHx-

Smoking: Never / Ex____yrs. ago / current____per day

Alcohol:____units/week

HVS swab: *result/date*

Signature.....

Chlamydia/GC swab: *result /date*

Name.....

Examination: Date:

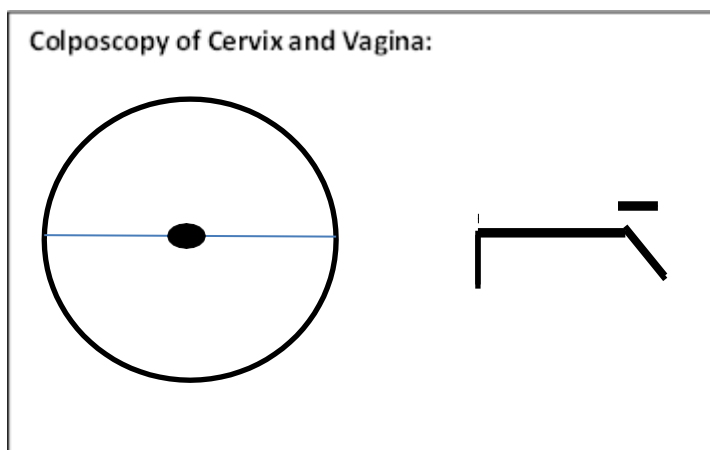
Clinician:

Findings: O/E PA:

PS: V&V -

PV: Uterus- (*if appropriate*)

Adnexae



Investigations

USS: *result/date*

Cervical punch biopsy: *single /multiple*

Pipelle:

Clinical impression:

Follow up:

- ☐ Return for treatment: (Cautery / Ablation / LLETZ)
- ☐ Booked for Hysteroscopy: Urgent / Routine
- ☐ Telephone consultation in _____ weeks
- ☐ Write with results
- ☐ Advised change of contraceptive to _____
- ☐ Refer to general Gynae for _____
- ☐ Discharge to GP

Additional Information:

Signature.....

Name

Post.....