HPV detection methods

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Abstract. Given the causal relation between a persistent high-risk human papillomavirus (hrHPV) infection and the development of high-grade cervical intraepithelial neoplasia (CIN) and cervical cancer, hrHPV testing has been advocated in addition to cytology for the detection of clinically relevant cervical lesions. HrHPV testing is thought to improve cervical screening algorithms, the management of women with cytologically equivocal smears, and the management of women treated for high grade CIN. In this chapter we discuss different methods for HPV detection and genotyping and their respective applications.

Keywords: Human papillomavirus, polymerase chain reaction, reverse hybridization, hybrid capture, real-time PCR, cervical carcinoma, NASBA, DNA probes, epidemiology

1. Introduction

Cervical cancer is preceded by premalignant lesions termed cervical intraepithelial neoplasia (CIN). Cervical cancer screening largely relies on cytological detection of these premalignant lesions, which can be treated relatively easily and with minor side effects. Given the causal relation between a persistent high-risk human papillomavirus (hrHPV) infection and the development of high-grade CIN and cervical cancer, hrHPV testing has been advocated in addition to cytology for the detection of clinically relevant cervical lesions. HrHPV testing is thought to improve cervical screening algorithms [1], the management of women with cytologically equivocal smears [2,3], and the management of women treated for high grade CIN [4]. In this chapter we elaborate on the different methods for HPV detection and their respective applications.

1.1. HPV phylogeny

HPVs are associated with various benign and malignant epithelial proliferative diseases. Over 100 geno-
determined yet [8]. However, some of them could be considered as potentially oncogenic due to their phylogenetic relationship with known hrHPV types. For example HPV type 67, which initially was isolated from a vaginal intraepithelial neoplasia (VIN) [12], clusters in the A9 group that contains only hrHPVs such as HPV16 [9].

1.2. The need for HPV genotyping

Many HPV tests rely on cocktails of probes representing 13 or 14 of the most common hrHPV types. It has been suggested that the added value of less frequent hrHPV types to such probe cocktails would be small, probably irrelevant for screening programs, and resulting in a substantial decrease in screening specificity [13,14]. However, the increasing need to distinguish individual hrHPV types is illustrated by several current developments in the field of HPV research. Firstly, there is growing evidence that certain hrHPV types confer increased risks for high-grade CIN and cervical carcinomas. This is discussed in more detail in the next paragraph. Secondly, 20–30% of HPV infections involve multiple HPV types and in those cases typing is necessary to determine the contribution of the individual HPV types. Thirdly, there is an increasing interest in prophylactic HPV vaccination (reviewed in [15]), which entails studies of geographical distribution of HPV types and their associated diseases as a basis for prophylactic vaccination programs. Once vaccination has started, HPV typing remains necessary to monitor the changes in prevalence of the type(s) represented in the vaccines as a measure for vaccine efficacy.

HPV genotyping may also have some advantages and perhaps some prognostic value in monitoring of women who have been treated for CIN3 or cervical cancer, although the treatment itself does not depend on the typing result. Demonstrating the same hrHPV type in the post-treatment specimen as in the primary CIN3 lesion may indicate a recurrent hrHPV infection due to incomplete removal of the lesion or disability of the respective woman to clear a particular hrHPV type [16,17] and might require more intense follow-up or more aggressive treatment.

Finally, the number of known HPV types is still increasing. Although novel HPV types are often isolated from malignancies, their oncogenic potential can only be deduced from a combination of in vitro and epidemiological studies, the latter of which should include typing.

1.3. HrHPV types and risk of high-grade cervical lesions

We showed that in the Dutch screened population [18], HPV types 16 and 33 are more prevalent in women with a cytological reading of moderate dyskaryosis or worse and underlying CIN2 or worse, than in women with normal cytology. This suggests that infection with these types confers an increased risk for development of high-grade cervical lesions [19].

Moreover, HPV type distribution in women with cervical carcinoma as compared with cytologically normal women showed that HPV18 is mainly a risk factor for the development of adenocarcinoma (AdCa) and its precursor adenocarcinoma in situ (ACIS), whereas HPV16 is associated with both squamous cell carcinoma (SCC) and AdCa [20]. In line with the abovementioned findings, an increased risk posed by HPV16 and HPV18 for cervical (pre)cancers of the squamous and adeno-histotypes was shown in a prospective screening cohort of 20810 women followed for up to 10 years [21]. In addition, an increased risk for cervical precancer posed by HPV16 but not by HPV18 was shown in a prospective trial of women with equivocal or mild cervical abnormalities [22]. A likely explanation for increased prevalence of HPV18 in invasive carcinoma but not in high-grade precursor lesions, is an association of HPV18 with a cytopathological effect high in the endocervical canal that is likely to be missed by cytology.

These findings strongly suggest that in particular women who are HPV16 or HPV18 positive should be monitored very closely, even if their smears are cytologically normal.

2. HPV detection methods

Because HPV cannot be cultured, all HPV tests currently in use rely on the detection of viral nucleic acids. HPV detection methods can be divided in target-amplification methods and signal-amplification methods. In the following section, technical aspects of the different currently available HPV detection methods are described.

2.1. Target amplification techniques

2.1.1. Consensus primer polymerase chain reaction (PCR)

In most of the PCR-based HPV detection systems, a broad spectrum of HPV types is amplified by consen-
2.1.2. Detection and/or genotyping of consensus PCR systems.

Amplicor [29] systems. Alternatively, they may contain mismatches that are accepted under low-stringency PCR conditions as in the MY09/11 [23] or SPF [27] primers, or sets of overlapping primers as is the case in the PGMY [28] and Amplicor [29] systems.

2.1.2. Detection and/or genotyping of consensus PCR products

For group-specific detection of HPVs without high-resolution typing, an enzyme immuno assay (EIA) can be applied conveniently using cocktails of e.g. HR- or LR-HPV probes [30,31]. Typing of PCR products was traditionally done by means of dot blotting or Southern blotting and hybridization with type-specific oligonucleotides. More recently, reverse hybridization techniques were introduced. These methods rely on the hybridization of labelled consensus PCR products to HPV-type specific oligos immobilized on filters. Examples are reverse line blot (RLB) analysis following MY09/11 [23] or GP5+/6+ [25] consensus PCR, or a line probe assay (LiPa) following SPF PCR [27]. Detection of the hybridized PCR product is done by a colorimetric reaction [23,27] or by chemiluminescence [25], the latter allowing repeated usage of the same filter [25]. Instead of filters, glass microarrays of HPV type-specific probes can also be used [32]. Recently, a quantitative and high-throughput method was developed [33] based on Lumines Narrow suspension array technology. This method relies on detection of consensus primer PCR (GP5+/6+) products with type-specific oligonucleotide probes coupled to fluorescence-labelled polystyrene beads and allows detection of up to 100 different HPV types simultaneously.

Two non-hybridization typing methods following consensus PCR are sequence analysis of the PCR product [34,35] and restriction fragment length polymorphism (RFLP) analysis [36]. RFLP implies the digestion of consensus PCR products with restriction endonucleases, and comparison of the digestion pattern with those of known HPV types. These techniques are useful if unknown types of HPV are present in the specimens, but they have several drawbacks as compared with hybridization methods. For example, RFLP and sequence analysis are not suitable for the detection of infections with multiple HPV types: these will usually give an uninterpretable mix-up of digestion/sequence patterns. In addition, these two methods are less suitable for high-throughput analyses because they are relatively laborious. Finally, RFLP and sequence analysis are less sensitive than hybridization methods because more PCR product is needed to generate a positive signal.

As opposed to typing after consensus PCR, it is also possible to type during the reaction, in real time. Various real-time PCR techniques are available (see also Section 3.2.2), the best known being molecular beacons, the fluorescent 5′ exonuclease assay (e.g. Taqman) and fluorescence energy resonance transfer assays (e.g. LightCycler).

At present the only real-time assay that allows consensus HPV-PCR with simultaneous typing is a molecular beacons assay [37]. However, the multiplicity of this technique is limited because the current generation of real-time thermocyclers do not allow for more than six differentially labelled probes. Hence, for typing of multiple HPV infections, reverse hybridization methods are preferred over real-time assays.

2.1.3. Type-specific PCR

If one is interested in a particular HPV type, type-specific PCR can be applied (described in [25,38] among others). Care should be taken when designing primers, because they may still react with other types if chosen in well-conserved regions. Confirmation of the specificity of type-specific PCRs, as with consensus primer PCRs, can be done by (regular or reverse) filter hybridization or by EIA, but also in real-time [39–41]. A great advantage of real-time PCR assays is the possibility to quantify the HPV in the specimen. Several studies have shown that the amount of hrHPV present in a cervical smear (the “viral load”) as measured by real-time PCR is predictive for the presence or development of high-grade cervical lesions [42–45]; reviewed in [46]. However, to obtain reliable quantification data DNA extraction is usually necessary, thus increasing the work load.

2.1.4. mRNA amplification

Several recent studies have shown that hrHPV testing can also be done through detection of the viral mRNA. The most relevant transcripts to look for are those encoding the viral oncoproteins E6 and E7. It is hypothesized that the presence of viral E6/E7 mRNA in a cervical smear has a better positive predictive value for high-grade cervical lesions than the presence of viral DNA, because the E6/E7 mRNA represents an active
infection with cell-transforming potential whereas the viral DNA may also be present in clinically irrelevant conditions.

Detection of viral mRNA in cervical smears can be done by reverse-transcriptase (RT-) PCR [47] or by nucleic acid sequence-based amplification (NASBA) [48]. For the latter assay type, a commercially available system was recently developed that detects E6/E7 transcripts from the five most common hrHPV types 16, 18, 31, 33 and 45 [49–53]. In hospital-based populations, hrHPV mRNA detection showed a better specificity for the detection of high-grade cervical lesions than hrHPV DNA detection [51,52]. However, this method still needs to be validated for population-based screening and triage of women with smears showing minor cellular abnormalities.

An alternative method relying on hrHPV transcription is the detection of HPV-human fusion transcripts that arise from hrHPV genomes integrated into the host cell genome. This method was designated “amplification of papillomavirus oncogene transcripts” (APOT) [54].

Although this method is rather laborious and therefore not suitable for large-scale hrHPV testing, it can give information that is not obtained using other mRNA amplification techniques. On the one hand, it reveals viral integration, an event that is specific for a real precancerous lesion as opposed to a productive viral infection [55]. On the other hand, it allows the identification of patient-specific fusion transcripts. This may be useful in monitoring of women who have been treated for high-grade lesions, because it will help to determine whether a high-grade lesion occurring after treatment is a recurrence or even a metastasis of the original lesion [56].

2.2. Signal-amplification techniques

2.2.1. Liquid-phase signal amplification techniques

The best known technique in this category is the commercially available, FDA-approved and clinically validated Hybrid Capture 2 (HC2) method [57]. This method uses a mixture of full-length RNA probes representing hrHPV types 16, 18, 31, 33, 35, 39, 45, 51, 52, 56, 58, 59, and 68. Prior to the test, the clinical samples are heat-alkaline-denatured. Hybridization of one or more of the probes to HPV DNA present in the samples is detected by peroxidase-labelled antibodies that recognize the RNA/DNA hybrid, and visualized by chemiluminescence. The analytical sensitivity of this method is 1 pg/ml of cloned HPV16 DNA, which corresponds to approximately $10^5$ HPV16 genome copies. This sensitivity is lower than that of most target-amplification methods. Because a mixture of probes is used, HC2 is at present not suitable for high-resolution typing. Some cross reactivity of the HC2 probes with HPV types not represented in the probe mix, including non-oncogenic HPVs, has been described [58].

2.2.2. Morphological signal-amplification techniques

In addition to the abovementioned methods, hrHPV detection can be performed by DNA in situ hybridization (ISH) to cytological slides [59,60] and histological preparations [61]. This can be achieved by fluorescent detection [61] or colored substrate deposition and bright field microscopy. The relatively small size of the HPV genome (7.8 kb) and thus of the probe precludes direct detection of hybrids in case of low viral genome copy numbers and therefore some type of signal amplification is generally necessary. A commercially available HPV ISH system uses an indirect biotin-streptavidin method (Ventana Inform HPV) which at present lacks sufficient sensitivity to detect high-grade cervical lesions [60]. Alternatively, tyramide signal amplification (TSA), also known as catalysed reporter deposition (CARD) can be used, both in fluorescent [61,62] and bright field [62] applications. Also for the CARD method, a commercially available system exists (Dako GenPoint). CARD greatly enhances sensitivity, but in general HPV ISH is too laborious to be used in high-throughput HPV testing.

3. Clinical specimens for HPV detection

The kind of clinical specimens available will determine the choice of the HPV detection method. Most PCR methods mentioned above can be done with any type of clinical material, including formalin-fixed, paraffin-embedded tissue and Pap-stained archival smears. The recent focus on liquid-based cytology (LBC) for cervical cancer screening has initiated numerous studies evaluating the feasibility of different HPV detection methods on cell suspensions destined for LBC, or cell suspensions remaining after LBC preparation. Some of the pros and cons of different types of clinical specimens are discussed below.
3.1. Clinical specimen types suitable for target amplification methods

3.1.1. Specimen types for PCR-based methods

PCR-based HPV DNA detection methods can generally be applied to all kinds of clinical specimens, provided the nucleic acids contained within are not heavily degraded or cross-linked and no putative PCR-inhibiting substances (e.g. organic solvents) are present. Notably, the nucleic acids in formalin-fixed, paraffin-embedded (FFPE) tissue are only amplified in a reliable manner when PCR products are generated that are smaller than 150 bp due to fixative-induced cross links [63]. On the other hand, this type of specimen generally does not contain large amounts of PCR-inhibiting substances, as demonstrated by the fact that a relatively crude extract can be used in the PCR without laborious extraction protocols [64].

As opposed to FFPE material, cervical smears collected in LBC media may show excellent preservation of cell morphology and integrity of nucleic acids. However, to achieve this preservation they usually contain relatively high amounts of organic solvents which will inhibit the PCR and need to be removed by careful extraction. An exception are smears collected in Digene Universal Collection Medium (UCM) [65,66], which can be used directly in the PCR after a simple dilution and freeze-thaw protocol (Hesselink et al., submitted).

Pap-stained archival cervical smears can also be used for PCR purposes, but they also need extraction of nucleic acids to remove remnants of dyes. In addition, the nucleic acids in this type of specimen may also be degraded due to fixation artefacts. For all above-mentioned samples, careful analysis of the nucleic acid quality is recommended to exclude false negative HPV test results due to DNA degradation and/ or PCR inhibition. This can be achieved e.g. by a PCR specific for a human single-copy gene [67].

For most reliable PCR results, it is recommended to standardize the amount of input DNA in the PCR. It is self-evident that this can only be done using purified nucleic acids.

3.1.2. Specimen types for mRNA amplification methods

Until recently, hrHPV mRNA testing on cervical smears was hampered by the lack of suitable collection media. The poor RNA quality of samples collected in solutions such as PBS precluded transcription analysis. However, use of LBC is increasing and this is accompanied by the introduction of preservation media that not only safeguard cellular morphology but also the integrity of DNA and mRNA. For example, recent studies have shown that cervical smears collected in PreservCyt LBC medium are of sufficient quality for HPV RT-PCR [68] and NASBA analysis [69].

Alternatively, cervical smears could be collected in a dedicated RNA-preservation buffer [51], but in general these contain chaotropic salts such as guanidinium isothiocyanate and thus destroy cellular morphology.

When hrHPV mRNA is to be detected in cervical biopsy specimens, these should preferably be snap-frozen to guarantee mRNA integrity. However, several groups claim successful mRNA extraction from FFPE material (reviewed in [70]). Provided the use of proper quality controls, this is of course a very interesting source of material, especially for retrospective studies or in situations where snap-freezing is not an option.

3.2. Clinical specimen types suitable for signal-amplification methods

The Digene HC2 method is FDA-approved for use on cervical smears and cervical biopsies collected in Digene’s Sample Transport Medium (STM). In addition, the FDA-approval of HC2 includes use on cervical smears collected in Cytyc’s ThinPrep medium. However, because this medium preserves cell integrity as opposed to STM, ThinPrep smears require an additional pre-treatment to obtain full cell lysis and release of nucleic acids [71]. With some modifications of the standard protocol, HC2 can also be used on the residual material of cervical smears collected in SurePath fluid (i.e. the suspension that remains after the LBC slides have been prepared) [72–74].

An intermediate between the commercially available LBC media that preserve cell morphology but require pre-treatment steps prior to HC2, and Digene’s STM that does not require pre-treatment but destroys cell morphology, is Digene Universal Collection Medium (UCM). This medium can be used for sample transport [66], does not require additional steps prior to HC2 analysis, and because it preserves cell morphology it also allows LBC [65].

4. Considerations regarding test sensitivity and specificity

4.1. Considerations for primary screening and triage

In general, the analytical sensitivities of HPV tests depend on the test system (target amplification ver-
sus signal amplification). The sensitivity of the HC2 method is approximately 5,000 copies of the HPV genome per reaction well according to the manufacturer. PCR-based techniques have a much higher analytical sensitivity than HC2, with a slight variation between the methods (reviewed in [46]). For most PCR assays, less than 100 to 1000 HPV genome equivalents in a reaction tube are sufficient to generate a positive PCR signal and to enable typing.

Despite these high analytical sensitivities, currently available hrHPV tests miss a subset of high-grade CIN lesions [75,76]. This may be due to several factors. For PCR-based methods, modification or loss of primer binding sites due to viral integration or naturally occurring sequence variants may be a problem. For HC2 where whole-genome probes are used this is generally not an issue, but then the lower analytical sensitivity of HC2 might preclude detection of lesions with low viral loads. The ideal hrHPV test used in primary screening should combine the high analytical sensitivity of PCR assays with the possibility to detect all configurations and variations of the hrHPV genome, including integrated genomes. Because the intact E6/E7 region of the hrHPV genome is necessary for cellular transformation [77] and is invariably retained upon integration [78], a PCR system detecting this region would be the method of choice [79]. It is self-evident that amplification primers should then be chosen in such a way that they cover all known HPV genomic variations.

On the other hand, a very sensitive hrHPV test consequently has a low clinical specificity, and as a primary screening tool it would result in a substantial increase in referrals for colposcopy and repeat smears. In the Dutch screening population, approximately 5.0% of the women is hrHPV DNA-positive by the GP5+/+6+-system [18]. Because even women with hrHPV DNA-positive normal cytology have a 210 times greater risk to develop CIN 3 compared to women with hrHPV negative normal cytology [80], all 5.0% hrHPV DNA-positive women would need follow-up. However, only 10% of them will actually develop CIN3 [81]. This calls for additional stratification of women who have a positive hrHPV DNA test, which can be achieved for example by viral load analysis or HPV mRNA analysis (see Sections 2.1.3 and 2.1.4).

### 4.2. Considerations for situations not involving triage

For several research questions, clinical specificity of the hrHPV test is not an issue as it is in population-based screening and triage. Methods with high analytical sensitivity are especially suitable for studies in which any HPV infection should be detected, irrespective of its clinical relevance. Examples are epidemiological studies, and studies to determine vaccination efficacy. It is self-evident that HPV genotyping is desired in such studies.

### 5. Conclusions

As seen in the previous paragraphs, the aim of the study and the clinical material available will determine the choice of the HPV detection method. In general, consensus PCR followed by reverse hybridization is very sensitive and gives the most extensive typing information for many kinds of clinical specimens, including those containing multiple infections.

In general, it is to be expected that future amplification systems will become faster and more automated.

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