Can HPV Selfy be considered as a clinically validated HPV test for use in cervical cancer screening?

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In primary cervical cancer screening, it is crucial to use only hrHPV tests that are clinically validated according to international guidelines in order to reduce the risks of missing relevant disease and of over-treatment. In the recent J Transl Med [1] paper, Avian et al. concluded that the HPV Selfy assay (Ulisse BioMed, Trieste, Italy) fulfils international validation criteria for hrHPV testing on clinician-collected cervical samples (Meijer guidelines) [2] as well as by extension on self-collected vaginal samples (VALHUDES) [3]. Our perception is that the study by Avian et al. has certain limitations that are worthy of consideration and which may call into question certain conclusions.

Validation requires an appropriately composed study population comprising a sufficient number of diseased subjects, derived from a continuous screening population or from a clearly described selection of CIN2+ cases and < CIN2 controls [4]. Avian et al. compiled cervical specimens for testing with HC2 (standard comparator test) and with the new HPV Selfy (index test) [1], but it remains unclear how the study population was composed. With 98 CIN2+ and 791 ≤ CIN1 subjects it was obviously not a continuous screening population, so more granularity on this would have been welcome. Additionally, detail on how non-disease was defined, which is essential for the evaluation of clinical specificity, was lacking.

The reported absolute sensitivity for CIN2+ of the HC2 comparator test was 82.7%, which was substantially lower than the sensitivities observed in validation studies following the VALGENT or Meijer protocols included in a meta-analysis (Fig. 1) [5]. This may raise suspicion of a certain degree of histological over-classification. Nonetheless, we verified the data matrices in Table 2 in Avian et al. [1] and confirm the correctness of the non-inferiority statistics (Table 1).

The claim that HPV Selfy on self-samples was non-inferior to clinician-collected samples was flawed by critical statistical errors. The number of subjects with discordant self+/clinician− and self−/clinician+ results (b and c cells in Table 4, in Avian et al. [1]) in the recommended formula for comparison of matched proportions were switched yielding reported p values < 0.05. Correct data entry would have generated non-inferiority p values 0.35 and 0.81 for sensitivity and specificity, respectively. The corresponding relative sensitivity and relative specificity for CIN2+ and 95% confidence intervals (not reported by authors) were 0.92 (95% CI 0.81–1.00) and 0.97 (95% CI 0.95–0.99), respectively, indicating non-significantly lower sensitivity and significantly lower specificity of HPV Selfy on self-versus clinician-collected samples.

Collaborations between science and industry are instrumental to advance clinical research, however contractual independency of researchers and autonomy of publication enhance scientific credibility. We observe that sixteen of thirty six authors (including the first and last) of the JTM paper are affiliated with the manufacturer of the assay. In the 2020 list of validated HPV assays
Fig. 1  Sensitivity of the HC2 assay (standard comparator used in validation of hrHPV assays) in studies included in the 2020 list of HPV assays validated for cervical screening [5] that applied the Meier [2] or VALGENT [4] validation protocols (on top) or included in the study of Avian et al. [1] (at the bottom).

| Study               | Index test        | Design | Estimate (95% CI) |
|---------------------|-------------------|--------|-------------------|
| In 2021 list        |                   |        |                   |
| Heideman, 2011      | Cobas 4800        | Meijer | 0.92 (0.82, 0.96) |
| Carozzi, 2011       | Abbott RT hrHPV test | Meijer | 0.98 (0.92, 0.99) |
| Lloveras, 2013      | Cobas 4800        | Meijer | 0.98 (0.91, 1.00) |
| Ejegod, 2014        | BD Onclarity      | Meijer | 0.94 (0.89, 0.97) |
| Polman, 2017        | HPV-Risk          | VALGENT| 0.96 (0.90, 0.98) |
| Ostrbenk, 2018      | Anyplex HR        | VALGENT| 0.96 (0.90, 0.98) |
| Dhillon, 2021       | Alinity           | VALGENT| 0.96 (0.91, 0.98) |
| Subtotal (I² = 0.0%, p = 0.66) |                   |        | 0.96 (0.94, 0.97) |
| Current study       |                   |        |                   |
| Avian, 2022         | HPV Selfy         | Mixed  | 0.83 (0.74, 0.89) |

Heterogeneity between groups: p < 0.001
Table 1  Computation of the relative specificity to exclude cervical intra-epithelial neoplasia of grade 2 or worse of Selfy on self-samples (SS) vs clinician-taken samples (clin) and non-inferiority statistics

| Correct statistic | Selfy clin— | Selfy clin+ |
|-------------------|-------------|-------------|
| Selfy SS—         | 708         | 16          |
| Selfy SS+         | 37          | 30          |
|                   | 745         | 46          |
| Specificity Selfy SS | = 724/791 = | 91.5%       |
| Specificity Selfy clin | = 745/791 = | 94.2%       |
| Relative specificity SS/clin | 0.97 | (95% CI 0.95–0.99)* |
| T non inferiority | —0.86       |             |
| p non-inferiority | 0.81        |             |

| Wrong statistic (b and c cells switched in the abcd matrix) | Selfy clin— | Selfy clin+ |
|------------------------------------------------------------|-------------|-------------|
| Selfy SS—                                                  | 708         | 16          |
| Selfy SS+                                                  | 37          | 30          |
|                                                             | 724         | 46          |
| Specificity Selfy clin                                      | = 745/791 = | 94.2%       |
| Specificity Selfy SS                                        | = 724/791 = | 91.5%       |
| Relative specificity clin/SS                                | 1.03        | (95% CI 1.01–1.05)* |
| T non inferiority                                           | 6.60        | <0.0001     |
| p non-inferiority                                           |             |             |

In italics: non-inferiority statistic reported by Avian et al. [1] which was due to erroneous switching the values 37 and 16. In fact this reported statistic reflects that Selfy on clin samples is not inferior to SS samples.

[5], assays evaluated by test developers were downgraded to "partially validated" if all other validation criteria were fulfilled. This principle may also apply on the HPV Selfy assessment [1]. We recommend test developers, HPV experts and collaborating epidemiologists or statisticians to design validation studies according to internationally established protocols and evaluation methodologies. Journal editors should take this advice into account as well.

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MA wrote the initial manuscript, all other authors critically revised the manuscript. All authors read and approved the final manuscript.

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Declarations

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Competing interests
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