Evaluating cytology for the detection of invasive cervical cancer

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Abstract

Objective: To assess the sensitivity, the number needed to screen (NNS) and the positive predictive value (PPV) of cervical cytology for the diagnosis of cancer by age in a screening population.

Methods: A retrospective cohort of women with invasive cervical cancer nested within a census of cervical cytology. All (c. 8 million) women aged 20–64 years with cervical cytology (excluding tests after an earlier abnormality). From April 2007 to March 2010, 3372 women had cervical cancer diagnosed within 12 months of such cytology in England. The sensitivity of cervical cytology to cancer, NNS to detect one cancer and predictive values of cytology were calculated for various ‘referral’ thresholds. These were calculated for ages 20–24, 25–34, 35–49 and 50–64 years.

Results: The sensitivity of at least moderate dyskaryosis [equivalent to a high-grade squamous intraepithelial lesion (HSIL) or worse] for cancer of 89.4% [95% confidence interval (CI) 88.3–90.4%] in women offered screening was independent of age. At all ages, women with borderline-early recall or mild dyskaryosis on cytology (equivalent to ASC-US and LSIL, respectively, in the Bethesda system) had a similar risk of cervical cancer to the risk in all women tested. The PPV of severe dyskaryosis/invasive and glandular neoplasia cytology (equivalent to squamous cell carcinoma and adenocarcinoma/adenocarcinoma in situ, respectively, in the Bethesda System) were 34% and 12%, respectively; the PPV of severe dyskaryosis (HSIL: severe dysplasia) was 4%. The NNS was lowest when the incidence of cervical cancer was highest, at ages 25–39 years, but the proportion of those with abnormal cytology who have cancer was also lowest in younger women.

Conclusions: The PPV of at least severe dyskaryosis (HSIL: severe dysplasia) for cancer was 4–10% of women aged 25–64 years, justifying a 2-week referral to colposcopy and demonstrating the importance of failsafe monitoring for such patients. The sensitivity of cytology for cervical cancer was excellent across all age groups.

Keywords: cervical cancer, cervical cytology, predictive value of tests, sensitivity, specificity, pap test

Introduction

Cervical screening aims to prevent cervical cancer through the diagnosis and treatment of premalignant cervical lesions. Although screening can lead to the early diagnosis of invasive cervical cancer, which is not its primary goal, and the value of cytology to detect cancer (rather than pre-cancerous lesions) has not been studied.

This paper considers the results of a census of cervical cytology; we show the distribution of cytology results, by age, in the general (screening) population and women with cervical cancer. We explore the sensitivity and positive predictive value (PPV) of cer-
cervical cytology at different thresholds for the detection of invasive cervical cancer in the general screening population, as well as the number of women needed to be screened (NNS) to detect one case of invasive cervical cancer.

During the study period (2006–2010) the technology used for screening in England changed. Liquid-based cytology (LBC) was introduced between 2003 and 2008 and will have resulted in cytology slides taken during the study period reported in a combination of conventional cytology and one of the two systems of LBC. In addition, six large cytology laboratories in England were using HPV testing to triage borderline and mild dyskaryosis (equivalent to ASC-US and LSIL, respectively, in the Bethesda system) during most of this period.

Materials and methods

For the results of cytology in women with cervical cancer, we used data from the National Audit of Invasive Cervical Cancers in England.1–3 We studied a retrospective cohort of women who had had cervical cytology taken in the 12 months prior to diagnosis, between April 2007 and March 2010. All cervical cytology was read using the British Society for Clinical Cytology (BSCC) terminology in laboratories subject to accreditation and quality assurance. Data on their screening histories were abstracted from cervical cytology records held on the Exeter Call/Recall System. For results in the general population we used an extract from the Exeter database (taken in October/November 2010) including annual attendances to the screening programme from April 2007 to March 2010.4 This resulted in a census of cytology results. The information in this extract included the women’s age, test result and category of screening invitation (for example routine recall, early recall after an abnormality or surveillance after treatment). In both women with cancer and the general population, we excluded cytology that was taken because of an earlier abnormal result. Note that by considering cytology in a 3-year window and excluding repeat tests, few women will have more than one test in this study. For women with cervical cancer, we define the ‘index test’ as the first non-recall (i.e. not a follow-up test) test result within 12 months of diagnosis. Twelve months was chosen to allow for diagnosis after early (6-month) recall triggered by borderline or mild dyskaryotic cytology (equivalent to ASC-US and LSIL, respectively, in the Bethesda system) while ensuring that the cancer was already present at the time of cytology. Sensitivity analyses taking the index test as the first within 9 and 18 months of diagnosis were also carried out.

The BSCC terminology can be compared broadly with the Bethesda System (TBS) as follows:4 Borderline changes (early recall) includes atypical squamous cells (ASC) and atypical glandular cells (AGC). Borderline, high-grade dyskaryosis not excluded, equivalent to ASC, cannot exclude a high-grade squamous intraepithelial lesion (ASC-H). The term dyskaryosis equates to a squamous intraepithelial lesion (SIL); mild dyskaryosis corresponds to a low-grade squamous intraepithelial lesion (LSIL); high-grade dyskaryosis (equating to HSIL) is defined as moderate or severe dyskaryosis. Separate categories exist for severe dyskaryosis/?invasive and ?glandular neoplasia for squamous cell carcinoma and cervical glandular intraepithelial neoplasia (CGIN) / adenocarcinoma, respectively.

Borderline cytology results were divided into two groups reflecting the associated differing risk of disease. ‘Borderline-high risk’ comprises tests for which immediate referral to colposcopy is recommended (because the high-grade disease could not be ruled out). ‘Borderline low-risk’ comprises tests for which a repeat test at 6 months is recommended.6 In England, although text reports distinguish borderline changes in which high-grade cannot be excluded, result codes on national records do not record subtypes of borderline. Because we did not have access to cytology reports, we used cytology test action codes of ‘suspend’ (i.e. referral to colposcopy) and ‘early-recall’ (i.e. repeat cytology in 6 months) to classify women as ‘borderline-high risk’ and ‘borderline low-risk’, respectively. Borderline samples with a positive HPV triage test and those with changes in endocervical cells are included in the borderline-high risk group as these samples would have triggered an immediate referral to colposcopy. Cytology test action codes were available for all women with cancer and a random sample (i.e. controls from the National Audit) of the general population. The proportion of controls classified as ‘borderline-high risk’ was applied to the census of cytology results to estimate the proportion of the general population.

We estimated the number needed to screen (NNS), the positive predictive value (PPV) and sen-
sitivity for cervical cancer according to three different groupings of cytology test results. This allowed us to assess which women would be likely to benefit from a 2-week referral to colposcopy for further investigation. These were defined as follows: Level 1 includes severe dyskaryosis or worse cytology results; Level 2 includes borderline-high risk, moderate dyskaryosis or worse cytology results; and Level 3 includes all cytological tests that resulted in a referral to colposcopy based on the screening programme protocols prior to diagnosis. Box 1 details the BSCC and Bethesda terminology included in each referral threshold Level.

Statistical analysis

Sensitivities were calculated as the proportion of women with cancer who had a positive test when considering each cytology result. Specificities were

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**Box 1. Referral threshold Levels* for colposcopy with their respective BSCC and Bethesda terminology**

| Level 1 | Level 2 | Level 3 |
|---------|---------|---------|
| BSCC    | Bethesda | BSCC    | Bethesda | BSCC    | Bethesda |
| Severe dyskaryosis | HSIL | Borderline, high-grade dyskaryosis not excluded | ASC-H | Third consecutive inadequate test |
| Severe dyskaryosis | Squamous cell carcinoma | Borderline change in endocervical cells | Atypical endocervical, endometrial or glandular. | First or second mild; second or third consecutive borderline |
| ?Invasive | | | | Un satisfactory for evaluation |
| ?Glandular neoplasia | AGC, favour neoplastic/AIS | High-grade dyskaryosis, moderate or severe | HSIL | Borderline, high-grade dyskaryosis not excluded |
| Adenocarcinoma: endocervical endometrial extrauterine not otherwise specified (NOS) | | | | |
| Severe dyskaryosis | Squamous cell carcinoma | | | |
| ?Invasive | | | | |
| ?Glandular neoplasia | AGC, favour neoplastic/AIS | High-grade dyskaryosis, moderate or severe | HSIL | Squamous cell carcinoma |
| Adenocarcinoma: endocervical endometrial extrauterine not otherwise specified (NOS) | | | | |

*Note: threshold levels are not mutually exclusive. BSCC, British Society for Clinical Cytology; LSIL, Low-grade Squamous Intraepithelial Lesion; HSIL, High-grade Squamous Intraepithelial Lesion; AIS, adenocarcinoma in situ; ASC-H, atypical cells of undetermined significance, cannot exclude a high-grade squamous intraepithelial lesion; ASC-US, atypical cells of undetermined significance; AGC, atypical glandular cells.
calculated as the proportion of women without cancer who had a test result less severe than the relevant cut-off. The PPVs were calculated by dividing the number of cancers diagnosed with a given index test result by the number of cytology tests (in the general population during the same period) with that test result. It was not possible to calculate the PPV for Level 3 cytology as the action code of each cytology test was not available in our extract of the Exeter database. The NNS is the total number of cytology tests divided by the total number of cancers diagnosed after cytology with that result or worse. To calculate the 95% confidence intervals (CIs) for the NNS, a 95% CI for the PPV was calculated assuming the number of cancers diagnosed from the number of cytology tests carried out has a binomial distribution, and the inverse was taken. \(^7\) Confidence intervals for borderline-low risk and borderline-high-risk took into account (by simulation) that the number of cytology tests in each of these categories in the general population was estimated. Analyses were carried out in STATA 12 (StataCorp., College Station, TX, USA).

**Results**

**Cytology results**

There were 3372 women diagnosed with cervical cancer, who had a routine (i.e. one not following a previous action code of early recall or suspend) cytology test within 12 months of diagnosis, between 1\(^{st}\) April 2007 and 31\(^{st}\) March 2010. Over the same period, there were 8,214,754 routine cytology tests in England (Figure 1). The majority of screening age cytology tests were negative (92.2%), 1.3% were Level 2 (moderate dyskaryosis or worse including borderline-high risk), and 0.6% were Level 1 (severe dyskaryosis or worse) (Table 1). The percentage that was negative increased with age (Supplementary Table S1), from 82.0% in women aged 20–24 years to 94.9% in women aged 50–64 years. The proportion of tests that were inadequate remained consistent across age groups, ranging from 2.6% to 3.1% and remained similar over the study period. The percentage that were Level 2 was higher in young women (20–24 years: 3.7%, 25–34 years: 2.4%) than the older age groups (35–49 years: 0.9%, 50–64 years: 0.4%) (Table 2).

**NNS, sensitivity and predictive values**

The NNS to identify one cancer in women aged 20–64 years with Level 2 cytology was 2726 (95% CI: 2630–2826) and the NNS was lowest for women aged 25–39 (1913, 95% CI: 1810, 2024). Trends in the NNS with age are shown in Figure 2a. The NNS is lowest when the incidence of cervical cancer is highest, at ages 25–39 years.

In 78% of women aged 20–64 years with cervical cancer with at least one test within 12 months of diagnosis, the first such test was Level 1 and 89% were Level 2. The sensitivity of cytology to cervical cancer was largely independent of age (Figure 2b).

Overall, the PPV for cervical cancer of a severe dyskaryosis/?invasive report was 33.5% (95% CI: 31.3–35.8%), the PPV of a report of ?glandular neoplasia was 12.4% (95% CI: 11.4–13.5%) and 3.6% (95% CI: 3.4–3.8%) for a report of severe dyskaryosis (Table 1). The PPV of moderate dyskaryosis was much less: 0.66% (95% CI: 0.57–0.75%) (i.e. 1 in 153 women with moderate dyskaryosis had cervical cancer), which was similar to the PPV of a borderline-high risk (0.61%, 95% CI: 0.52–0.71%). The overall PPV of all Level 2 cytology was 2.77% (95%
Table 1. Result of the first routine cytology test in the last 12 months in women aged 20–64 years, with cervical cancer (‘Cancers’) and in the general population (‘Cytology tests’) and the positive predictive value (PPV) of the test result to cervical cancer

| Cytology test result | % cancers diagnosed with test result or worse | Cytology tests | % of all tests | PPV   |
|----------------------|--------------------------------------------|---------------|---------------|-------|
| Negative             | 106                                        | 7 570 265     | 92.2          | 0.00% |
| Inadequate           | 58                                         | 221 956       | 2.7           | 0.03% |
| Borderline – low risk| 118                                        | 203 322       | 2.5           | 0.06% |
| Mild dyskaryosis     | 76                                         | 110 333       | 1.3           | 0.07% |
| Borderline – high risk| 164                                    | 26 672       | 0.3           | 0.61% |
| Moderate dyskaryosis | 215                                        | 32 823       | 0.4           | 0.66% |
| Severe dyskaryosis   | 1589                                       | 43 930       | 0.5           | 3.62% |
| ?Glandular neoplasia | 459                                        | 3701         | 0.0           | 12.40%|
| ?Invasive squamous carcinoma | 587                                    | 1752         | 0.0           | 33.50%|
| Total                | 3372                                       | 8 214 754    | 0.04%         |       |
| Level 1              | 2635                                       | 49 383       | 0.6           | 5.34% |
| Additional value of moderate dyskaryosis and borderline-high risk cytology | 379 | 59 495 | 0.7 | 0.64% |

*First routine cytology test in the last 12 months.

Table 2. Positive predictive value (PPV) of high grade (Level 2) and severe dyskaryosis or worse (Level 1) cytology to cervical cancer by age at diagnosis

| Age group | % cancers diagnosed with test result | % of all Cytology tests | PPV (%)
|-----------|------------------------------------|-------------------------|--------|
| Level 2   |                                    |                         |        |
| 20–24     | 97                                 | 9599                    | 3.7    | 1.0 |
| 25–34     | 1246                                | 57 006                  | 2.4    | 2.2 |
| 35–49     | 1293                                | 34 825                  | 0.9    | 3.7 |
| 50–64     | 378                                 | 7447                    | 0.4    | 5.1 |
| Level 1   |                                    |                         |        |
| 20–24     | 76                                 | 3785                    | 1.5    | 2.0 |
| 25–34     | 1069                                | 27 484                  | 1.2    | 3.9 |
| 35–49     | 1160                                | 14 936                  | 0.4    | 7.8 |
| 50–64     | 330                                 | 3178                    | 0.2    | 10.4 |

*First routine cytology test in the last 12 months.

How good is cytology at detecting invasive cervical cancer?

The PPVs of mild dyskaryosis (0.07%, 95% CI: 0.05–0.09%) and of borderline-low risk (0.06%, 95% CI: 0.05–0.07%) were similar to the prevalence of cancer in all screened women (0.041%, 95% CI: 0.040–0.042%). In other words, the chance of the cancer being diagnosed in the next year in a woman with mild dyskaryosis is only slightly higher than in a randomly selected woman attending screening (1 in 2436).

Women’s first screen

When data for young women were split according to whether it was the woman’s first screen or subsequent screen, the proportion of Level 2 results was slightly higher for first screens (age 20–24 years: first screen 4.0%, subsequent screens 3.6%; age 25–29: first screen 3.2%, subsequent screens 2.7%). However, the PPV of a Level 1 cytology result was substantially higher for first screens than subsequent screens, particularly at age

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20–24 years [first screen 4.51% (95% CI: 3.45–5.78%), subsequent screens 0.69% (95% CI: 0.40–1.10%)] (Table 3). This reflects the fact that prevalent occult cancers are most likely detected on the first screen.

**Results by stage at diagnosis**

FIGO (International Federation of Gynaecologists and Obstetricians) stage was recorded for 91.6% of cancers diagnosed aged 20–64 years. Over half of those without cytology in the 12 months prior to diagnosis had stage 2+ (57.5%) compared to only 11.0% of women with cytology in the 12 months prior to diagnosis (Table 4).

The results were very similar when cytology within 9 or 18 months of diagnosis was considered in place of 12 months (Supplementary Table S2).

**Discussion**

We have shown that in a screening population, regardless of age, the predictive value of cytology for cervical cancer is extremely high for reports of severe dyskaryosis/?invasive carcinoma (34%) and ? glandular neoplasia (12%), and high for reports of severe dyskaryosis (3.6%). Collectively these three groups account for a high proportion (78%) of invasive cervical cancers, and thereby confirm that the current guidelines for a 2-week referral to colposcopy for severe dyskaryosis or worse cytology results are appropriate. Additionally, our results demonstrate the need for failsafe monitoring for women with severe dyskaryosis or worse cytology.

The PPV of severe dyskaryosis or worse (Level 1) cytology was substantially higher on the first cervical screening test in young women than among those who had been screened previously, reflecting the large number of prevalent cancers diagnosed at first screen, although it is important to note that in England 72% of women aged 20–29 years diagnosed with cervical cancer on their first cytology test had stage 1A cancer.8 These results explain why the number needed to screen is lower and the PPV of cytology to cancer is higher among women under the age of 35 years than among older women. The sensitivity of cytology for cervical cancer is excellent; in fact it is much better than the reported sensitivity for CIN3 or worse.9 The sensitivity of cytology to cancer was similar across all age groups.

When interpreting these findings it is important to bear in mind that results refer to the use of cervical cytology for the early diagnosis of cancer, and not the standard use of cytology (i.e. screening for the detection of precursor disease). As such, the NNS...
How good is cytology at detecting invasive cervical cancer?

Table 3. Predictive value of cytology to cervical cancer by age at diagnosis

| Cytology test result* | Tests and cancers between April 07/March 10 | Tests and cancers between April 07/March 10 |
|-----------------------|--------------------------------------------|--------------------------------------------|
|                       | % cancers diagnosed with test result or worse | Cytology tests | % of all tests | PPV | % cancers diagnosed with test result or worse | Cytology tests | % of all tests | PPV |
| **Age 20–24 – first screen** | | | | | | | | |
| Negative              | 0                                          | 100                                      | 63 167 | 83.2 | 0.00% | 0                                          | 100                                      | 148 763 | 81.5 | 0.00% |
| Inadequate            | 0                                          | 100                                      | 2029 | 2.7 | 0.00% | 0                                          | 100                                      | 5480 | 3.0 | 0.00% |
| Borderline – low risk/mild dyskaryosis | 7                                          | 100                                      | 7694 | 10.1 | 0.09% | 7                                          | 100                                      | 21 693 | 11.9 | 0.03% |
| Borderline – high risk/ moderate dyskaryosis | 15                                         | 91                                        | 1698 | 2.2 | 0.88% | 6                                          | 77                                        | 4116 | 2.3 | 0.15% |
| Severe dyskaryosis or worse | 59                                         | 73                                        | 1307 | 1.7 | 4.51% | 17                                         | 57                                        | 2478 | 1.4 | 0.69% |
| Total                 | 81                                         | 75 895                                   | 0.107% | | | 30                                         | 182 530                                   | 0.016% | |
| **Age 25–29 – first screen** | | | | | | | | |
| Negative              | 4                                          | 100                                      | 365 199 | 86.4 | 0.00% | 13                                         | 100                                      | 743 270 | 88.0 | 0.00% |
| Inadequate            | 2                                          | 99                                        | 10 800 | 2.6 | 0.02% | 5                                          | 97                                        | 21 423 | 2.5 | 0.02% |
| Borderline – low risk/mild dyskaryosis | 10                                         | 98                                        | 33 370 | 7.9 | 0.03% | 30                                         | 96                                        | 56 991 | 6.7 | 0.05% |
| Borderline – high risk/ moderate dyskaryosis | 28                                         | 94                                        | 7040 | 1.7 | 0.40% | 65                                         | 89                                        | 12 071 | 1.4 | 0.54% |
| Severe dyskaryosis or worse | 226                                        | 84                                        | 6330 | 1.5 | 3.57% | 310                                        | 73                                        | 10 674 | 1.3 | 2.90% |
| Total                 | 270                                        | 422 739                                   | 0.064% | | | 423                                        | 844 429                                   | 0.050% | |
| **Age 30–34 – first screen** | | | | | | | | |
| Negative              | 2                                          | 100                                      | 100 522 | 89.2 | 0.00% | 11                                         | 100                                      | 915 341 | 91.2 | 0.00% |
| Inadequate            | 5                                          | 98                                        | 3417 | 3.0 | 0.15% | 4                                          | 98                                        | 26 843 | 2.7 | 0.01% |
| Borderline – low risk/mild dyskaryosis | 3                                          | 94                                        | 6200 | 5.5 | 0.05% | 31                                         | 97                                        | 43 336 | 4.3 | 0.07% |
| Borderline – high risk/ moderate dyskaryosis | 14                                         | 92                                        | 1249 | 1.1 | 1.12% | 70                                         | 92                                        | 9163 | 0.9 | 0.76% |
| Severe dyskaryosis or worse | 100                                        | 81                                        | 1308 | 1.2 | 7.65% | 433                                        | 79                                        | 9172 | 0.9 | 4.72% |
| Total                 | 124                                        | 112 696                                   | 0.110% | | | 549                                        | 1 003 855                                  | 0.055% | |

will be much higher and predictive values in this paper much lower than those obtained in a screening context.

The main strength of this study is in the use of cytology results for the entire population of England together with the linked screening histories for
women diagnosed with cervical cancer. Thus, it has wide-ranging validity and is essentially unbiased. The number of borderline cytology tests that were high risk in the screening population had to be estimated from the audit dataset. However, as a result of the introduction of HPV triage in England, the new BSCC terminology guidelines no longer include details on whether or not high-grade disease can be excluded from borderline results, relying instead on the HPV test result. Further, the introduction of LBC testing in England resulted in vigorous training for laboratory and primary care staff just prior to and during the study period, which may have affected the PPV of cytology.

Despite the fact that vaginal cytology was first described as a method of detecting cervical cancer, we are not aware of any studies showing the PPV of modern cervical cytology to cancer. Results presented here demonstrate the usefulness of cytology with a threshold of borderline-high risk or worse (Level 2) for the diagnosis of cervical cancer, regardless of age. In fact, these cytology results were observed in 89% of cancers.

There has been concern that cytology in the presence of invasive cancer is unreliable because the cytology may be inadequate (or even negative). In this study, we found no evidence (in any age group) to suggest this, and additionally show that the proportion of cytology tests that are inadequate in the presence of cancer remains low across all age groups. As some false-negative cytology does occur, it remains important that GPs follow-up women with gynaecological symptoms of possible cervical cancer, and that they make a specialist referral if symptoms persist or worsen.

A substantial benefit of cervical screening comes from early diagnosis of occult cancers. The NNS to detect one cancer using cervical cytology (2700 per cancer) is considerably higher than the NNS to diagnose one colorectal cancer using faecal occult blood tests (516 per cancer). However, the incidence of cervical cancer has been reduced by 46% since the introduction of the national screening programme in England in 1988, and it is difficult to estimate what the incidence would be in the absence of screening.

This study reports cytology read using BSCC terminology in laboratories subject to accreditation and quality assurance, and the conclusions may not generalize to other systems for reporting cytology or to countries with less quality assurance.

### Conclusion

The PPV of severe dyskaryosis or worse cytology for cervical cancer is 4–10% in women aged 25–64 years, clearly justifying a 2-week referral to colposcopy and emphasizing the importance of failsafe monitoring for such patients. The sensitivity of cytology for cervical cancer is excellent across all age groups.

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Contributors: P.D.S. had full access to all of the data in the study and confirms that it is an honest, accurate and transparent account of the study being reported, that no important aspects of the study have been omitted; and that any discrepancies are disclosed. P.D.S. conceived and designed the study. R.L., A.C. and P.D.S. analysed the cytology data.

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Table 4. Screening category by FIGO stage in women aged 20–64 years

| Screening Category | % of Cancers Diagnosed at Stage 1A | % of Cancers Diagnosed at Stage 1B | % of Cancers Diagnosed at Stage 2+ |
|--------------------|-----------------------------------|-----------------------------------|-----------------------------------|
| Not screened       | 93.0                              | 345.0                             | 593.0                             |
| Missed             | 147.0                             | 153.0                             | 92.0                              |
| Cytology detected  | 1836.0                            | 1262.0                            | 326.0                             |
| Total              | 2076.0                            | 1760.0                            | 1011.0                            |

*Has had a cytology test within a year of diagnosis, but no ‘suspend’ action code within a year of diagnosis.
A.W.L., T.H., W.H. and N.D. interpreted the data analysis. All authors contributed to the drafting and revising of the manuscript and have approved the final version. P.D.S. affirms that the manuscript is an honest, accurate and transparent account of the study being reported; that no important aspects of the study have been omitted.

**Ethical approval**

The collection of data as part of audit has been covered since 2003 by section 251 of the NHS Act 2006 re-enacted Section 60 of the Health and Social Care Act 2001 approval (PIAG 1-08(a)/2003). The analysis of anonymised data in this context is considered service evaluation that does not require research ethics approval according to the UK guidelines (NRES).13

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**Supporting Information**

Additional Supporting Information may be found in the online version of this article:

**Table S1.** Result of the first non-recall cytology test in the last 12 months by age group in women with cervical cancer and in the general population and predictive value of the test result to cervical cancer.

**Table S2.** Sensitivity analysis: result of the first non-recall cytology test in the last 9 and 18 months in women aged 20–64 years with cervical cancer (‘Cancers’) and in the general population (‘Cytology tests’) and positive predictive value (PPV) of the test result to cervical cancer.