

1 **A Protocol for Revealing Oral Neutrophil Heterogeneity by Single-Cell Immune Profiling**
2 **in Human Saliva**

3

4 Running Title: *Oral Neutrophil Profiling by sc-RNAseq*

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30 **ABSTRACT**

31 Neutrophils are the most abundant white blood cells in the human body responsible for fighting
32 viral, bacterial and fungi infections. Out of the 100 billion neutrophils produced daily, it is
33 estimated that 10 % of these cells end up in oral biofluids. Because saliva is a fluid accessible
34 through non-invasive techniques, it is an optimal source of cells and molecule surveillance in
35 health and disease. While neutrophils are abundant in saliva, scientific advancements in neutrophil
36 biology have been hampered likely due to their short life span, inability to divide once terminally
37 differentiated, sensitivity to physical stress, and low RNA content. Here, we devise a protocol
38 aiming to understand neutrophil heterogeneity by improving isolation methods, single-cell RNA
39 extraction, sequencing and bioinformatic pipelines. Advanced flow cytometry 3D analysis, and
40 machine learning validated our gating system model, by including positive neutrophil markers and
41 excluding other immune cells and uncovered neutrophil heterogeneity. Considering specific cell
42 markers, unique mitochondrial content, stringent and less stringent filtering strategies, our
43 transcriptome single cell findings unraveled novel neutrophil subpopulations. Collectively, this
44 methodology accelerates the discovery of salivary immune landscapes, with the promise of
45 improving the understanding of diversification mechanisms, clinical diagnostics in health and
46 disease, and guide targeted therapies.

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60 1. INTRODUCTION

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62 Rapid advances in mapping single-cell transcriptional states have been made through the
63 Human Cell Atlas (HCA)¹ and the NIH BRAIN initiatives, providing important insights into
64 human health². By considering transcript and protein compositions, functional categorization of
65 cells is reshaping our models of ontology and helping discover new cell populations and
66 elucidating heterogeneity. Immunity works through orchestrated cellular actions with specialized
67 cellular tasks. In an effort to catalogue immune repertoire, studies are surveying specific tissues
68 and biofluids revealing unique signals, differentiation patterns, cell activation states and diversity.
69 While a large number of immune cell populations, especially lymphocytes and macrophages, have
70 been widely investigated by single cell analysis there is a lack of investigation in neutrophils, the
71 most abundant myeloid cells³⁻⁶. Investigating neutrophil cell diversity is an emerging field with
72 the potential to reveal novel cell functions and applications⁷⁻¹⁰.

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74 Neutrophils are produced at a rate of 10^{11} cells/day, comprising 50 to 70 % of all leukocytes
75¹¹ and highly active to respond to infection, injury, migrating to sites initiating inflammation^{12,13}.
76 Human neutrophils mature in the bone marrow from committed myeloid precursors and through
77 subsequent stages of differentiation they develop into segmented adult mature cells¹⁴.
78 Characterization of neutrophil diversity by transcriptomic data has been limited due to their
79 complex nature including short lifespan, lower RNA content, high levels of intracellular
80 phosphatases and nucleases, and sensitivity to handling during experimentation, resulting in false
81 negative and low heterogeneity in single cell studies. Neutrophils are very sensitive to physical
82 stress such as higher centrifugation speeds, and cold temperatures commonly used in processing
83 of cells for sorting and RNA-seq. Moreover, the levels of neutrophils in published datasets could
84 have been artificially reduced by quality control steps in data processing of RNAseq data which
85 bias against their properties of neutrophils^{15,16}.

86

87 Conventional protocols in blood employ peripheral blood mononuclear layer PBMC as a
88 sample source for single-cell transcriptomic studies of immune cells. In the context of neutrophils,
89 this method allows the detection of homogenous normal density neutrophils (NDNs) found in the
90 granulocyte layers at the interface between red blood cells and the gradient layers, but fails to

91 detect other populations, including activated NDNs that are immunosuppressive with
92 proinflammatory functions¹⁷. These cells are called low density neutrophils (LDN) and are only
93 found in chronic diseases at the mononuclear layer (PBMCs) between gradient and plasma, which
94 are not found in samples from healthy subjects¹⁸. Another level of difficulty is the fact that cells
95 in PBMCs contain 10-20 times more RNA than neutrophils¹⁹, thus sequencing data from
96 neutrophils gets masked or filtered out through conventional PBMC protocols¹⁹. Consequently,
97 in cell heterogeneity studies, neutrophil detection has not followed the trends of other immune
98 cells with regard to diversity, which we believe is related to these technical issues. Since the results
99 from neutrophil single cell transcriptomics reveal either low gene expression levels or decreased
100 functional diversity^{15,16,20,21}, there is a need to overcome technical difficulties and further
101 understand neutrophil heterogeneity.

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103 In addition to blood, neutrophils are also found in high abundance in mucosal tissues and
104 biofluids such as urine, tears, and saliva. The oral cavity is, however, an excellent and non-invasive
105 source to access biological materials for sampling and single cell studies. Buccal swabs and saliva
106 are the two most common oral sampling methods used for biomedical research. Saliva is rich in
107 mucins, host cells, and most types of proteomic markers found in the body are also detected here.
108 The immune salivary landscape is poorly defined, and leukocytes, lymphocytes and squamous
109 epithelial cells are also present in the oral cavity²². The cellular content is derived from mucosal
110 blood vessels, crevicular fluids, oral sweat, and salivary glands. It is estimated that 1/10 of all
111 neutrophils produced daily in the bone marrow are destined for oral tissues²³. The function,
112 diversity and heterogeneity of neutrophils remains elusive, and while knowledge from bulk assays
113 has been developed suggesting heterogeneity, unbiased single-cell investigations are needed to
114 capture the full repertoire.

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116 Processing single cells in biofluids presents unique challenges including preservation of
117 activation states and relationship to the microbiome that require the development of specialized
118 methods, including buffers, cell handling and protocol to successfully derive scientific insights to
119 specific cells and their populations. In addition to biological limitations specific to neutrophils,
120 there are no established protocols aiming to improve their detection rate and functional analysis.
121 In this study, we establish that neutrophils require unique handling strategies to retain their RNA

122 integrity during experimental procedure than those commonly used for preparation of other cell
123 types in RNA-seq studies. We further demonstrate a protocol that uncovers the diversity of
124 neutrophil heterogeneity. A modified Smart-seq2 method for cDNA synthesis, and bioinformatics
125 pipeline for scRNA-seq data analysis of low mRNA-containing cells is shown, profiling of a large
126 number of cell populations and genome-wide features by changes of buffers and cell handling of
127 neutrophils. We detected novel neutrophil subpopulations and leveraged recent technological
128 advancements in bioinformatics to design filtering criteria to define the signatures. In combination
129 with machine learning profiling, and 3D flow cytometry analysis, we have validated our findings
130 and explored clusters based on surface markers and density. Altogether, our protocol and
131 feasibility studies demonstrated signatures of cells contributing to the human oral neutrophil
132 heterogeneity. We suggest that our methods can be applied to heterogeneity in salivary immune
133 cells, especially oral neutrophils, as an important feature to health and disease investigations,
134 providing insights into possible mechanisms of cell diversification and their functions.

135 **1.1. Development of the Protocol:**

136 Our experimental workflow (Fig. 1) begins with harvesting primary cells by collection of
137 saliva through oral rinse from three healthy donors to obtain cells present in the buccal cavity,
138 using 0.9 % sterile saline solution, followed by a series of filtration using 40, 20, and 10 μm cell
139 strainers to eliminate food particles and larger epithelial cells to obtain > 98 % enriched saliva
140 neutrophil. After each filtration step the change in the cell population is verified by microscopic
141 imaging of unstained cells by 3D- holotomographic microscope (Fig. 1A). Unstained/unlabeled
142 neutrophils are then FACS sorted for high-FSC and high-SSC to obtain cell samples for unbiased
143 transcriptomic gene-signature profiling and the possible discovery of new sub-population(s) of
144 neutrophil (Fig. 1B), which is the primary goal of this study. Single cells were FACS sorted into
145 separate wells containing lysis buffers in a 384 well FrameStar plate and stored at -80 °C for
146 downstream processing. scRNA-seq was performed by a modified Smart-seq2 protocol following
147 various quality-control (QC) steps between each step²⁴. Single cell wells passing QC assay criteria
148 were selected for NexteraXT library preparation and sequencing on an Illumina NovaSeq 6000
149 platform. Bioinformatics analysis was performed on the generated sequencing data to assess
150 sequence quality and batch effects, to map reads to the reference human transcriptome and quantify

151 expression, to identify distinct transcriptional phenotypes using unsupervised clustering, and to
152 evaluate expression of known neutrophil markers from the literature (Fig. 1D). Enriched saliva
153 neutrophil samples were also used for FACS sorting to validate expression of known neutrophil
154 markers including CD66b⁺CD11b⁺CD14⁻ (Fig. 1C).

155 **1.1.1. Sample Handling for Saliva Neutrophil Enrichment:**

156 Fresh primary human saliva was harvested and prepared from healthy donors. Each donor
157 rinsed five times with ~10 mL of 0.9 % NaCl solution (sterile) for 30-60 sec. with a gap of 3 min
158 between each rinse to collect 50 mL total volume. The cells were then pelleted down through
159 centrifugation at 160 x g at room temperature (RT) for 5-10 min. From 50 mL total volume, 40
160 mL of supernatant were discarded by aspirating carefully without disturbing the cell pellet. It is of
161 utmost importance to maintain each step of primary saliva neutrophil isolation at RT as the cells
162 tend to lose viability faster at lower temperature (e.g. at 4 °C). Cells were then resuspended in the
163 remaining ~10 mL of 0.9 % saline solution and filtered through a 40-micron nylon mesh filter
164 using gravity to remove food particles and eliminate mucus present in the collected oral rinse. 40
165 µm filtered oral rinse was sequentially filtered through 20 µm and 10 µm pluriStrainer nylon mesh
166 filters to remove epithelial cells. All filtration steps were done by gravity flow or centrifugation at
167 160 x g at RT for 1 min. Vacuum suction for pluriStrainers should be avoided due to increased
168 contaminants. Viability and cell density were determined through a trypan blue exclusion using
169 the Countess automated cell counter. Cells obtained by this method from a single healthy donor in
170 50 mL oral rinse of resting saliva was ~1.8 x 10⁵ cells. The purity of saliva neutrophils isolated by
171 this enrichment method was verified by both microscopy (Fig. 2) and flow cytometry (Fig. 3). To
172 compare the identification of neutrophil subpopulation by flow cytometry and scRNA-seq, we
173 looked at the heterogeneity of the immune cell in whole saliva by computational analysis of flow
174 cytometry data using FLOCK²⁵ unsupervised analysis for identifying both known and novel cell
175 populations from the flow cytometry data (Fig. 4). The expression profile of neutrophil-specific
176 genes from the scRNA-seq data also verified the purity of the FACS-sorted single cell as
177 neutrophils (Fig. 5 and Fig. 6).

178

179 **1.1.2. Flow Cytometry Sorting:**

180 To ensure unbiased transcriptomic patterns, cells were not labeled with antibody/cell-
181 surface-marker prior to sorting. The viability of cells was checked before each sorting and only
182 samples containing > 70 % viable cells were sorted. Cells were sorted using BD FACS Aria II (BD
183 Biosciences) with custom PMT2 using a 130-micron (10 PSI) nozzle size. A nozzle size of 100-
184 micron is desirable due to a flow pressure of 20 PSI and would also be suitable to use. The flow
185 pressure of 70 PSI generated by a 70-micron nozzle during sorting affects both the viability and
186 RNA quality of sorted neutrophil cells. Cell populations were selected by software gating for high-
187 FSC/high-SSC Scatter-gate followed by doublet-discrimination (DD) (FSC-DD and SSC-DD
188 gates) to isolate single cells from any multicellular clusters/clump. Except for control wells, single
189 cells were sorted into each well of a Framestar 384-well plate containing 2 μ L of pre-dispensed
190 lysis buffer having ERCC (External RNA Controls Consortium) spike-in RNA standards (final
191 concentration of 2 % Triton X-100, 2 U/ μ L RNase inhibitor, 1:20,000 dil. ERCC and nuclease free
192 water). Blank or 'No Template Control (NTC)' wells contained 2 μ L nuclease free water only;
193 'spike-in' control well contained lysis/FACS sort buffer with ERCC but no cell; and 'UHR' control
194 well contained 2 μ L lysis/sort buffer + 10 pg of Universal Human RNA (UHR). Except for the
195 sorting-sample tube, all other reagents and FACS sorting plates were maintained at 4 °C on ice by
196 using chill blocks. The lysis/sort buffer must be prepared fresh. The sorted plates were then frozen
197 on dry ice and stored at -80 °C for downstream processes, including cell lysis, cDNA synthesis
198 and PCR amplification according to the SMART-seq2 protocols. Out of three 384 wells, we
199 generated 1084 single-cell cDNA wells and selected for QC-Pass criteria/parameters set to a single
200 cDNA library plate (Supplementary Fig. 2).

201 To compare the effect of pressure and electrostatic charge on cells while sorting by BD
202 FACS Aria II, we also tested single cell sorting using other microfluidics based cell sorters,
203 including the On-chip Sort-microfluidic chip cell sorter (On-chip Biotechnologies Co., Ltd, Japan)
204 and the WOLF Cell sorter (NanoCollect Biomedical Inc., CA, USA). With the WOLF cell sorter,
205 the volume requirement was very high and not compatible for use with the Smart-Seq II reaction.
206 On the other hand, the volume needed by On-chip Sort is minimal allowing for its use with the
207 Smart-seq protocols. In addition, On-Chip-SPiS single-cell dispenser captures high quality images
208 providing a visual confirmation of single deposition into the wells (Supplementary Fig. 1). We

209 compared the purity of bulk sorted cells and quality of RNA of both systems and found that cell
210 viability and cDNA obtained from On-Chip sorters were superior. However, due to its inherent
211 pipetting system, the timing of this procedure was longer than neutrophil handling would allow.
212 BD FACS Aria II workflow provided cell sorting with the volume needed, during a time frame
213 compatible with maintaining good neutrophils viability and RNA quality.

214 **1.1.3. Cell lysis, cDNA Synthesis and Quality Control:**

215 Detailed procedure of cell lysis and cDNA for Smart-seq2 protocol was previously
216 published²⁶. In addition to modifications²⁴ e.g. increasing cDNA PCR-amplification cycle from
217 18 to 21 cycles to compensate for lower mRNA levels in neutrophil^{15,16,20} and modifying Template
218 Switch Oligo (TSO) primer by 5'-biotinylation to reduce non-specific amplification caused by
219 TSO concatemers^{24,27}. Before preparing the library for scRNA-seq, we carried out quality control
220 assays of the cDNA samples by quantification using a PicoGreen dsDNA quantitation assay kit
221 and qRT-PCR for ACTB expression using a TaqMan gene expression assay. A Hamilton Microlab
222 STAR liquid handling system was used to select samples that passed the QC parameters for both
223 PicoGreen dsDNA assay (RFU > 0.3) and qRT-PCR for ACTB expression (Ct < 35)
224 (Supplementary Fig. 2).

225 **1.1.4. Preparation of sequencing library and sequencing:**

226 We employed the Illumina Nextera XT DNA library preparation kit and performed
227 multiplexed paired-end sequencing of barcoded libraries using an Illumina MiSeq in order to
228 determine the required sequencing parameters for sequencing on the pooled libraries on a NovaSeq
229 6000 sequencing system. Samples in each well were barcoded by using a unique combination of
230 Nextera XT Index Kit v2 Set A and Set D to identify sequence data from each single cell. To
231 ensure initial quality of the sequencing libraries, we performed a MiSeq run on a pooled library of
232 16 randomly selected Nextera XT libraries created from the selected cDNA plates from each donor
233 in order to determine the sequence quality and coverage needed to discover subpopulations among
234 saliva neutrophils. Prior to loading the MiSeq, an Agilent 2100 BioAnalyzer High Sensitivity DNA
235 chip run was performed and insert size of the sequencing library was determined to be between

236 250–500 bp. For the MiSeq-MO (medium output) run, 270 pM of 16-plex pool library was spiked
237 with 1 % PhiX174. The 150 base paired-end MiSeq run of 16 single-cell data showed an average
238 Q30 of Read 1 (75%), Read 2 (69%), i7 (83%) and i5 (79%) pass filter rate and a sequencing depth
239 of $1.5\text{--}2.0 \times 10^6$ reads per cell. It has been previously shown that read-depth of $1.5\text{--}2.0 \times 10^6$ is
240 adequate for the detection of saturating levels of RNA expression in single cells ²⁸. This
241 information suggested that we could perform a sequence run of NovaSeq 6000 with a single pool
242 of 384 Nextera XT libraries without over saturation of the RNA-seq read depth and gene counts
243 from each single cell.

244 Based on the MiSeq run results, a total of 3 NovaSeq 6000 SP flow cell 2x150 XP workflow
245 runs were performed on each of the 3 donors with 3 library pools consisting of 384 Nextera XT
246 libraries from both single cells and controls. Each pool was loaded at 300 pM with a 1 % PhiX174
247 spike-in. Quality of 3 NovaSeq 384-plex pooled libraries were analyzed by both Agilent HS DNA
248 chip and qPCR-based NGS library quantification using KAPA Library Quantification Kit -
249 Illumina. FACS sorted single-cell plates were evaluated by PicoGreen dsDNA quantification assay
250 (for cDNA concentration) and qRT-PCR TaqMan assay (for expression of housekeeping gene
251 ACTB) for cDNA quality. Each selected cDNA library was used to generate an Illumina Nextera
252 XT library and combined into a 384-plex pool for sequencing on the Illumina NovaSeq 6000
253 system.

254 **1.1.5. Cell Morphology and Immunofluorescence 3D Holotomographic Microscopy:**

255 Cell diversity of healthy human resting saliva and elimination of each cell type during each
256 filtration steps of 40-, 20-, and 10-micron by our saliva neutrophil enrichment protocol was
257 analyzed by 3D- holotomography imaging of unstained cells and cells stained with fluorophore-
258 conjugated cell surface CD-markers using 3D Cell Explorer microscope (Nanolive's 3D Cell
259 Explorer-*fluor*; Model CX-F). Cells were collected after each filtration step of neutrophil
260 enrichment protocol. Unstained and stained cells suspended in 0.9 % saline solution and staining
261 buffer respectively were imaged at 60X magnification using class 1 low power Laser ($\lambda=520$ nm,
262 sample exposure 0.2 mW/mm²) and a depth of field of 30 μ m. For fluorescent imaging, cells were
263 stained by a flow cytometry staining protocol and were imaged by the fluorescent module of 3D
264 Cell Explorer-*fluor* in DAPI, GFP (or FITC), and OFP (or TRITC) fluorescent channels. Exposure

265 of 100/100/100 ms, gains of 35/35/35 and intensities of 35/50/50 were used to capture images in
266 DAPI/GFP/OFP-channels, respectively (Fig. 2A).

267 **1.1.6. Validation of Neutrophil Morphology by Microscopy and Giemsa Staining:**

268 To confirm the identity of unstained neutrophils done by 3D Holotomography microscope,
269 based on their intracellular structure, we stained the same samples with Giemsa stain solution and
270 imaged by Zeiss AxioVision microscope (Carl Zeiss Microscopy, LLC, NY, USA) at 40X
271 objective using Zen Blue software. For staining, enriched neutrophils obtained after 10 μ m
272 filtration step were spread into a cell monolayer on the charged side of the slide by CytoSpin 4
273 centrifugation. Around 25,000 cells in 50-200 μ L volumes of cell suspension are loaded in
274 Cytofunnel with white filter cards and caps (Shandon EZ Single Cytofunnel, cat. no. A78710003)
275 for each slide and centrifuged at 700 RPM for 7 min at medium acceleration. Boundaries were
276 drawn around the cell monolayer by using a hydrophobic pen and allowed to dry at RT inside the
277 hood for ~30 mins. After drying, the cells were methanol fixed by incubating for 5-7 min at RT.
278 The slides were carefully washed twice in PBS⁽⁻⁾ and dried by draining the PBS⁽⁻⁾ completely. At
279 this step, the slides were dried and stored overnight at 4°C. After drying, Giemsa stain solution
280 (1:20 dilution) was added and incubated at RT for 30-60 mins. After incubation, the slides were
281 washed carefully by draining the stain and slowly dipping the slide in an angular position in
282 deionized water to prevent cells from getting washed away. A second wash was performed by
283 dipping the slide in fresh deionized water for 2 min. Slides were dried in the hood and then
284 mounted with coverslip using aqueous based mounting media such as CytoSeal 60 and observed
285 in brightfield under 40X objective of microscope and images. Images were captured and processed
286 by using Zen Blue software (Fig. 2B) and Nanolive 3D-Cell explorer microscope (Fig. 2C; section
287 1.9.10).

288 **1.1.7. Immunofluorescence for Microscopy and Flow Cytometric Analysis:**

289 Cells from saliva samples were further processed for immunofluorescence. Briefly, cells
290 were fixed by 4 % paraformaldehyde (PFA) on ice for 30 min. After incubation, fixed cells were
291 washed at least twice by adding a staining buffer and centrifuged at 160 x g for 10 mins at RT to

292 pellet cells. Because neutrophils don't form a clear pellet, the supernatant is carefully aspirated at
293 slow speed from the top. After cell counting, 1 μ L of Fc-block per million cells was added,
294 followed by incubation for 15 mins at RT. The volume was reconstituted to 200 μ L cell
295 suspension/tube. Anti-human monoclonal antibodies were added to each tube according to
296 manufacturer instructions. In the master mix, cell surface markers were added and incubated at RT
297 for 1 hr in dark. Suggestive cell surface markers: PerCP/Cy5.5-CD11b (cat. no. 101227,
298 BioLegend, CA), Brilliant Violet 650-CD15 (cat. no. 323033, clone W6D3, BioLegend, CA),
299 Pacific Blue-CD66b (cat. no. 305111, clone G10F5, BioLegend, CA), APC/Cy7-CD3 (cat. no.
300 300425, clone UCHT1, BioLegend, CA), PE-CD19 (cat. no. 302207, clone HIB19, BioLegend,
301 CA), APC-CD14 (cat. no. MHCD1405, Invitrogen, CA), FITC-CD18 (cat. no. MHCD1801,
302 Invitrogen, CA). Cell counts were obtained by trypan blue exclusion in the Countess cell counter
303 (Invitrogen, CA). Fixed-unstained/stained samples for flow analyses were run on a BD FACS Aria
304 II (BD Biosciences) to obtain raw data in FCS format, which were later analyzed by FlowJo
305 v10.6.1 (BD Biosciences) for 2D analysis (Fig. 3) and by FLOW Clustering without K (FLOCK
306 v1)²⁵ for computational analysis using all markers simultaneously (Fig. 4), which is explained in
307 detail in section 1.1.11. Each sample was analyzed for 'ungated' total cell populations and 'high-
308 SSC gated' populations used for single-cell sorting of saliva neutrophil in this study.

309 **1.1.8. scRNA-Seq data processing and Analysis:**

310 Single-cell RNA-seq data were processed according to the procedure described in detail in
311 Krishnaswami et. al. 2016²⁴. Briefly, raw sequencing files were demultiplexing using Illumina
312 barcodes, while sequencing primers and low-quality bases were removed using the Trimmomatic
313 package²⁹. Trimmed reads were then aligned using HISAT³⁰ in two steps: first to a reference of
314 ERCC sequences, and then the remaining reads were mapped to GRCh38 (GENCODE). StringTie
315³⁰ was then used to assemble the resulting alignments into gene expression values (TPM) estimated
316 using GENCODE v25 annotation (Ensembl 87; 10-2016).

317

318 Expression values for non-control cells were imported into Scanpy for PCA, UMAP, and
319 cluster analysis³¹. Cells were filtered for quality using two different approaches. The first liberal
320 filtering consisted of removing cells with less than 50 genes per cell or with greater than 10% of

321 the total gene counts being from mitochondrial genes. Genes were required to be present in at
322 least 4 cells, with greater than 50 total counts. From these genes, the top 2500 variable genes were
323 selected (Fig. 5). The second stricter filtering consisted of removing cells with less than 400 genes
324 per cell or with greater than 10% of the total gene count being from mitochondrial genes. In fact,
325 genes were required to be present in at least 4 cells, with greater than 50 total counts. From these
326 genes, the top 2000 variable genes were selected (Fig. 6).

327

328 Unsupervised clustering was performed by first performing PCA to determine principal
329 components, then a neighborhood graph was constructed using those components. Next Louvain
330 clustering was performed using the neighborhood graph. Using the Louvain clustering solution,
331 marker gene determination was performed using logistic regression³¹. The outputs of this
332 computational pipeline produce a set of unbiased cell type clusters, a gene expression matrix with
333 the expression levels of genes in individual single cells grouped into cell type clusters, and a set
334 of marker genes for each cell type cluster.

335 **1.1.9. Publicly available healthy control data**

336 In addition to salivary neutrophils, RNA-seq data of immune-cell types and PBMC
337 obtained from blood were acquired from the Gene Expression Omnibus (GEO) database under
338 accession code [GSE64655](#)³². This data was analyzed by Scanpy in a similar fashion described
339 above. Expression patterns for targeted marker genes, those with known differential expression
340 patterns in the oral and blood neutrophils, were visualized (Supplementary Fig.6).

341 **1.1.10. Microscopic Analysis:**

342 Raw images obtained for both unstained and stained cells were processed by software
343 (STEVE software v1.6.3496, Nanolive) with similar parameters for the brightfield images which
344 was also used to obtain digital acid-stained and RI 3D-rendering images. Parameters for processing
345 of raw images captures in each fluorescent channel (i.e. DAPI, FITC and TRITC) to determine the
346 background cutoff pixel was determined by comparing with images of unstained negative control
347 samples. These parameters were used consistently for all images processed for each fluorescent
348 channel.

349 For unbiased identification of cell types based on CD-markers, unstained cells were imaged
350 and stained via digital acid-staining (STEVE software v1.6.3496, Nanolive) according to the
351 refractive index (RI) of the intracellular structures (Fig. 2A). Four major cell types i.e. epithelial
352 cells, neutrophils, monocytes, and lymphocytes were identified based on their size and nuclear
353 structure and quantified by manually counting from multiple images obtained from three healthy
354 donors (Fig. 2C). To further verify the neutrophil enrichment and confirm the elimination of other
355 cell types, we did Giemsa staining of 10 μ m filtered unstained samples (Fig. 2B insert).

356 To rule out the biased staining of cell types with cell surface markers, stained samples used
357 in flow cytometry analyses were further imaged for selected CD-markers conjugated with
358 fluorophores detectable in the three fluorescent channels (i.e. DAPI, FITC, and TRITC channels)
359 available in Nanolive's 3D Cell Explorer microscope. Classical neutrophil positive marker CD-
360 66b conjugated with pacific blue was imaged in the DAPI channel, whereas classical neutrophil
361 negative markers CD14 and CD19 conjugated with FITC and PE, respectively were imaged in the
362 FITC and TRITC channels (Fig. 2A). Four major CD-marker positive cells i.e. CD-66b-PacBlue
363 for neutrophils, CD14-FITC for monocytes, and CD19-PE for B-lymphocytes were identified
364 based on obtaining positive signals in the respective channels.

365 **1.1.11. Flow cytometry Data Analysis:**

366 The FCS files from BD FACS Aria II flow cytometry were transformed using FCSTrans³³
367 on R programming language. FCSTrans applies a logical transformation on fluorescence channels
368 and a linear transformation on scattering parameters followed by a min-max linear rescaling
369 applied across all the channels to scale the range to [0, 4095]. After FCSTrans transformation, area
370 parameters of measured channels were selected and saved to tab-delimited text files for
371 downstream data clustering analysis. Auto-gating is applied using an unsupervised clustering
372 approach – FLOCK²⁴ (Flow Clustering Without K). FLOCK is publicly available on ImmPort
373 Galaxy (<https://immportgalaxy.org/>). FLOCK identified 23 cell subsets across the 6 stained
374 samples (two replicates for each of the three filtrations: 40 μ m, 20 μ m and 10 μ m) by using all
375 measured parameters simultaneously in density-based clustering analysis. Then each identified
376 cell subset was visually examined on dot plots of all 2D marker combinations for manually
377 annotating the phenotype (e.g., CD3⁻CD19⁻CD14⁻CD11b⁺CD15⁺CD66⁺ neutrophils).

378 Frequencies of the 23 FLOCK-identified cell subsets were quantified by their percentages
379 (with the total number of cells as the parent). Mean fluorescent intensity (MFI) values of each cell
380 population for each marker were also calculated. Figure 4A visualizes the MFI values of a 40 µm
381 sample (Tube-10_Specimen_001_002) using a heatmap for indicating the phenotype of each
382 identified cell population. Based on visual examination of the 2D dot plots, 5 of the 23 FLOCK-
383 identified cell subsets are salivary neutrophils. For each neutrophil subpopulation identified by
384 FLOCK, mean percentage is calculated for duplicate files in each filtration step. Figure 4B shows
385 the bar graph for mean percentages across the five FLOCK-identified neutrophil subsets,
386 indicating that the major/abundant neutrophil subpopulations consistently increased their
387 percentages as filtration size decreased. For both manual annotation and interpretation of the 5
388 neutrophil subpopulations, 2D dot plots (Fig. 4C) of the 40 µm sample with each neutrophil
389 subpopulation highlighted in a different color were generated to visualize the phenotype
390 differences of these subpopulations.

391 **1.2. Comparison to other methods:**

- 392 ● The key strength of our protocol are as follows:
- 393 ● The protocol overcomes the methodological limitations that produced the false-negative
394 expression of neutrophils in many published studies due to low mRNA content of neutrophils
395 and the different experimentation conditions needed in comparison with PBMCs.
- 396 ● Saliva is an easily accessible and readily available clinical sample which makes this protocol
397 non-invasive to patients requiring deep sequencing for diagnosis.
- 398 ● We suggested important experimental conditions for neutrophil that are overlooked by
399 researchers, different from processing of PBMCs. Neutrophils prefer room temperature and
400 lower centrifugation speeds of 160 x g to 300 x g.
- 401 ● This study presented processing time of primary neutrophils and the rate of RNA degradation.
402 This information is important to consider a sample for further downstream steps of an NGS
403 workflow.
- 404 ● For deep sequencing, the samples should be collected fresh and processed within 4 hours.
405 Storage of samples at -150 °C (liquid N2) or -20 °C after collection by DMSO-cryopreservation

406 and methanol-fixation protocols established in other immune cells are not suitable for deep
407 sequencing of neutrophils^{16,34}.

- 408 ● This protocol demonstrates stringency levels needed for filtering raw sequence data during
409 bioinformatics analysis. It suggests inclusion of mitochondrial gene expression data during
410 analysis that is considered an exclusion criterion for single cell sequence data processing.

411 **1.3. Limitations of this protocol:**

- 412 ● This study is aimed for unbiased sequencing of saliva neutrophils in which unlabeled
413 morphology was the gating determinant. While we coupled with specific pipelines able to
414 detect other cells, it may lead to lack of specificity in the sorting scheme.
- 415 ● Neutrophils are complex cells that are fragile and get easily activated during handling¹⁵. Thus,
416 personnel need to be trained, and reagents prepared carefully to yield replicable results.
- 417 ● Neutrophils contain 10-20 times lower amounts of RNA per cell than PBMC¹⁹. Therefore, if
418 another PBMC cell is sorted along with neutrophil by chance during FACS sorting, the
419 probability of amplification of non-targeted cell mRNA during cDNA synthesis is higher,
420 which may lead to erroneous sequencing results.
- 421 ● Due to the small amount of mRNA in neutrophils it may be necessary to optimize the PCR
422 cycles required to obtain sufficient cDNA for the NGS-workflow. We amplified the single cell
423 neutrophil cDNA with 21 cycles, similar to single nucleus protocols²⁴ because of low amounts
424 of RNA, compared with 18 cycles for other cell types²⁶. Some low-copy number transcripts
425 may still be difficult to detect in neutrophils. However, increasing the number of PCR cycles
426 could introduce some amplification bias in the library by compressing expression values for
427 high-copy number transcripts.
- 428 ● Small noncoding RNAs (ncRNAs) and other short sequence mRNAs lacking polyA tails would
429 not be detected. The low amounts of RNA contained in the neutrophils may also prevent the
430 detection of some ncRNAs.
- 431 ● Since the neutrophils are found to have generally lower RIN values compared to other cell
432 types, it is possible that the RNA of the samples are partly degraded and may result in an
433 increasingly 3' biased library preparation as suggested by Chen et al., and therefore losing
434 valuable reads from your data is a possibility³⁵.

435 **1.4. Future Applications:**

436 *Comparing Health and Disease.* Viral, autoimmune, metabolic, and chronic inflammatory diseases
437 require novel and non-invasive methods to monitor cellular phenotypes from humans, comparing
438 health versus disease states. This protocol provides the experimental conditions and time needed
439 for processing of neutrophils for NGS-workflow to obtain their transcriptomic signatures.

440

441 *Revealing Oral-Systemic Axis.* Emerging evidence demonstrates that markers expressed in
442 biofluids such as saliva are representative of systemic changes. A protocol for the unbiased
443 evaluation of single cells in saliva could yield a better understanding of systemic health through
444 oral sampling.

445

446 *Longitudinal Monitoring.* Sampling saliva is non-invasive and easy to perform. Thus, continuous
447 monitoring of cells, biomarkers and gene expression patterns in saliva provides an effective system
448 for longitudinal survey. In addition to research studies, this system would also be optimal for the
449 development of novel diagnostic systems and drug delivery.

450 **2. MATERIALS:**

451 **2.1. Reagents:**

- 452 ● *Cell sample:* We have successfully isolated human saliva neutrophils by series of the filtration
453 processes and obtained enrichment of neutrophils. We have assessed the purity of the enriched
454 neutrophil by microscopic and flow analysis. **CAUTION:** An Institutional Review Board
455 approval or patient consent form may be required for sample collection from healthy donors.
- 456 ● HL-60 cell line (ATCC, cat. no. CCL-240)
- 457 ● Iscove's Modified Dulbecco's Medium (IMDM) (Gibco, cat. no. 12440-061)
- 458 ● Roswell Park Memorial Institute (RPMI) 1640 (Gibco, cat. no. 11875-093)
- 459 ● Fetal Bovine Serum (FBS) (Gibco, cat. no. 26140-095)
- 460 ● Penicillin-Streptomycin (10,000 U/mL; Gibco, cat. no. 15-140-122)
- 461 ● Dimethyl sulfoxide (DMSO) (ATCC, cat. no. 4-X)
- 462 ● Centrifuge tubes (50 mL; Denville Scientific Inc., cat.no. C1060-P)

- 463 ● 0.9% Saline solution sterile (Teknova, cat. no. 50-843-140)
- 464 ● pluriStrainer 20 µm-sterile cell strainer (pluriSelect, cat. no. 43-50020-03)
- 465 ● pluriStrainer 10 µm-sterile cell strainer (pluriSelect, cat. no. 43-50010-03)
- 466 ● 40 µm sterile cell strainer (Fisher Scientific, cat. no. 22363547)
- 467 ● Connector ring (pluriSelect, cat. no. 41-50000-03)
- 468 ● RNaseZap RNase decontamination solution (Ambion, cat. no. AM9780)
- 469 ● Nuclease free water (Ambion, cat. no. AM9932)
- 470 ● β-Mercaptoethanol (14.3 M; Sigma, cat. no. M6250-100 mL) **CAUTION:** This is a
- 471 combustible liquid. Avoid contact with skin and eyes. Avoid inhalation of vapor or mist and
- 472 handle it while you are wearing appropriate personal protective equipment (PPE). It is toxic if
- 473 swallowed or if inhaled. It is very hazardous in case of skin contact (permeator) and ingestion.
- 474 Severe overexposure can result in death. It causes skin irritation, and it may cause an allergic
- 475 skin reaction. It also causes serious eye damage.
- 476 ● Tris buffer (pH 7.0, 1 M; Ambion buffer kit, cat. no. 9010)
- 477 ● Magnesium chloride (MgCl₂) (1 M; Ambion buffer kit, cat. no. 9010)
- 478 ● EDTA (pH 8.0, 0.5 M; Ambion buffer kit, cat. no. 9010)
- 479 ● RNase inhibitor, cloned (40 U/µl; Ambion, cat. no. AM2682)
- 480 ● Triton X-100 (Sigma-Aldrich, cat. no. T8787-100ML). **CAUTION:** Harmful if swallowed and
- 481 can cause serious eye damage. Handle it while wearing appropriate PPE.
- 482 ● dNTP mix (10 mM each; Thermo Fisher, cat. no. 18427-088)
- 483 ● ProtoScript II reverse transcriptase (New England Biolabs, cat. no. M0368X). Includes:
- 484 ● ProtoScript II buffer (5X)
- 485 ● 1,4-Dithiothreitol (DTT) 0.1 M
- 486 ● ProtoScript II Reverse Transcriptase (200 U/µl)
- 487 ● Betaine (BioUltra, ≥ 99.0 %; Sigma-Aldrich, cat. no. 61962)
- 488 ● PicoGreen dsDNA quantitation assay kit (Invitrogen, cat. no. P7589)
- 489 ● KAPA HiFi HotStart ReadyMix (2X; KAPA Biosystems, cat. no. KK2602)
- 490 ● KAPA Library Quantification Kit - Illumina (KAPA Biosystems, cat. no. KK4835)
- 491 ● Ethanol- molecular biology grade (Sigma-Aldrich, cat. no. E7023-500 ml)
- 492 ● Agencourt AMPure XP beads (Beckman Coulter, cat. no. A63881)

- 493 ● Adapter oligos (See cDNA Synthesis on section 3.4). Locked Nucleic Acid (LNA)-modified
 494 TSO were ordered from QIAGEN (<https://www.qiagen.com/>). All other oligos were ordered
 495 from IDT (<https://www.idtdna.com>). All oligos were HPLC-purified. The identity of LNA-
 496 modified TSO compounds is also confirmed by MS.
- 497 ● Biotin TSO Custom LNA Oligonucleotide (1 μmole synthesis; QIAGEN, cat. no.
 498 339413; GeneGlobe ID: YCO0078131; 5'-biotin-
 499 AAGCAGTGGTATCAACGCAGAGTACrGrG+G-3')
 - 500 ● oligo-dT (1 μmole, 57 bases; 5'-AAG CAG TGG TAT CAA CGC AGA GTA CTT
 501 TTT TTT TTT TTT TTT TTT TTT TTT TVN-3')
 - 502 ● ISPCR oligo (1 μmole, 23 bases; 5'-AAG CAG TGG TAT CAA CGC AGA GT -
 503 3')
 - 504 ● UltraPure BSA (50 mg/ml; Ambion, cat. no. AM2616)
 - 505 ● Trypan blue (0.4 %; Sigma-Aldrich, cat. no. T8154)
 - 506 ● ERCC RNA spike-in mix 1 (Ambion, cat. no. 4456740)
 - 507 ● Universal Human RNA (UHR) control (Takara/CloneTech, cat. no. 636538, discontinued
 508 product)
 - 509 ● PhiX Control v3 Library (Illumina, cat. No. FC-110-3001)
 - 510 ● RNase-free PBS, pH 7.4 (Ambion, cat. no. AM9625)
 - 511 ● 0.5 % RNase-free BSA (Ambion, cat. no. AM2616)
 - 512 ● Mouse IgG1κ (BD Pharmingen, cat. no. 554121)
 - 513 ● Fluorochrome conjugated anti-human monoclonal antibodies:
 - 514 ● PerCP/Cy5.5-CD11b (BioLegend, cat. no. 101227)
 - 515 ● Brilliant Violet 650-CD15 (clone W6D3; BioLegend, cat. no. 323033)
 - 516 ● Pacific Blue-CD66b (clone G10F5; BioLegend, cat. no. 305111)
 - 517 ● APC/Cy7-CD3 (BioLegend, cat. no. 300425, clone UCHT1)
 - 518 ● PE-CD19 (clone HIB19; BioLegend, cat. no. 302207)
 - 519 ● APC-CD14 (Invitrogen, cat. no. MHCD1405)
 - 520 ● FITC-CD18 (Invitrogen, cat. no. MHCD1801)
 - 521 ● LIVE/DEAD Fixable Aqua Dead Cell Stain Kit (Invitrogen, cat. no. L34965)
 - 522 ● Cell Staining Buffer (BioLegend, cat. no. 420201)
 - 523 ● Paraformaldehyde (PFA) (16 % w/v; Electron Microscopy Sciences, cat. no. 15710)

- 524 ● Giemsa stain solution (LabCam, cat. no. LC148407)
- 525 ● CytoSeal 60 (Thermo Scientific. cat. no. 8310-4)
- 526 ● Yellow fluorescent polystyrene microspheres, 10 μm (Spherotech, cat. no. FP-10052-2)
- 527 ● PerfeCTa qPCR FastMix II, ROX (2X, Quanta Biosciences, cat. no. 95119-05K)
- 528 ● TaqMan gene expression assay, ACTB (60X, Applied Biosystems, cat. no. 4351368, assay ID-
- 529 Hs01060665_g1).
- 530 ● RNeasy Mini Kit (50) (Qiagen, cat. no. 74104)
- 531 ● AllPrep DNA/RNA/Protein Mini Kit (50) (Qiagen, cat. no. 80004)
- 532 ● Quant-iT PicoGreen dsDNA assay kit (Invitrogen, cat. no. P11496)
- 533 ● Agilent RNA 6000 pico kit (Agilent Technologies, cat. no. 5067-1513)
- 534 ● Agilent high sensitivity DNA kit (Agilent Technologies, cat. no. 5067-4626)
- 535 ● Nextera XT DNA library preparation kit, 96 samples (Illumina, cat. no. FC-131-1096)
- 536 ● Nextera XT Index Kit v2 Set A (Illumina, cat. no. FC-131-2001)
- 537 ● Nextera XT Index Kit v2 Set D (Illumina, cat. no. FC-131-2004)
- 538 ● MiSeq Reagent Kit v2 (300-cycles) (Illumina, cat. no. MS-102-2002)
- 539 ● CytoOne T75 filter cap TC flask (USA Scientific, cat. no. CC7682-4875)
- 540 ● E-Gel General Purpose Agarose Gels, 1.2 % (Invitrogen, cat. no. A03076)
- 541 ● 1 Kb Plus DNA Ladder (Invitrogen, cat. no. 10787018)
- 542 ● Blue/Orange Loading Dye (6X, Promega, cat. no. G190A)
- 543 ● Countess cell counting chamber slides (Invitrogen, cat.no. C10283)

544 **2.2. Equipment:**

- 545 ● On-chip Sort the microfluidic chip cell sorter (Model: HSG; On-chip Biotechnologies Co., Ltd,
- 546 Tokyo, Japan; cat.no. 362S2001G)
- 547 ● Single Particle isolation System “On-chip SpiS” (On-chip Biotechnologies Co., Ltd, Tokyo,
- 548 Japan)
- 549 ● Microfluidic chip for sorting (2D Chip-Z1001; On-chip Biotechnologies Co., cat. no. 1002004)
- 550 ● BD FACS-ARIA II Flow sorter with an automated cell deposit unit (BD Biosciences)
- 551 ● BD Falcon tube with a cell strainer cap (Becton Dickinson, cat. no. 352235)
- 552 ● Falcon polystyrene conical tube (50 mL, BD Biosciences, cat. no. 352095)

- 553 ● Inverted fluorescence microscope Olympus IX70 (Olympus Corporation)
- 554 ● Zeiss AxioVision microscope (Carl Zeiss Microscopy, LLC, NY, USA)
- 555 ● 3D Cell Explorer microscope (3D Explorer-fluo; Model CX-F, Nanolive SA, Switzerland)
- 556 ● Hemocytometer (Hausser Scientific, cat. no. 1483)
- 557 ● Countess automated cell counter (Invitrogen, cat. no. C10281)
- 558 ● 96-well black Fluorac microplate (VWR, cat. no. 82050728)
- 559 ● FrameStar Clear 384-well Skirted PCR Plates (Phenix Research Products, cat. no. MPC-
- 560 384HS4NH-C)
- 561 ● FrameStar 384 Well Skirted PCR Plates- black frame with white well (Phenix Research
- 562 Products, cat. no. MPC-384HS4-WW)
- 563 ● 96-well plates (twin.tec PCR plate 96 LoBind, skirted, colorless; Eppendorf, cat. no.
- 564 0030129512)
- 565 ● 8-strip, nuclease-free, 0.2 mL, thin-walled PCR tubes with caps (Eppendorf, cat. no.
- 566 951010022)
- 567 ● Microcentrifuge DNA LoBind Safe-Lock tubes (1.5 mL; Eppendorf, cat. no. 022431021)
- 568 ● ErgoOne Multichannel pipette (USA Scientific)
- 569 ● ErgoOne single channel pipettes set, 1–10 µL; 2–20 µL; 20-200 µL (USA Scientific)
- 570 ● accu-jet pro Pipette Controller (BrandTech, cat. no. 26332)
- 571 ● TipOne ultra low retention filter tips- 10 µL; 200 µL; 1250 µL (USA Scientific)
- 572 ● DynaMag-96 side skirted magnetic rack (Thermo Fisher, cat. no. 12027)
- 573 ● MicroAmp clear adhesive film (Applied Biosystems, cat. no. 4306311)
- 574 ● MicroAmp optical adhesive film (Applied Biosystems, cat. no. 4311971)
- 575 ● GeneAmp PCR System 9700 (Applied Biosystems, cat. no. 4307808)
- 576 ● QuantStudio 6 Flex Real-Time PCR System (Applied Biosystems, cat. no. 4485694)
- 577 ● FlexStation 3 multi-mode microplate reader (Molecular Devices, San Jose, CA, USA)
- 578 ● NanoDrop ND-1000 Spectrophotometer (Thermo Fisher)
- 579 ● Cytospin 4 Cytocentrifuge (Thermo Scientific, cat. no. A78300003)
- 580 ● Microlab STAR Liquid Handling System (Hamilton Company, Reno, NV, USA)
- 581 ● Agilent BioCel 1200 system (Agilent Technologies, Santa Clara, CA, USA, cat. No. G5500-
- 582 90011) Integrates following devices:
- 583 ● Bravo Automated Liquid Handling Platform (Agilent Technologies)

- 584 ● Direct Drive Robot (DDR)
- 585 ● Microplate Centrifuge (Eppendorf)
- 586 ● BioRAPTR FRD Microfluidic Workstation (Beckman Coulter)
- 587 ● Thermo CUBE 300A Thermoelectric Recirculating Chiller (Solid State Cooling
- 588 System, NY, USA)
- 589 ● Auxiliary Barcode reader
- 590 ● Lid Hotel Station
- 591 ● Labware MiniHub
- 592 ● Labware Stacker
- 593 ● PlateLoc Thermal Microplate Sealer
- 594 ● Microplate Labeler
- 595 ● Agilent 2100 Bioanalyzer (Agilent Technologies)
- 596 ● Refrigerated centrifuge (Eppendorf, Model: Centrifuge 5804 R)
- 597 ● DNA sequencing instrument. CRITICAL: A compatible Illumina DNA sequencing instrument
- 598 (MiSeq, NextGen 500, HiSeq 2000, HiSeq 2500, NovaSeq 6000) is necessary to complete
- 599 sequencing of the Nextera XT libraries, as the barcodes and sequencing adapters are designed
- 600 for the Illumina sequencing platform.
- 601 ● 64-bit computer running Linux with 4 GB of RAM (16 GB preferred)

602 **2.3. Equipment software:**

- 603 ● VENUS (for Microlab STAR Liquid Handling System)
- 604 ● VWorks Automation Control software (for Agilent BioCel 1200 system)
- 605 ● BioRapTR 3.3.2 (for BioRAPTR FRD Microfluidic Workstation)
- 606 ● SoftMax Pro (for Flexstation 3)
- 607 ● BD FACSDiva Software v8.0.2 (for BD FACS-ARIA II Flow sorter)
- 608 ● FlowJo v10.6.1 (BD Biosciences)
- 609 ● 2100 Expert Software (For Agilent 2100 Bioanalyzer System)
- 610 ● STEVE FULL v1.6.3496 (for Nanolive 3D Cell Explorer microscope; Nanolive SA,
- 611 Switzerland)

- 612 • Zen 3.0 -blue edition (for Zeiss AxioVision microscope; Carl Zeiss Microscopy, LLC, NY,
613 USA)

614 **2.4. Sequence data analysis software:**

Software function	Software name	Web link	References
For sequence quality assessment	FASTX	http://hannonlab.cshl.edu/fastx_toolkit/download.html	
	fastQC	http://www.bioinformatics.babraham.ac.uk/projects/fastqc/	
	RSeQC (alternative to FASTX and fastQC)	http://rseqc.sourceforge.net/	36,37
For sequence trimming	Trimmomatic	http://www.usadellab.org/cms/uploads/supplementary/Trimmomatic/Trimmomatic-0.33.zip	
	Cutadapt (alternative to Trimmomatic)	https://cutadapt.readthedocs.org/en/stable/	38
For sequence alignment	HISAT2	http://daehwankimlab.github.io/hisat2/download/	30
	SAM tools	http://sourceforge.net/projects/samtools/files/samtools/	
For RNA expression analysis	StringTie	https://ccb.jhu.edu/software/stringtie/#install	
	RSEM (alternative to HISAT2/ Stringtie)	http://deweylab.biostat.wisc.edu/rsem/	

For data analysis	R	https://cran.r-project.org/	
	Python	https://www.python.org/	
	Jupyter	https://jupyter.org/install	
	Pandas	http://pandas.pydata.org/	
	Matplotlib	http://matplotlib.org/	
	Scanpy	https://pypi.org/project/scanpy/	
	Bedtools	http://bedtools.readthedocs.org/en/latest/	
	IGV	http://www.broadinstitute.org/igv/	

615

616 **2.5. Reagent setup:**

617 **2.5.1. Low Tris-EDTA (TE) buffer:**

618 Add Tris buffer, pH 7.0 (10 mM final concentration) and EDTA, pH 8.0 (0.1 mM final
619 concentration)

620 **2.5.2. Lysis/FACS Sort Buffer:**

Components of Lysis/ sort buffer	Final concentration	Vol. for 96-well (μL)	Vol. for 384-well (μL)
Triton X-100 (10 % v/v)	0.20 %	4.00	16.900
RNase inhibitor (40 U/μL)	2 U/μL	10.00	42.250
ERCC spike-in, Dilution-1 (1:2,000 dilution of stock)	1:20,000,000	0.20	0.845

Nuclease free water		185.00	785.005
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621

622 **2.5.3. Lysis Mix (Oligo dT/dNTP addition):**

Components of Lysis mix	HALF reaction (μL/rxn)	Final concentration
Anchored oligo-dT-primer (25 μM)	0.25	2.50 μM
dNTP (25 mM)	0.25	2.50 mM
Total volume	0.50	-
Lysis buffer volume/well (From STEP-9)	2.00	-
Total reaction volume/well -Lysis mix	2.50	-

623

624 **2.5.4. Reverse Transcriptase (RT) reaction mix:**

Components of Reverse transcriptase (RT) reaction	HALF reaction (μL/rxn)	Final concentration
ProtoScript II buffer (5X)	1.000	
DTT (100 mM)	0.250	4.55 mM
Betaine (5 M)	1.000	0.50 M
MgCl ₂ (100 mM)	0.300	
Template Switch Oligo (TSO) 5'-Biotinylated (100 μM)	0.055	0.05 μM
Nuclease free water	0.015	-
RNase inhibitor (40 U/μL)	0.130	5.20 U
NEB ProtoScript II reverse transcriptase (200 U/μL)	0.250	50.0 U
RT master mix total	3.000	

Lysis reaction (From Step-18) (2 μ L lysis/sort buffer + 0.50 μ L Lysis mix)	2.500	
Total volume/well -RT reaction master mix	5.500 μL	

625

626 **2.5.5. PCR-preamplification reaction master mix for cDNA synthesis master-mix:**

Components of PCR-preamplification reaction	HALF reaction (μ L/rxn)	Final concentration
KAPA HiFi HotStart ReadyMix (2X)	6.250	0.50 X
ISPCR primer (10 μ M)	0.125	0.05 μ M
Nuclease free water	1.125	-
KAPA master mix total	7.500	-
1st strand cDNA (From STEP-20)	5.500	-
Total volume/well -PCR reaction master mix	13.00	-

627

628 **2.5.6. PicoGreen dsDNA assay mix:**

Components of PicoGreen working solution	Volume (μ L/rxn)
TE-buffer (1X) volume	24.375
PicoGreen stock volume	0.125
Total volume/well (200-fold diluted PicoGreen stock sol.)	24.50
cDNA volume (From STEP-24)	00.50
Total volume/well -PicoGreen reaction mix	25.00

629

630 **2.5.7. λ -DNA standard for PicoGreen dsDNA assay:**

- 631 • First, prepare working stock of Lambda gDNA (10 ng/ μ L) by adding 2 μ L of stock (100 ng/ μ L)
 632 and 18 μ L of TE buffer
 633 • Then make 2-fold serial dilution from 10 ng/ μ L Lambda gDNA as per the table below.
 634

Final λ -DNA concentration	TE volume (μ L/tube)	Serial dilution- λ -DNA (μ L/tube)	Total vol. (μ L)
0 ng/ μ L	10 μ L	0 μ L	10 μ L
0.15625 ng/ μ L	10 μ L	10 μ L	20 μ L
0.3125 ng/ μ L	10 μ L	10 μ L	10 μ L*
0.625 ng/ μ L	10 μ L	10 μ L	10 μ L*
1.25 ng/ μ L	10 μ L	10 μ L	10 μ L*
2.50 ng/ μ L	10 μ L	10 μ L	10 μ L*
5.00 ng/ μ L	10 μ L	10 μ L	10 μ L*
10.00 ng/ μ L	18 μ L	2 μ L λ -DNA stock	10 μ L*
<i>* These tubes had 10 μL removed from the 20 μL volume for the next dilution</i>			

635

636 **2.5.8. qRT-PCR master-mix:**

Components of qRT-PCR TaqMan assay master mix	Volume (μ L/rxn)	Final concentration
PerfeCTa FastMix II, ROX (2X)	5.000	1X
ACTB- primer-probe mix (60X) [TaqMan gene expression assay ID: Hs0160665_g1]	0.166	1X
Nuclease free water	2.300	-
Total volume/well	7.500	-
cDNA; 1:10 diluted (From STEP-24)	2.500	-

Total volume/well -TaqMan assay master mix	10.000	-
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637

638 **2.5.9. Nextera library dilution for QC:**

Dilution	Diluted/undiluted Pooled library vol. (μL)	Dilution solution (μL)
1:12	2 μL (1:1 dilution)	22 μL
1:120	5 μL (1:12 dilution)	45 μL
1:1200	5 μL (1:120 dilution)	45 μL
1:12000	5 μL (1:1200 dilution)	45 μL

639

640 **2.5.10. KAPA qPCR-library quantification master-mix:**

Components of KAPA qPCR-library quantification	Volume (μL/rxn)	Final concentration
KAPA SYBR Fast qPCR master mix (2X) + Primer premix (10X)	6.00	
Sample to be assayed	4.00	
Total volume/well -KAPA qPCR master mix	10.00	

641

642 **2.6. Equipment Setup**

643 **2.6.1. Sorting**

644 For high-throughput single cell sorting with accuracy by flow cytometry using FACS
645 (Fluorescent-Activated Cell Sorting), the operator should have knowledge of doublet-
646 discrimination gating strategy and setting instrument for sorting single cell events by determining

647 the 384-well plate targeting accuracy of the droplet stream prior to sorting single cells. The
648 accuracy of a single cell being sorted to each well of a 384-well microplate for cDNA synthesis
649 can be confirmed by targeting the bottom of each microplate well with 10- μ m yellow fluorescent
650 polystyrene microspheres and observing the wells by inverting plate and imaging the microspheres
651 on an inverted fluorescent microscope. Obtaining a target accurately on 16 wells of both ends of
652 a 384-well microplate typically gives > 95 % accuracy for a single cell being sorted in each well
653 of the plate. The saliva neutrophils obtained by the enrichment protocol is split into two separate
654 tubes: one tube where cells are neither fixed nor stained for unbiased sorting (Fig. 1B) and the
655 second tube where cells are stained with CD-marker (CD11b, CD66b, CD14) and viability dye
656 (Aqua LIVE/DEAD) to check the percentage of viable cells and neutrophil population in the
657 sorting gate (Fig. 1C). If selective sorting of the target cell is desired the cells can be stained with
658 the desired marker specific to the target cell and the sort gate can be set accordingly.

659 **CRITICAL:** On the day of sorting, if the same individual operates the equipment and also
660 perform the saliva neutrophil enrichment experiment, then for sorting of unfixed saliva neutrophils
661 the equipment should be set up and kept ready before starting the saliva neutrophil
662 isolation/enrichment protocol. This is because setting up of instrument including calibration,
663 droplet delay, targeting etc. that takes almost 1 hour which will degrade the quality of RNA (i.e.
664 RIN) of isolated/enriched saliva neutrophil when left at room temperature (Supplementary Fig. 4).

665 **2.6.2. BioCel 1200 system**

666 For full automation of high-throughput single cell cDNA synthesis and library preparation,
667 a laboratory automation platform such as Agilent BioCel 1200 system should be set up and
668 validated prior to processing the sorted plates of single cells. The compatible plastic consumables
669 (i.e. microplates, tips etc.) should be identified and equipment should be optimized for them. The
670 protocols for accurately dispensing reagents should be created and saved in run control software
671 comparable to Agilent VWorks software. A dummy run of the protocol should be performed to
672 test the accuracy of automated protocols for each experimental run in which reagents can be
673 substituted with buffer or diluted glycerol to match the viscosity of reagents to be used.
674 Nomenclature for barcoding the plates of each experimental step needs to be decided and standardized
675 (e.g. Data_Expt_Plate#_BatchNo_ExperimenterInitial).

676

677 **2.6.3. 3D-Cell Explorer Holotomography (Nanolive) microscope**

678 3D-Cell explorer microscopes are set up as per manufacturer's instruction. Among the four
679 filters provided for Blue (DAPI), Green (FITC), and Red (OFP) fluorescent channels, only Red
680 channel has two interchangeable filters (i.e. TRITC or Cy5) and the correct filter needs to be put
681 based on the fluorochrome of the stained cells. The correct hardware configuration needs to be set
682 and the right refractive index (RI) of the mounting media selected on STEVE software before
683 imaging. A designated storage folder with enough data storage capability needs to be created for
684 saving the captured image and post-processed images.

685 **3. PROCEDURE:**

686 **3.1. Oral rinse collection • TIMING: 1-1.5 hrs**

687 **CRITICAL:** Maintain each step of primary saliva neutrophil isolation at room temperature
688 (RT) as the cells tend to lose viability faster at a colder temperature (e.g. 4 °C) that is generally
689 used for PBMCs or any other primary cell type isolation.

- 690 1) Label the sterile 50 mL centrifuge tubes for collection of oral rinse and set the centrifuge to
691 RT.
- 692 2) Before collecting the first oral rinse, each donor must clean their oral cavity and wait for 3 min.
- 693 3) Each donor must rinse their oral cavity five times with ~10 mL of 0.9 % NaCl solution (sterile)
694 for 30-60 sec each time with a gap of 3 min between each rinse to collect 50 mL total volume.
695 (**CRITICAL:** The collection tube should be immediately proceeded for the enrichment process,
696 as neutrophil RNA keeps degrading over time.)

697

698 **3.2. Saliva Neutrophil Isolation/enrichment • TIMING: 1-1.5 hrs**

- 699 4) Pellet down the cells by centrifuging at 160 x g at RT for 5-10 min. (Alternatively, if there is
700 high mucus content in the sample, the cell suspension can be passed through 40 µm sterile cell
701 strainer before centrifugation).? TROUBLESHOOTING
- 702 5) Discard 40 mL of supernatant from 50 mL total vol. by aspirating carefully without disturbing
703 the cell pellet. (CRITICAL: Neutrophils do not form a solid pellet)
- 704 6) Resuspended the cells in the remaining ~10 mL of 0.9 % saline solution and passed through
705 40 µm sterile cell strainer using gravity to remove any food particles or mucus present in the
706 collected oral rinse.? TROUBLESHOOTING
- 707 7) Then sequentially filter the 40 µm filtered oral rinse through pluriStrainer nylon mesh filters
708 20 µm and 10 µm to remove epithelial cells. (CRITICAL: All filtration steps are to be done by
709 gravity flow or centrifuge at 160 x g at RT for 1 min. Avoid using vacuum suction as this
710 suck's in smaller epithelial cells through the 20 µm & 10 µm pluriStrainer. Check under an
711 optical microscope after each filtration process to ensure removal of larger cell and food
712 particles. If necessary, filter again. Check cell viability and density after each filtration step.
713 The cell viability tends to increase with reduction of epithelial cells, which are mostly dead
714 and add to reduced cell viability before 20 and 10 µm filtration).
- 715 8) Count the cells to check the viability and cell density by staining 10 µL aliquot of filtered cell
716 suspension with trypan blue (10 µL) and loading into Countess cell counting chamber slides
717 and using Countess automated cell counter. (Expected total number of cells obtained from a
718 healthy donor in 50 mL oral rinse of resting saliva is $\sim 1.8 \times 10^5$ cells having viability of 70-
719 80 %. This procedure provides > 95 % enriched saliva neutrophils). (CRITICAL: Proceed
720 immediately to the downstream experimental step requiring unfixed cells. Otherwise, for
721 downstream experiments requiring fixed cells, proceed with cell fixation by 4 % PFA).?
722 TROUBLESHOOTING

723 **3.3. FACS sorting • TIMING: 2-3 hrs**

724 (CRITICAL: The FACS equipment for sorting should be prepared and kept ready
725 beforehand to minimize the exposure of saliva neutrophil in its non-native environment.) [NOTE:
726 All provided timing is for single 384-well plate]

727 On verifying the desired purity (i.e. > 95 %) by observing under optical microscope (Fig.
728 2) and determining > 70 % cell viability by trypan blue exclusion method (using Countess cell
729 counter) of enriched neutrophils (Supplementary Fig. 5), proceed immediately with FACS sorting.
730 As we performed unbiased cell sorting, we did not stain the cells with any antibody/marker and
731 used the cell suspension in saline solution for FACS sorting. Alternatively, cell pellets can be
732 resuspended in PBS⁽⁻⁾. (**OPTIONAL**: Cell viability and purity can be checked by flow cytometry
733 using viability dye.)

734 9) In a 1.5-ml Eppendorf tube prepare the desired volume of lysis buffer by adding the reagents
735 as in **section 2.5.2.** and then place it on ice. (**CRITICAL**: For each experiment, the lysis buffer
736 should be made fresh.)

737 10) Prepare 384-well thin-walled PCR plates for sorting by adding 2 μ L of lysis/FACS sort buffer
738 to each well, except NTC-control well(s). (**CRITICAL**: Maintain the lysis buffer added plates
739 at 4 °C using chill blocks)

740 11) Prepare the FACS instrument with 100/130-micron nozzle size for daily FACS setup, testing,
741 and droplet delay optimization, plate targeting. (**CRITICAL**: Failure to optimize the droplet
742 breakoff may sort satellite droplet instead of the droplet of interest by placing a charge on the
743 satellite. Follow the FACS manufacturer's instructions for the droplet stream optimization for
744 timing delay.)

745 12) Using the FACSDiva software, prepare the gating strategy for doublet discrimination gating to
746 prevent the sorting of cell doublets or multiple cell clumps. Load a small amount of sample
747 into the instrument to confirm the set gating and rearrange the gates if needed. Adjust the
748 voltage of the instrument for each channel if needed.

749 13) Confirm the FACS setting parameters for single-cell sorting by targeting the plate using 10-
750 μ m yellow fluorescent polystyrene microspheres or similar fluorescent beads and observing
751 under a fluorescent microscope for accurate targeting. (**CRITICAL**: Achieving an accuracy of
752 a minimum 95 % single microsphere sorting is recommended. To obtain this we suggest
753 practice sorts for single microsphere before actual sort day.)

754 14) Proceed with FACS sorting for a single cell. The overall event rate for 100-micron nozzle is
755 kept at 1000–2,000 events per second on the FACS instrument (minimum 1200 events for 130-
756 micron nozzle setup). Sort 1 cell in each except the Control wells (**CRITICAL/OPTIONAL**:

757 Final confirmation of single-cell sorting can be performed by sorting a single cell in a slide
758 and observing under microscope.)? **TROUBLESHOOTING**
759 15) Seal the sorted plates with MicroAmp Thermo-Seal lid, and immediately proceed with lysis
760 and reverse transcription. **PAUSE POINT**: Otherwise, immediately freeze the plate on dry ice
761 for storage at -80°C .

762

763 **3.4. cDNA synthesis by Smart-seq2 • TIMING: 1 day all steps**

764 We performed cDNA synthesis by using modified Smart-seq2 protocol, previously
765 published by our team. Cell lysis, cDNA synthesis and Nextera XT library preparation can be
766 performed using any of the currently available methods for single cells³⁹⁻⁴¹. All liquid dispensing
767 steps are performed using the BioCel-1200 system incorporated with Bravo and BioRapTR
768 fluidics systems. For dispensing of master mixes by BioCel we used Half-reaction volumes that
769 have been shown below.

770 **3.4.1. Single cell lysis • TIMING: ~15 min**

771 16) Perform cell lysis on each single cell well by adding lysis mix containing 25 μM Oligo dT
772 (0.25 μL) and 25 mM dNTP mix (0.25 μL) to each reaction mix (see **section 2.5.3**). To the
773 control wells, add 10 $\text{pg}/\mu\text{L}$ UHR (1 μL) to UHR-Control well; nuclease-free water (2 μL) to
774 each NTC-control well; nothing to ERCC spike-in control wells. (**CRITICAL**: If FACS sorted
775 plates are taken out of -80°C storage, thaw the plates on ice in chill blocks.)

776 17) Denature by incubating at 72°C for 3 min and immediately putting the plate on ice.

777 18) Centrifuge at $700 \times g$ for 10 sec at RT to spin down the samples to the bottom of the well.
778 Immediately put the plate back on ice. At this step, the oligo-dT primer is hybridized to the
779 poly(A) tail of mRNA strands.

780 **3.4.2. Reverse transcriptase (RT) reaction • TIMING: 3 hrs**

781 19) Add reagents for RT-reaction as on **section 2.5.4** to each well containing 2.50 μL of lysed cell
782 soup (STEP 18) by adding 3 μL of RT-master mix.

783 20) Perform first-strand cDNA synthesis of RT-reaction in Thermocycler by following reaction
784 cycle:

785 42 °C for 90 min : RT and template-switching
786 10 cycles of
787 50 °C for 2 min : RNA-secondary structure unfolding
788 42 °C for 2 min : Completion of RT and template-switching
789 70 °C for 15 min : Final heat inactivation of enzyme
790 4 °C hold : Temperature for safe storage.

791

792 3.4.3. PCR-preamplification • **TIMING: 3 hrs**

793 21) Prepare PCR-preamplification master mix for cDNA synthesis by addition of ISPCR primer
794 as in **section 2.5.5.**

795 22) Run PCR-preamplification reaction in a thermocycler by using the following reaction cycle:

796 98 °C for 3 min : Denaturation
797 21 cycles of
798 98 °C for 20 min : Denaturation
799 67 °C for 15 sec : Annealing
800 72 °C for 6 min : Extension
801 72 °C for 5 min : Final extension
802 4 °C for Infinite : HOLD temperature

803 **PAUSE POINT:** PCR run plate can be stored at -20 °C for short term or at -80 °C for long
804 term storage

805 3.4.4. PCR Purification of cDNA synthesis product • **TIMING: ~45 min**

806 23) Using BRAVO protocol, perform the purification of the cDNA synthesis product by adding
807 AMPure XP beads (1:1 ratio) to the RT-reaction mix from above. Incubate the mix for 5 min
808 at RT and then place it on a magnetic rack for 2 min. Carefully remove the supernatant by
809 pipetting and wash the beads twice with 80 % ethanol (molecular biology grade) for 30 sec.

810 Dry the beads on a magnetic rack for 10 sec. Finally, elute the biotinylated-cDNA with 12.5
811 μL of Low Tris-EDTA (TE) buffer (10 mM Tris + 0.1 mM EDTA) by incubating for 10 min
812 at RT followed by 2 min on a magnetic rack. (**CRITICAL**: Low TE buffer and AMPure XP
813 beads should be at RT.)

814 24) Collect purified cDNA by pulling the supernatant onto a newly labeled thin-wall PCR plate.
815 **PAUSE POINT**: Seal the plate and store it at $-80\text{ }^{\circ}\text{C}$. Otherwise, proceed with the QC step for
816 analysis of cDNA quality as in the next step.

817 **3.5. Quality control analysis of purified cDNA • TIMING: 1-4 hrs**

818 (**CRITICAL**: Quality of purified cDNA library can be analyzed by four methods: Agilent
819 high-sensitivity DNA chips, PicoGreen dsDNA assay, qRT-PCR for expression of housekeeping
820 gene, and qRT-PCR for expression of cell-type specific gene.) Agilent high-sensitivity DNA chips
821 are used for a few randomly picked cDNA samples and analyzed on the 2100-Bioanalyzer system
822 to check the fragment size of DNA. PicoGreen dsDNA analysis is performed on the whole plate
823 to accurately quantify the cDNA concentration in each well. Finally, TaqMan assay is performed
824 in qRT-PCR to check the expression of housekeeping genes such as β -actin (ACTB) to make sure
825 that each well has an eukaryotic cell. (**OPTIONAL**: TaqMan assay for a known cell-type specific
826 marker gene(s) can be used to confirm the target cell sorted into each well. We didn't use any
827 neutrophil specific marker as we performed unbiased sorting.)

828 **3.5.1. QC1: Quality check of cDNA library by Agilent high-sensitivity DNA kit • TIMING:** 829 **1 hr**

830 25) cDNA library size distribution and quality are checked by Agilent high-sensitivity DNA chips
831 on 2100-Bioanalyzer system for randomly picked cDNA samples from the plate by following
832 the manufacturer's protocol for Agilent high-sensitivity DNA kit.

833 26) Undiluted cDNA sample (1.0 μL) is loaded on each of the Agilent high-sensitivity DNA chip
834 and then run on the 2100-Bioanalyzer system to obtain the raw data.

835 27) The sample run data of each chip was analyzed by 2100 Expert software to obtain the
836 electropherogram. Sample free of < 500 bp fragments and showing a peak at ~1.5-2 kb is
837 considered a good library.? [TROUBLESHOOTING](#)

838 **3.5.2. QC2: Picogreen dsDNA quantitation assay • TIMING: 2 hrs**

839 28) Quant-iT PicoGreen dsDNA assay kit (Invitrogen, cat. no. P11496) is used for quantification
840 of dsDNA following the manufacturer's protocol. Working concentration of PicoGreen
841 solution from stock concentration is prepared by 200-fold dilution in TE-buffer (1X) as in
842 **section 2.5.6** and 24.50 μL is dispensed in each well of a 384 Black flat bottom plate using
843 BioRapTR.

844 29) On the other hand, prepare Lambda (λ) gDNA standard of 10 ng/ μL working concentration
845 from the provided stock solution in TE-buffer (1X). Prepare λ -gDNA of varying concentrations
846 by serial dilution (10.00 ng/ μL , 5.00 ng/ μL , 2.50 ng/ μL , 1.25 ng/ μL , 0.625 ng/ μL , 0.3125
847 ng/ μL , 0.15625 ng/ μL , and 0.00 ng/ μL) to obtain the standard curve, which is used to
848 determine the cDNA concentration of each sample well by plotting the RFU (Relative
849 Fluorescence Unit) value of each sample in the standard curve.

850 30) To each of the reaction well transfer 0.50 μL of purified cDNA samples using Bravo to obtain
851 a final concentration of 1:50 dilution. To the standard wells, by using a pipette manually load
852 0.50 μL of Lambda (λ) gDNA standard of varying concentration prepared by serial dilution
853 (see **section 2.5.7**).

854 31) Seal the plate, mix the reaction components and centrifuge briefly to bring the reaction mix to
855 the bottom.

856 32) Incubate for 2-5 min at RT by protecting from light.

857 33) Read plates in FlexStation 3 or any available Fluorescent microplate reader using standard
858 Fluorescein wavelengths of Ex/Em of 480/520 nm

859 34) The RFU-values obtained from each sample well is plotted against the standard curve
860 generated from the lambda-DNA standards (obtained by serial dilutions) to obtain the cDNA
861 concentration of each single-cell well and the control wells (i.e. NTC, ERCC and UHR
862 controls) by using the SoftMax Pro Software for FlexStation 3 (Molecular Devices).

863 35) The SoftMax Pro software generated file is saved in .txt format. The cDNA concentration for
864 each sample well is then incorporated in the final project template file in .xlsx format. Samples
865 with DNA concentration > 0.30 ng/ μ L are considered good suitable for downstream
866 processes.? [TROUBLESHOOTING](#)

867 **3.5.3. QC3: qRT-PCR for housekeeping gene expression • TIMING: 4 hrs**

868 36) For qRT-PCR TaqMan assay, first dilute the cDNA sample to 1:10 in a new 384 well
869 FrameStar plate by adding 9 μ L Low TE buffer or nuclease free water to 1 μ L of cDNA using
870 Bravo.

871 37) Prepare required volume of qRT-PCR master mix as in **section 2.5.8** to be dispensed to each
872 well of the PCR-plate by BioRapTR. (**CRITICAL**: Consider the “dead volume” of BioRapTR
873 while calculating the total required volume of qRT-PCR Master mix.)

874 38) Dispense 7.50 μ L RT-Master mix with BioRPTR using designated Reservoir and Tip to each
875 well.

876 39) Using BRAVO, add 2.5 μ L of diluted cDNA template of each sample for a total reaction
877 volume of 10 μ L/well. For NTC-control wells, added 2.5 μ L of nuclease-free water.

878 40) Load the reaction plate on qRT-PCR machine (QuantStudio 6 Flex Real-Time PCR system)
879 and run the following reaction cycle:

880	95 °C for 2 min	: Denaturation
881	<u>50 cycles of</u>	
882	95 °C for 10 sec	: Denaturation
883	60 °C for 30 sec	: Annealing
884	4 °C for Infinite	: HOLD temperature

885
886 41) On completion of the qRT-PCR reaction cycles, analysis of the generated raw data is done by
887 QuantStudio Real Time PCR Software V1.3. The "CT Settings" under the "Analysis" is
888 changed from its “Default Settings” by changing the ‘Threshold’ to 0.01 and 'Baseline Start
889 and End' to 2 and 10 respectively. Then “Analysis Settings" is applied to obtain the final Ct-
890 values.

891 42) The Ct-values are then exported in the .XLS format to be incorporated in the final project
892 template file. Samples with Ct-values < 35 is considered good quality and suitable for
893 downstream processes.

894 **3.5.4. QC4: qRT-PCR for target cell specific gene expression • TIMING: 4 hrs**

895 (OPTIONAL: Expression of genes known to be expressed in the target cell sorted can be
896 checked by qRT-PCR to confirm the single cell sorted in each well.)

897 43) Taqman assay or preferred qRT-PCR method targeting the marker gene specific to the
898 target cell sorted for the study can be performed to verify the single cell sorted in each well

899 **3.5.5. cDNA-library plate preparing by HitPicking of QC-Pass wells • TIMING: 2 hrs**

900 44) The cDNA concentration obtained from PicoGreen assay (From STEP 35) along with the Ct-
901 values for ACTB expression obtained from qRT-PCR Taqman Assay (From STEP 42) for each
902 sample well in sample plate is pasted on the project template file. Project template file is an
903 excel file prepared to keep track of each sample well of each sort plate.

904 45) The samples having cDNA concentration > 0.3 ng/μL and ACTB Ct-values < 35 are
905 considered double QC-Pass and selected for Hamilton transfer to a new cDNA-library plate.

906 46) For Hamilton transfer, a “Hamilton input file” in .CSV format is generated for each library
907 plate of each donor (viz. H-SN1_Lib#1 etc.). This file is loaded in the “VENUS” software and
908 Hamilton transfer protocol is run after placing the desired tips and plates at their designated
909 location set on the protocol.

910 47) On completion of the Hamilton transfer to combine HitPicked cDNA-library plate from
911 two/three single-cell cDNA plates, the plates are sealed, barcoded and stored at -80 °C for
912 downstream processes.

913

914 **3.6. Illumina Nextera XT Library preparation of HitPicked cDNA-library plates:**

915 Illumina Nextera XT library is prepared for the HitPicked cDNA library plate by using
916 ‘Nextera XT DNA library preparation kit’ and each sample is barcoded by using ‘Nextera XT

917 index kit Set A and Set D' following the manufacturers protocol. We used 1/8th reaction protocol
918 for automated/robotic dispensing system, where 1/8th the volume of each reagent is used as
919 mentioned in manufacturers protocol for 96-well reaction plate. The required target DNA quality
920 for Nextera Library preparation is 1 ng of input DNA with 260/280 ratio of 2.0 - 2.2

921 **3.6.1. Normalization of cDNA library plate • TIMING: ~1 hr**

922 48) Before starting the Nextera XT 1/8th reaction protocol, 'Normalized cDNA Library' plate is
923 prepared to obtain 0.2 ng/μL cDNA concentration in all wells.

924 49) cDNA library plate stored at -80 °C is taken out and thawed on a chill block in ice.

925 50) From each sample well 1 μL of cDNA is transferred to a new Framestar 384-plate by BRAVO
926 and the desired volume of low TE-buffer is dispensed by BioRapTR to obtain 0.2 ng/μL cDNA
927 concentration in each well. Care is taken so that total volume per well should not exceed 100
928 μL. If the calculated volume for any well exceeds 100 μL, the final dispensing volume is
929 calculated for 100 μL.

930 **3.6.2. Nextera XT Tagmentation reaction • TIMING: 10 min**

931 51) 0.625 μL of diluted cDNA (0.2 ng/μL) from cDNA normalization plate is added to 1.250 μL
932 of Tagment DNA Buffer (TD, 2X) and 0.625 μL of Amplification Tagment Mix (ATM) in a
933 Framestar 384-well microplate to obtain 2.5 μL total tagmentation reaction mix volume.

934 52) The plate is sealed, mixed by brief centrifuge and loaded on the thermocycler to run the
935 tagmentation reaction by incubation at 55 °C for 10 min.

936 53) On completion of the reaction, immediately add 0.625 μL of NT buffer to neutralize the
937 Tagmentation reaction to obtain 3.125 μL Total Neutralized Tagmentation volume/well.

938

939 **3.6.3. Nextera XT PCR reaction with Set A and Set D barcoding kits • TIMING: 1 hr**

940 54) To the 3.125 μ L of Tagmentation volume, 1.875 μ L NPM PCR master mix and 1.250 μ L of
941 Index Primer mix (0.625 μ L Index Primer i5 + 0.625 μ L Index Primer i7) is added to obtain
942 total volume of 6.25 μ L Nextera PCR reaction/well.

943 55) Seal and centrifuge FrameStar Plate at 4 $^{\circ}$ C, 500 x g (2,000 RPM) for 30 sec to mix, keep on
944 ice till running the thermocycler reaction.

945	72 $^{\circ}$ C for 3 min	: Extension
946	95 $^{\circ}$ C for 30 sec	: Denaturation
947	<u>16 cycles of</u>	
948	95 $^{\circ}$ C for 10 sec	: Denaturation
949	55 $^{\circ}$ C for 30 sec	: Annealing
950	72 $^{\circ}$ C for 60 sec	: Extension
951	72 $^{\circ}$ C for 5 min	: Final extension
952	4 $^{\circ}$ C for Infinite	: HOLD

953

954 **PAUSE POINT:** Seal the plate and store at -80 $^{\circ}$ C until ready for library purification and
955 cleanup.

956 **3.6.4. Nextera XT Library purification and cleanup • TIMING: ~45 min**

957 56) Purify each sample individually as is Step 23-24 but use 0.9:1 ratio of AMPure XP beads to
958 Nextera library (i.e. 5.625 μ L beads + 6.25 μ L of Nextera Library). Elute with 6.25 μ L of Low
959 TE buffer into a new FrameStar plate "Purified Nextera XT".

960 **PAUSE POINT:** Seal the plate and store at -80 $^{\circ}$ C until ready for PicoGreen QC or
961 normalization of the purified library.

962

963 **3.6.5. QC5: Nextera XT Library QC by PicoGreen assay • TIMING: 2 hrs**

964 57) Use 1 μ L of the purified Nextera XT reactions for Picogreen dsDNA assay as in Step 28-35.

965 **3.6.6. Normalization of Nextera XT Library plate • TIMING: ~1 hr**

966 58) Prepare 1.0 ng/μL normalization plate of purified Nextera library based on the PicoGreen
967 quantification above.

968 59) Using BioRaPTR dispense the desired amount of Low-TE buffer to a new Framestar plate and
969 then using BRAVO to add 1 μL of purified Nextera XT library sample to obtain 1.0 ng/μL
970 purified Nextera XT sample in each well.

971 **3.6.7. 16 sample pooling of NexteraXT samples for MiSeq run • TIMING: 30 min**

972 (OPTIONAL: As the sequencing of NovaSeq run for 384-plex pool library is expensive,
973 we sequenced randomly picked 16-plex pool library from each NexteraXT library plate in MiSeq-
974 Nano to confirm the sequence quality, determine the coverage needed and required depth of the
975 transcriptome. We used Illumina MiSeq Reagent kit v2 (300 cycle) for the MiSeq-Nano low output
976 run.)

977 60) Pool 16 samples (from Step- 59) by pipetting 3 μL of all normalized NexteraXT samples (1.0
978 ng/μL) for a 3 ng pool into a 1.5 mL Eppendorf LoBind tube.

979 61) Reverse pipette to determine the total volume and then add 90 % of that volume AMPure beads
980 to purify the MiSeq-pooled library as in Nextera XT library cleanup on Step 56.

981 62) Elute with Low-TE buffer using 10-fold lower volume than the original volume of pooled
982 library determined by reverse pipetting.

983 63) For QC check on Agilent high-sensitivity DNA chips, prepare 1:1 dilution (take out 1 μL),
984 1:10 dilution (1 μL library + 9 μL Low TE), and 1:20 dilution (2 μL of 1:10 dilution + 2 μL of
985 Low-TE)

986 64) Run each of the 3 samples in triplicate on Agilent high-sensitivity DNA chip. Calculate the
987 average fragment size and the average pool concentration in pmol/L (pM) and nM of all
988 replicates from DNA chip run report.

989 **PAUSE POINT:** Store the MiSeq-pooled library at -20 °C until ready for sequencing run.

990 **3.6.8. 384 sample pooling of NexteraXT samples for NovaSeq run • TIMING: 30 min**

991 65) For each 384-well library plate, combine all 384 normalized Nextera XT libraries from each
992 well to a single well by using BRAVO. Pipette the 384-plex pool to an Eppendorf tube and
993 label the tube with sample and library name (e.g. H-SN1_Lib#1, H-SN2_Lib#1, and H-
994 SN3_Lib#1 in this study).

995 **3.6.9. Pooled Nextera XT Library Cleanup • TIMING: ~45 min**

996 66) Clean the 384-plex pool Nextera XT library by AMPure XP bead purification as in Step 23-24
997 using manual protocol for fewer pooled library sample tubes. BRAVO protocol for AMPure
998 XP bead purification can be used for more samples.

999 **3.6.10. QC6: QC check of 384-plex Nextera XT Library (Agilent DNA Chip) • TIMING: 1**
1000 **hr**

1001 67) Use 1 μ L of the 384-plex pool NexteraXT Library samples to check the quality on 2100-
1002 Bioanalyzer using Agilent high-sensitivity DNA chips. The DNA concentration of the sample
1003 is assumed to be within the range recommended by Agilent for the high-sensitivity DNA chips.

1004 68) The Bioanalyzer report for each 384-plex pool library is saved and needed to submit samples
1005 to the sequencing core.

1006 **PAUSE POINT:** The Pooled library can be stored at -20 °C for short term and at -80 °C
1007 for long term till ready for sequencing run.

1008 **3.6.11. QC7: KAPA Library Quantification Kits - Illumina • TIMING: 2-3 hrs**

1009 69) Calculate and prepare volumes enough for 3 replicates of NTC, each sample dilutions, and
1010 each standard.

1011 70) Six pre-diluted DNA Standards of concentration 20, 2, 0.2, 0.02, 0.002, 0.0002 pM
1012 respectively are provided in the kit.

- 1013 71) Before starting, prepare four different dilution sets of the pooled NexteraXT library (from Step
 1014 66) by adding required volume of dilution solution (10 mM Tris-HCL + 0.05 % Tween 20) as
 1015 in **section 2.5.9**.
- 1016 72) Prepare qPCR master mix as in **section 2.5.10** by combining the 1 mL of Illumina Primer
 1017 Premix (10X) and the 5 mL bottle of KAPA SYBR Fast qPCR master mix (2X) provided in
 1018 the kit after thawing properly. Vortex briefly to mix well and store at -20 °C till ready to use.
- 1019 73) For half reaction volume, add 6 µL of qPCR master mix and 4 µL of sample or standards to
 1020 appropriate well. Add 4 µL nuclease free water to NTC wells
- 1021 74) Seal plate, gently vortex to mix and spin down sample to bottom of the well. Protect plate from
 1022 light until ready to run.
- 1023 75) Run the sample plate on QuantStudio 6 Flex or any other qPCR machine by selecting the
 1024 “Standard Curve” experimental method and “SYBR Green” detector. Run the instrument in
 1025 “Fast” mode using the following thermocycler protocol:
- | | | |
|------|---------------------|----------------|
| 1026 | 95 °C for 5 min | : Denaturation |
| 1027 | <u>35 cycles of</u> | |
| 1028 | 95 °C for 20 sec | : Denaturation |
| 1029 | 60°C for 45 sec | : Annealing |
| 1030 | 4 °C for Infinite | : HOLD |
- 1031
- 1032 76) On completion of qPCR run, analyze the data and evaluate the Slope and R2. If the auto set
 1033 Ct-threshold if the acceptable range of slope (-3.58 to -3.10) and/or R2 (~0.99) is not obtained,
 1034 manually adjust Ct by setting threshold of 0.2 and set start cycle to 2 and end cycle to 3.
- 1035 77) The average Ct value of each DNA Standard is plotted against its known concentration (pM)
 1036 to generate a standard curve which is used to determine the concentration (pM) of diluted
 1037 libraries. Finally, the working concentration of each library is calculated from the
 1038 concentration of diluted libraries.
- 1039 78) To further check the fragment size, the qPCR amplified product from three replicate wells were
 1040 combined and ran on 1.2 % E-Gel (Invitrogen, cat. no. A03076) for 30 min using 1 kb plus
 1041 ladder (Invitrogen, cat. no. 10787018) and 6X loading dye (Promega, cat. no. G190A).
- 1042 79) On verification of the quality and fragment size of the pooled NexteraXT library samples,
 1043 proceed to the cDNA sequencing step.

1044 **3.7. cDNA Sequencing**

1045 **3.7.1 cDNA Sequencing: kit selection, run parameters, and yield • TIMING: ~24 hrs**

1046 80) The purified pooled-NexteraXT library is subjected to paired-end sequencing on a suitable
1047 Illumina NGS platform (MiSeq, HiSeq 2500, NextSeq 500, and NovaSeq 6000) with the aim
1048 to generate 1-2 million reads per sample having a read length of 100-150 bases. The sequencing
1049 data generated by the HGS platform is in fastq format.

1050

1051 The following sections have been briefly explained in this paper. For details on “RNA-seq
1052 analysis” of the fastq data files, please refer to Step 26 of previous publication from our group²⁴.

1053 **3.7.2. RNA-seq analysis: sequence quality assessment and preprocessing • TIMING:**
1054 **Variable**

1055 81) *Sequence quality assessment*: Sequence quality is assessed by evaluating the fastq sequence
1056 files (from Step 80) from each cell (i.e. single cell saliva neutrophil) using the fastQC tool for
1057 sequence yield, base quality, GC profile, k-mer distribution, contamination and other desired
1058 parameters.

1059 82) *Sequence duplication*: Sequence duplication is determined in the input data. Tools such as
1060 fastx_collapser are used to calculate the absolute number of identical reads (i.e. duplicates) in
1061 the input sample fastq sequences (from Step 80). Use correct base quality score offset (-Q).
1062 Process multiple files by repeating each sequence file at a time, as the program accepts only
1063 one sequence file as input.

1064 83) *Sequence trimming*: Sequence trimming of input paired-end fastq reads (from Step 80) is
1065 performed by Trimmomatic program to remove adapter/primer sequences and low-quality end
1066 bases.

1067

1068 **3.7.3. RNA-seq analysis: sequence mapping and gene expression analysis • TIMING:**
1069 **Variable**

1070 84) *Prepare the reference genome*: Prepare the reference genome index for alignment using the
1071 build function in HISAT2 program and the reference genome fasta file. Here we used GRCh38
1072 downloaded from Ensembl /Gencode.

1073 85) *Calculating expression values*: Calculate the gene expression values (transcripts per million or
1074 TPM) by mapping the paired-end reads that passed trimming (from Step 83) to the reference
1075 index using HISAT2 and then evaluating the alignments using StringTie to estimate levels of
1076 expression per gene models in the annotation file. Here we used gencode.v25.annotation.gtf.

1077 86) *Calculate and plot overall mapping statistics*: Calculate the number of reads that are mapped
1078 to the genome, to the ERCC spike-in transcripts as well as that remained unmapped using SAM
1079 tools.

1080 **4. TIMING**

1081
1082 Steps 1–3, Oral rinse collection: 1–1.5 hr
1083 Steps 4–8, Saliva Neutrophil Isolation/enrichment: 1–1.5 hr
1084 Steps 9–15, FACS sorting: 2–3 hr
1085 Steps 16–18, Single cell lysis: ~15 min
1086 Steps 19–20, Reverse transcriptase (RT) reaction: 3 hr
1087 Steps 21–22, PCR-preamplification: 3 hr
1088 Steps 23–24, PCR Purification of cDNA synthesis product: ~45 min
1089 Steps 25–43, QC analysis of purified cDNA: 1-4 hrs depending on QC method(s) chosen
1090 Steps 25–27, QC1 (OPTIONAL) - Agilent high-sensitivity DNA kit: 1 hr
1091 Steps 28–35, QC2- Picogreen dsDNA quantitation assay: 1 hr
1092 Steps 36–42, QC3- qRT-PCR for housekeeping gene expression: 4 hr
1093 Step 43, QC4 (OPTIONAL) - qRT-PCR for target cell specific gene expression: 4 hr
1094 Steps 44–47, cDNA-library plate preparing by HitPicking of QC-Pass well: 2 hr
1095 Steps 48–56, Illumina Nextera XT Library preparation of HitPicked cDNA-library plates: ~3
1096 hr

1097 Steps 48–50, Normalization of cDNA library plate: 1 hr
 1098 Steps 51-53, Nextera XT Tagmentation reaction: ~10 min
 1099 Steps 54-55, Nextera XT PCR reaction: 1 hr
 1100 Steps 56, Nextera XT Library purification and cleanup: ~45 min
 1101 Steps 57, QC5: Nextera XT Library QC by PicoGreen assay: 2 hr
 1102 Steps 58-59, Normalization of Nextera XT Library plate: 1 hr
 1103 Steps 60-65, (OPTIONAL) 16 sample pooling of NexteraXT samples for MiSeq run: 30 min
 1104 Step 66, Pooled Nextera XT Library Cleanup: ~45 min
 1105 Steps 67-68, QC6: 384-plex Nextera XT Library QC by Agilent DNA Chip: 1 hr
 1106 Steps 69-79, QC7: KAPA Library Quantification Kits - Illumina: 2-3 hr
 1107 Steps 80, cDNA Sequencing: kit selection, run parameters, and yield: Variable
 1108 Steps 81-83, RNA-seq analysis: sequence quality assessment and preprocessing: variable
 1109 Steps 84-87, RNA-seq analysis: sequence mapping and gene expression analysis: variable
 1110

1111 **5. TROUBLESHOOTING**

1112

Step	Problem	Possible reason	Solution
4	Missing Cell Pellet	3 min gap not given after rinsing oral cavity	Restart and provide 3 min before collecting the first rinse
		Low cell density, thus not forming proper pellet	Centrifugation speed can be increased up to 300 x g but check its effect on your cells i.e. viability and RIN.
6	Cell pellet visible on step 4 but not on step 6	Cells lost while discarding supernatant	Neutrophils don't form solid pellets. Avoid vacuum aspiration to discard supernatant. Gently and slowly pipette out the supernatant from the center of the tube while holding the tube vertically.
		High Mucus content in sample	Mucus present may attach cells. While pipetting out mucus be very slow.
			Pass cell suspension through a 40 µm nylon mesh filter before the first centrifugation on step 4.

8	Lower cell viability	Lower temperature set on centrifuge	Check the centrifuge temperature set.
		Cells came in contact with cold temperature	Make sure sample tubes are not placed on an ice bucket during and after sample collection.
	low cell count	Saline volume per rinse and no. of time rinse collected in Oral rinse collected protocol not followed properly	Saline sol. Per oral rinse should be 10 mL and collection for five times to make total vol of 50 ml
		Volume of saline used per rinse is more than 10 ml	Measure 10 ml of saline solution separately each time for five times
	Cloudy filtered sample or cell debris visible under microscope	Vacuum aspiration or high centrifuge speed used for filtration	Depending on the saliva viscosity uses gravity for filtration. Avoid vacuum aspiration for quick filtration. Centrifuge at 160 x g for not more than 1 min.
14	Poor recovery of single cell from FACS	Single cell targeting of FACS is compromised	Optimize FACS conditions; sort fluorescent beads onto a glass slide and observing under a fluorescent microscope
			Determination of sorting gates by using antibody stained samples (see Fig. 1C).
			Sort a single cell onto a glass slide and visualize under an optical microscope.
27 & 35	No product after cDNA preamplification	Cell were dead or damaged during FACS	Keep the cells suspended in 0.9 % saline solution or PBS ⁽⁻⁾ maintained always at room temperature.
			Check cell viability using viability dye on the flow cytometer (e.g.: Aqua LIVE/DEAD from Invitrogen).
			Check the time of cells from the isolation step (see Fig. 6).

			Negatively stained with viability dye can be used and viable cells can be gated for sorting following DD-gating strategy.
			Check the nozzle size and nozzle pressure used during FACS sorting.
	No cell in the well (after FACS)	cell targeting was not accurate	Optimize single-cell targeting into wells of the 384-well microtiter plate before FACS.
	Low quality RNA obtained	RNA degraded	Minimize the time between cell harvest and sorting below 4 hr to maintain minimum acceptable RIN value of 3. Saliva neutrophil's RIN value goes down with every hour (see Fig. 6).
27	Amount of cDNA is much larger than usual (i.e. samples contain > 500bp fragments)	Contamination occurred during reagent mastermix preparation	Keep the tools and workstation RNases free. Perform all experiments in a clean and dedicated area free from RNases and DNases.
35	Low/High RFU-values than expected	No/ more than single cell in the well	Cell targeting was not accurate. Optimize single cell targeting into wells before FACS.

1113

1114 **6. ANTICIPATED RESULTS**

1115 This protocol enables isolation of immune cells and enrichment of human primary salivary
1116 neutrophils, for cell isolation, flow cytometry analysis, sorting and scRNA-seq workflow (Fig. 1).
1117 Our protocol shows that the repertoire of a myeloid derived cell can be evaluated at a single cell
1118 level after saliva collection. The use size exclusion allowed > 98 % pure enriched neutrophils with
1119 viability compatible with the protocols. In order to develop the protocol that is consistent, oral
1120 samples were collected from healthy subjects throughout the project for all the experiments
1121 including microscopy (Fig. 2), flow cytometry (Fig. 3 and Fig. 4), and scRNA-seq (Fig. 5 and 6).
1122 We found that neutrophils were sensitive to cold temperature and physical stress employed in cell
1123 isolation procedures for NGS workflow. Chen et. al has shown that the integrity of total RNA is a
1124 critical parameter for RNA-seq analysis and degraded RNA heavily influences the gene expression
1125 profiles³⁵. Here we have shown that neutrophil's RIN (RNA Integrity Number) values⁴² decreases

1126 by one with every passing hour and after 4 hours of harvest the RNA is degraded enough (RIN <
1127 3) which is not suitable for transcriptomic profiling, though they do not lose their viability
1128 completely in the time period (Supplementary Fig. 4). After RNA sequencing, our liberal filtration
1129 criteria of raw data allowed to obtain transcriptomic signature of neutrophils similar to other cell
1130 types and identification of eight novel sub-populations of neutrophils from healthy human saliva
1131 as compared to four sub-populations identified by stringent filter criteria typically used for analysis
1132 of RNA-seq data (Fig. 5 and Fig. 6). In addition, both manual gating analysis and the FLOCK-
1133 based automated gating analysis of the flow cytometry data confirmed diverse neutrophil
1134 subpopulations and a more stringent analysis revealed five sub-populations based on markers and
1135 density (Fig. 4).

1136
1137 Through microscopy, saliva presented four major different cell types (i.e. epithelial cells,
1138 neutrophils, monocytes, and lymphocytes) were initially identified based on histochemical
1139 morphology by cytospin and GIEMSA staining. For verification of the purity of neutrophils the
1140 enriched samples from healthy donors were also stained (Fig. 2B) and quantified from a minimum
1141 of 10 slides per donor which shows ~98 % neutrophil purity. This was confirmed by
1142 immunofluorescence of CD66b, CD14, CD19 markers (Fig. 2A and 2B). Our robust
1143 holotomography microscopy preserves live cells while imaging. At least 45 images per sample
1144 were taken, and cell count was represented by percentage of the total count. This morphological
1145 data was also compared through flow cytometry. While epithelial cells are quite abundant, their
1146 morphology was distinct from immune cells. In our serial filtration strategies, epithelial cells did
1147 not show significant reduction after 40 or 20 μm filtration ($p = \text{ns}$), but were significantly reduced
1148 after 10 μm filtration (40 versus 20 μm , $p = 0.0037$; 40 versus 10 μm , $p = 0.0197$). Similarly,
1149 neutrophil enrichment was feasible after 40 μm and 20 μm filtered samples ($p = \text{ns}$), reaching
1150 significant increase abundance after 10 μm filtration ($p = 0.0025$). This was also true for 10 μm
1151 filtered samples when compared to 40 μm filtered samples ($p = 0.0233$). No significant increase
1152 or decrease has been observed in monocyte, lymphocyte, and unidentified cell types (Fig. 2C).
1153 Morphological analysis was also confirmed by immunofluorescence analysis.

1154
1155 We further compared gating strategies through viability, expression marker and cell size.
1156 When utilizing gating for Aqua LIVE/DEAD fixable dye (Fig. 3A and 3C) neutrophils were

1157 positive in high abundance, but minimum monocyte marker expression is detachable. We
1158 hypothesized that by gating the cells on their size we would further exclude the other immune cell.
1159 High-SSC gating demonstrated high purity of neutrophils and exclusion of other immune cells
1160 (Fig. 3B and 3D). Comparatively, live-cell gating and high-SSC gating were quantified and plotted
1161 for the type of cells (Fig. 3E and 3F). Thus, the employment of high-SSC high-FSC gating to sort
1162 unlabeled cells allows for selection of viable cells, prevents activation and cell death of oral
1163 neutrophils, minimizing the possibility of sorting monocytes or lymphocytes.

1164

1165 Flow cytometry analysis (Fig. 3) revealed that oral neutrophils were positive for specific
1166 neutrophil markers (CD15, CD66b, CD11b) and negative for monocyte and lymphocytes markers
1167 (CD14, CD19, CD3). In fact, when staining for most common blood immune-cell markers i.e.
1168 CD11b, CD66b, CD15, CD14, CD19, and CD18 oral neutrophils were positive with different
1169 levels of expression. Oral neutrophils were identified on CD14⁻CD19⁻CD3⁻
1170 CD15⁺CD66b⁺CD11b⁺Aqua⁻ (Fig. 4) whereas monocytes and lymphocytes were identified for
1171 CD14⁺CD11b⁺ and CD19⁺CD3⁺ respectively in live-cell and high-SSC gated populations (Fig.
1172 3C and 3D). CD15⁺ neutrophil density increases from 38.30 % of live-cell gated population for 40
1173 μ m filtered un-enriched samples to 60.72 % in 10 μ m filtered enriched neutrophil. In addition, this
1174 increase in neutrophil was also seen to be associated with more CD14⁺ monocytes in 10 μ m filtered
1175 samples (Fig. 3C). When using the high-SSC gating (instead of live-cell gating), an increase of
1176 CD15⁺ neutrophil density from 30.10 % of high-SSC gated population in 40 μ m filtered sample to
1177 47.35 % in 10 μ m filtered enriched sample. Though this increase in density of neutrophil is not
1178 very high, the CD14⁺ monocytes were completely eliminated on 10 μ m filtered enriched samples
1179 (Fig. 3D). Similar to monocytes, CD3⁺ T cells and CD19⁺ B cells were totally eliminated even in
1180 40 μ m filtered samples when selected for high-SSC gate instead of live-cell gate. Similarly, when
1181 the frequency of each immune-cell types in the live-cell gated population (Fig. 3E) and high-SSC
1182 gated population (Fig. 3F) compared to the total ungated cell population, we see an increase in the
1183 neutrophil population and decrease in monocytes, B cells and T cells.

1184

1185 To characterize heterogeneity of neutrophils in a data-driven way, we employed an
1186 unsupervised data clustering method²⁴ (FLOCK, <http://importgalaxy.org>) to understand cell
1187 phenotype differences and cell surface markers expression levels (Fig. 4). We first applied the

1188 FLOCK method to identify cell populations from the 40 μm filtered sample, before applying the
1189 identified cluster centroids across all 6 samples for cross-sample identification and comparison of
1190 the 23 cell populations. The advantage of unsupervised clustering analysis is being data-driven,
1191 without requiring or being limited by predefined cellular phenotype. Therefore, manual annotation
1192 of each identified data cluster for identifying the cellular phenotype and interpreting the phenotype
1193 difference between the identified cell populations is usually required. Percentages of identified cell
1194 populations, mean fluorescence intensities (MFI) for each marker, expression profiles across the
1195 markers (levels 1 to 4, from negative, low, positive to high) as well as 2D dot plots of samples
1196 with cell populations highlighted in different colors are automatically generated or calculated by
1197 the FLOCK procedure. Heatmap of MFI (Fig. 4A) and bar graphs of population percentages (Fig.
1198 4B) are commonly used to visualize the characteristics of the FLOCK-identified cell populations.
1199 Based on the MFI heatmap in figure 4A, one can easily identify the 5 salivary neutrophil
1200 subpopulations by selecting those with the phenotype of high-FSC/high-SSC/ $\text{CD11b}^+\text{CD14}^-$
1201 $\text{CD15}^+\text{CD66}^+\text{CD3}^-\text{CD19}^-$. From the FACS Aria II data, the FLOCK-identified neutrophil
1202 phenotype included:

1203

- 1204 ● Pop10: $\text{CD14}^-\text{CD19}^-\text{CD3}^-\text{CD15}^+\text{CD66b}^+\text{CD11b}^+\text{Aqua}^-\text{CD18}^-$
- 1205 ● Pop12: $\text{CD14}^{\text{int}}\text{CD19}^-\text{CD3}^-\text{CD15}^+\text{CD66b}^+\text{CD11b}^+\text{Aqua}^-\text{CD18}^-$
- 1206 ● Pop14 and Pop15: $\text{CD14}^{\text{int}}\text{CD19}^-\text{CD3}^-\text{CD15}^{\text{hi}}\text{CD66b}^{\text{hi}}\text{CD11b}^+\text{Aqua}^-\text{CD18}^{\text{int}}$
- 1207 ● Pop17: $\text{CD14}^{\text{int}}\text{CD19}^{\text{int}}\text{CD3}^-\text{CD15}^{\text{hi}}\text{CD66b}^{\text{hi}}\text{CD11b}^+\text{Aqua}^-\text{CD18}^{\text{int}}$

1208

1209 When examining the percentage values of the 5 neutrophil subpopulations, we found that
1210 the abundant/known neutrophil subpopulations (Pop10 and Pop12, with different expression levels
1211 of CD14) increased frequency as the filtration size decreased, which confirmed the finding in Fig.
1212 3. However, FLOCK also identified three other rare salivary neutrophil subpopulations (Pop14,
1213 Pop15, and Pop17) that were not in the region of the “classical” neutrophils. These rare neutrophil
1214 subpopulations have larger size and complexity (based on FSC.A and SSC.A, Fig. 4C) as well as
1215 slightly higher expression on CD15/C66b/CD18 than the classical neutrophils (Fig. 4C). The
1216 frequencies of these rare neutrophil cell populations did not increase as the filtration size
1217 decreased. Limited by the small number of markers measured in flow cytometry, this finding

1218 further emphasizes the necessity of performing a single cell RNA-seq assay to elucidate
1219 transcriptional profiles of these interesting neutrophil subpopulations.

1220

1221 Because we determined the optimal gating for oral neutrophils that maintain their viability
1222 and purity, we pursued sorting single neutrophils for RNA-seq. After raw sequencing files were
1223 demultiplexing using Illumina barcodes, and processed²⁹, we further evaluated expression values
1224 for non-control cells were imported into Scanpy for PCA, UMAP, and cluster analysis³¹. The
1225 liberal filtering 8 neutrophil subpopulations and top 2500 variable genes were selected (Fig. 5). In
1226 fact, half of the subpopulations presented differential expression of surface markers when
1227 compared to other clusters. In contrast the strict filtering showed 4 subpopulations and top 2000
1228 variable genes were selected (Fig. 6). Unsupervised clustering determined the differential
1229 expression of each cell population provided individual single cells grouped into cell type clusters,
1230 and a set of sensitive and specific marker genes. Logistic regression showed the comparison of
1231 gene levels in each cluster (Fig.5B and 6B). Oral neutrophils highly expressed the following gene
1232 markers CD11c, CD14, CD16a, CD16b, CD32, CD55, CD62L, CD141, Lysozyme, BEST1,
1233 FTH1, with moderately levels of CD10, CD11b, CD18, CD31, CD50, CD63, CD85D and low
1234 levels of CD11a, CD13, CD19, CD43, CD170, CD172A, CHEMR23 (Fig. 5E and 6E). These gene
1235 signatures are important to understand future neutrophil functions.

1236

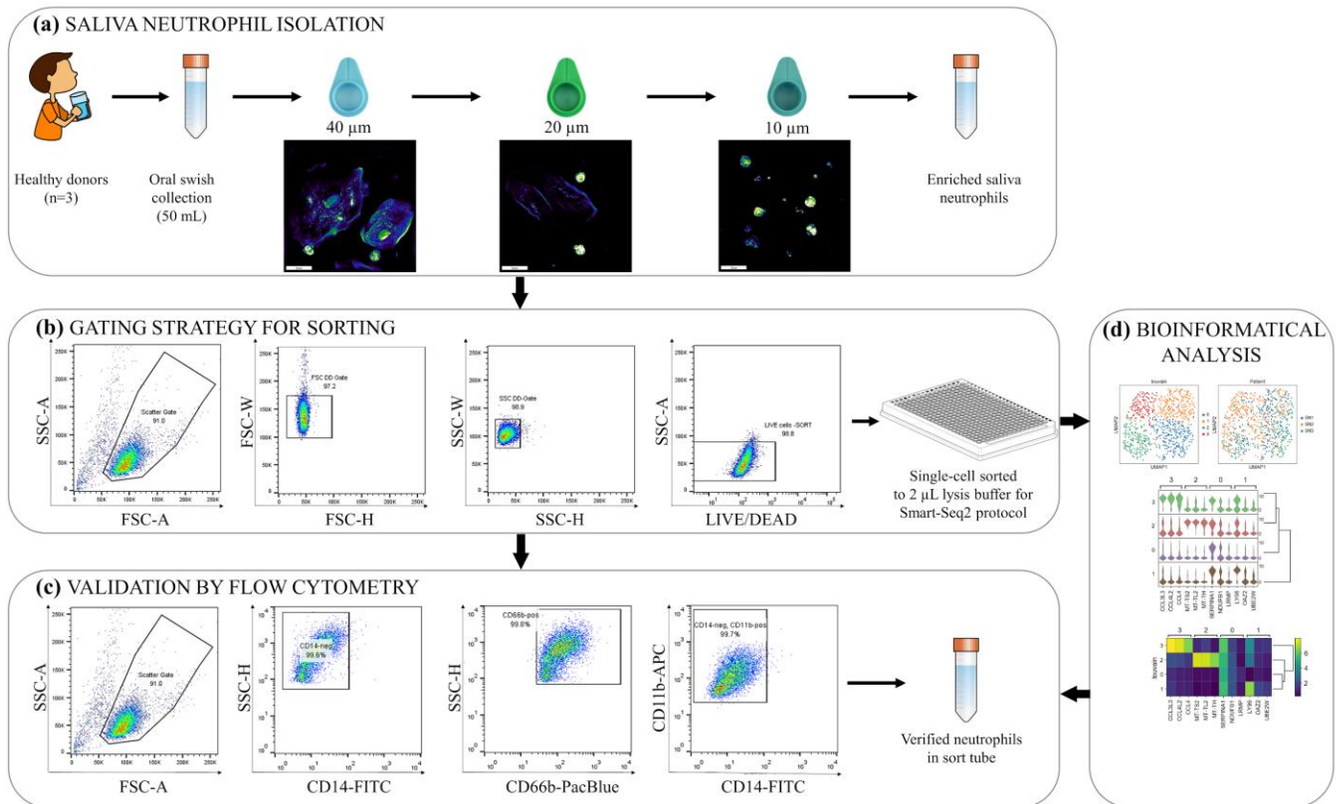
1237 Previous single-cell sequencing of immune cells was unsuccessful in generating desirable
1238 transcriptomic profile/signature of neutrophils among PBMCs⁴³. We have evaluated a dataset that
1239 is publicly available (Supplementary Fig. 6). Evaluation following our oral immune-cell pipeline
1240 applied to the publicly available dataset accession number GSE64655. Similar to oral neutrophils,
1241 blood derived neutrophils presented strong RNA expression levels of CD55, CD16b, CD15, and
1242 not CD66b. Blood neutrophils however presented much higher expression levels on CD10, CD11a
1243 and CD11b. We believe that the cell collection procedures of PBMCs such as the Ficoll-Hypaque
1244 method are not suitable for neutrophils as the NDNs found in the granulocyte layers at the interface
1245 of red blood cells and the gradient layer are excluded during the sample collection. In health and
1246 disease this is extremely important. SLE is characterized by neutrophil subsets known as low-
1247 density granulocytes (LDGs). When compared to other immune cell subtypes, LDGs showed the
1248 highest expression of interferon gamma genes, CD10 with subpopulations that were specifically

1249 positive correlation with disease severity and coronary patterns ⁴⁴. In addition, neutrophils patterns
1250 were also described in the emerging COVID-19 infectious disease. Bronchoalveolar single cell
1251 analysis demonstrated that myeloid cells such as macrophages were highly inflamed expressing
1252 high levels of CD68, while neutrophils from infected lungs were highly expressed (FCGR3B or
1253 CD16b) ⁴⁵. Interestingly, similar to lung neutrophils, oral neutrophils highly expressed Cd16b as
1254 part of our cell clustering system. Whereas similar to SLE cells, oral cells expressed moderate
1255 levels of CD10, revealing that oral neutrophils present shared markers with systemic cells. Future
1256 studies should implement similar protocols for systemic neutrophils compared to oral cells to
1257 uncover methods and heterogeneity.

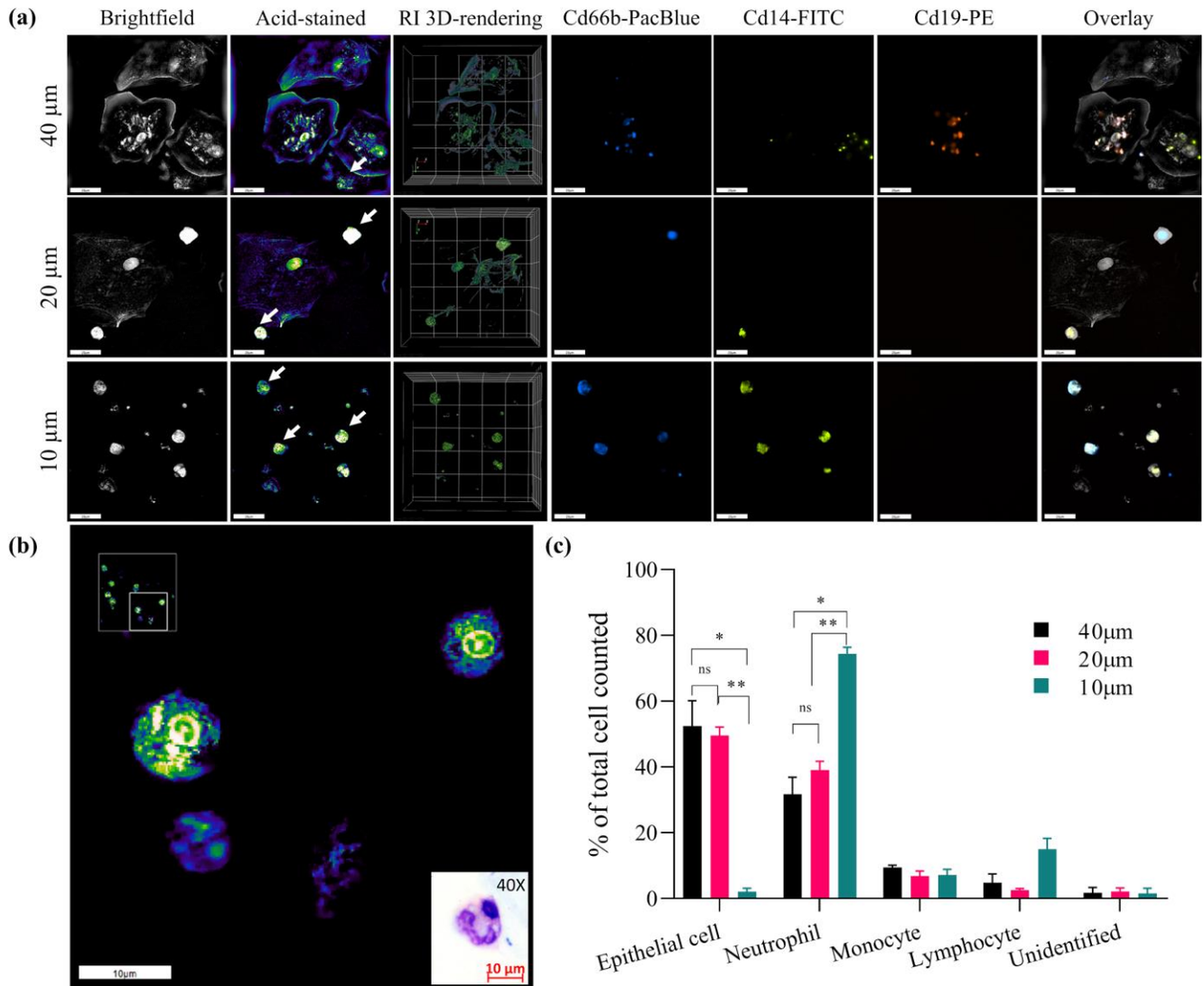
1258

1259 Altogether, we developed a protocol to survey neutrophil cell heterogeneity by improving
1260 isolation methods, flow cytometry evaluation, single-cell RNA extraction, sequencing and
1261 bioinformatic pipeline. Our findings suggest novel transcriptomic signatures with identification of
1262 novel sub-populations. By combining flow cytometry with machine learning, we validated our
1263 model and gating strategies which led to exclusion of other immune cells, while enriching for
1264 neutrophils. Ultimately, this methodology advances methods to understand oral immune cell
1265 landscape and heterogeneity.

1266



1269
 1270 **Figure 1. Saliva single-cell pipeline to discover novel neutrophil populations.** Schematics of
 1271 single-cell isolation and sorting strategies for RNA-seq analysis prior to flow cytometry validation.
 1272 (a) Saliva neutrophil collection in 0.9 % saline solution and filtration 40, 20, and 10-micron cell
 1273 strainers allows for high purity *ex vivo* neutrophils. (b) High-SSC flow cytometry strategy for
 1274 sorting, followed by FSC- and SSC-doublet discrimination (DD) gating strategy and live cell
 1275 gating provided optimal sorting strategies for single cell RNA-seq. Smart-seq2 sequence library
 1276 preparation provides full reads of neutrophil transcriptome. (c) Protein validation of neutrophils
 1277 markers is achieved in tandem (e.g. CD66b⁺CD11b⁺CD14⁻). (d) Bioinformatic evaluation of RNA
 1278 raw sequences is developed to identify novel neutrophil populations and heterogeneity.

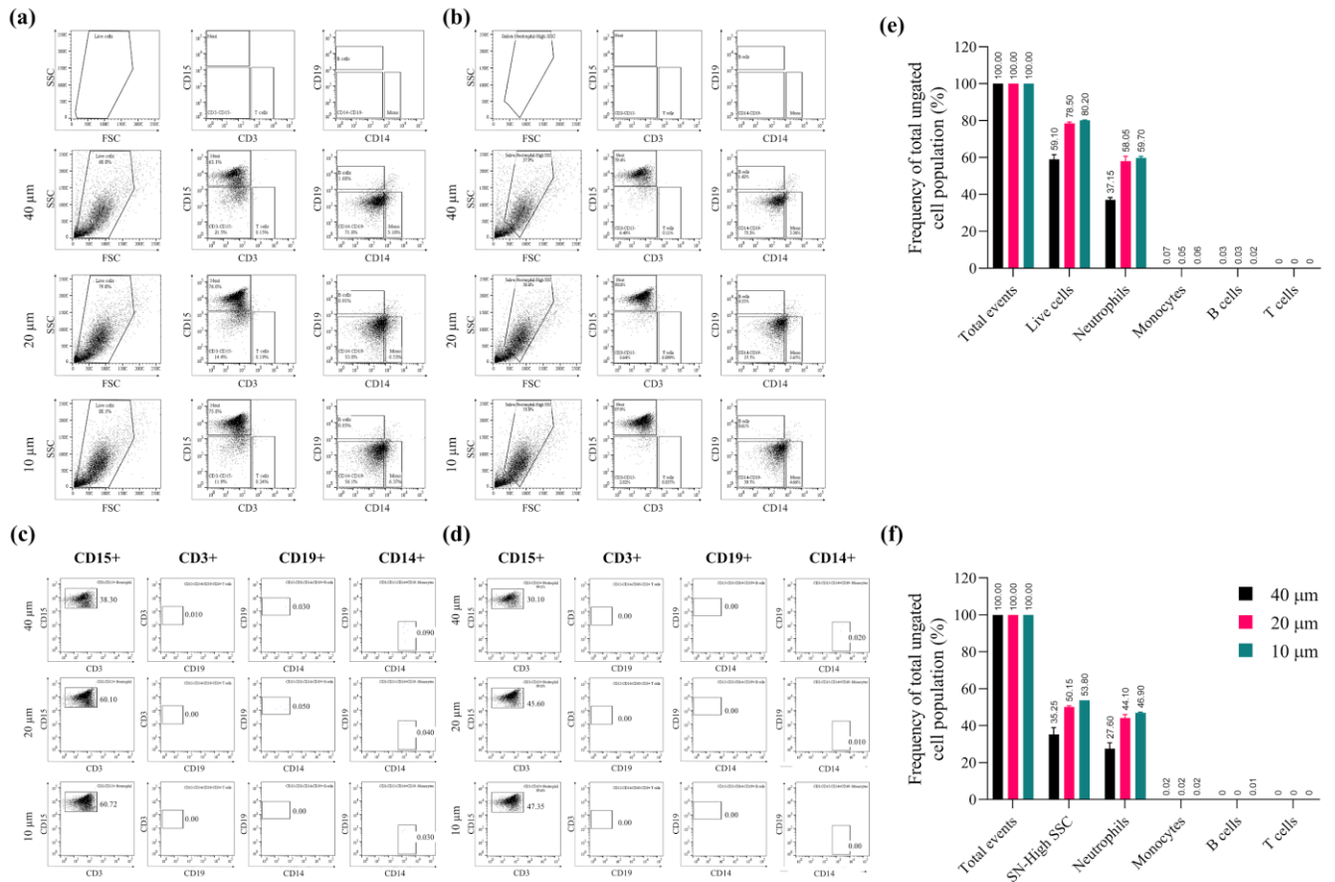


1280

1281 **Figure 2. Microscopic visualization and quantification of neutrophils in saliva.** (a) Representative
 1282 images of unstained and stained samples obtained by holotomography microscope (Nanolive 3D
 1283 Cell Explorer) at 60X magnification shows reduction of oral epithelial cells and enrichment of
 1284 neutrophils from 40-micron to 10-micron filtration. Brightfield, digitally-acid stained images and
 1285 RI (refractive index) 3D rendering images were used to identify each unstained cell type based on
 1286 their intracellular structure. Classic markers for immune cells including neutrophils (CD66b⁺),
 1287 monocytes (CD14⁺) and lymphocytes (CD19⁺) were employed to confirm the identity of cells and
 1288 enrichment protocol. (b) By zooming the images captured at 60X objective we looked for

1289 morphological cues to (i) demonstrate classic horse-shoe shaped nuclei of banded-neutrophils after
 1290 10-micron filtration and (ii) their vitality. Giemsa staining imaged by Zeiss AxioVision at 40X
 1291 objective demonstrates classic neutrophil morphology. (c) Quantification of cell types
 1292 demonstrated decrease of oral epithelial cells ($p = 0.0037$) and neutrophil enrichment ($p = 0.0025$).
 1293 Data includes cell counts from minimum 10 nanolive-images obtained from each healthy donor (n
 1294 $= 3$). White arrowhead indicates the neutrophils, ($*p \leq 0.05$; $**p \leq 0.01$; $***p \leq 0.001$). Please see
 1295 *Supplementary Fig. 5 for cell viability*.

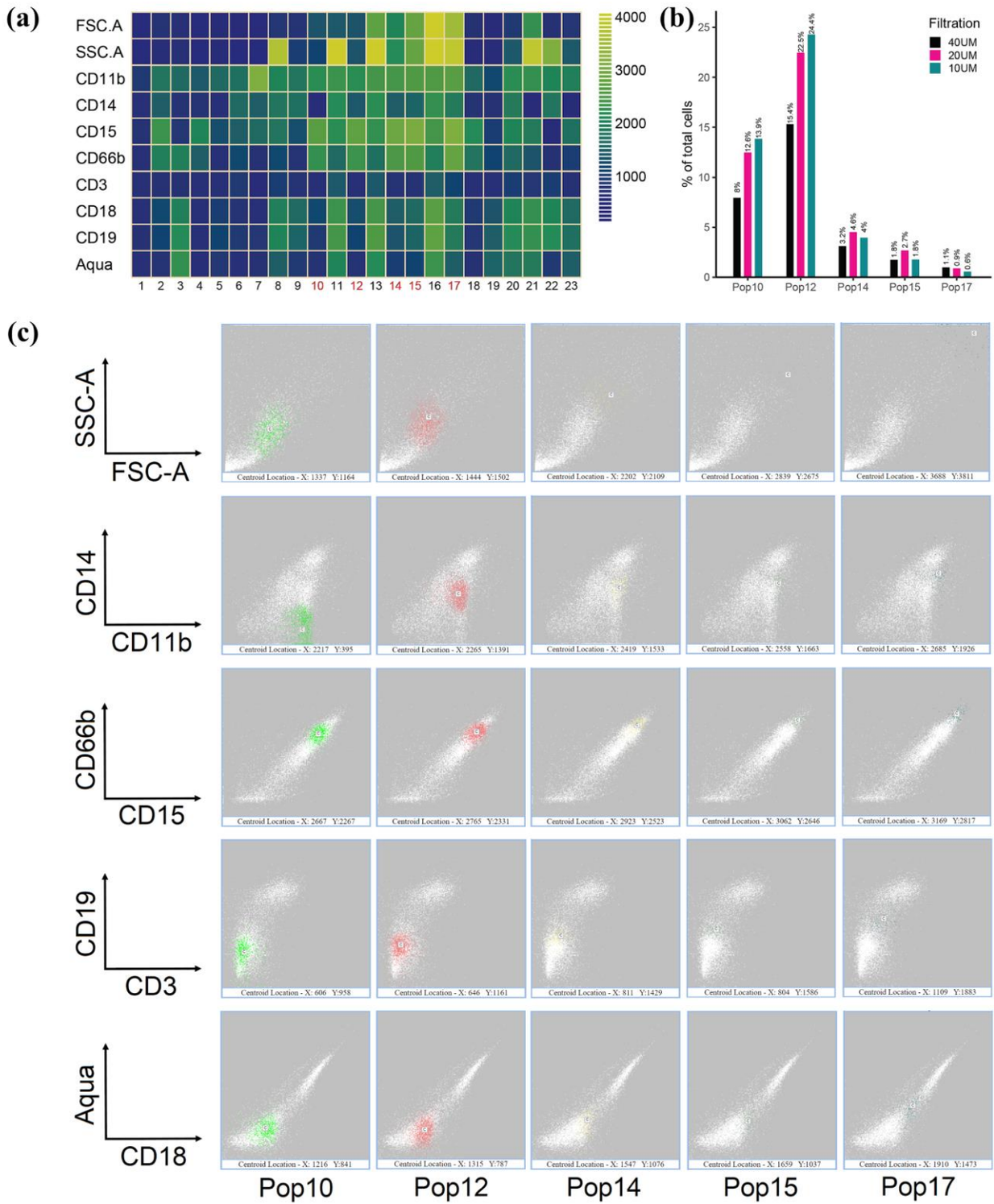
1296 **Figure 3**



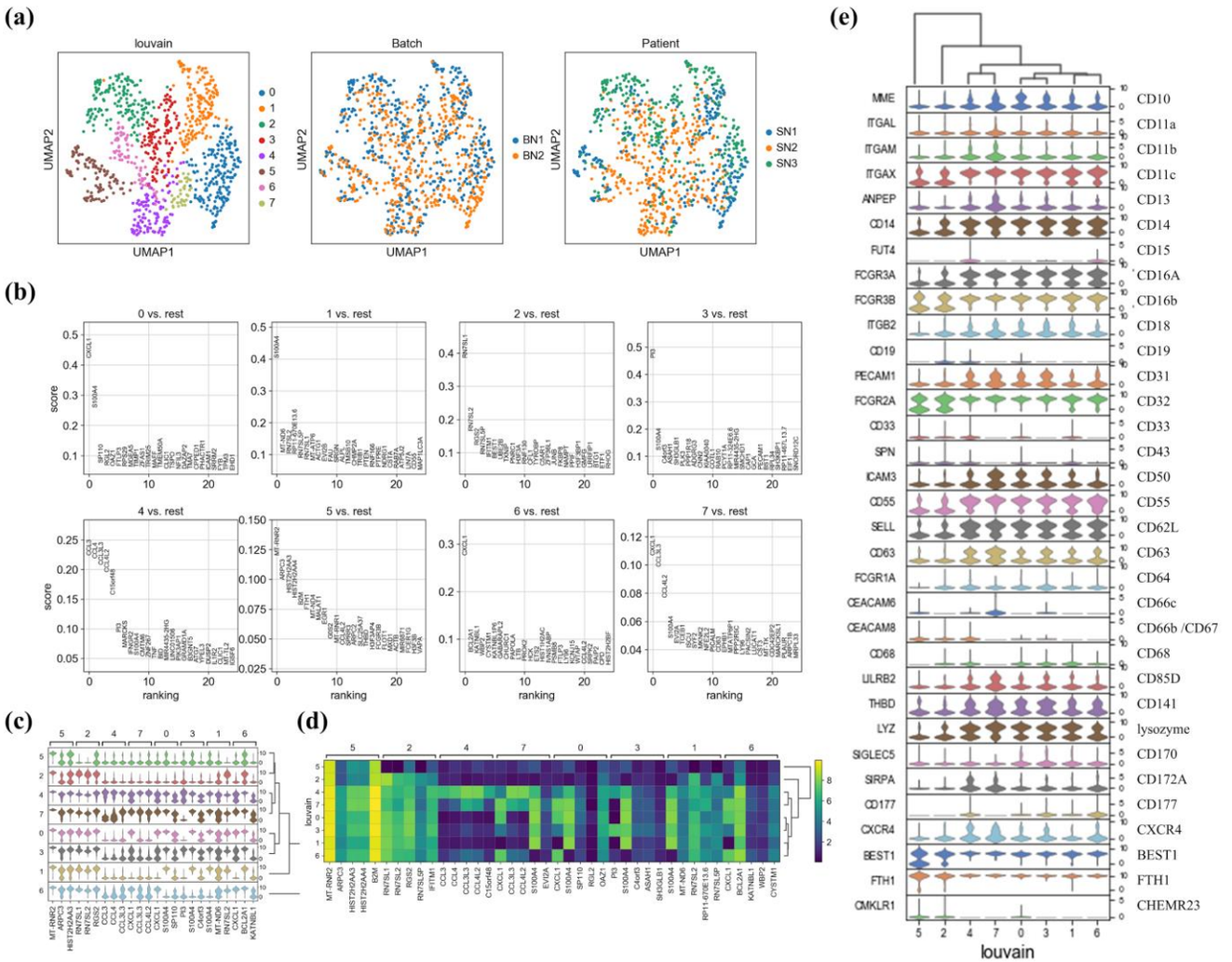
1297

1298 **Figure 3. Flow cytometry analysis reveals the immune landscape of salivary cell populations.**
 1299 Enriched saliva neutrophil is selected by live cells and high-SSC gating for saliva neutrophils.
 1300 Change in population density for all cells from subsequent 40 μm to 20 μm to 10 μm filtration is
 1301 compared among (a) live cell gated cells, in comparison to (b) high SSC cells. Immune cell types

1302 present in the healthy human saliva is determined by presence of classical CD-markers in human-
1303 blood immune cells (i.e. CD19⁺CD3⁺ as lymphocytes; CD14⁺CD11b⁺ as monocytes; and
1304 CD11b⁺CD15⁺ as neutrophils). (c) To look at the frequency of cell type in live cell gating on
1305 filtration from 40 μ m to 10 μ m, in comparison to total ungated events. (d) Change in frequency of
1306 cell type in saliva neutrophil high-SSC gate on filtration from 40 μ m to 20 μ m to 10 μ m, in
1307 comparison to total ungated events. Qualification of live cells (e) and high-SSC (f) were plotted.
1308 Cells were reduced/eliminated on subsequent 40 μ m (black), 20 μ m (magenta) and 10 μ m (teal)
1309 filtration steps used for saliva neutrophil enrichment protocol. Results for all measurements are
1310 mean \pm SD. Mean values are shown above each bar.



1313 **Figure 4. Unsupervised data clustering analysis identifies 23 cell populations from FACS**
1314 **Aria II flow cytometry data, including 5 subsets of salivary neutrophils. (a)** Heatmap of mean
1315 fluorescence intensity (MFI) of the 23 cell populations identified by FLOCK from the 40 μm
1316 sample with population ID on the x-axis and parameters measured on y-axis. IDs of neutrophil
1317 subpopulations are highlighted in red based on phenotype definition:
1318 $\text{FSC-A}^+\text{SSC-A}^+\text{CD11b}^+\text{CD14}^-\text{CD15}^+\text{CD66b}^+\text{CD3}^-\text{CD19}^-\text{Aqua}^-$. **(b)** Bar graph showing frequency
1319 changes of neutrophil subpopulations with population ID on the x-axis and percentages of total
1320 cells on y-axis (mean of the two replicates of each filtration). Frequencies of the two abundant
1321 neutrophil subpopulations (Pop10 and Pop12) increased from 40 μm to 10 μm filtration, while
1322 frequencies of the three rare ones did not. **(c)** 2D dot plots showing the position of each identified
1323 neutrophil subpopulation (represented by different colors) from the 40 μm sample, with other cells
1324 in the sample in white background. Each column corresponds to a neutrophil subpopulation with
1325 pairwise combination of measured parameters on rows: FSC-A vs. SSC-A (cell size and
1326 complexity), CD15 vs. CD66b (classical neutrophil markers), CD14 vs. CD11b (classical
1327 monocyte markers), CD3 vs. CD19 (classical lymphocyte markers), and CD18 vs. Aqua
1328 (live/dead).

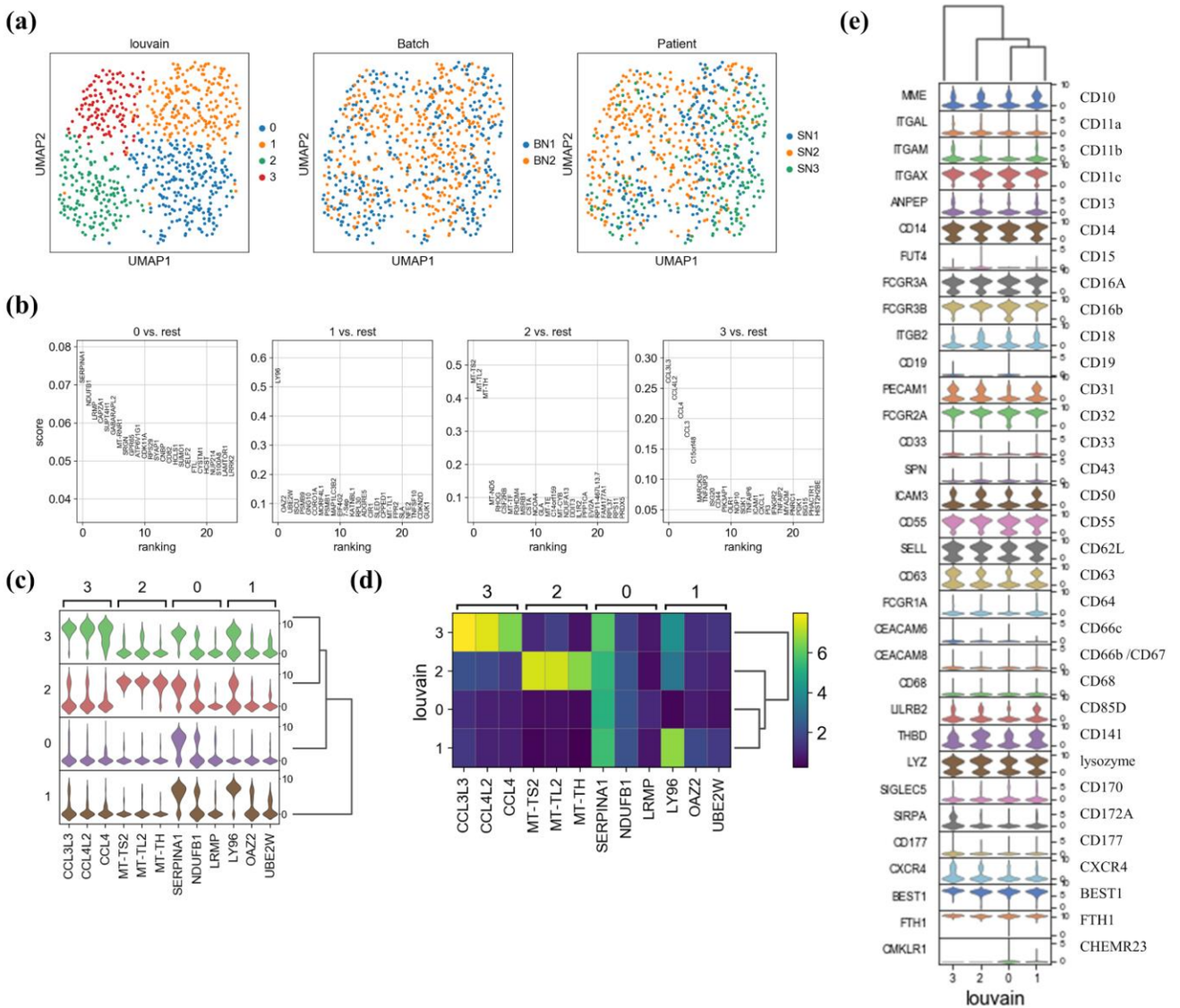


1330

1331 **Figure 5. Minimal scRNA-Seq filtering criteria determined more subsets of novel oral**
 1332 **neutrophils than standard filtering criteria suggesting the use of less stringent filter criteria**
 1333 **for neutrophils.** “Minimal Filtering” criteria were established including inclusion of cells with
 1334 greater than 50 genes per cell and cluster analysis of 2500 top variable genes. (a) Louvain
 1335 clustering at a resolution 0.8 identified eight clusters (0 to 7) among three healthy donor’s saliva
 1336 neutrophils (SN1, SN2, and SN3) without any “batch effect” among single-cell sorted plates run
 1337 on BioCel 2100 robotics in two batches (BN1 and BN2). (b) Marker genes were determined by
 1338 logistic regression performed by comparing each target cluster with the rest and distance among
 1339 each cluster, here the gene ranks on the x-axis while the regression score is given on the y-axis (c)

1340 heatmap of marker genes showing expression levels per cluster **(d)** is shown here. **(e)** Expression
 1341 levels of already known/published human neutrophil markers from both saliva and blood are given
 1342 for each louvain cluster. The gene names are represented on the left side of cluster-map and their
 1343 commonly used synonyms are shown on the right side.

1344 **Figure 6.**

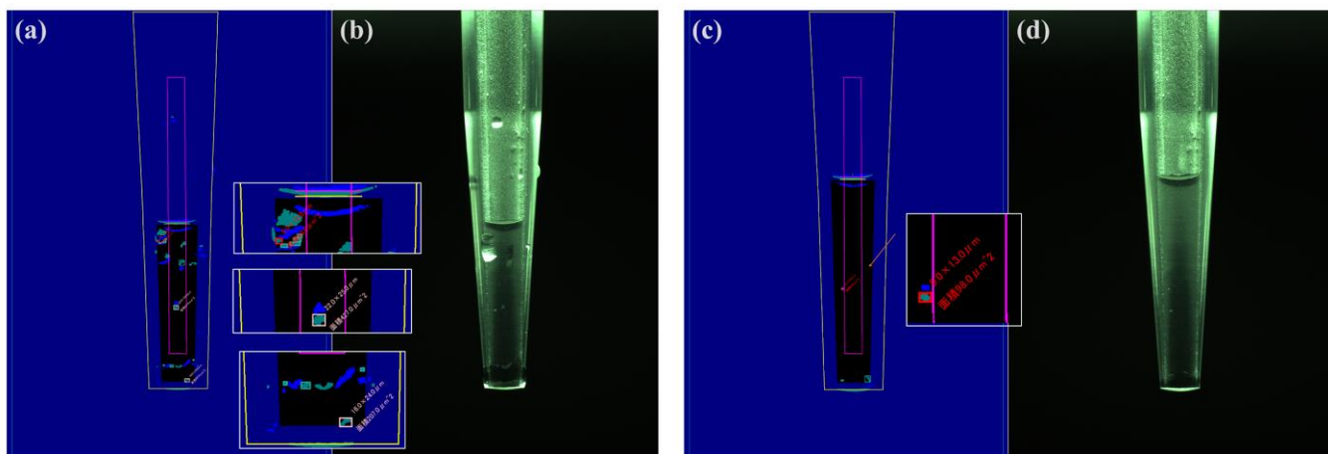


1345
 1346 **Figure 6. Stringent/standard scRNA-Seq filtering criteria determined fewer novel oral**
 1347 **neutrophil subsets. “Stringent Filtering”** criteria were established, including inclusion of cells

1348 with greater than 400 genes per cell and cluster analysis of 2000 top variable genes. (a) Louvain
1349 clustering at a resolution 0.8 identified eight clusters (0 to 3) among three healthy donor's saliva
1350 neutrophils (SN1, SN2, and SN3) without any "batch effect" among single-cell sorted plates run
1351 on BioCel 2100 robotics in two batches (BN1 and BN2). (b) Marker genes were determined by
1352 logistic regression performed by comparing each target cluster with the rest and distance among
1353 each cluster, here the gene ranks on the x-axis while the regression score is given on the y-axis (c)
1354 heatmap of marker genes showing expression levels per cluster (d) is shown here. (e) Expression
1355 levels of already known/published human neutrophil markers from both saliva and blood are given
1356 for each louvain cluster. The gene names are represented on the left side of cluster-map and their
1357 commonly used synonyms are shown on the right side.

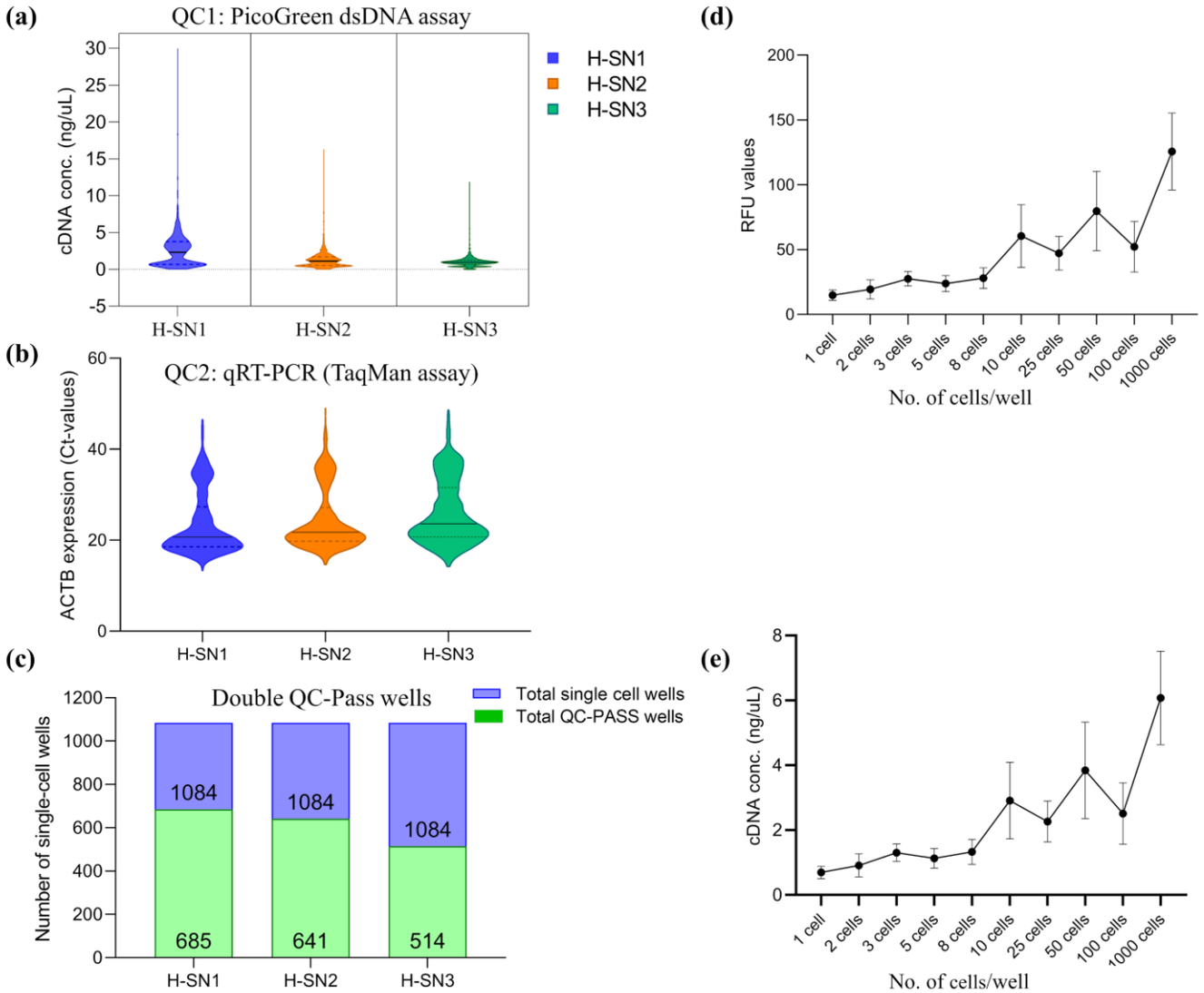
1358 8. SUPPLEMENTARY FIGURES

1359 Supplementary Figure 1.



1360
1361 **Supplementary Figure 1. Imaging of single cells sorting of oral neutrophils.** Software
1362 generated image of automated pipetting system by On-Chip-SPiS single-cell dispenser showing
1363 (a) selection of single cell images (b) showing the rejected sample dispensing image due to
1364 presence of multiple cells in the dispensing pipette tip and their sizes (Range of cell sizes are user
1365 input parameters). (c) Software generated image of (d) showing the allowed sample dispensing
1366 image due to presence of a single cell in the dispensing pipette tip having the desired cell size.
1367 (Insert shows the zoomed image of the large or doublet cells identified by software.)

1368 **Supplementary Figure 2.**

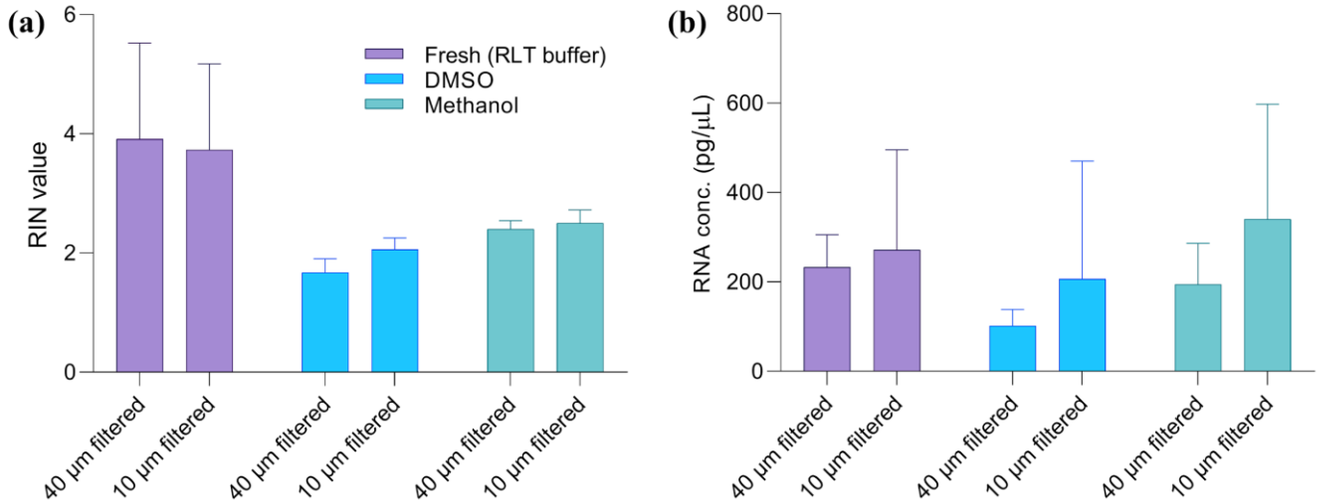


1369

1370 **Supplementary Figure 2. Quality control (QC) analysis of each single cell well.** PicoGreen
 1371 dsDNA assay for cDNA concentration and qRT-PCR TaqMan assay for ACTB expression. **(a)**
 1372 Violin plot representing the cDNA concentration (ng/μL) by PicoGreen dsDNA assay of each
 1373 single cell well from three 384-well plates of three healthy donors. **(b)** Violin plot representing the
 1374 Ct-values for ACTB-expression by TaqMan assay of each single cell well from three 384-well
 1375 plates of three healthy donors. Concentration of cDNA increases with increase in the number of
 1376 cells/wel. **(c)** Superimposed graph showing the consistent QC-Pass wells (green) from total 1084
 1377 of single-cell wells (blue) of three healthy donors. QC-Pass wells are selected and combined into

1378 a single 384-well HitPicked cDNA library plate (named SN1, SN2, and SN3 respectively for each
1379 donor). (d) Mean RFU and (e) cDNA concentration obtained by PicoGreen dsDNA assay for
1380 varying numbers of cells sorted in each well.

1381 **Supplementary Figure 3.**

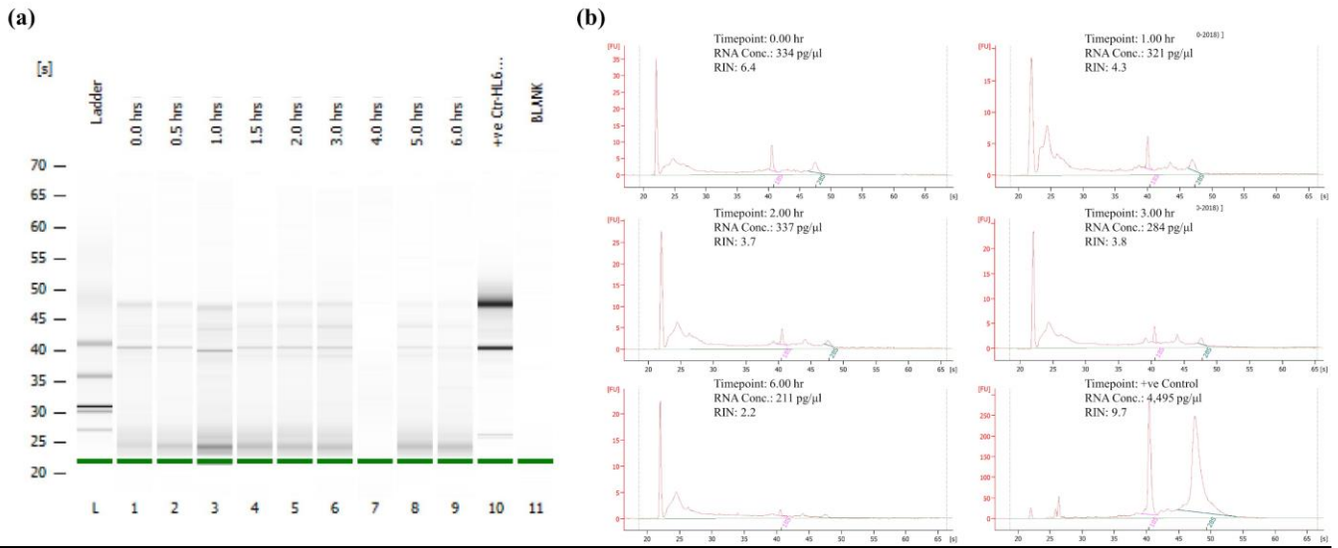


1382

1383 **Supplementary Figure 3. RNA quality degrades on storing saliva neutrophil by DMSO**
1384 **cryopreservation or methanol fixation storage methods.** RNA from enriched saliva neutrophils
1385 were either isolated fresh by immediately using RLT buffer (violet) of QIAGEN RNA isolation
1386 kit or stored at -80 °C by DMSO cryopreservation (blue) following *Wohnhaas et al.* 2019 and
1387 methanol fixation (teal) following *Chen et al.* 2018. Stored cells were recovered after a week and
1388 RNA was isolated using the same kit and (a) RIN and (b) RNA concentration was determined by
1389 Bioanalyzer. The result suggests using fresh saliva neutrophil samples and not DMSO or methanol
1390 cryopreserved samples for RNA-seq pipeline.

1391

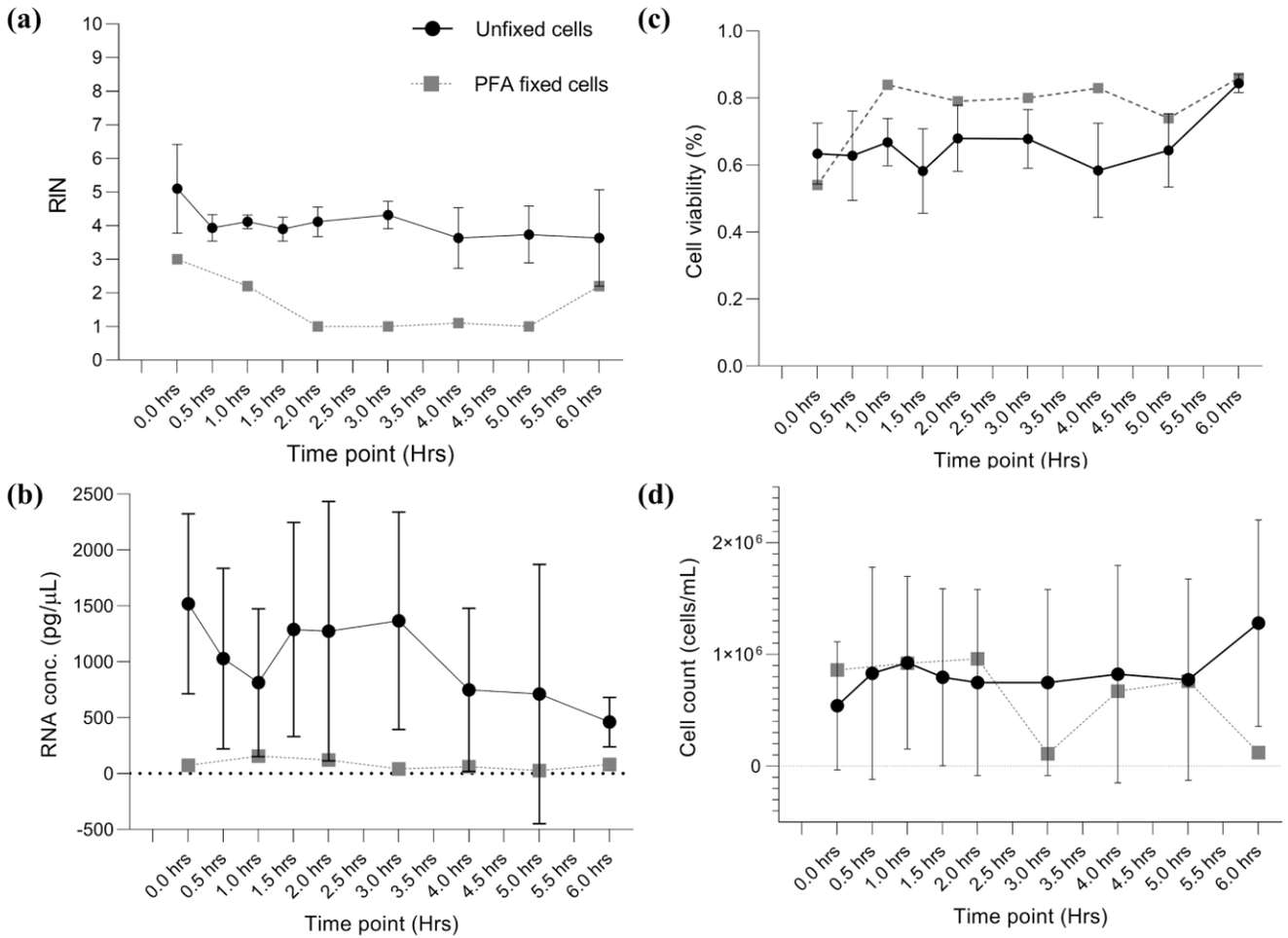
1392 **Supplementary Figure 4.**



1393

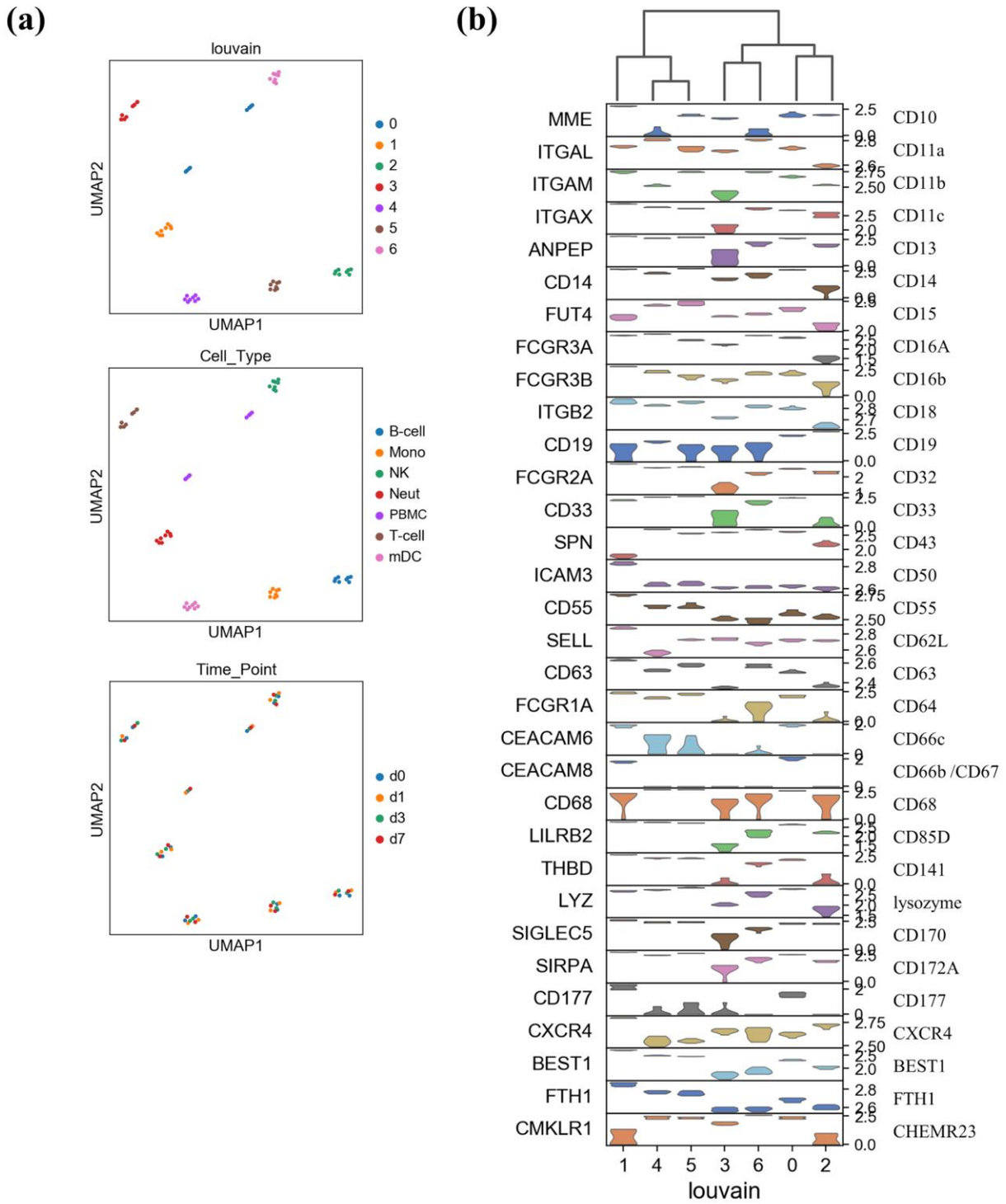
1394 **Supplementary Figure 4. RNA degradation patterns of oral neutrophils. (a)** Representative
1395 image of Bioanalyzer-Gel image showing the degradation of RNA over time. Lane 1 to 9 are
1396 enriched saliva neutrophil samples collected at different time points (denoted above each lane). As
1397 a positive control, total RNA of the HL-60 cell line is used in lane 10 and lane 11 is left blank as
1398 negative control. **(b)** Representative image of Bioanalyzer-electropherogram for time-point 0, 1,
1399 2, 3, and 6 hours and total RNA of cultured HL60 cell lines is used as a positive control. Cells
1400 were maintained at room temperature in 0.9 % saline solution/ PBS⁽⁻⁾.

1401 **Supplementary Figure 5.**



1402

1403 **Supplementary Fig. 5. Comparing cell viability and RNA quality of fixed (4 % PFA) and**
 1404 **unfixed saliva neutrophils. (a)** Quality of RNA with decrease in RNA integrity number (RIN)
 1405 and **(b)** total RNA concentration (ng/ μ L) of cells in enriched saliva neutrophils over time. **(c)**
 1406 Saliva neutrophils maintain consistent cell-viability to 70-80 % and **(d)** total cell-count even after
 1407 6 hours' time period. Cells were maintained at room temperature in 0.9 % saline solution/ PBS⁽⁻⁾.



1411 of blood cells following our oral immune-cell pipeline applied to the publicly available dataset
1412 accession number GSE64655. **(a)** Louvain clustering with resolution of 1 from a graph produced
1413 using 5 neighbors (n) identified seven clusters (0 to 6) among two healthy donor's blood that
1414 correlates with immune cell types B cells, monocytes (Mono), NK cells, neutrophils (Neut),
1415 PBMCs, T cells and dendritic cells (mDC), collected on day 0, 1, 3, and 7 post-vaccination. Cluster
1416 1 represents the neutrophil population. **(b)** Gene expression level of previously identified human
1417 neutrophil markers. The gene names were represented on the left side of cluster-map and their
1418 commonly used names were shown on the right side.

1419 **9. SUPPLEMENTARY INFORMATION**

1420 **9.1. Supplementary Information:**

1421 **9.1.1. Cultured cell lines:**

1422 As a model neutrophil cell type to optimize the protocols for the successful processing of
1423 human primary saliva neutrophils and troubleshoot unexpected problems we used HL-60 cells
1424 and/or differentiated HL-60 (dHL-60) cells. HL-60 cell line was obtained from ATCC (cat. no.
1425 CCL-240) and was maintained on Iscove's Modified Dulbecco's Medium (IMDM) media
1426 containing 10 % fetal bovine serum (FBS) and 1 % penicillin. HL-60 cell lines were differentiated
1427 by culturing them in differentiation media containing RPMI 1640, containing 10 % FBS and 1 %
1428 penicillin and 1.25 % Dimethylsulfoxide (DMSO) to obtain dHL-60 cells. HL-60 and dHL-60
1429 cells were harvested from the culture flask by centrifuging at 160 x g for 10 min at 4 °C. The cell
1430 pellet is washed once in Ca²⁺ and Mg²⁺ -ions free PBS (PBS⁽⁻⁾) and then resuspended into a sorting
1431 buffer for FACS sorting or staining buffer for flow cytometry staining.

1432

1433 **9.1.2. Cryopreservation of neutrophil sample by DMSO or methanol fixation method**
1434 **degrades RNA and is not suitable for NGS-workflow:**

1435 Neutrophils have a short lifespan and thus need to be processed within 4-6 hours of sample
1436 collection for NGS-workflow. Obtaining samples from patients for NGS-workflow and processing
1437 them within this short duration is always not possible as the sample collection center is usually not
1438 located at the same place where they are processed. Previous studies have shown that DMSO
1439 cryopreservation³⁴ and methanol fixation of PBMCs¹⁶ are methods of choice for storage of
1440 samples and processing later for single-cell RNA sequencing. Therefore, we decided to check the
1441 quality of RNA in neutrophils by using their storage protocols and compared with fresh saliva
1442 neutrophils. The saliva neutrophil samples were collected from two healthy donors (HSN1 and
1443 HSN2) as mentioned in section 1a and cells were collected after 40 μm and 10 μm filtration steps.
1444 RNA was isolated from fresh samples by using RLT-buffer (from Qiagen DNA/RNA/Protein
1445 isolation kit); from DMSO cryopreserved samples stored at -150 °C (in liquid N₂) and from
1446 methanol fixed samples stored at -20 °C for 2 weeks. We checked the degradation of RNA by
1447 using Agilent RNA 6000 pico chip in Bioanalyzer.

1448 The data obtained from Bioanalyzer suggests that for saliva neutrophils fresh samples are
1449 best for scRNA-seq studies having RIN values between 4 and 6 as compared to DMSO-
1450 cryopreserved or methanol fixed samples having RIN values < 3 (Supplementary Fig. 3).

1451 **9.2. Source Data:** Flow cytometry data will be completely deposited to ImmunoImport and single
1452 cell transcriptomics will be deposited to GSEA. Accession number pending.

1453 **10. AUTHOR CONTRIBUTIONS STATEMENTS:**

1454 Writing original draft. SC, MF. Writing review & editing: all authors. Oversaw the project and
1455 assist with the writing of the manuscript: MF, MN, RS, YQ. Prepared Graphs: SC, BA, AM, YQ,
1456 AD; Assisted with sample processing: SC, MN, SL. Assisted with sample analysis: SC, MN, BA,
1457 SL; Project Administration: MF; Funding Acquisition: MF, RS, YQ. All authors reviewed the
1458 manuscript.

1459

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1469 The authors declare no potential conflicts of interest with respect to the authorship and/or
1470 publication of this article.

1471 **14. REFERENCES:**

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