Detection of minimal residual disease in lymph predicts recurrence in HPV-negative head and neck cancer patients

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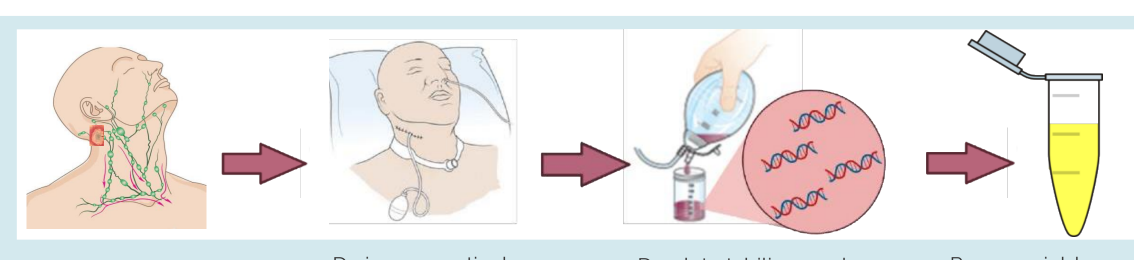
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Introduction

Locoregional cancer relapse remains a major cause of failure in head and neck squamous cell carcinoma (HNSCC), particularly for HPV-negative patients whose 2-year locoregional failure rate is up to 50%1. Methods to measure minimal residual disease (MRD) using ctDNA have emerged but have thus far had limited applications for detection of locoregional recurrence, especially after surgery. There is an unmet need for an accurate diagnostic test that predicts the risk of recurrence prior to adjuvant therapy selection. We present a novel proximal assay ("Droplet") for MRD profiled in lymphatic exudate collected via surgical drains ("lymph") and compare its performance to plasma-based MRD profiling and standard pathologic features for recurrence prediction.

Methods and Materials



Lymphatic fluid flows through all tumoradjacent lymph nodes.

Drains are routinely placed after tumor resection to prevent fluid build-up and speed healing.

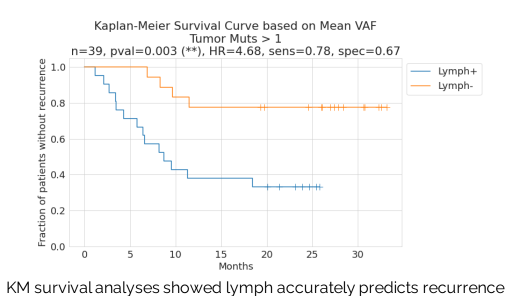
Droplet stabilizes and extracts lymphatic fluid from the drain material.²

Process yields a novel lymph-based analyte for MRD analysis using ctDNA

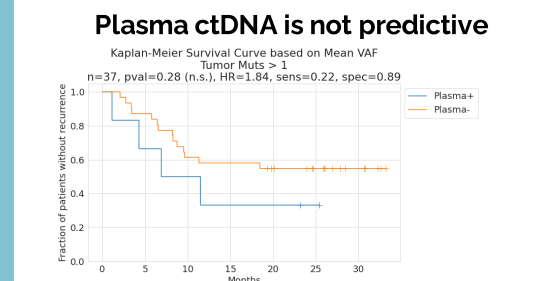
Lymph, plasma, and blood were collected from 46 HPVnegative HNSCC patients postoperatively at 24 hours along with resected tumor. Cell-free DNA was extracted from lymph and plasma and sequenced using the TruSight Oncology 500 panel to a depth of >100 million reads at Droplet Biosciences. Somatic mutations were identified by exome sequencing (200x) tumor and blood. Five patients had <2 somatic mutations in tumor and were excluded. Two patients were censored due to lack of clinical data, yielding 18 patients with disease recurrence (REC) and 21 with no evidence of disease (NED) with >1 year of follow-up. Two plasma samples were not available. Tumor-specific variants were force-called in lymph and plasma using a custom pipeline. Patients were considered MRD positive if the mean variant allele fraction (mVAF) was greater than 0.015% (the estimated limit of detection). Mann-Whitney U test was used for group comparisons. The Kaplan-Meier (KM) estimator with log-rank test and Cox proportional-hazards model were used for survival analyses. Logistic regression

models were performed with 5-fold cross-validation.

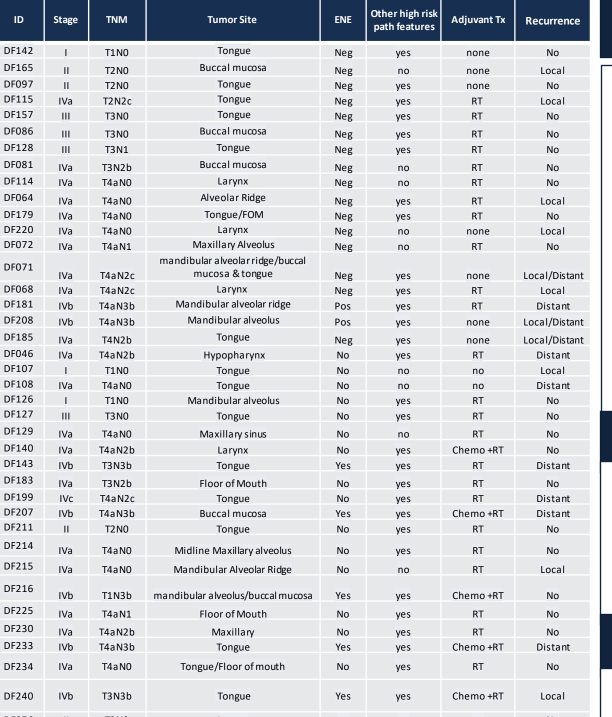
Lymph ctDNA predicts recurrence



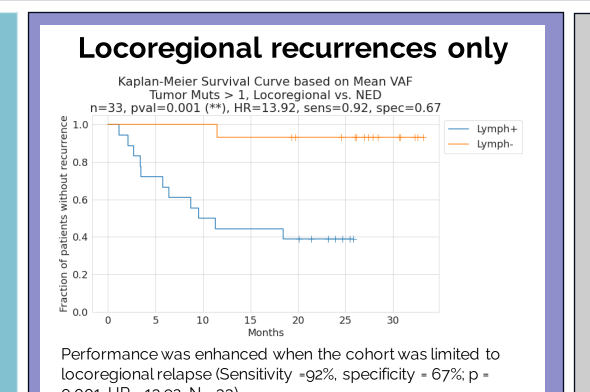
(sensitivity = 78%, specificity = 67%; p = 0.003, Hazard ratio (HR) = 4.68).

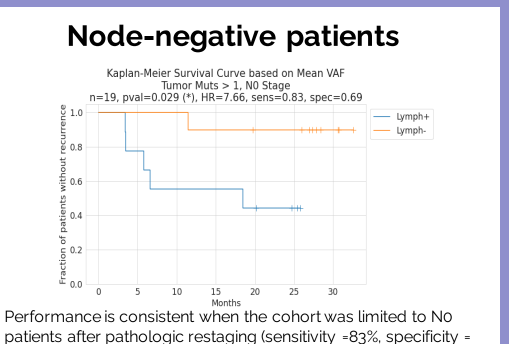


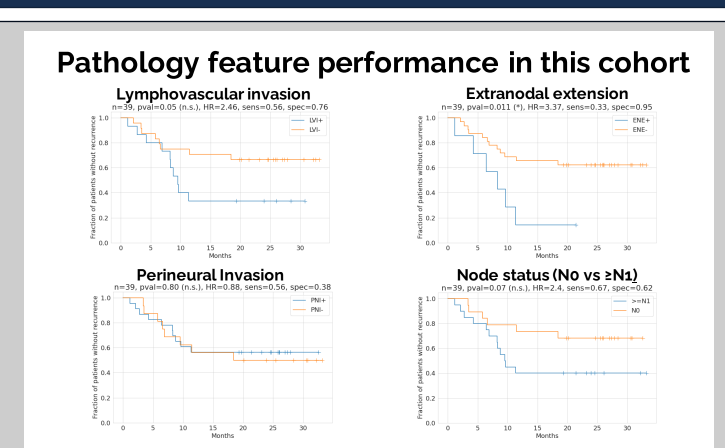
KM survival analyses indicated that plasma obtained 24 hrs after surgery does not predict recurrence (sensitivity = 22%, specificity = 89%; p = 0.28, HR = 1.84).



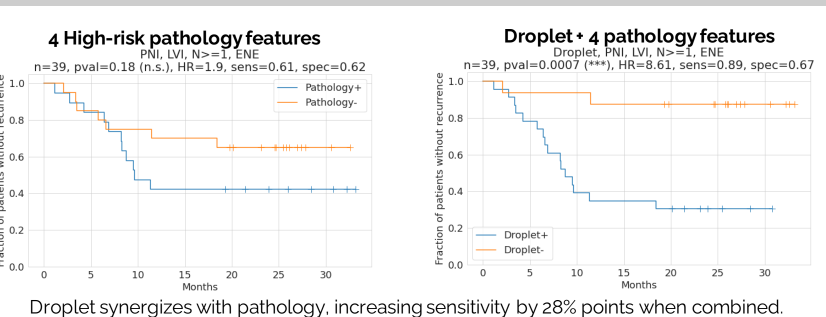
Results







The Droplet test outperformed pathology features (extranodal extension, perineural invasion, lymphovascular invasion, and nodal disease status) as well as a logistic regression model of all 4 (SN = 61%) SP = 62%; p = 0.18, HR = 1.9). A model incorporating Droplet plus the 4 high-risk pathology features showed superior performance over either lymph alone or pathology alone (SN = 89%, SP = 67% p = 0.0007, HR = 8.61).



Conclusions

- Postoperative ctDNA analysis from surgical lymphatic fluid represents a novel MRD approach in HPV-negative HNSCC.
- Patients who have increased ctDNA in lymph recur significantly more often and earlier than patients who have low or undetectable levels of lymph ctDNA, including patients with locoregional relapse and patients with No disease.
- · Lymph significantly outperforms plasma for prediction of recurrence.
- The Droplet assay gives superior prediction of recurrence than a multi-feature pathology model.
- The observed synergy between lymph MRD testing and traditional pathology suggests that incorporating postoperative lymph analysis has the potential to:
 - Augment traditional pathology
 - 2. Provide more personalized adjuvant treatment
- Validation in a large, prospective multi-institutional cohort of patients is ongoing

References

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