



CHS37 Obtain cervical cytology samples from individuals

OVERVIEW

This standard covers taking cervical cytology samples for screening. The aim of cervical screening is to identify conditions that might otherwise develop into invasive cancer so that preventative treatment can be provided. The sample taker should take care to explain the aim, procedure and possible outcomes of the screening programme and acknowledge and address women's concerns and anxieties. Women and their families should have confidence in the screening programme. The nature, urgency and efficacy of any follow-up actions are dependent upon the reliability of the screening results. It is important therefore that the sample taking procedure results in high quality samples which can be screened effectively. This standard is for qualified practitioners who have completed a recognised training programme for taking samples for cervical screening. This standard does not cover the taking of vault samples. Users of this standard will need to ensure that practice reflects up to date information and policies. Version No 1

KNOWLEDGE AND UNDERSTANDING

You will need to know and understand:

- 1.The legislation which relates to your actions within the healthcare environment including:
 - 1.confidentiality and information sharing
 - 2.the provision of services
 - 3.human rights
 - 4.anti-discriminatory practice
- 2.The statutory and professional standards and codes of practice for your area of work and how to interpret and apply these
- 3. The importance of working within your own sphere of competence and seeking advice when faced with situations outside your sphere of competence
- 4. The importance of applying standard precautions and appropriate health and safety measures and the potential consequences of poor practice
- 5.Local protocols for patient identification
- 6.Local policy and protocol for arranging and working with a chaperone
- 7.Legislation and legal processes relating to consent
- 8. The principles of valid consent, including implied consent and expressed consent
- 9. The relevant national and organisational policies and guidelines on consent
- 10. Why it is essential to maintain the individual's confidentiality

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- 11.The national cervical screening programme guidelines including eligibility for screening
- 12. The importance of complying with the national cervical screening guidelines and the potential consequences of poor practice
- 13. The epidemiology and natural history of cervical cancer
- 14. The risk factors for cervical cancer
- 15. Human papilloma virus (HPV) and its links with cervical cancer
- 16. The effectiveness and limitations of cervical screening
- 17. National and local guidelines for the clinical management of women on the national cervical screening programme
- 18.Colposcopy procedures
- 19. Sources of authoritative information on cervical screening, including evidence based information
- 20. The current and emerging issues, research and developments for cervical screening
- 21.Information and educational resources for practitioners, and methods for accessing them
- 22. The appearance of normal female genital anatomy
- 23. The appearance of benign/harmless abnormalities of female genital anatomy
- 24.An in-depth understanding of the:
 - 1.position of the cervix
 - 2.anatomy and physiology of the cervix
 - 3.cellular structure of the cervix
 - 4.development of and changes in the transformation zone
- 25. The physical manifestations of common genital tract infections
- 26. The presentation and appearance of genital tract malignancy
- 27.How to communicate with individuals to explain procedures and reassure, including those with special needs
- 28. The aim, risks, benefits and limitations of cervical screening and how to explain these to individuals
- 29.Customs and health beliefs of local minority ethnic groups and the implications for taking cervical cytology samples
- 30. Signs of anxieties in women and how to respond to these
- 31. The meaning and implications of different test results, including normal, inadequate and abnormal results, and how to explain these to women
- 32. National and local information materials and how to access these
- 33.Materials and equipment required for taking cervical cytology samples
- 34. Contraindications for the use of equipment e.g. latex allergy
- 35. The factors to take into account in selecting specula and samplers
- 36. The correct storage and type of container for cervical cytology samples
- 37. The national and local call and recall system and fails afe procedures
- 38. The types of information required in taking a history, and why each is required
- 39. The steps you would take to clarify and confirm any relevant information which is ambiguous or missing from the woman's history
- 40. Specific preparation for women prior to taking cervical cytology samples
- 41. Circumstances when it would be inappropriate to take a cervical cytology sample
- 42. Appropriate use of the speculum to visualise the cervix
- 43. How to avoid unnecessary discomfort or distress to women during consultation and sample taking for cervical screening
- 44. The laboratory screening process and the implications for the type, quantity and quality of samples required
- 45.Potential problems in obtaining cervical cytology samples and what you should do in response to these

- 46. The possible adverse reactions that may occur when taking cervical cytology samples and how to respond to these
- 47.Local procedures for reporting incidents and concerns
- 48. The full and complete labelling of samples and the implications of incorrect or incomplete labelling
- 49. The requirements for safe and secure transfer of samples to the laboratory for screening
- 50. When and how to refer women for further opinion, examination or intervention
- 51. The importance of, and procedures and timescales for, emergency referral to an appropriate clinician when malignancy is suspected
- 52. Arrangements and approximate timing for notifying women of their test results
- 53. The importance of recording information clearly, accurately and in a systematic manner
- 54. The importance of confirming the woman's address and arrangements for her to receive her test results
- 55. The information that is required to ensure accurate labelling of samples
- 56. The relevant test request form and the information required
- 57. The information which must be recorded in the woman's notes

PERFORMANCE CRITERIA

You must be able to do the following:

- 1.comply with the national cervical screening programme guidelines
- 2.receive the woman and check her identification details in accordance with local protocols
- 3.clearly explain who you are and your role in the cervical screening programme
- 4.communicate with individuals in a supportive and approachable manner consistent with their:
 - 1.understanding
 - 2.culture and background
 - 3.preferred language and ways of communicating
- 5.ensure that the woman is due for cervical screening and has received a copy of the national leaflet explaining the screening programme
- 6.establish any specific requirements to achieve effective consultation and sample taking
- 7.clearly explain the screening process and possible outcomes, including benefits and limitations
- 8.offer, and arrange if appropriate, the services of a chaperone in accordance with local policy
- 9.answer any questions from individuals accurately and promptly consistent with your professional role and responsibilities, and refer any questions that you cannot answer to the appropriate person
- 10.obtain the woman's valid consent to cervical screening and the sample taking procedure
- 11.take and record a relevant history
- 12.complete the appropriate documentation fully, clearly and accurately

- 13.stop the procedure if the necessary preparations have not been followed by the individual or staff
- 14.ensure the correct positioning of the woman to minimise discomfort and facilitate the sample taking procedure
- 15.carry out the sample taking procedure correctly to obtain an adequate sample
- 16.act with discretion, sensitivity and respect throughout the consultation and sample taking procedure
- 17.ask the woman to indicate if she wishes to stop the procedure at any time
- 18.ensure the woman's privacy, dignity and confidentiality at all times, taking into account the woman's ethnic and religious background which might influence certain aspects of taking cervical cytology samples
- 19.apply standard precautions for infection control and take other appropriate health and safety measures
- 20.recognise and take appropriate action in response to:
 - 1.any condition or behaviour which may signify adverse reactions to the sample taking procedure
 - 2.inability to visualise or sample the cervix
 - 3.any signs of genital tract infection
 - 4.clinical suspicion of malignancy
- 21.preserve and store the sample with due regard for sample quality, safety and security
- 22.carefully remove and dispose of the speculum
- 23.label the sample clearly and correctly, and attach the relevant documentation
- 24.store the sample in the correct place for collection or transport
- 25.report any problems or concerns to the relevant person
- 26.26. check that the woman understands:
 - 1.how and when she will receive her result
 - 2.possible results and follow up
 - 3.how to obtain help, support, advice and further information if required
- 27.keep accurate, complete and legible records of the sample taking procedure and outcomes in accordance with local policies and procedures

ADDITIONAL INFORMATION

This National Occupational Standard was developed by Skills for Health. This standard links with the following dimension within the NHS Knowledge and Skills Framework (October 2004): Dimension: HWB7 Interventions and treatments