



COMPARISON OF HIV DRUG-RESISTANT MUTANT DETECTION BY NGS WITH AND WITHOUT UNIQUE MOLECULAR IDENTIFIERS (UMI)

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Sensitive, Scalable and Affordable Assays are needed for HIV Drug Resistance Testing

| | Sanger Sequencing | Next Generation Sequencing (NGS) |
|--------------------|--|---|
| Sensitivity | Cannot detect minor variants (>20%) | High sensitivity for minor variant detection (?<20%) |
| Scalability | Low throughput Limited potential for automation | Sample multiplexing - potential for automation |
| Cost | High cost per sample (>100 USD) | Reduced cost per sample (<50 USD) |
| Validation | Widely Validated; adapted into kits | Sensitivity, specificity and reproducibility unknown |

(Table adapted from Inzaule et al. The Lancet. 2016)

The effects of sequencing error and variation of sampling depth on the accuracy of mutant detection by NGS are not well defined.

Unique Molecular Identifiers (UMIs) may be used to reduce sequencing artifacts and quantify the true sampling depth.

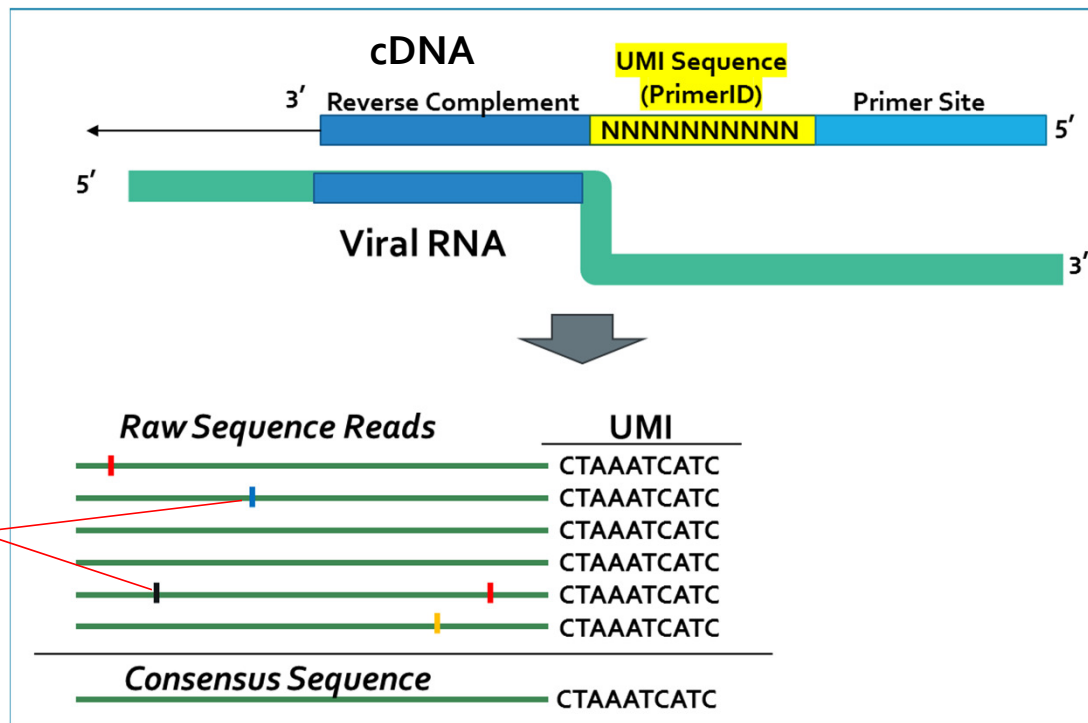


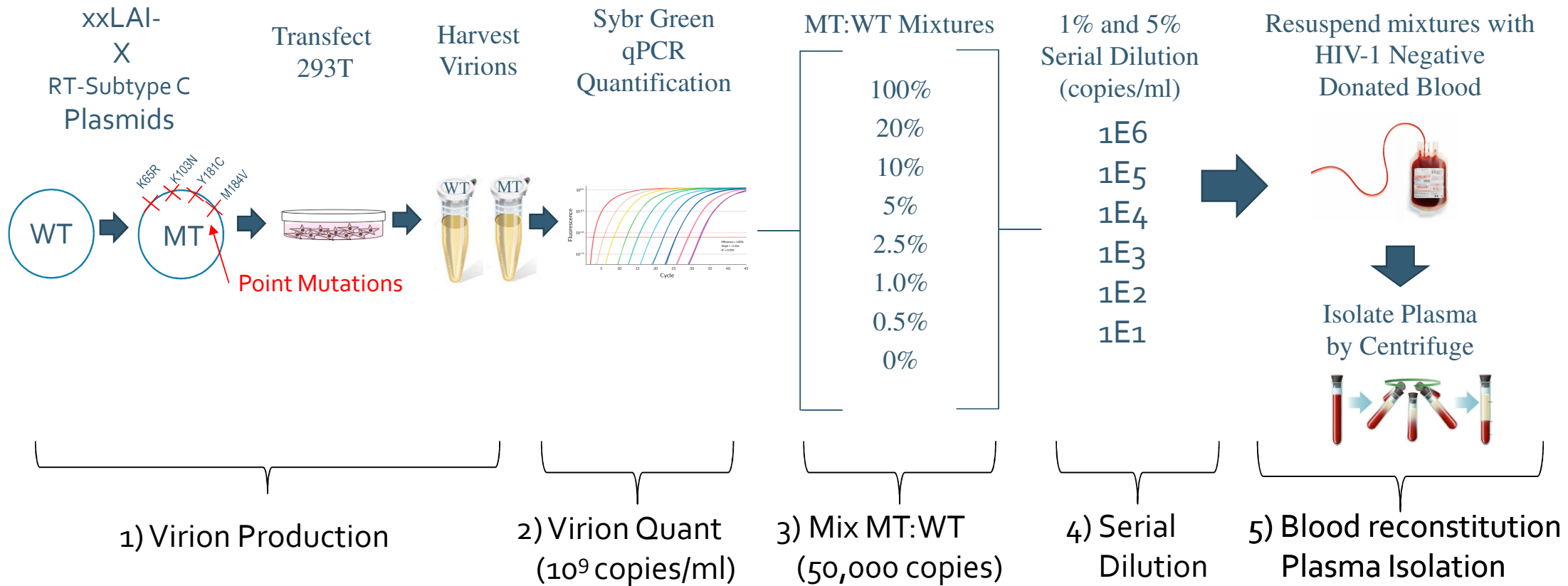
Figure adapted from <https://tcs-dr-dept-tcs.cloudapps.unc.edu/> (Jabara, et al. PNAS. 2011 and Zhou et al. JVI 2015)

OBJECTIVE

Perform a comparative assessment of UMI- and non-UMI-based NGS assays for mutant detection and confident analysis of low frequency HIV DR.

- 1) The sensitivity and specificity**
- 2) The limits of detection**
- 3) The accuracy and precision with clinical samples**

METHODS: Design of the Recombinant HIV Drug Resistant Mixture Panel



METHODS: NGS Library Prep and Data Analysis

All NGS libraries were constructed from the RT region of HIV-1 using the ultrasensitive single-genome sequencing method (Boltz et al. 2016).

non-UMI-based NGS Analysis

- PASEq v1.4 (<https://www.paseq.org>)
- 10,000 raw read coverage with >30 Miseq quality filter
- Minor allele frequencies were reported from the "Amino acid Variants .CSV" file

UMI-based NGS Analysis

- Zhou method was used for consensus building. (Zhou et al. JVI 2015)
- 80% homologous UMI binned reads required for consensus; allowing for 1 mismatched codon
- 298 and 54 UMI consensus sequences were required to call 1% and 5% minor variants respectively (95% confidence)

Similar false positive mutation rates in UMI and non-UMI-based Wildtype samples (Specificity)

| False positive Mutation Frequencies | non-UMI-based NGS (n=8) | UMI-based NGS (n=17) |
|--|--------------------------------|-----------------------------|
| Ave Number | 482 | 68 |
| Mean | 0.06% | 0.08% |
| Median | 0.02% | 0.05% |
| Min | 0.01% | 0.01% |
| Max | 0.46% | 0.46% |

Higher false negatives result with non-UMI NGS at lower viral RNA input and UMI NGS at high RNA input

| Expected (% MT) | Viral Load (cp/ml) | non-UMI-based NGS % Detected | | | UMI-based NGS % Detected | | | # UMI |
|-----------------|--------------------|------------------------------|----------------|----------------|--------------------------|----------------|----------------|-------|
| | | K103N | Y181C | M184V | K103N | Y181C | M184V | |
| 5% | 1.00E+06 | 3.8 | 3.8 | 3.8 | 0.6 | 1.5 | 1.5 | 4308 |
| | 1.00E+05 | 4.0 | 4.0 | 4.0 | 1.3 | 3.2 | 3.2 | 24991 |
| | 1.00E+04 | 3.9 | 4.0 | 4.0 | 4.0 | 4.7 | 4.7 | 2149 |
| | 1.00E+03 | 4.8 | 4.8 | 4.8 | 3.1 | 3.1 | 3.1 | 261 |
| | 1.00E+02 | 0.8 | 0.9 | 0.9 | <BCO | <BCO | <BCO | 16 |
| | 1.00E+01 | 0 ^a | 0 ^a | 0 ^a | <BCO | <BCO | <BCO | 0 |
| 1% | 1.00E+06 | 0.9 | 0.9 | 0.9 | 0 ^a | 0 ^a | 0 ^a | 5010 |
| | 1.00E+05 | 0.9 | 0.8 | 0.8 | 0 ^a | 0.7 | 0.7 | 20848 |
| | 1.00E+04 | 0.8 | 0.8 | 0.8 | 0.6 | 0.7 | 0.7 | 2325 |
| | 1.00E+03 | 1.6 | 1.6 | 1.6 | 0.8 | 0.8 | 0.8 | 266 |
| | 1.00E+02 | 0 ^a | 0 ^a | 0 ^a | <BCO | <BCO | <BCO | 19 |
| | 1.00E+01 | 0 ^a | 0 ^a | 0 ^a | <BCO | <BCO | <BCO | 3 |

Below Biological Cutoff (<BCO)

0^a = false negative detection; <BCO = below biological cutoff (insufficient UMI counts)

non-UMI NGS false negative detection with inefficient PCR from polymorphic primer binding sites

HIV RNA Input = 10,000 copies/sample

| # of PCR FWD primer mismatches | Expected Allele Frequency (% MT) | non-UMI-based NGS detected allele frequency (%) | UMI-based NGS detected allele frequency (%) | # UMI |
|--------------------------------|----------------------------------|---|---|-------|
| 0 | 5.0 | 7.0 | 4.0 | 1213 |
| | 1.0 | 1.5 | 0.9 | 1447 |
| 2 | 5.0 | 7.7 | 10.7 | 215 |
| | 1.0 | 0 ^a | <BCO | 248 |

0^a = false negative detection; <BCO = below biological cutoff (insufficient UMI counts)

METHOD: Sensitivity and specificity of non-UMI NGS analysis vs UMI-based consensus building.

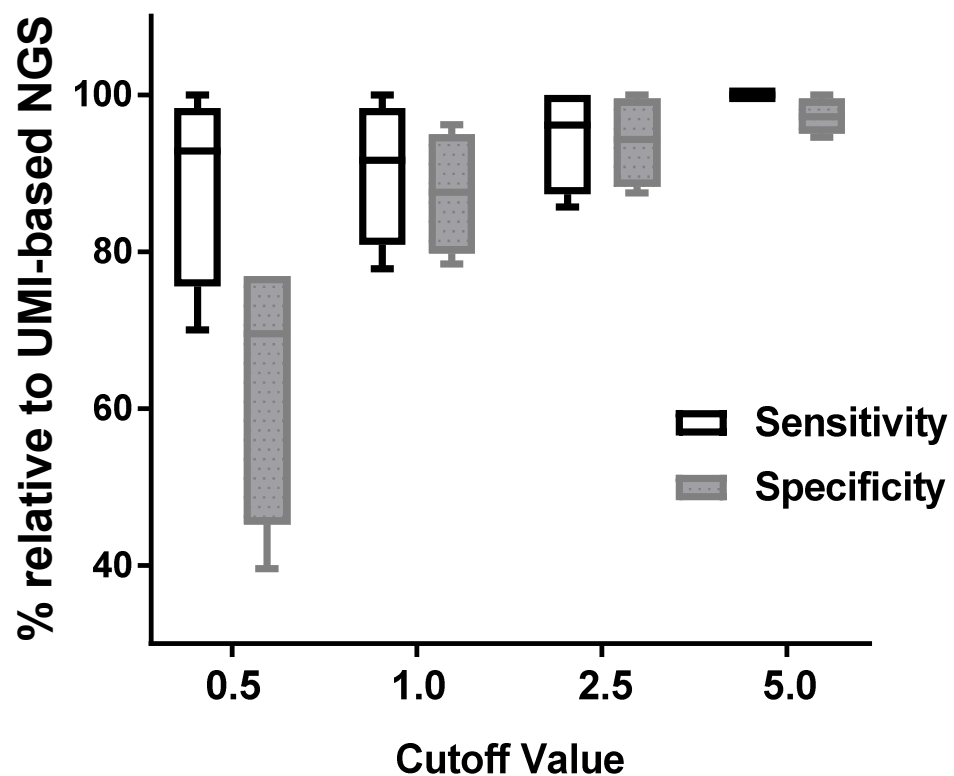
- We re-analyzed without UMIs, a UMI generated clinical dataset.
 - 62 plasma samples from viremic donors with HIV acute infection.

| | | UMI-based NGS (gold standard) | |
|-------------------|---|----------------------------------|------------------------------|
| | | True Positive MT Codon(s) | True Negative MT Codon(s) |
| Non UMI-based NGS | Test Positive (MT codon(s) detected) | A (True Positive) | B (False Positive) |
| | Test Negative (no MT codon(s) detected) | C (False Negative) | D (True Negative) |

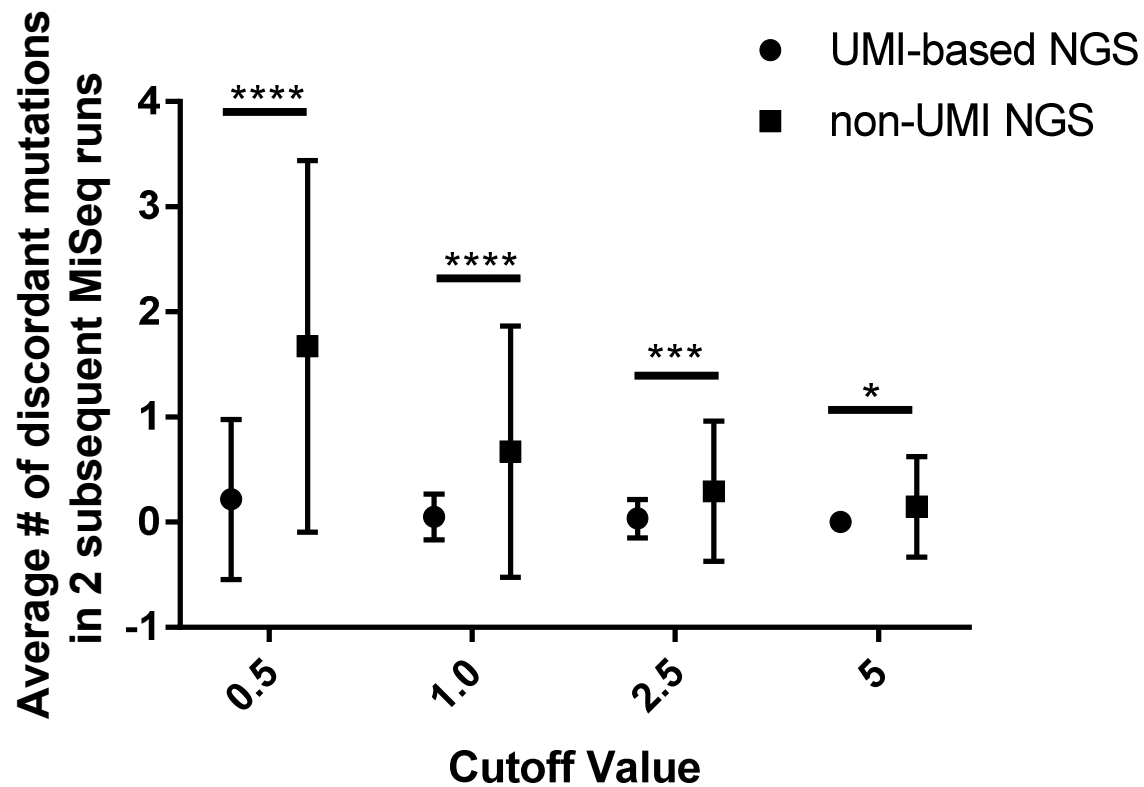
Sensitivity = proportion of samples with mutant codons that were detected with non UMI
$$\text{Sensitivity} = A/(A+C)$$

Specificity = proportion of samples without mutant codons that were negative with non UMI
$$\text{Specificity} = D/(B+D)$$

non-UMI-based NGS had similar sensitivity but reduced specificity in the re-analyzed clinical dataset.



Lower reproducibility of non-UMI-based NGS analysis



Note: The same samples were run on MiSeq twice

SUMMARY

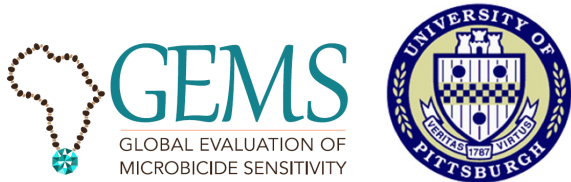
- Both UMI and non-UMI methods detected mutants at 0.5% in an isogenic background at high template input (50,000 copies), indicating that both methods are generally robust.
- False negative error rates are higher in non-UMI-based NGS with limited sampling depth (<1000 cp/ml or PCR bias).
- Non-UMI mutant detection at <5% was not reproducible for clinical samples.

IMPLICATIONS

- UMI-based consensus building should be used if calling mutations at frequencies below 5%.
- This is particularly true for samples that are likely to have low template input or inefficient PCR.
 - Low plasma HIV RNA.
 - Dried blood spots!
 - Polymorphic primer binding sites (always a concern).



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