



This is a repository copy of *Point-of-care oral cytology tool for the screening and assessment of potentially malignant oral lesions*.

White Rose Research Online URL for this paper:  
<https://eprints.whiterose.ac.uk/154613/>

Version: Accepted Version

---

**Article:**

McRae, M., Modak, S., Simmons, G. et al. (10 more authors) (2020) Point-of-care oral cytology tool for the screening and assessment of potentially malignant oral lesions. *Cancer Cytopathology*, 128 (3). pp. 207-220. ISSN 1934-662X

<https://doi.org/10.1002/cncy.22236>

---

This is the peer reviewed version of the following article: McRae, M.P., Modak, S.S., Simmons, G.W., Trochesset, D.A., Kerr, A.R., Thornhill, M.H., Redding, S.W., Vigneswaran, N., Kang, S.K., Christodoulides, N.J., Murdoch, C., Dietl, S.J., Markham, R. and McDevitt, J.T. (2020), Point-of-care oral cytology tool for the screening and assessment of potentially malignant oral lesions. *Cancer Cytopathology*, which has been published in final form at <https://doi.org/10.1002/cncy.22236>. This article may be used for non-commercial purposes in accordance with Wiley Terms and Conditions for Use of Self-Archived Versions.

**Reuse**

Items deposited in White Rose Research Online are protected by copyright, with all rights reserved unless indicated otherwise. They may be downloaded and/or printed for private study, or other acts as permitted by national copyright laws. The publisher or other rights holders may allow further reproduction and re-use of the full text version. This is indicated by the licence information on the White Rose Research Online record for the item.

**Takedown**

If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing [eprints@whiterose.ac.uk](mailto:eprints@whiterose.ac.uk) including the URL of the record and the reason for the withdrawal request.



[eprints@whiterose.ac.uk](mailto:eprints@whiterose.ac.uk)  
<https://eprints.whiterose.ac.uk/>

# Title

Point-of-Care Oral Cytology Tool for the Screening and Assessment of Potentially Malignant Oral Lesions

# Running Title

Point-of-Care Oral Cytology Tool

# Authors

1. Michael P. McRae, Ph.D.  
Department of Biomaterials, Bioengineering Institute, New York University, New York, NY, USA  
michael.mcrae@nyu.edu
2. Sayli S. Modak, M.S.  
Department of Biomaterials, Bioengineering Institute, New York University, New York, NY, USA  
sayli.modak@nyu.edu
3. Glennon W. Simmons, B.S.  
Department of Biomaterials, Bioengineering Institute, New York University, New York, NY, USA  
glennon.simmons@nyu.edu
4. Denise A. Trochesset, D.D.S.  
Department of Oral and Maxillofacial Pathology, Radiology & Medicine, New York University  
College of Dentistry, New York, NY, USA  
dat5@nyu.edu
5. A. Ross Kerr, D.D.S., M.S.D.  
Department of Oral and Maxillofacial Pathology, Radiology & Medicine, New York University  
College of Dentistry, New York, NY, USA  
ark3@nyu.edu
6. Martin H. Thornhill, M.B.B.S., B.D.S., Ph.D.  
Department of Oral & Maxillofacial Medicine, Surgery and Pathology, School of Clinical Dentistry,  
University of Sheffield, Sheffield, UK  
M.Thornhill@sheffield.ac.uk
7. Spencer W. Redding, D.D.S., M.Ed.  
Department of Comprehensive Dentistry and Mays Cancer Center, The University of Texas Health  
Science Center at San Antonio, San Antonio, TX, USA  
redding@uthscsa.edu
8. Nadarajah Vigneswaran, B.D.S., Dr.Med.Dent., D.M.D.

1  
2  
3 Department of Diagnostic and Biomedical Sciences, The University of Texas Health Science  
4 Center at Houston, Houston, TX, USA  
5 Nadarajah.Vigneswaran@uth.tmc.edu  
6

7  
8 9. Stella K. Kang, M.D., M.S.

9 Departments of Radiology, Population Health New York University School of Medicine, New York,  
10 NY, USA

11 stella.kang@nyulangone.org

12  
13 10. Nicolaos J. Christodoulides, Ph.D.

14 Department of Biomaterials, Bioengineering Institute, New York University, New York, NY, USA  
15 nicolaoschristo19@gmail.com  
16

17  
18 11. Craig Murdoch, Ph.D.

19 Department of Oral & Maxillofacial Medicine, Surgery and Pathology, School of Clinical Dentistry,  
20 University of Sheffield, Sheffield, UK

21 c.murdoch@sheffield.ac.uk

22  
23 12. Steven J. Dietl, M.Eng.

24 SensoDx, LLC, Victor, NY, USA  
25 sdietl@sensodx.com  
26

27  
28 13. Roger Markham, Ph.D.

29 SensoDx, LLC, Victor, NY, USA

30 rmarkham@sensodx.com

31  
32 14. John T. McDevitt, Ph.D.

33 Chair, Department of Biomaterials, Bioengineering Institute, New York University, New York, NY,

34 USA

35 mcdevitt@nyu.edu  
36

37 Corresponding Author: John T. McDevitt, Ph.D., Chair, Department of Biomaterials, Bioengineering

38 Institute, New York University, 433 First Avenue, Room 820, New York, NY 10010-4086, USA. Email:  
39 mcdevitt@nyu.edu. Phone: 212-998-9204.  
40  
41

## 42 43 44 Funding Support

45  
46  
47 Funding for this work was provided by the National Institutes of Health (NIH) through the National Institute  
48 of Dental and Craniofacial Research (Award Number 1RC2DE020785-01, 5RC2DE020785-02,  
49 3RC2DE020785-02S1, 3RC2DE020785-02S2, 4R44DE025798-02, R01DE024392). The content of this  
50  
51 paper is solely the responsibility of the authors and does not necessarily represent or reflect the official  
52  
53  
54

views of the NIH or the US government. Segments of this work are supported by Renaissance Health Service Corporation and Delta Dental of Michigan.

## Conflict of Interest Disclosures

Principal Investigator, John T. McDevitt, has an equity interest in SensoDx, LLC. He also serves on the Scientific Advisory Board of SensoDx. Michael P. McRae has served as a consultant for SensoDx.

## Author Contributions

**Michael P. McRae:** Conceptualization, investigation, data curation, formal analysis, methodology, software, validation, visualization, writing - original draft, and writing - review and editing. **Sayli S. Modak:** Investigation, validation, and writing - review and editing. **Glennon W. Simmons:** Project administration, investigation, methodology, data collection, and writing - review and editing. **Denise A. Trochesset:** Conceptualization, writing - review and editing. **A. Ross Kerr:** Conceptualization and writing - review and editing. **Martin H. Thornhill:** Conceptualization and writing - review and editing. **Spencer W. Redding:** Conceptualization and writing - review and editing. **Nadarajah Vigneswaran:** Conceptualization and writing - review and editing. **Stella K. Kang:** Conceptualization and writing - review and editing. **Nicolaos J. Christodoulides:** Conceptualization, writing - original draft, and writing - review and editing. **Craig Murdoch:** Investigation and writing - review and editing. **Steven J. Dietl:** Methodology and writing - review and editing. **Roger Markham:** Methodology and writing - review and editing. **John T. McDevitt:** Conceptualization, funding acquisition, project administration, resources, supervision, writing - original draft, and writing - review and editing.

## Acknowledgements

The authors thank the University of Texas Health Science Center at San Antonio (UTHSCSA) (Stephanie Rowan, Chih-Ko Yeh, Stan McGuff, Frank Miller), University of Texas Health Science Center at Houston (UTHSCH) (Jerry Bouquot, Nagi Demian, Etan Weinstock, Nancy Bass), New York University / Bluestone Center for Clinical Research (Joan Phelan, Patricia Corby, Ismael Khouly), Sheffield Teaching Hospitals NHS Foundation Trust and the University of Sheffield (Paul Speight, Christine Freeman, Anne Hegarty, Katy D'Apice) for assistance in obtaining clinical samples. The authors also thank Rho, Inc. (Chapel Hill,

1  
2  
3 North Carolina) (Julie Vick) for assisting with patient data management and (Robert James) for statistical  
4 and data analysis support. Finally, the authors thank Timothy J. Abram for early contributions to the project,  
5 including assay development and database organization.  
6  
7  
8  
9

## 10 Precis

11  
12 A point-of-care oral cytology tool was developed for non-invasive detection and monitoring of potentially  
13 malignant oral lesions. Distributions of cell phenotypes identified by machine learning and a cytology-on-a-  
14 chip approach provide useful information in the assessment of oral lesions with improved interpretability,  
15 calibration, and generalizability relative to conventional methods.  
16  
17  
18

## 19 Abstract

20  
21 **BACKGROUND:** Effective detection and monitoring of potentially malignant oral lesions (PMOL) are  
22 critical to identifying early stage cancer and improving outcomes. In this study, the authors describe  
23 cytopathology tools including machine learning algorithms, clinical algorithms, and test reports developed  
24 to assist pathologists and clinicians with PMOL evaluation. **METHODS:** Data were acquired from a multi-  
25 site clinical validation study of 999 subjects with PMOLs and oral squamous cell carcinoma (OSCC) using  
26 a cytology-on-a-chip approach. A machine learning model was trained to recognize and quantify the  
27 distributions of four cell phenotypes. A least absolute shrinkage and selection operator (lasso) logistic  
28 regression model was trained to distinguish PMOLs and cancer across a spectrum of histopathologic  
29 diagnoses ranging from benign, to increasing grades of oral epithelial dysplasia (OED), to OSCC using  
30 demographics, lesion characteristics, and cell phenotypes. Cytopathology software was developed to assist  
31 pathologists in reviewing brush cytology test results, including high-content cell analyses, data visualization  
32 tools, and results reporting. **RESULTS:** Cell phenotypes were accurately determined through an automated  
33 cytological assay and machine learning approach (99.3% accuracy). Significant differences in cell  
34 phenotype distributions across diagnostic categories were found in three phenotypes (Type 1 'mature  
35 squamous', Type 2 'small round', and Type 3 'leukocytes'). The clinical algorithms resulted in acceptable  
36 performance characteristics (AUC = 0.81 for benign vs. mild dysplasia and 0.95 for benign vs. malignancy).  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53

54 **CONCLUSION:** These new cytopathology tools represent a practical solution for rapid PMOL assessment  
55  
56  
57  
58  
59  
60

1  
2  
3 with the potential to facilitate screening and longitudinal monitoring in primary, secondary, and tertiary  
4 clinical care settings.  
5

6  
7 **KEY WORDS:** squamous cell carcinoma; oral epithelial dysplasia; point-of-care testing; single-cell  
8 analysis; artificial intelligence; cytology; biomarkers.  
9

10  
11  
12 **Text Pages:** 23

13  
14 **Tables:** 0

15  
16 **Figures:** 7

17  
18 **Supporting Files for Publication:** Suppl. Methods, Suppl. Figures (6 total), Suppl. Video Content  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60



## Introduction

Cancers of the lip, oral cavity, and pharyngeal subsites are estimated to affect over 500,000 people globally each year.<sup>1</sup> The National Cancer Institute's Surveillance, Epidemiology, and End Results (SEER) program estimates 53,000 new cases and 10,860 deaths attributed to oral and pharyngeal cancer (OPC) in 2019 in the US alone, of which approximately 50% involve oral cavity subsites. Collectively, OPCs represent approximately 3% of all cancers.<sup>2</sup> Approximately two-thirds of OPCs are diagnosed at Stage III or IV when the 5-year survival rate is just 45% and 32%, respectively.<sup>3</sup> For the remaining third of OPCs detected at early stages,<sup>4</sup> survival increases to 84%.<sup>2</sup> Despite steady improvements in overall survival rates for OPC over the last four decades, identifying OPCs at an early stage remains a challenge for oral health care providers.<sup>5</sup> The current diagnostic paradigm of procuring a biopsy is based on remote lab services which can take days/weeks to provide results, and this further prolongs anxiety for patients. A point-of-care (POC) solution could provide immediate feedback within the same visit. Thus, there is a strong need for technology-driven solutions that can precisely and rapidly diagnose the entire spectrum of oral epithelial dysplasia (OED) and oral squamous cell carcinoma (OSCC) using minimally invasive sampling at the POC.

A successful diagnostic adjunctive test for primary care settings should be able to discriminate potentially malignant oral lesions (PMOLs) that are at "risk" (i.e., malignant lesions or those with an elevated risk for undergoing malignant transformation) from more common benign lesions with no malignant potential, thus improving the referral efficiency to secondary or tertiary care (e.g., reducing over-referral of patients with benign lesions and improving the early identification and prompt referral of malignant or high-grade dysplastic PMOLs for oncologic care). Numerous adjunctive tests are available to assist in the diagnosis of PMOLs. In a meta-analysis of oral cancer adjuncts, vital staining and visualization adjuncts (e.g., autofluorescence and tissue reflectance) demonstrated insufficient accuracy to be recommended for use as lesion triage tools by general dentists.<sup>6</sup> Cytology, however, has demonstrated greater sensitivity and specificity relative to the other adjuncts, suggesting its potential as a surrogate for gold-standard histopathology. This evidence to support the accuracy of cytology is largely based on accuracy studies performed in secondary and tertiary care settings. Although cytology is unable to replace histopathologic diagnosis based on tissue architecture, this relatively inexpensive, easy to perform, and minimally-invasive method may be useful for triaging lesions in any setting: primary care settings such as a dental office, low-



1  
2  
3 resource/remote settings, and secondary/tertiary settings. Incisional biopsy followed by histopathologic  
4 examination represents the current standard of care for diagnosing PMOLs. However, incisional biopsy of  
5 PMOLs, particularly in those that are large non-homogeneous leukoplakias, leads to underestimation of the  
6 severity of OED up to 30% of the time because the biopsy sample (typically 5 mm in diameter) may not be  
7 representative of the variable pathology across the field of the entire PMOL.<sup>7</sup> Brush cytology could enable  
8 a wider sampling of PMOLs that encompass larger areas or are multifocal with the potential to reduce  
9 sampling errors encountered with incisional biopsies.  
10

11  
12  
13  
14  
15  
16  
17 Previously, we have demonstrated the conceptual basis and the efficacy of chip-based cell capture,  
18 multispectral fluorescence measurements, and single-cell analysis approaches yielding high content  
19 diagnostic information related to oral lesions.<sup>8-10</sup> This compact and integrated lesion diagnostic adjunct  
20 approach has been studied previously through a multi-site clinical validation effort that has led to the  
21 development of one of the largest oral cytology databases ever assembled for PMOLs.<sup>11,12</sup> These efforts  
22 included the development of an “enhanced gold standard” adjudication process<sup>12</sup> that was used to correlate  
23 brush cytology measurements with six levels of histopathological diagnosis, ranging from benign, to OED,  
24 to OSCC. The same approach showed strong promise for OSCC surveillance in Fanconi Anemia patients<sup>13</sup>  
25 and for the development of a cytology based numerical risk index for cancer progression.<sup>14</sup> Overall, these  
26 past efforts have revealed that microfluidic-based cell capture systems with integrated imaging and  
27 embedded diagnostic algorithms can yield diagnostic accuracies that rival and exceed the capabilities of  
28 previously developed adjunct devices. These tools were developed previously to serve as adjunctive aids  
29 capable of distinguishing between high risk and low risk oral lesions with the goal of improving the pipeline  
30 of referrals from primary care settings to secondary and tertiary treatment centers. Thus, these models  
31 were intended for assisting primary care providers in making binary referral decisions and considered  
32 hundreds of complicated image-based cytomorphometric features with minimal clinical interpretability (i.e.,  
33 “black box”).  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48

49 This manuscript targets the development of a Point of Care Oral Cytology Tool (POCOCT), the first  
50 precision oncology technology capable of high content cell analysis for near patient testing. The POCOCT  
51 platform comprises a minimally invasive brush cytology test kit, disposable assay cartridge, instrument,  
52 clinical algorithms, and cloud-based software services that automate the quantification and analysis of  
53  
54  
55  
56

1  
2  
3 cellular and molecular signatures of dysplasia with results available in a matter of minutes as compared to  
4 days for traditional labor intensive lab-based pathology methods. This paper features the development of  
5 new diagnostic models using the same database described above with the goal of greatly simplifying the  
6 diagnostic algorithms and their interpretation through the classification and quantification of cellular  
7 phenotypes, resulting in more informative and transparent models for cytopathologists. Likewise, this work  
8 explores the utility of cell phenotype identification through machine learning, their implementation in  
9 diagnostic models with interpretable predictors and responses, and the practical application of these  
10 software tools in a cytopathology service.  
11  
12  
13  
14  
15  
16  
17  
18

## 19 Materials and Methods

### 22 Oral Cytology Data

23  
24 Data used in this study originated from the 999-patient multisite prospective non-interventional  
25 study evaluating the cytology-on-a-chip system for the measurement of cytological parameters on brush  
26 cytology samples to assist in the diagnosis of PMOL.<sup>11,12</sup> Briefly, both histopathological and brush  
27 cytological samples for 714 subjects from three patient groups were measured: (1) subjects with PMOL  
28 who underwent scalpel biopsy as part of the standard of care for microscopic diagnosis, (2) subjects with  
29 recently diagnosed malignant lesions, and (3) healthy volunteers without lesions. Histopathological  
30 assessment of scalpel biopsy specimens classified lesions into six categories (benign, mild-, moderate- or  
31 severe-dysplasia, carcinoma-in-situ, and OSCC), including healthy controls without lesions. While  
32 traditionally the grading of OED has been considered subjective and lacking intra- and inter-observer  
33 reproducibility,<sup>15,16</sup> this new study implemented an “enhanced gold standard” adjudication.<sup>12</sup> Here, two  
34 adjacent serial histologic sections were independently scored by two pathologists. In the event that the  
35 pathologists disagreed, a third independent adjudicating pathologist reviewed both sections. If the  
36 adjudicator did not agree with either of the initial two pathologists, a third stage consensus review was  
37 conducted to attain a final diagnosis. This “enhanced gold standard” process was able to achieve 100%  
38 consensus agreement compared to an initial pre-adjudication 69.9% agreement rate.  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52

53 Brush cytology specimens were collected and processed using protocols published previously.<sup>11,12</sup>  
54 Cytopathological assessment of brush cytology specimens implemented a cytology-on-a-chip approach  
55  
56  
57  
58  
59  
60

1  
2  
3 which measured morphological and intensity-based cell metrics as well as the expression of six molecular  
4 biomarkers ( $\alpha\beta6$ , EGFR, CD147, McM2, Geminin, and Ki67), resulting in a total of 13 million cells analyzed  
5 with over 150 image-based parameters. The molecular biomarkers were selected based on their capacity  
6 to distinguish benign, dysplastic, and malignant oral epithelial cells through prior immunohistochemistry  
7 studies.<sup>9,17,18</sup> Specific details on the molecular biomarker selection, patient characteristics, sample  
8 collection and processing, cytology assay, and cytological parameters were published previously<sup>11</sup> and are  
9 summarized in the **Supplemental Methods**.

### 17 **Cell Identification Model Training and Validation**

18 A cell phenotype classification model was explored for its ability to discriminate and quantitate the  
19 frequency and distributions of four cell phenotypes: Type 1: cells presenting as polygonal in shape with a  
20 low nuclear-cytoplasmic ratio (NC ratio) which represent mature squamous epithelial cells; Type 2: cells  
21 presenting as small round cells representing immature parabasal cells; Type 3: cells presenting as  
22 mononuclear leukocytes; Type 4: cells represented by lone (naked) nuclei without cell membrane and  
23 cytoplasm. To recognize these cell types, a machine learning algorithm was trained on 144 cellular/nuclear  
24 features from single-cell analyses, including morphological and intensity-based measurements. Prior to  
25 model development, principal component analysis (PCA) was performed on the training set. The PCA  
26 method is an unsupervised statistical learning technique for exploratory data analysis which improves data  
27 visualization by reducing the dimensionality of complex datasets<sup>19</sup> and has been used for phenotypic  
28 identification in flow cytometric data.<sup>20</sup> Detailed methods for the training and validation of the cell  
29 identification model are provided in the **Supplemental Methods**.

### 43 **Numerical Index and Diagnostic Models for Assessing PMOL**

44 A numerical index was developed for the purpose of discriminating benign vs. dysplasia/malignant  
45 lesions (OED-spectrum model 2|3). Detailed methods for the training and validation of the numerical index  
46 and detailed definition of predictors are provided in the **Supplemental Methods**. Briefly, subjects were  
47 dichotomized into “case” and “non-case” outcomes according to their lesion determination (non-case for  
48 benign lesions and case for [mild, moderate, severe] dysplasia and malignant lesions). Due to relatively  
49 few numbers of moderate and severe dysplasia patients (total of 21), these lesion determinations were  
50  
51  
52  
53  
54  
55  
56

1  
2  
3 combined. Lasso logistic regression was selected for its ability to reduce the number of predictors in high-  
4 dimensional datasets to improve prediction performance and generalizability.<sup>21-24</sup> Non-zero lasso logistic  
5 regression coefficients were retained for the following predictors: percentage of non-mature squamous  
6 cells, percentage of small round cells, percentage of leukocytes, age, sex, smoking pack years, lesion major  
7 axis diameter, clinical impression of lichen planus, and lesion color (red, white, or red/white). Diagnostic  
8 performance was characterized by area under the curve (AUC), sensitivity, and specificity. Median  
9 numerical indices were compared for each diagnostic classification using a two-sided Wilcoxon rank sum  
10 test at a significance level of  $p = 0.05$ . Internal calibration was performed by sorting and grouping the  
11 predicted responses (i.e., numerical index) into deciles and measuring the observed proportions of  
12 dysplasia/malignant lesions in each decile. The Hosmer-Lemeshow goodness of fit statistic was used to  
13 assess the model fit.<sup>21</sup>

14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24 Following this same method, diagnostic algorithms for mild vs. moderate dysplasia (OED-spectrum  
25 model 3|4), low vs. high risk (4|4), moderate vs. severe dysplasia (4|5), healthy control (no lesion) vs.  
26 malignant (0|6), and benign vs. malignant (2|6) were also developed, and AUC, sensitivity, and specificity  
27 were reported as mean and 95% confidence interval values for the cross-validated test set.

## 28 29 30 31 32 33 **Cytopathology Software**

34  
35 Measurements of individual cells, such as morphometric appearance and biomarker staining  
36 intensity, were recorded using the open-source software CellProfiler.<sup>25</sup> All model development and data  
37 analyses were completed with MATLAB R2017b (MathWorks, Natick, MA, USA) software. A graphical user  
38 interface for visualizing cytopathology results was developed in MATLAB R2017b. The results summary  
39 report tool was developed with Python 3.6.3. Figures of the cytopathology software interface and results  
40 summary were compiled from a test on the integrated POCOCT instrument.

## 41 42 43 44 45 46 **Level of Integration**

47  
48 Data originating from our 999-patient NIH Grand Opportunity (GO) study and used in the cell  
49 identification and diagnostic models were collected using non-integrated cytology-on-a-chip flow cell  
50 prototypes, syringe pumps, research microscope stations, and a collection of commercial and open-source  
51 software packages (see **Supplemental Methods** for more details).<sup>11</sup> More recently, we have integrated the  
52 cytology-on-a-chip technology into a POC device comprising integrated instrument, microfluidic cartridges  
53  
54  
55  
56

1  
2  
3 with on-board blister packs, and dedicated software. Likewise, sample processing steps have been  
4 significantly reduced. Cell identification and diagnostic models developed on the non-integrated platform  
5 were translated to the POC instrument, and software screenshots and results reports presented here were  
6 completed with this integrated POC platform.  
7  
8  
9

## 10 Results

### 11 Cell Identification Model

12  
13  
14  
15  
16  
17 A cell identification tool to assist in the accurate and precise estimation of histopathological  
18 endpoints for the entire spectrum of OED and OSCC was developed. Figure 1 shows the diagnostic  
19 categories and rates for oral cancer and dysplasia based on WHO classification<sup>26</sup> found during mass  
20 screening,<sup>27</sup> showing 5-year malignant transformations<sup>28</sup> and 5-year cancer recurrence.<sup>29</sup> The literature  
21 presents a range of 5-year transformation and recurrence rates, and the ones listed here are representative  
22 of those reported previously.<sup>30</sup>  
23  
24  
25  
26  
27  
28

29 The POCOCT platform (Figure 2) comprises a minimally invasive brush cytology test kit, disposable  
30 assay cartridge, instrument, clinical algorithms, and cloud-based software services to automate the  
31 quantification and analysis of cellular and molecular signatures of dysplasia and OSCC. The cell  
32 identification tool automatically classified four distinct cell phenotypes (Figure 3A). Type 1 'mature  
33 squamous' or 'mature keratinocytes' were broad/flat cells, approximately 50-100  $\mu\text{m}$  in diameter, had a low  
34 NC ratio, and demonstrated a relatively low cytoplasm staining intensity (Phalloidin-Alexa Fluor® 647).  
35  
36  
37  
38 Type 2 'small round' cells were small (12-30  $\mu\text{m}$  in diameter) highly circular cells with high NC ratio and a  
39 brightly stained cytoplasm representing immature basaloid keratinocytes. Type 3 'leukocytes' appeared as  
40 small, brightly stained pink objects 6-23  $\mu\text{m}$  in diameter representing mononuclear leukocytes. Type 4 'lone  
41 nuclei' represented by lone or naked nuclei without a cytoplasm appeared as brightly stained blue objects  
42 approximately 5-12  $\mu\text{m}$  in diameter.  
43  
44  
45  
46  
47  
48

49 The PCA scatter plot of the first two principal components revealed a glimpse of the internal data  
50 structure and variance (Figure 3A). Here, populations according to each cell type were clearly observed.  
51  
52 Further, over 90% of the variance was explained by the first 20 principal components from a total of 144,  
53 with 30% and 14% variance explained in the first and second principal components, respectively. Despite  
54  
55  
56  
57  
58  
59  
60

Types 2 and 3 having similar cytomorphology, the features with the largest association with the first principal component were NC ratio and mean cytoplasm intensity, suggesting that cell size and cellular actin content/distribution play a dominant role in explaining the variance among these cell phenotypes.

The cross-validated *k-nearest neighbors* (*k*-NN) algorithm resulted in overall accuracy of 96.9% and accuracy of 100%, 90.1%, 96.0%, and 99.0% for Types 1 (mature), 2 (small), 3 (leukocytes), and 4 (lone nuclei), respectively. An additional label ('unknown') was added for cells that had four or less similar neighbors. After accounting for this 'unknown' cell type, the overall accuracy was 99.3%. When applied to the study population, cell phenotype distributions showed significant differences across all diagnostic categories (Figure 3B). The proportion of Type 1 (mature) cells decreased with more advanced disease. In contrast, the proportions of Type 2 (small) and Type 3 (leukocytes) cells increased with disease progression. Median values for Type 1 (mature) and Type 2 (small) cells were significantly different between all lesion determinations. For Type 3 (leukocytes), all lesion determinations had significantly different median values except for benign vs. dysplasia ( $p = 0.0539$ ).

The same cell identification model development process was completed on recently developed integrated instrumentation, cartridges, and cloud-based analysis tools. Images from two samples, one each from benign and malignant lesions, were collected with the POCOCT platform, and cell phenotype labels were overlaid on each recognized cell object (Figure 3C). Here, the benign lesion sample contained mostly Type 1 (mature) cells, while the malignant sample contained a mixture of primarily Type 2 (small), Type 3 (leukocytes), and Type 4 (lone nuclei).

#### **Numerical Index and Diagnostic Models for Assessing PMOL**

Expanding on this capability, a numerical index for discriminating benign and dysplasia/malignant lesions was developed using the cell phenotypes as predictors. Figure 4A shows the ROC curve

representing discrimination performance of the multivariate model. The numerical index is a score between 0 and 100 that can be interpreted literally as the probability of dysplasia/malignancy. The diagnostic accuracy of the model is defined by the cutoff score that maximizes its AUC (benign vs. dysplasia/malignant numerical index cutoff of 36). Predictors for the model were retained as follows: cell phenotype distributions (Types 1, 2, and 3), age, sex, smoking pack years (i.e., packs per day times years of smoking), lesion size (maximum diameter), clinical impression of lesion as lichen planus, and lesion color (white, red, or both)

(Figure 4B). Minimal differences were observed between training and test error (28% and 27% misclassification rate on the training and test sets, respectively) which suggests no evidence of overfitting. The numerical index showed significant differences between all lesion diagnostic categories studied ( $p < 0.01$ ) except for mild vs. moderate/severe dysplasia ( $p = 0.1519$ ) (Figure 4C); however, significant differences were observed in a dichotomous model for mild vs. moderate dysplasia (i.e., 3|4) ( $p = 0.04$ ). Model calibration shows the numerical index relative to the observed proportions of dysplasia/malignant subjects when sorted and grouped into deciles (Figure 4D). A non-significant result of the Hosmer-Lemeshow goodness of fit test suggests that there is no evidence of a poor fit ( $p = 0.6259$ ).

Models were also developed for dichotomous classification across the OED spectrum, and Figure 5 summarizes the diagnostic performance of these models. The clinical algorithms resulted in AUCs ranging 0.81 (95% CI 0.76–0.86) for benign vs. mild dysplasia (3|4) to 0.97 (0.94–1.00) for healthy control (no lesion) vs. malignancy (0|6). While previous work demonstrated AUCs of 0.836 for the binary low vs. high risk (4|4) split and 0.883 for moderate vs. severe dysplasia (4|5),<sup>11</sup> these new optimized models here presented resulted in improved AUCs of 0.88 (0.84–0.93) and 0.92 (0.88–0.96) for the same diagnostic splits, respectively.

### Cytopathology Software

A cytopathology interface tool was developed to assist pathologists in reviewing the brush cytology test results, enabling rich content cellular analyses on single- and multi-cell levels (Figure 6 and

**Supplemental Figures**). This interface enables the pathologist users to access data stored and processed on cloud-based services, view results summaries, explore cytology results through data visualization tools, and generate automated oral cytopathology reports (Figure 7) which provide the adjunctive referral recommendations and summarize important information from cytology, including total cell count, cell

phenotype distributions (Types 1, 2, and 3), and mean values for NC ratio, molecular biomarker fluorescence intensity, and cell circularity. The ability to assess cumulative data on this cloud-based cytopathology platform may improve pathologist decision making (e.g., through learning about their own histopathologic assessment vs. the POCOCT and, ultimately, the surgical pathology).

## Discussion

This work demonstrates an evolution of the POCOCT technology towards a rapid and simple brush cytology analysis for POC or in a remote laboratory setting. We have demonstrated that (1) cell phenotypes can be accurately determined through the automated cytological assay and machine learning approach; (2) significant differences in cell phenotype distributions across diagnostic categories are found in three phenotypes (Types 1, 2, and 3); and (3) these cell phenotypes are valuable predictors for distinguishing lesion diagnostic categories in a multivariate lasso logistic regression model. The compilation of these results suggests that the observed cellular phenotypic variations within cytological samples are equated with disease severity and, thus, may be useful in the evaluation of PMOLs. Although cell phenotyping can be completed by a pathologist by manually identifying cells in a cytological sample, this is a lengthy process subject to human errors. Providing a means to automate metrics, such as the distributions of cell phenotypes, may increase adoption of this POCOCT approach through a cytopathology service and allow for pathologists to complete more efficient and more effective recommendations.

The optimized numerical index for evaluating PMOLs developed here represents a simple, practical, and effective approach that is directly applicable to clinical implementation and interpretation. While previous models relied on complicated high-dimensional cytological parameters, the classification and quantitation of cell phenotypes greatly simplifies the predictive algorithm and its interpretation, substantially improves performance for diagnostic splits relative to these earlier efforts,<sup>11,14</sup> and supports the translation of research methodologies from laboratory-based microscopy stations to an integrated POC instrument. With a total of 9 predictors, the practical model developed here represents a sparse solution (i.e., reduction of over 150 variables to 9) with greater potential generalizability without sacrificing any diagnostic performance. Further, excellent model calibration performance and significant differences between the diagnostic endpoints demonstrates strong potential for the numerical index as a continuous indicator of PMOL risk. While previous work was primarily focused on delivering binary results for referral decisions,<sup>11</sup> this new work involves a cytopathology interface tool, developed to assist pathologists in reviewing the brush cytology test results, and a numerical index, enabling rich content cellular analyses on single- and multi-cell levels. This interface enables the pathologist to access data stored on cloud-based services, view results summaries, explore cytology data through data visualization tools, and generate a



1  
2  
3 report that provides recommendations. Accurate diagnostic models spanning the entire OED spectrum also  
4 demonstrate the potential for the POCOCT to be used for multiple applications, such as screening PMOLs  
5 in primary care and the surveillance of patients with a history of OED and OSCC in secondary or tertiary  
6 care settings.  
7  
8  
9

10  
11 Although light-based adjuncts offer clinicians a new perspective to view a lesion at the POC, their  
12 diagnostic utility remains unproven.<sup>5</sup> Rashid and Warnakulasuriya reviewed the performance of light-based  
13 adjuncts in discriminating low and high risk lesions (VELscope [sensitivity/specificity: 30–100 / 15–100],  
14 ViziLite Plus [0–100 / 0–78], and Microlux DL [78 / 71]) and concluded that there is insufficient evidence to  
15 validate their efficacy as screening adjuncts.<sup>31</sup> Despite the numerous adjunctive tests available to assist in  
16 the diagnosis of PMOLs today, only cytology shows potential as a surrogate for gold standard  
17 histopathology.<sup>32</sup> Several commercial cytopathology services exist today including OralCDx (CDx  
18 Diagnostics, Inc.), OralCyte (ClearCyte Diagnostics, Inc.), Cyt ID (Forward Science), and ClearPrep OC  
19 (Resolution Biomedical). OralCDx, for example, provides an oral brush sample collection kit for their  
20 BrushTest.<sup>33</sup> Despite the ease of collection, samples need to be shipped to a commercial laboratory for  
21 analysis, resulting in delays between sample collection and test results. Further, the test often returns an  
22 ambiguous “atypical” result for which the positive predictive value for dysplasia or carcinoma has been  
23 determined to be only 30-40%.<sup>34</sup> Additionally, prior studies of cytology adjuncts demonstrated  
24 methodological gaps by only performing matched gold-standard histopathology on a subset of lesions with  
25 a higher index of suspicion for malignancy, and not for lesions with a lower index of suspicion which are  
26 frequently encountered in primary care settings.<sup>35,36</sup> A clinically validated POC cytology service capable of  
27 distinguishing the degree of OED in PMOL and stratifying the risk of malignant progression as a numerical  
28 index in near real-time would fulfill a significant unmet need mitigating unnecessary referrals to experts,  
29 leading to a more efficient process in surveillance clinics and reducing the patient distress related to waiting  
30 for test results.  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48

49 One limitation is that previous studies of the POCOCT, and cytology adjuncts in general, primarily  
50 focused on PMOL evaluation in secondary care settings where the prevalence of dysplastic and malignant  
51 lesions may be substantially higher than in the primary care. Additionally, while expert clinicians in  
52 secondary and tertiary care settings have extensive training and experience in the recognition and risk  
53  
54  
55  
56

1  
2  
3 stratification of PMOLs, primary care clinicians may have difficulty distinguishing PMOLs from normal/non-  
4 neoplastic lesions. Thus, the POCOCT technology may potentially have a larger impact in primary care  
5 settings where there is a strong need to accurately interrogate the PMOLs detected there and generate a  
6 dichotomous outcome to indicate if referral of patients to higher care settings for expert evaluation and  
7 possible biopsy is required and if such referral should be urgent.  
8  
9  
10  
11  
12

13 This manuscript provides a key step towards the development of new tools that could pave the way  
14 for new capabilities in the area of 'precision lesion diagnostics'. Helping to push forward this theme, we  
15 have demonstrated the utility of temporal changes in numerical index in a pilot study of Fanconi Anemia  
16 (FA) patients.<sup>13</sup> These efforts showed strong potential for patient-specific temporal changes in the lesion  
17 numerical index to track early signs of disease for this high risk population. Plans are now in place to (1)  
18 evaluate the POCOCT's precision lesion diagnostic capabilities through a prospective longitudinal study of  
19 malignant transformation and cancer recurrence and (2) move the POCOCT into a clinical trial to assess  
20 the POCOCT's diagnostic performance vs. routine care in primary care clinics.  
21  
22  
23  
24  
25  
26  
27  
28

## 29 Conclusion

30  
31 In summary, we have demonstrated the utility of a POC-amenable cytology platform that has the  
32 potential to screen and monitor oral lesions across the entire diagnostic spectrum of OED. Cell phenotype  
33 distributions provided additional information in the assessment of PMOL. Further, a practical model  
34 comprised of patient information, lesion characteristics, and cell types from cytology showed similar  
35 performance characteristics to more complicated models previously developed. Cytopathology software  
36 may assist expert pathologists and non-expert care providers in reviewing and understanding the brush  
37 cytology test results. We developed data visualization tools to provide high content cellular analyses on  
38 single- and multi-cell levels with full transparency of test results data for pathologists. Additionally, oral  
39 cytopathology results summarize the test's most important predictors through indications of potential lesion  
40 progression for care providers and patients. Along with recently developed instrumentation and cartridges,  
41 this simple and sensitive system could provide non-invasive triage for PMOLs detected in primary,  
42 secondary, and tertiary care settings.  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

1  
2  
3 Future work may expand the utilization of molecular biomarkers and explore the identification of  
4 additional rare cell phenotypes to further improve performance. Future clinical studies may also be directed  
5 to determine: whether brush cytology could enable a wider sampling of large/multifocal lesion areas relative  
6 to incisional biopsies via multiple site-precise samplings; the effect of inflammation on the cytological  
7 analysis; whether the system can identify candida and distinguish clinical leukoplakia from neoplastic vs.  
8 non-neoplastic conditions; its placement in existing monitoring algorithms for PMOLs. Clinical trials are  
9 needed to assess the POCOCT's ability to identify early stage cancer relative to existing protocols and to  
10 validate the POCOCT as a substitute for biopsy. Future publications will describe and validate the integrated  
11 POC hardware (i.e., instrument, cartridge, and assay). To accelerate the translation and expand the  
12 adoption of the POCOCT platform, a cytopathology service for secondary and tertiary care oral cytology  
13 applications is now in development. Scaling and distribution of this versatile cytology approach is now  
14 underway with potential to serve diagnostic and surveillance applications in primary, secondary, and tertiary  
15 care settings.  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

## References

1. Shield KD, Ferlay J, Jemal A, et al. The global incidence of lip, oral cavity, and pharyngeal cancers by subsite in 2012. *CA Cancer J Clin.* 2017;67(1):51-64.
2. National Cancer Institute Surveillance, Epidemiology, and End Results Program. Cancer stat facts: oral cancer and pharynx cancer. <https://seer.cancer.gov/statfacts/html/oralcav.html>. Accessed May 10, 2019.
3. Neville BW, Damm DD, Allen CM, Chi AC. Epithelial Pathology. In: Neville BW, Damm DD, Allen CM, Chi AC, eds. *J Oral Maxillofac Pathol.* 4th ed. St. Louis, MO: Elsevier Health Sciences; 2015:331-421.
4. Siegel RL, Miller KD, Jemal A. Cancer statistics, 2018. *CA Cancer J Clin.* 2018;68(1):7-30.
5. Huber MA. Adjunctive Diagnostic Techniques for Oral and Oropharyngeal Cancer Discovery. *Dent Clin North Am.* 2018;62(1):59-75.
6. Lingen MW, Abt E, Agrawal N, et al. Evidence-based clinical practice guideline for the evaluation of potentially malignant disorders in the oral cavity: A report of the American Dental Association. *J Am Dent Assoc.* 2017;148(10):712-727.
7. Lee J-J, Hung H-C, Cheng S-J, et al. Factors associated with underdiagnosis from incisional biopsy of oral leukoplakic lesions. *Oral Surg Oral Med Oral Pathol Oral Radiol Endod.* 2007;104(2):217-225.
8. Weigum SE, Floriano PN, Christodoulides N, McDevitt JT. Cell-based sensor for analysis of EGFR biomarker expression in oral cancer. *Lab Chip.* 2007;7(8):995-1003.
9. Weigum SE, Floriano PN, Redding SW, et al. Nano-bio-chip sensor platform for examination of oral exfoliative cytology. *Cancer Prev Res.* 2010;3(4):518-528.
10. McDevitt JT, Weigum SE, Floriano PN, et al. A new bio-nanochip sensor aids oral cancer detection. *SPIE Newsroom.* 2011.
11. Abram TJ, Floriano PN, Christodoulides N, et al. 'Cytology-on-a-chip' based sensors for monitoring of potentially malignant oral lesions. *Oral Oncol.* 2016;60:103-111.

12. Speight PM, Abram TJ, Floriano PN, et al. Interobserver agreement in dysplasia grading: toward an enhanced gold standard for clinical pathology trials. *Oral Surg Oral Med Oral Pathol Oral Radiol*. 2015;120(4):474-482.
13. Abram TJ, Pickering CR, Lang AK, et al. Risk stratification of oral potentially malignant disorders in Fanconi Anemia patients using autofluorescence imaging and cytology-on-a chip assay. *Transl Oncol*. 2018;11(2):477-486.
14. Abram TJ, Floriano PN, James R, et al. Development of a cytology-based multivariate analytical risk index for oral cancer. *Oral Oncol*. 2019;92:6-11.
15. Bosman FT. Dysplasia classification: pathology in disgrace? *J Pathol*. 2001;194(2):143-144.
16. Warnakulasuriya S, Reibel J, Bouquot J, Dabelsteen E. Oral epithelial dysplasia classification systems: predictive value, utility, weaknesses and scope for improvement. *J Oral Pathol Med*. 2008;37(3):127-133.
17. Vigneswaran N, Beckers S, Waigel S, et al. Increased EMMPRIN (CD 147) expression during oral carcinogenesis. *Exp Mol Pathol*. 2006;80(2):147-159.
18. Torres-Rendon A, Roy S, Craig GT, Speight PM. Expression of Mcm2, geminin and Ki67 in normal oral mucosa, oral epithelial dysplasias and their corresponding squamous-cell carcinomas. *Br J Cancer*. 2009;100(7):1128-1134.
19. Jolliffe I. *Principal component analysis*. 2nd ed. New York: Springer; 2011.
20. Lugli E, Pinti M, Nasi M, et al. Subject classification obtained by cluster analysis and principal component analysis applied to flow cytometric data. *Cytometry A*. 2007;71A(5):334-344.
21. Hosmer DW, Lemeshow S. *Applied Logistic Regression*. 2nd ed. New York: John Wiley & Sons, Inc.; 2004.
22. LaValley MP. Logistic Regression. *Circulation*. 2008;117(18):2395-2399.
23. Hastie T, Tibshirani R, Friedman JH. *The Elements of Statistical Learning: Data Mining, Inference, and Prediction*. 2nd ed. New York: Springer; 2009.
24. Wang D, Zhang W, Bakhai A. Comparison of Bayesian model averaging and stepwise methods for model selection in logistic regression. *Stat Med*. 2004;23(22):3451-3467.

- 1  
2  
3 25. Carpenter AE, Jones TR, Lamprecht MR, et al. CellProfiler: image analysis software for identifying  
4 and quantifying cell phenotypes. *Genome Biol.* 2006;7(10):R100.  
5  
6  
7 26. El-Naggar AK, Chan JK, Grandis JR, Takata T, Slotweg PJ. *WHO classification of tumours of the*  
8 *head and neck.* 4th ed. Lyon: IARC Press; 2017.  
9  
10  
11 27. Bouquot JE. Common oral lesions found during a mass screening examination. *J Am Dent Assoc.*  
12 1986;112(1):50-57.  
13  
14  
15 28. Sperandio M, Brown AL, Lock C, et al. Predictive value of dysplasia grading and DNA ploidy in  
16 malignant transformation of oral potentially malignant disorders. *Cancer Prev Res.* 2013;6(8):822-  
17 831.  
18  
19  
20 29. Brands MT, Smeekens EAJ, Takes RP, et al. Time patterns of recurrence and second primary  
21 tumors in a large cohort of patients treated for oral cavity cancer. *Cancer Med.* 2019;8(12):5810-  
22 5819.  
23  
24  
25 30. Mehanna HM, Rattay T, Smith J, McConkey CC. Treatment and follow-up of oral dysplasia — A  
26 systematic review and meta-analysis. *Head Neck.* 2009;31(12):1600-1609.  
27  
28  
29 31. Rashid A, Warnakulasuriya S. The use of light-based (optical) detection systems as adjuncts in the  
30 detection of oral cancer and oral potentially malignant disorders: a systematic review. *J Oral Pathol*  
31 *Med.* 2015;44(5):307-328.  
32  
33  
34 32. Lingen MW, Tampi MP, Urquhart O, et al. Adjuncts for the evaluation of potentially malignant  
35 disorders in the oral cavity: Diagnostic test accuracy systematic review and meta-analysis—a report  
36 of the American Dental Association. *J Am Dent Assoc.* 2017;148(11):797-813.e752.  
37  
38  
39 33. CDx Diagnostics: The Painless Test for Common Oral  
40 Spots <https://www.cdxdiagnostics.com/brushtest/>. Accessed May 10, 2019.  
41  
42  
43  
44 34. Svirsky JA, Burns JC, Carpenter WM, et al. Comparison of computer-assisted brush biopsy results  
45 with follow up scalpel biopsy and histology. *Gen Dent.* 2002;50(6):500-503.  
46  
47  
48 35. Sciubba JJ. Improving detection of precancerous and cancerous oral lesions: computer-assisted  
49 analysis of the oral brush biopsy. *J Am Dent Assoc.* 1999;130(10):1445-1457.  
50  
51  
52 36. Poate TWJ, Buchanan JAG, Hodgson TA, et al. An audit of the efficacy of the oral brush biopsy  
53 technique in a specialist Oral Medicine unit. *Oral Oncol.* 2004;40(8):829-834.  
54  
55  
56

## Figure Legends

Figure 1. Diagnostic categories for oral cancer and dysplasia based on WHO classification with 5-year malignant transformations and 5-year cancer recurrence rates. While 10% of US adults may present to their dentist for a routine care visit with an abnormal oral cavity lesion, about 83% of these lesions are diagnosed clinically as having no malignant potential, and 17% have unknown significance and meet the clinical criteria for PMOL. About 17% of PMOLs are histopathologically diagnosed with OED or OSCC. OED is about 15 times more common than OSCC, yet only a fraction of patients with dysplastic PMOLs undergo malignant transformation.

Figure 2. The POCOCT assay platform allows for the analysis of cellular samples obtained from a minimally invasive brush cytology sample. The cell suspension collected in this manner allow for the simultaneous quantification of cell morphometric data and expression of molecular biomarkers of malignant potential in an automated manner using refined image analysis algorithms based on pattern recognition techniques and advanced statistical methods. This novel approach turns around cytology results in a matter of minutes as compared to days for traditional pathology methods, thereby making it amenable to POC settings. The POC testing is expected to have tremendous implications for disease management by enabling dental practitioners and primary care physicians to circumvent the need for multiple referrals and consultations before obtaining assessment of molecular risk of PMOL.

Figure 3. A cell type identification model was developed to automatically classify cell Types 1-4. Panel A (left) shows the four distinct cell phenotypes that were identified: Type 1 ('mature squamous cells'), Type 2 ('small round cells'), Type 3 ('leukocytes'), and Type 4 ('lone nuclei'). Principal component analysis (right) shows cell phenotypes clustered into distinct groups with substantial separation between cell phenotype labels, demonstrating strong promise for an effective cell phenotype recognition algorithm. Boxplots in Panel B show the study population distributions of mature squamous cells (left), small round cells (center), and leukocytes (right), representing the predicted mean cell type percentages across six biomarker assays ( $\alpha\beta6$ , CD-147, EGFR, geminin, Ki-67, and MCM2) within each lesion class: normal (n=121), benign (n=241), dysplasia (n=59), and malignant (n=65). The results shown include only patients with definitive

1  
2  
3 lesion determinations and patients with evaluable data for all six biomarkers. Panel C shows limited field of  
4 view cytology pseudocolor images (fluorescence images acquired with a monochrome camera and digitally  
5 assigned to red, green, and blue color channels) of benign (left) and malignant (right) lesions with the cell  
6 phenotype model output labels overlaid as follows: "M" for mature squamous cells, "S" for small round cells,  
7 "W" for leukocytes, and "L" for lone nuclei (Unknown type "U" not shown). Fluorescent staining shows the  
8 cytoplasm (red), nuclei (blue), and Ki-67 biomarker (green).

9  
10  
11  
12  
13  
14  
15  
16  
17 Figure 4. Algorithm results of the dichotomous benign vs. dysplasia/malignant lesion model from 241 benign  
18 lesion and 124 dysplasia and malignant lesion subjects for six molecular biomarker assays on the POCOCT  
19 system. Panel A shows the ROC curve for the model. The lasso logistic regression coefficients are provided  
20 in Panel B. The predictors are as follows: "1-%TYPE 1" (percent of cells that are non-mature squamous  
21 cells), "%TYPE 2" (percent of cells that are small round cells), "%TYPE 3" (percent of cells that are  
22 leukocytes), "AGE", "SEX", "PACKYR" (pack years), "LSIZEMAX" (lesion diameter of the major axis),  
23 "LICHENFN" (clinical impression of lichen planus), and "LESIONCOLOR" (red, white, or red/white). The  
24 boxplot in Panel C shows cross-validated algorithm response ("numerical index") for the lasso logistic  
25 regression on the test set averaged over all biomarker assays. Distribution of scores are represented for  
26 benign (n=241), mild dysplasia (n=38), moderate/severe dysplasia (n=21), and malignant lesions (n=65).  
27 Panel D shows a model calibration plot of the predicted responses (numerical index) sorted and grouped  
28 into deciles vs. the observed proportions of dysplasia and malignant lesions.

29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41 Figure 5. Diagnostic models for the OED spectrum. Results are shown for the cross-validated clinical  
42 algorithms for benign vs. dysplasia (2|3), mild vs. moderate dysplasia (3|4), low vs. high risk (4|4), moderate  
43 vs. severe dysplasia (4|5), healthy control (no lesion) vs. malignant (0|6), and benign dysplasia vs.  
44 malignant (2|6) models. Model responses for each subject were averaged over all biomarker assays to  
45 inform diagnostic performance. AUC, sensitivity, and specificity are mean and 95% confidence interval  
46 values for the cross-validated test set.



1  
2  
3 Figure 6. Cytopathology interface tool provides pathologists with cloud access to test results summaries  
4 and detailed data visualizations (A), scatter plots (B), and histograms (C) for over 150 different cytology  
5 parameters. With this tool, pathologists can view all cells within the field of view, zoom in for more detail,  
6  
7 and isolate individual cells of interest.  
8  
9

10  
11  
12 Figure 7. Oral cytopathology test results. The algorithm result is a numerical index between 0 and 100 with  
13 a cutoff of 36 that distinguishes benign and dysplasia/malignant (“atypical”) lesions (left). Other informative  
14 cytopathology results are displayed on a reference range, including total cell counts, cell phenotype  
15 distributions, mean values for NC ratio, molecular biomarker fluorescence intensity, and cell circularity.  
16  
17 Images and outlines of the cells are provided for additional test context (right).  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

# Supplemental Figures

Data Explorer

Tools

**View Test Results** Showing test results for 190816-BICR56-EGFR-2

Select

**Test Information**

Test ID	CID
Test Date	2019-09-11
Biopsy Date	N/A
Lesion Location	N/A
Total Cells	286
Total Objects	429
Mature Squamous Cells	62 (22%)
Small Round Cells	210 (73%)
WBCs	14 (5%)
Lone Nuclei	23
Unknown	120
NC Ratio	0.2612
Cell Circularity	0.3046
EGFR	0.0465
Age	56
Sex	Female
Pack Years	12
Lesion Size	7
Lichen Planus	No
Lesion Color	Red and White
Algorithm Result	57.0189

Generate Report

**Image View**

Raw Image  Mature Squamous

R  Small Round

G  Leukocytes

B  Lone Nuclei

Cell Outlines  Unknown

The image shows a field of cells with red cytoplasm, blue nuclei, and some green spots. The cells are densely packed and vary in size and shape, consistent with the 'Small Round Cells' category mentioned in the test information.

Figure S1. Screenshot of cytopathology interface showing BICR 56 cancer cells magnified view with all three fluorescent labels (red: phalloidin, green: ...)

40 EGFR, blue: DAPI).

41  
42  
43  
44  
45  
46  
47

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47

Data Explorer

Tools

**View Test Results** Showing test results for 190816-BICR56-EGFR-2

Select

**Test Information**

Test ID	CID
Test Date	2019-09-11
Biopsy Date	N/A
Lesion Location	N/A
Total Cells	286
Total Objects	429
Mature Squamous Cells	62 (22%)
Small Round Cells	210 (73%)
WBCs	14 (5%)
Lone Nuclei	23
Unknown	120
NC Ratio	0.2612
Cell Circularity	0.3046
EGFR	0.0465
Age	56
Sex	Female
Pack Years	12
Lesion Size	7
Lichen Planus	No
Lesion Color	Red and White
Algorithm Result	57.0189

Generate Report

**Image View**

Raw Image  Mature Squamous

R  Small Round

G  Leukocytes

B  Lone Nuclei

Cell Outlines  Unknown

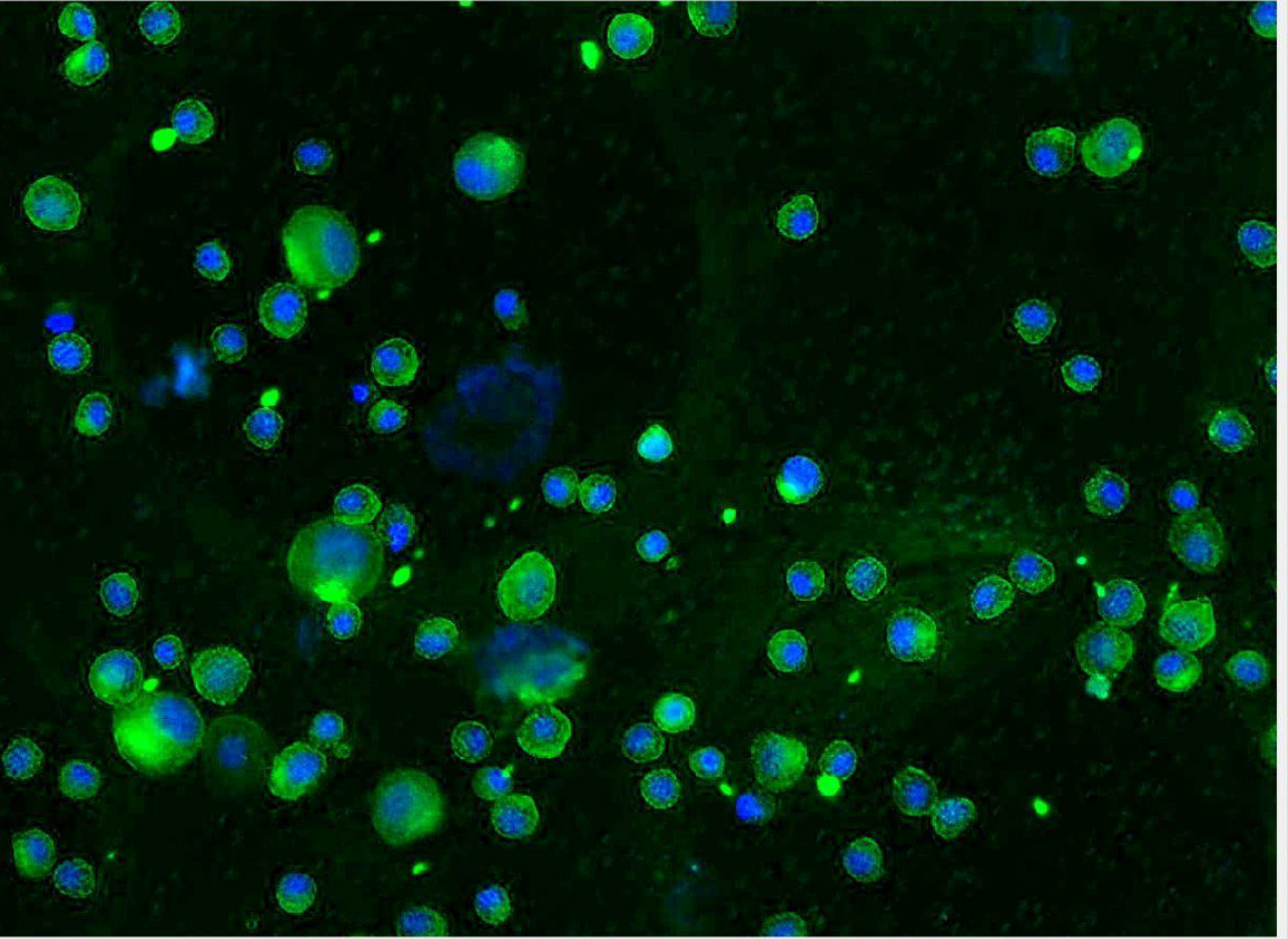


Figure S2. Screenshot of cytopathology interface showing BICR 56 cancer cells magnified view with green (EGFR) and blue (DAPI) fluorescent labels.

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47

Data Explorer

Tools

**View Test Results** Showing test results for 190816-BICR56-EGFR-2

Select

**Test Information**

Test ID	CID
Test Date	2019-09-11
Biopsy Date	N/A
Lesion Location	N/A
Total Cells	286
Total Objects	429
Mature Squamous Cells	62 (22%)
Small Round Cells	210 (73%)
WBCs	14 (5%)
Lone Nuclei	23
Unknown	120
NC Ratio	0.2612
Cell Circularity	0.3046
EGFR	0.0465
Age	56
Sex	Female
Pack Years	12
Lesion Size	7
Lichen Planus	No
Lesion Color	Red and White
Algorithm Result	57.0189

Generate Report

**Image View**

Raw Image

R

G

B

Cell Outlines

Mature Squamous

Small Round

Leukocytes

Lone Nuclei

Unknown

Figure S3. Screenshot of cytopathology interface showing BICR 56 cancer cells with cell phenotype labels overlaid (M: mature squamous, S: small

37 round, W: leukocytes, L: lone nuclei, U: unknown).

38

39

40

41

42

43

44

45

46

47

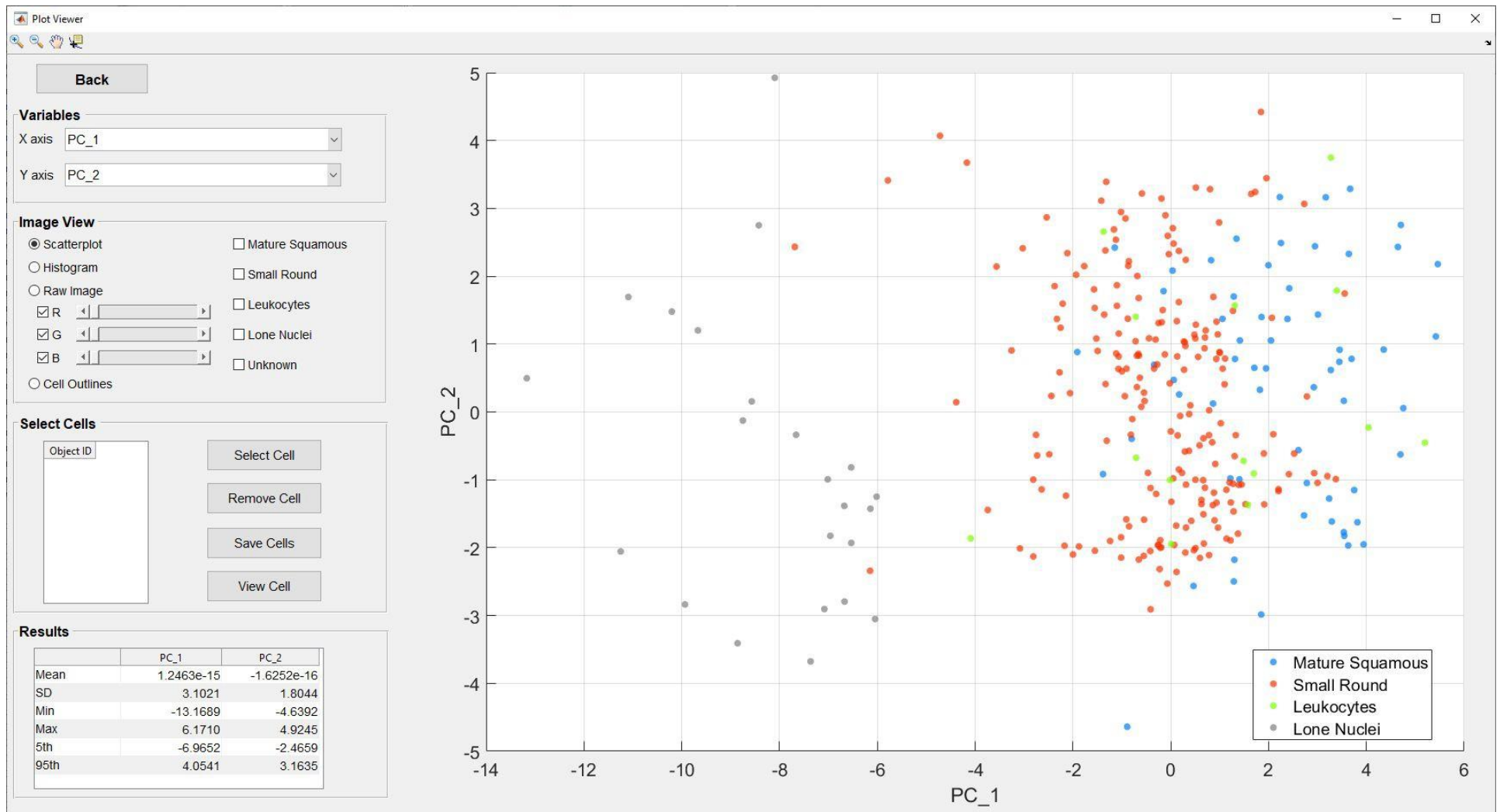


Figure S4. Screenshot of cytopathology interface showing a principal component scatter plot from a sample of BICR 56 cancer cells.

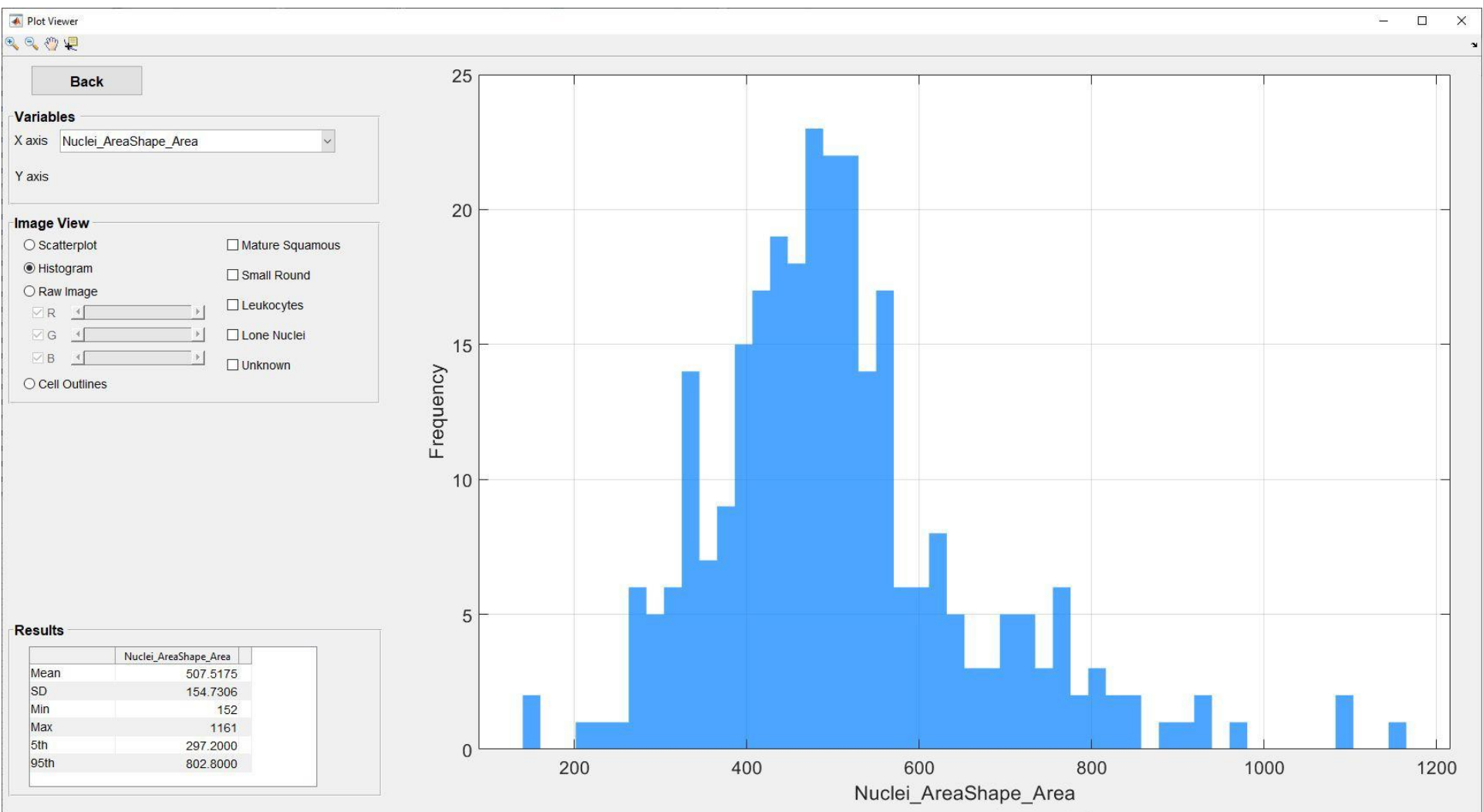


Figure S5. Screenshot of cytopathology interface showing histogram of nuclear area measurements from a sample of BICR 56 cancer cells.



1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47

Data Explorer

Tools

**View Test Results** Showing test results for 190827-Healthy-EGFR

Select

**Test Information**

Test ID	CID
Test Date	2019-09-11
Biopsy Date	N/A
Lesion Location	N/A
Total Cells	115
Total Objects	298
Mature Squamous Cells	89 (77%)
Small Round Cells	22 (19%)
WBCs	4 (3%)
Lone Nuclei	88
Unknown	95
NC Ratio	0.0764
Cell Circularity	0.4093
EGFR	0.0212
Age	30
Sex	Female
Pack Years	0
Lesion Size	2
Lichen Planus	No
Lesion Color	White
Algorithm Result	25.3827

Generate Report

**Image View**

Raw Image

R

G

B

Cell Outlines

Mature Squamous

Small Round

Leukocytes

Lone Nuclei

Unknown

Figure S6. Screenshot of cytopathology interface showing brush biopsy sample of healthy control cells with cell phenotype labels overlaid (M: mature

37 squamous, S: small round, W: leukocytes, L: lone nuclei, U: unknown, not shown).

38  
39  
40  
41  
42  
43  
44  
45  
46  
47

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

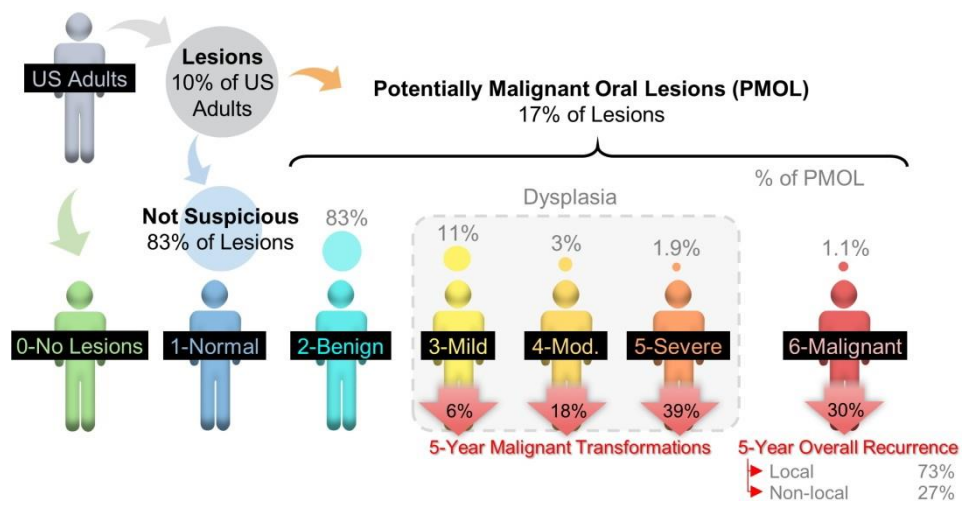


Figure 1. Diagnostic categories for oral cancer and dysplasia based on WHO classification with 5-year malignant transformations and 5-year cancer recurrence rates. While 10% of US adults may present to their dentist for a routine care visit with an abnormal oral cavity lesion, about 83% of these lesions are diagnosed clinically as having no malignant potential, and 17% have unknown significance and meet the clinical criteria for PMOL. About 17% of PMOLs are histopathologically diagnosed with OED or OSCC. OED is about 15 times more common than OSCC, yet only a fraction of patients with dysplastic PMOLs undergo malignant transformation.

171x101mm (300 x 300 DPI)

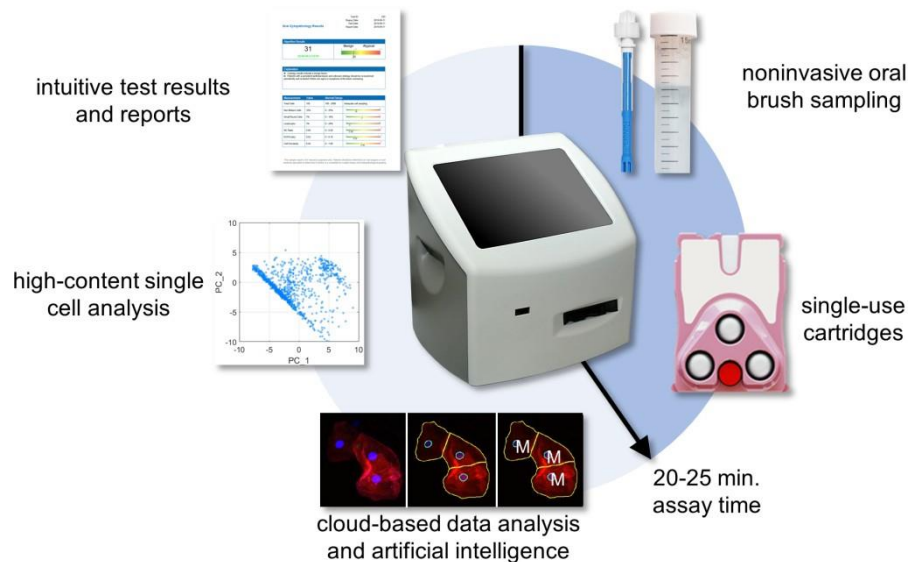


Figure 2. The POCOCT assay platform allows for the analysis of cellular samples obtained from a minimally invasive brush cytology sample. The cell suspension collected in this manner allow for the simultaneous quantification of cell morphometric data and expression of molecular biomarkers of malignant potential in an automated manner using refined image analysis algorithms based on pattern recognition techniques and advanced statistical methods. This novel approach turns around cytology results in a matter of minutes as compared to days for traditional pathology methods, thereby making it amenable to POC settings. The POC testing is expected to have tremendous implications for disease management by enabling dental practitioners and primary care physicians to circumvent the need for multiple referrals and consultations before obtaining assessment of molecular risk of PMOL.

171x96mm (300 x 300 DPI)

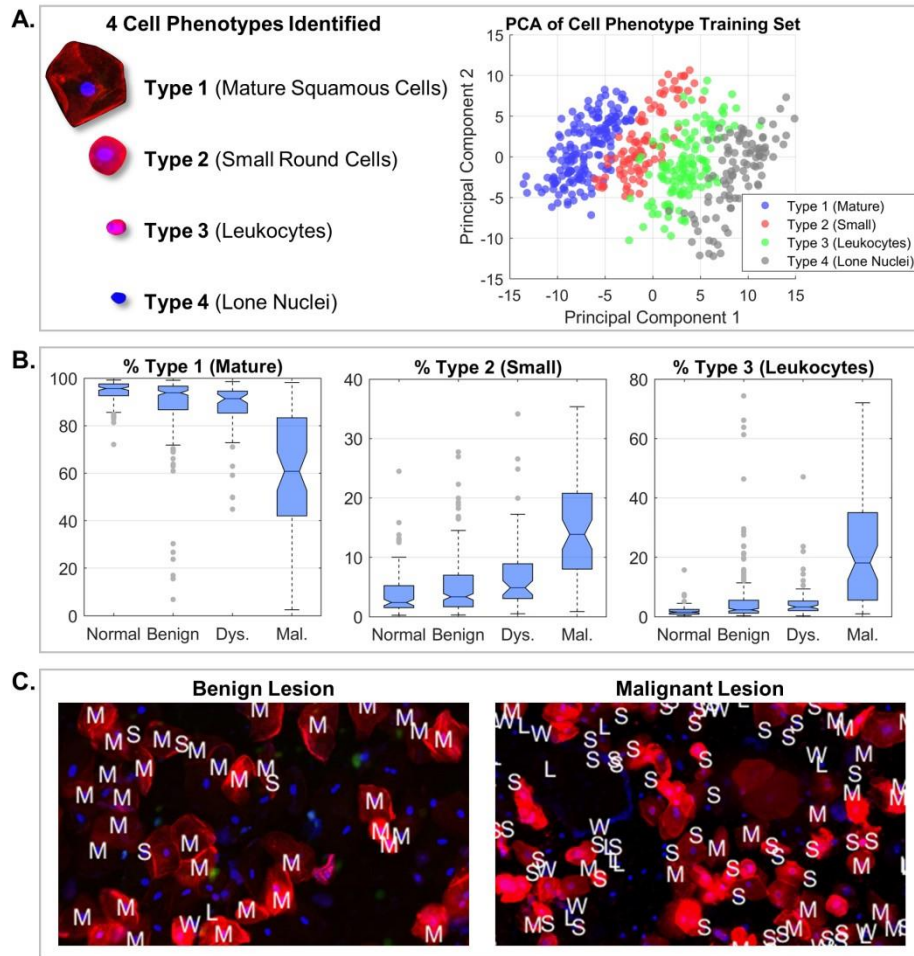


Figure 3. A cell type identification model was developed to automatically classify cell Types 1-4. Panel A (left) shows the four distinct cell phenotypes that were identified: Type 1 ('mature squamous cells'), Type 2 ('small round cells'), Type 3 ('leukocytes'), and Type 4 ('lone nuclei'). Principal component analysis (right) shows cell phenotypes clustered into distinct groups with substantial separation between cell phenotype labels, demonstrating strong promise for an effective cell phenotype recognition algorithm. Boxplots in Panel B show the study population distributions of mature squamous cells (left), small round cells (center), and leukocytes (right), representing the predicted mean cell type percentages across six biomarker assays (avβ6, CD-147, EGFR, geminin, Ki-67, and MCM2) within each lesion class: normal (n=121), benign (n=241), dysplasia (n=59), and malignant (n=65). The results shown include only patients with definitive lesion determinations and patients with evaluable data for all six biomarkers. Panel C shows limited field of view cytology pseudocolor images (fluorescence images acquired with a monochrome camera and digitally assigned to red, green, and blue color channels) of benign (left) and malignant (right) lesions with the cell phenotype model output labels overlaid as follows: "M" for mature squamous cells, "S" for small round cells, "W" for leukocytes, and "L" for lone nuclei (Unknown type "U" not shown). Fluorescent staining shows the cytoplasm (red), nuclei (blue), and Ki-67 biomarker (green).



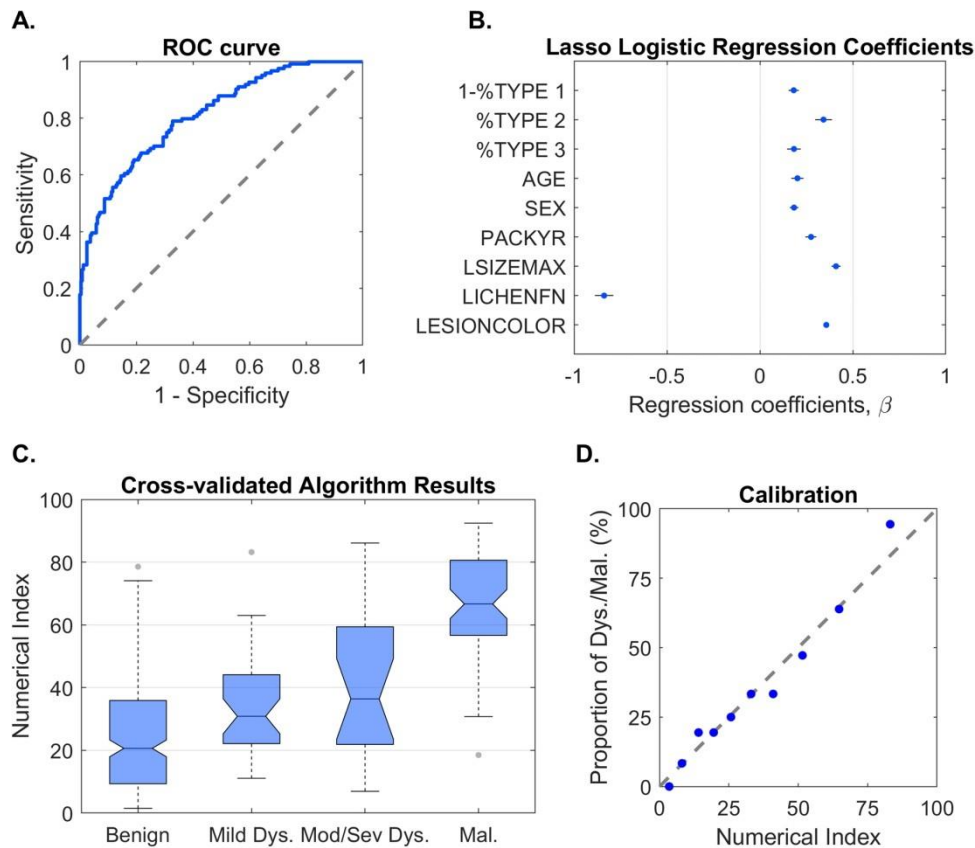


Figure 4. Algorithm results of the dichotomous benign vs. dysplasia/malignant lesion model from 241 benign lesion and 124 dysplasia and malignant lesion subjects for six molecular biomarker assays on the POCOCT system. Panel A shows the ROC curve for the model. The lasso logistic regression coefficients are provided in Panel B. The predictors are as follows: "1-%TYPE 1" (percent of cells that are non-mature squamous cells), "%TYPE 2" (percent of cells that are small round cells), "%TYPE 3" (percent of cells that are leukocytes), "AGE", "SEX", "PACKYR" (pack years), "LSIZEMAX" (lesion diameter of the major axis), "LICHENFN" (clinical impression of lichen planus), and "LESIONCOLOR" (red, white, or red/white). The boxplot in Panel C shows cross-validated algorithm response ("numerical index") for the lasso logistic regression on the test set averaged over all biomarker assays. Distribution of scores are represented for benign (n=241), mild dysplasia (n=38), moderate/severe dysplasia (n=21), and malignant lesions (n=65). Panel D shows a model calibration plot of the predicted responses (numerical index) sorted and grouped into deciles vs. the observed proportions of dysplasia and malignant lesions.






Model	Non-case / Case	Sensitivity	Specificity	AUC
2 3		0.69 (0.64–0.74)	0.77 (0.72–0.81)	0.81 (0.76–0.86)
3 4		0.79 (0.74–0.83)	0.85 (0.81–0.89)	0.88 (0.84–0.93)
4 4		0.78 (0.73–0.82)	0.87 (0.83–0.90)	0.88 (0.84–0.93)
4 5		0.82 (0.78–0.86)	0.88 (0.84–0.91)	0.92 (0.88–0.96)
2 6		0.89 (0.85–0.92)	0.90 (0.85–0.93)	0.95 (0.91–0.98)
0 6		0.94 (0.89–0.97)	0.92 (0.87–0.95)	0.97 (0.94–1.00)

Figure 5. Diagnostic models for the OED spectrum. Results are shown for the cross-validated clinical algorithms for benign vs. dysplasia (2|3), mild vs. moderate dysplasia (3|4), low vs. high risk (4|4), moderate vs. severe dysplasia (4|5), healthy control (no lesion) vs. malignant (0|6), and benign dysplasia vs. malignant (2|6) models. Model responses for each subject were averaged over all biomarker assays to inform diagnostic performance. AUC, sensitivity, and specificity are mean and 95% confidence interval values for the cross-validated test set.

171x99mm (300 x 300 DPI)



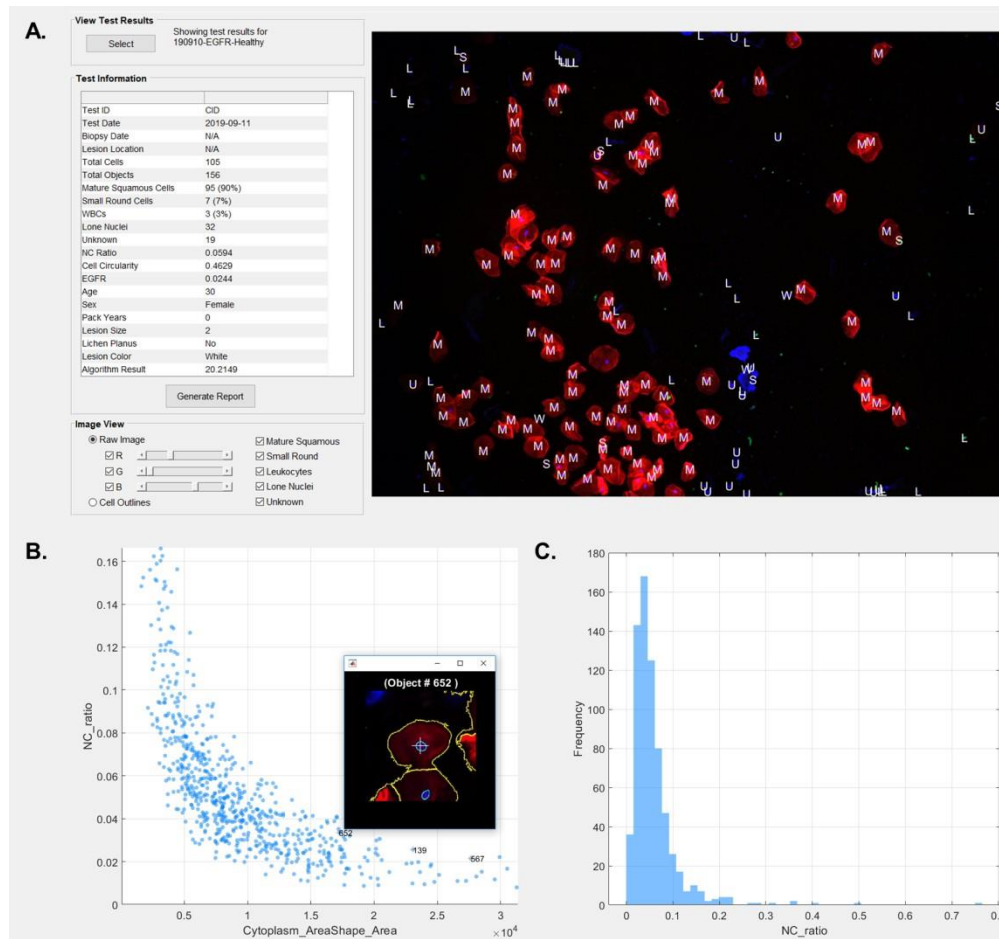


Figure 6. Cytopathology interface tool provides pathologists with cloud access to test results summaries and detailed data visualizations (A), scatter plots (B), and histograms (C) for over 150 different cytology parameters. With this tool, pathologists can view all cells within the field of view, zoom in for more detail, and isolate individual cells of interest.

171x160mm (300 x 300 DPI)

1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

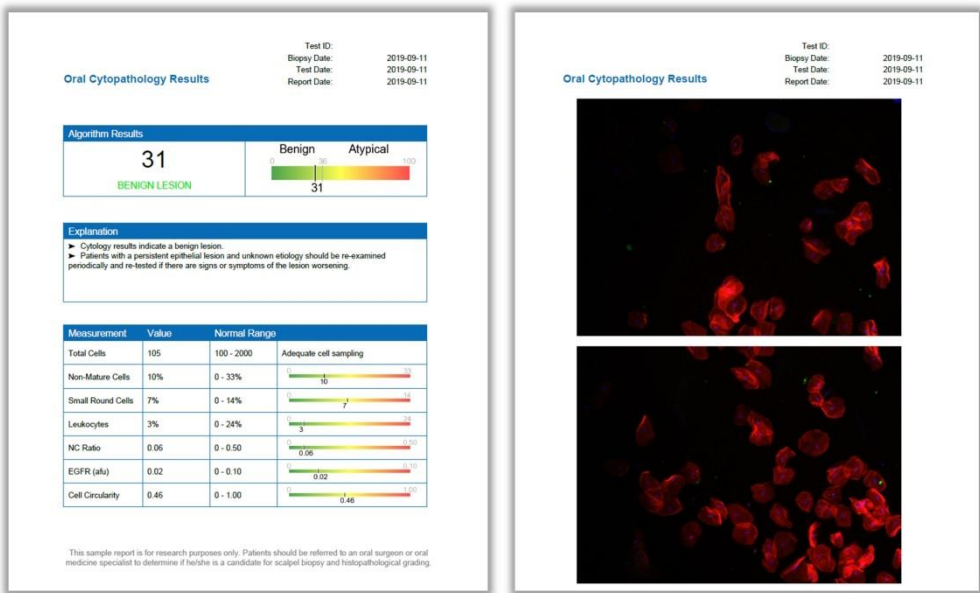


Figure 7. Oral cytopathology test results. The algorithm result is a numerical index between 0 and 100 with a cutoff of 36 that distinguishes benign and dysplasia/malignant (“atypical”) lesions (left). Other informative cytopathology results are displayed on a reference range, including total cell counts, cell phenotype distributions, mean values for NC ratio, molecular biomarker fluorescence intensity, and cell circularity. Images and outlines of the cells are provided for additional test context (right).

171x106mm (300 x 300 DPI)



## Supplemental Methods

### Biomarker Selection Rationale

Six molecular biomarkers were selected ( $\alpha\beta6$ , CD147, EGFR, geminin, Ki67, and MCM2) based on their capacity to distinguish benign, dysplastic, and malignant oral epithelial cells through prior immunohistochemistry studies.<sup>1-3</sup> These markers fall into three groups based on their localization: cell membrane, cytoplasm, and nucleus. **Table S1** summarizes the molecular biomarkers used in the study.

**Table S1. Summary of molecular biomarkers**

Biomarker	Localization	Function
$\alpha\beta6$	CM	an integrin receptor undetectable in normal oral epithelium, but highly expressed in dysplasia and OSCC <sup>4,5</sup>
CD147	CM	a multifaceted molecule that facilitates tumor progression by several mechanisms <sup>6</sup>
EGFR	CM + C	a transmembrane glycoprotein whose overexpression may contribute to tumor progression <sup>7</sup>
Geminin	N + C	a marker of proliferation <sup>2</sup>
Ki67	N	a marker of proliferation that is overexpressed at initial stages of oral carcinogenesis <sup>7</sup>
MCM2	N	an essential component for DNA replication associated with deregulated expression in dysplastic and malignant epithelial cells <sup>8,9</sup>

\* CM: cell membrane; C: cytoplasm; N: nucleus

### Patient Recruitment

Data used in this study originated from the 999-patient multisite prospective non-interventional study evaluating the cytology-on-a-chip system for the measurement of cytological parameters on brush cytology samples to assist in the diagnosis of PMOL. Briefly, both histopathological and brush cytological samples for 714 subjects from three patient groups were measured: (1) subjects with PMOL who underwent scalpel biopsy as part of the standard of care for microscopic diagnosis, (2) subjects with recently diagnosed malignant lesions, and (3) healthy

volunteers without lesions. Only subjects with complete biomarker results were included in the analysis (N = 486). **Table S2** summarizes the patient characteristics of those subjects included in the analysis.

**Table S2. Patient characteristics and histopathological diagnoses**

Characteristics and Histopathological Diagnoses	N (%)
Total	486
Sex	
Male	211 (43.4)
Female	275 (56.6)
Age	
>60	165 (34.0)
≤60	320 (65.8)
Patient Group	
Healthy Volunteer	121 (24.9)
Subjects with Previously Diagnosed Malignant Lesion	36 (7.4)
Subject with a Potentially Malignant Lesion	329 (67.7)
Histopathological Diagnosis	
Normal	121 (24.9)
Benign	241 (49.6)
Mild Dysplasia	38 (7.8)
Moderate Dysplasia	12 (2.5)
Severe Dysplasia	9 (1.9)
Malignant	65 (13.4)

### Clinical Protocol

The clinical protocol for this study was published previously<sup>10</sup> and is summarized as follows. Patients in group 1 underwent brush sampling of the oral lesion and a brush sampling of

the contralateral, clinically normal mucosa. The brush cytology sample was taken immediately



1  
2  
3 before the same lesion underwent a scalpel biopsy. Patients in group 2 underwent brush biopsy  
4 of the known cancerous lesion, as well as the contralateral, clinically normal mucosa. For healthy  
5  
6  
7 volunteers in group 3, a brush biopsy of normal appearing tissue on the lateral or ventral surface  
8  
9 of the tongue and a brush biopsy of normal appearing tissue on the left or right buccal mucosa  
10  
11 were taken. Brush biopsy samples were taken using a soft Rovers Orcellex oral cytology brush  
12  
13 (Rovers Medical Devices B.V., Oss, The Netherlands). The brush was applied directly to the  
14  
15 lesion or control oral mucosa using mild pressure and rotated 360 degrees approximately 10-15  
16  
17 times in the same direction to obtain the cytologic sample.  
18

### 19 **Cytology-on-a-Chip Protocol**

21  
22 The following methods have been published previously<sup>11</sup> and are summarized here for  
23  
24 convenience. Immediately after brush cytology samples were collected, cells were harvested by  
25  
26 vortexing the brush head in minimum essential medium (MEM) culture media, followed by a PBS  
27  
28 wash, re-suspension in FBS containing 10% of the cryo-preservative dimethyl-sulfoxide (DMSO),  
29  
30 frozen, and stored in a -80 degrees C freezer.  
31

32  
33 Prior to processing on the device, patient samples were thawed rapidly in a 37 degrees C  
34  
35 water bath, washed with PBS, and fixed for one hour in 0.5% formaldehyde prepared fresh from  
36  
37 a 16% stock solution (Polysciences, Warrington, PA, #18814-20). After fixation, cells were  
38  
39 washed twice in PBS, re-suspended in 150  $\mu$ L 0.1% PBS with 0.1% BSA (PBSA), and stored at  
40  
41 40 degrees C until ready to process. Before sample delivery, the cell suspension was diluted in a  
42  
43 20% glycerol/0.1% PBSA solution to improve cell distribution across the membrane and to reduce  
4  
45 cell clumping.  
46

47  
48 Using a custom built manifold connecting external fluidic tubing to the inlet and outlet ports  
49  
50 of the microfluidic device, the assembly was positioned on a robotically controlled microscope  
51  
52 stage (ProScan II, Prior Scientific, Cambridge, UK) and connected to a peristaltic pump (SciQ  
53  
54 400, Watson Marlow, Wilmington, MA) and manually controlled 6-position injector valve (Vici,  
55  
56  
57  
58  
59  
60

1  
2  
3 Valco Instruments, Houston, TX). Antibody stock solutions were vortexed for 30 seconds and  
4 centrifuged at 14,000 rpm for 5 minutes before preparing working dilutions to avoid precipitates.  
5  
6

7 All assays contained Phalloidin and DAPI in the secondary antibody cocktail, but each  
8 was specific for a single molecular biomarker primary-secondary antibody pair. Working dilutions  
9 of antibodies were prepared in 0.1% PBSA with 0.1% Tween-20 (EMD Millipore, Billerica, MA, #  
10 655206). Primary monoclonal antibodies were raised from either mouse (EGFR [Life  
11 Technologies, Carlsbad, CA, #MS-378-P, 10 µg/mL]), rabbit (αβ6 [Abcam, Cambridge, MA, #  
12 #Ab124968, 6 µg/mL], Ki67 [Abcam #Ab15580, 29 µg/mL], and MCM2 [Abcam #Ab108935, 10  
13 µg/mL]), or goat (CD-147 [EMMPRIN] [R&D Systems, Minneapolis, MN, #AF972, 20 µg/mL].  
14 AlexaFluor-488 conjugated secondary antibodies were specific for F (ab')<sub>2</sub> fragments of mouse  
15 IgG (Life Technologies #A11017, 20 µg/mL for EFGR), rabbit IgG (Life Technologies #A11070,  
16 50 µg/mL for αβ6, 64 µg/mL for Ki67, and 23.5 µg/mL for MCM2), or goat IgG (Life Technologies  
17 #A11078, 40 µg/mL for CD147). A working concentration of 0.33 µM was used for Phalloidin-  
18 AlexaFluor-647 (Life Technologies #A22287) and 5 µM for DAPI (Life Technologies #D3571).  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32

33 In summary, the lab-on-a-chip sample processing was comprised of the following steps:  
34 1) the device was primed with PBS at a flow rate of 735 µL/min for 2 minutes, 2) the cell  
35 suspension in 20% glycerol/0.1% PBSA was delivered at 1.5 mL/min for 2 minutes, 3) cells were  
36 washed with PBS at 1 mL/min for 2.5 min, 4) the primary antibody solution was delivered through  
37 a 0.2 µm PVDF syringe filter at 250 µL/min for 2.5 min, 5) a wash step similar to step 3 was  
38 performed, 6) the secondary antibody solution was delivered under the same conditions as step  
39 4, 7) a final wash step was performed, and 8) automated image capture was performed.  
40  
41  
42  
43  
44  
45  
46  
47

## 48 **Sample Digitization**

49 More complete details on cytology sample digitization and a complete list of intensity and  
50 morphological parameters can be found in our previous publication.<sup>11</sup> Images were recorded with  
51 a motorized reflected fluorescence microscope (Olympus BX-RFAA) equipped with a CCD  
52  
53  
54  
55  
56  
57  
58  
59  
60

1  
2  
3 camera (Hamamatsu ORCA-03G) through a 10x objective (10x/0.30NA UPlanFI, Olympus). A  
4 total of 25 unique fields of view (FOVs) repeated for 3 different z-focal planes were automatically  
5 captured across a 20 mm<sup>2</sup> area using a robotic x-y-z microscope stage. Due to the complex three-  
6 dimensional morphology of oral squamous cells, multiple z-focal planes were captured and  
7 subsequently combined into a single, enhanced depth-of-field image to simplify the multi-spectral  
8 detection of the three fluorescent labels using ImageJ “stack focuser”.

9  
10 Combinations of custom macros and the open-source image analysis tools ImageJ<sup>12</sup> and  
11 Cell Profiler<sup>13</sup> were developed to automatically detect individual cells and define their nuclear and  
12 cytoplasmic boundaries as individual regions of interest (ROI). These ROIs were used to obtain  
13 intensity measurements associated with the three spectral channels and were used to define  
14 morphometric parameters. The DAPI and Phalloidin molecular labels served primarily to assist in  
15 the automated segmentation of individual nuclei and cytoplasm, respectively.  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26

## 27 28 **Cell Identification Model Training and Validation**

29  
30 A cell type classification model was explored for its ability to discriminate and quantitate  
31 the frequency and distributions of four cell types: Type 1 (mature squamous cells), Type 2 (small  
32 round cells), Type 3 (leukocytes), and Type 4 (lone nuclei). To recognize these phenotypes, a  
33 machine learning algorithm was trained on 144 cellular/nuclear features from single-cell analyses,  
34 including morphological and intensity-based measurements. A training set was manually  
35 compiled by randomly selecting and labeling cells, resulting in approximately 100-200 single-cell  
36 objects for each of the four cell types. All features were log-normalized and standardized for zero  
37 mean and unit variance. Principal component analysis (PCA) was performed on the training set,  
38 and a scatterplot of the first two principal components was generated to visualize the internal data  
39 structure and variance. A *k*-nearest neighbors (*k*-NN) classifier was trained on the standardized  
40 features using 10-fold cross-validation and configured to find the nearest 7 neighbors in feature  
41 space (Euclidean distance). Cross-validated predicted responses by the *k*-NN classifier were  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60

1  
2  
3 recorded, and accuracy was reported for the overall cross-validation set and individually for each  
4 of the four cell types.  $k$ -NN model responses with 4 or less out of 7 similar neighbors were labelled  
5  
6  
7 “unknown” type, and cross-validated accuracy was reported for the overall training set after  
8  
9 accounting for unknown object types.

10  
11  
12 The cell type classification model was retrained on the entire training dataset, and this  
13  
14 final model was applied to the study population and averaged across each of the six molecular  
15  
16 biomarker assays. Results are presented for only subjects with evaluable data for all biomarker  
17  
18 measurements ( $N = 486$ ). Boxplots were generated to show the distributions of cell phenotypes  
19  
20 across 4 diagnostic categories as follows: 121 normal/non-neoplastic, 241 benign, 59 dysplasia,  
21  
22 and 65 malignant. Median values of cell phenotypes were compared for all lesion determinations  
23  
24 using a two-sided Wilcoxon rank sum test at a significance level of  $p = 0.05$ . Cell phenotype  
25  
26 frequencies and distributions for each subject were retained for use in clinical algorithm  
27  
28 development.

29  
30  
31 The same cell type identification model development process was completed on recently  
32  
33 developed integrated instrument, cartridges, and cloud-based analysis tools. Images of benign  
34  
35 and malignant lesions were collected with this cloud POC cytology platform, and cell phenotype  
36  
37 labels were overlaid on each recognized cell object.

### 38 39 **Numerical Index and Diagnostic Models for Assessing PMOL**

40  
41 The analysis of dichotomous outcomes with mutually exclusive levels is common in clinical  
42  
43 diagnostics, and logistic regression is regarded as the standard method of analysis for these  
44  
45 situations attributed to its probabilistic interpretation and ability to function as a dichotomous  
46  
47 classifier. Clinical data are often challenged by high-dimensionality and highly correlated  
48  
49 predictors that may generate model coefficients with high variance. For these situations, a size  
50  
51 penalty as implemented by the lasso technique may be applied to shrink the effect sizes and  
52  
53 reduce coefficient variability. Additionally, the lasso technique performs automatic parameter  
54  
55 selection by eliminating predictors with less importance. In high-dimensional data sets, reducing  
56  
57  
58  
59  
60

1  
2  
3 the set of predictors often leads to better prediction performance and generalizability and has  
4 shown improvements over manual stepwise selection methods. This lasso logistic regression  
5 model is suited to our platform because it is inherently more intuitive than previous methods which  
6 consider hundreds of measurements from cytology that are difficult to interpret.  
7  
8

9  
10  
11 A lasso logistic regression approach was used to prevent overfitting, reduce coefficient  
12 variability, and retain a sparse model with improved generalizability and interpretability. Subjects  
13 were dichotomized into “case” and “non-case” outcomes according to their lesion determination  
14 (non-case for benign lesions and case for [mild, moderate, severe] dysplasia and malignant  
15 lesions). Only subjects with evaluable data for all biomarker measurements and PMOL status  
16 were considered (N = 365). Algorithm results were recorded for 241 benign lesion and 124  
17 dysplasia and malignant lesion subjects. Diagnostic performance was characterized by area  
18 under the curve (AUC), sensitivity, and specificity. The results from six molecular biomarker  
19 assays on the POCOCT system were pooled to obtain final estimates. A receiver operating  
20 characteristic (ROC) curve was plotted for the cross-validated test set. Non-zero lasso logistic  
21 regression coefficients were retained for the following predictors: percentage of non-mature  
22 squamous cells, percentage of small round cells, percentage of leukocytes, age, sex, smoking  
23 pack years, lesion major axis diameter, clinical impression of lichen planus, and lesion color (red,  
24 white, or red/white) (see **Table S3**). Boxplots of cross-validated algorithm results were generated  
25 for the test set responses for benign, mild dysplasia, moderate/severe dysplasia, and malignant  
26 lesions. Median numerical indices were compared for each diagnostic classification using a two-  
27 sided Wilcoxon rank sum test at a significance level of  $p = 0.05$ . Internal calibration was performed  
28 by sorting and grouping the predicted responses (i.e., numerical index) into deciles and measuring  
29 the observed proportions of dysplasia/malignant lesions in each decile. The Hosmer-Lemeshow  
30 goodness of fit statistic was used to assess the model fit.  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52

53  
54 Following this same method, diagnostic algorithms for mild versus moderate dysplasia  
55 (3|4), low versus high risk (4|4), moderate versus severe dysplasia (4|5), healthy control (no  
56  
57  
58  
59  
60

lesion) versus malignant (0|6), and benign dysplasia versus malignant (2|6) were also developed. Model responses for each subject were averaged over all biomarker assays to inform diagnostic performance. AUC, sensitivity, and specificity were reported as mean and 95% confidence interval values for the cross-validated test set.

**Table S3. Predictor definitions**

Abbreviation	Reference	Details
1-%TYPE 1	percentage of non-mature squamous cells	1 – (number of mature squamous cells / total cells), where ‘total cells’ is the number of cells Types 1-3
%TYPE 2	percentage of small round cells	number of small round cells / total cells, where ‘total cells’ is the number of cells Types 1-3
%TYPE 3	percentage of leukocytes	number of leukocytes / total cells, where ‘total cells’ is the number of cells Types 1-3
AGE	age	age in years
SEX	sex	male = 1, female = 0
PACKYR	calculated pack years	average cigarettes smoked per day times years smoked divided by 20
LSIZEMAX	lesion size in maximum dimension	lesion diameter along the long axis in mm
LICHENFN	clinical impression of lichen planus	binary measure completed by clinician at time of brush cytology sample collection indicating the presence (“1”) or absence (“0”) of the clinical features of lichen planus
LESIONCOLOR	lesion color (red, white, or red/white)	variable indicating lesion color; white = 0, red = 1, red and white = 2

## Supplemental Methods References

1. Vigneswaran N, Beckers S, Waigel S, et al. Increased EMMPRIN (CD 147) expression during oral carcinogenesis. *Exp Mol Pathol*. 2006;80(2):147-159.
2. Torres-Rendon A, Roy S, Craig GT, Speight PM. Expression of Mcm2, geminin and Ki67 in normal oral mucosa, oral epithelial dysplasias and their corresponding squamous-cell



- 1  
2  
3 3. Weigum SE, Floriano PN, Redding SW, et al. Nano-bio-chip sensor platform for  
4 examination of oral exfoliative cytology. *Cancer Prev Res.* 2010;3(4):518-528.  
5  
6
- 7 4. Li HX, Zheng JH, Fan HX, Li HP, Gao ZX, Chen D. Expression of alphavbeta6 integrin  
8 and collagen fibre in oral squamous cell carcinoma: association with clinical outcomes and  
9 prognostic implications. *J Oral Pathol Med.* 2013;42(7):547-556.  
10  
11
- 12 5. Ylipalosaari M, Thomas GJ, Nystrom M, et al. Alpha v beta 6 integrin down-regulates the  
13 MMP-13 expression in oral squamous cell carcinoma cells. *Exp Cell Res.*  
14 2005;309(2):273-283.  
15  
16
- 17 6. Yu YH, Morales J, Feng L, Lee JJ, El-Naggar AK, Vigneswaran N. CD147 and Ki-67  
18 overexpression confers poor prognosis in squamous cell carcinoma of oral tongue: a  
19 tissue microarray study. *Oral Surg Oral Med Oral Pathol Oral Radiol.* 2015;119(5):553-  
20 565.  
21  
22
- 23 7. Daniel FI, Fava M, Hoffmann RR, Campos MM, Yurgel LS. Main molecular markers of  
24 oral squamous cell carcinoma. *Applied Cancer Research.* 2010;30(3):279-288.  
25  
26
- 27 8. Williams GH, Romanowski P, Morris L, et al. Improved cervical smear assessment using  
28 antibodies against proteins that regulate DNA replication. *Proc Natl Acad Sci.*  
29 1998;95:14932-14937.  
30  
31
- 32 9. Scott IS, Odell E, Chatrath P, et al. A minimally invasive immunocytochemical approach  
33 to early detection of oral squamous cell carcinoma and dysplasia. *Br J Cancer.*  
34 2006;94(8):1170-1175.  
35  
36
- 37 10. Speight PM, Abram TJ, Floriano PN, et al. Interobserver agreement in dysplasia grading:  
38 toward an enhanced gold standard for clinical pathology trials. *Oral Surg Oral Med Oral*  
39 *Pathol Oral Radiol.* 2015;120(4):474-482.  
40  
41
- 42 11. Abram TJ, Floriano PN, Christodoulides N, et al. 'Cytology-on-a-chip' based sensors for  
43 monitoring of potentially malignant oral lesions. *Oral Oncol.* 2016;60:103-111.  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
54  
55  
56  
57  
58  
59  
60



1  
2  
3  
4  
5  
6  
7  
8  
9  
10  
11  
12  
13  
14  
15  
16  
17  
18  
19  
20  
21  
22  
23  
24  
25  
26  
27  
28  
29  
30  
31  
32  
33  
34  
35  
36  
37  
38  
39  
40  
41  
42  
43  
44  
45  
46  
47  
48  
49  
50  
51  
52  
53  
55  
56  
57  
58  
59  
60

12. Schneider CA, Rasband WS, Eliceiri KW. NIH Image to ImageJ: 25 years of image analysis. *Nat Methods*. 2012;9(7):671-675.

13. Carpenter AE, Jones TR, Lamprecht MR, et al. CellProfiler: image analysis software for identifying and quantifying cell phenotypes. *Genome Biol*. 2006;7(10):R100.

