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2022 Medical Research Day Posters

Oct 13th, 12:00 AM

# Comparing Roche Linear Array® to Next Generation Sequencing for HPV Genotype Identification

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### **Recommended Citation**

Konur, Luke; Winters, Ashley; Williams, Brock; Luo, Meng; Baddoo, Melody; and Cameron, Jennifer, "Comparing Roche Linear Array® to Next Generation Sequencing for HPV Genotype Identification" (2022). *Medical Research Day*. 44.

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## **Presenter Information**

Luke Konur, Ashley Winters, Brock Williams, Meng Luo, Melody Baddoo, and Jennifer Cameron

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Comparing Roche Linear Array® to Next Generation Sequencing for HPV Genotype Identification LSU Health

NEW ORLEANS School of Medicine

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Sample Number

1

2

3

4

5

6

7

٩

10

11

12

13

14

15

16

17

18

Positive by LA only

89

35, 52, 84

55, 62, 73

42.61.68

16, 51, 83, 84

45, 58, 70

62

62.89

54, 62, 84

58

42

61

Positive by NGS only

33

6.53

6, 16

81, 85

53

54, 61, 83

6, 31, 81, 83

16

16

44, 85

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**Objective** 

The Objective of this work is to compare the

performance of Next Generation Sequencing with

MY09/11 Primers against the Roche Linear Array.

**Genotype Results** 

HPV enotyp

18

31

33

42

45

51 52

53 54

54

58

61

62

68

70

71

73

81

83

84

85

89

Table 2: Results of LA, NGS, and both for each genotype found in this work. Red genotypes are undetectable by LA

Positive by both

16

71

53

6. 31. 53

18

31

16, 54

61, 83

16

62, 81

16, 18, 42

18, 53, 70

81

83

Sample Positive LA

by

Samples Positive by

oth LA and NGS

Samples Positive by NGS

## Introduction

#### **HPV in Cervical Cancer**



Figure 1: Progression of human cervical cancer after HPV infection<sup>1</sup>



Figure 2: Phylogenetic tree of Alpha HPV family<sup>4</sup>. HR: High-oncogenic risk; LR: Low-oncogenic risk.

#### Linear Array

- The Linear Array Genotyping Test (LA) from Roche is a colorimetric reverse line blot hybridization array that detects 37 genotypes of HPV and has recently been discontinued.5
- The LA has been the gold standard assay used in research for HPV genotyping over the past two decades. Cross-hybridization between
- genotypes has been reported and could result in misidentification of HPV genotypes in a patient sample.5

#### Next Generation Sequencing

- Newer high-throughput assays like Next Generation Sequencing (NGS) have broadened the variety of HPV types that can be detected.
- NGS outputs sequence data that can be aligned with a continually updated library of reference genomes, allowing no lag time between new genotype discovery and its addition to the sequence library.
- The use of the MY09/11 degenerate primer set allow for nonspecific amplification of HPV late gene 1(L1) regions and sequencing of amplicons through NGS.<sup>2</sup>
- These primers target a 450 base pair region that is highly conserved in the L1 gene of a wide range of genital HPV types.2

Human papillomavirus (HPV) is known to be a causative agent in most cervical neoplasia, thus identifying HPV in a patient's cervical swab is an important step in detecting risk for cervical cancer and preventing further development.<sup>2</sup>

Over 200 genotypes have been found to date, of which 30-40 have a role in Cervical Cancer, with high-risk types being mos prevalent in cervical dysplasia.<sup>3</sup>

> Table 1: Genotypes found by LA NGS, or both for each individual sample. Red genotypes are undetectable by LA

- · A total of 23 HPV types were found by LA and 16 HPV types were found through NGS.
- There was an average concordance between LA and NGS of 92.01% (including positive and negative results).
- NGS found 2 HPV types not detected by LA, HPV 44, and 85, highlighted in red in Tables 1 and 2
- LA found HPV 35, 45, 51, 52, 55, 58, 68, 73, 84, and 89 infections that were not detected by NGS.

#### Methods

- DNA extracts from cervical swabs, collected previously from patients in an HPV observational study.
- The samples were tested by the Linear Array Genotyping Test (Roche Diagnostics, according to manufacturer specifications. Next, HPV libraries were prepared using MY09/11 primers, the samples
- were sequenced using the Illumina MiSeq platform and sequence reads aligned to HPV reference genomes using custom Perl scripts.
- Best concordance between LA and NGS was found when a read count threshold of  $\geq$  20 reads was applied to NGS output.
- Data from NGS was then compared to Linear Array results using Excel and Graphpad Prism.

This work was supported by the Center of Biomedical Research Excellence (CoBRE) in Translational Viral Oncology (CTVO) funded by the NIH NIGMS, award # P20GM121288.

## **Concordance vs. Mismatch Count**

Mean Concordance Percentage vs Mismatch Number



Figure 1: Comparison of concordance for each HPV genotype against number of mismatches between HPV reference consensus sequence and MY09/11 primer sequence.

- First the sequence of the primer binding region for each HPV
- genotype was compared to the primer set, and the number of nucleotide mismatches summed(see example).
- Then, the concordance of each HPV genotype was plotted against this mismatch number. with concordance of types with same mismatch number averaged.
- Simple linear regression yielded no significant trend in this comparison.

Example Alignment for HPV 31 MYII MYII - 5'-GCM CAG GGW CAT AAY AAT GG-3' HPV 3I - 5'-GCT CAG GGA CA<mark>C</mark> AAT AAT GG-3'

MY09 - 5'-GAT CAG TWT CCV YTK GGA CG-3' HPV 31 - 5'-GAT CAG TTT CCA CTG GGT CG-3'

This sample alignment shows the primer sequence first with degenerate primers, then the reference genome consensus sequence. Mismatches are highlighted in yellow. Type 31 had a total of 4 mismatches, and it had a concordance of 94.44%. Degenerate primers: M = A/C; W = A/T; Y = C/T; Degenera K = G/T .

- Conclusion
- NGS with MY09/11 primers were able to identify two genotypes not detected by linear array, HPV 44 and 85.
- HPV 85 is classified as High-risk, and HPV 44 is classified as low risk, so these are important infections that are not detected by LA.
- Cross-hybridization has been reported possibly explaining why LA detected genotypes not found by NGS in individual samples.
- The lack of correlation between mismatch number and concordance could be due to low sample number in this work.
- Perhaps, in a larger sample size this correlation could be seen as a better match between primer sequence and genotype sequence should lead to better amplification, and therefore better concordance.
- Cross-hybridization could also be confounding this correlation. Overall, the NGS platform circumvents the cross-hybridization of LA as well as allows for the detection of a broader range of HPV types.

#### References

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ARRAY H Roche Figure 3: Roche LA genotyping kit reference guide and LA strip.6