DATA NOTE
Flow cytometry analysis of epithelial cell populations from touch samples using the BD Influx flow cytometry platform [version 1; referees: 1 approved with reservations, 1 not approved]

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Abstract
'Touch' or trace cell mixtures submitted as evidence are a significant problem for forensic laboratories as they can render resulting genetic profiles difficult or even impossible to interpret. Optical signatures that distinguish epidermal cell populations from different contributors could facilitate the physical separation of mixture components prior to genetic analysis, and potentially the downstream production of single source profiles and/or simplified mixtures. For this dataset, optical properties including forwards scatter (FSC), side scatter (SSC), and fluorescence emissions in the Allophycocyanin (APC) channel were measured in epithelial cell populations from touch samples collected from several different contributors on multiple days to assess inter- and intra-contributor variability.

This article is included in the Data: Use and Reuse collection.
Introduction
Flow cytometry has proven a viable approach for differentiating cell populations in many types of uncompromised (i.e. non-degraded) forensic mixture sample (Dean et al., 2015; Schoell et al., 1999; Verdon et al., 2015). However, application to ‘touch’ or trace epithelial cell mixtures remains a challenge since many cell surface features are lost or obscured during the process of keratinocyte differentiation, leaving few biochemical or structural features in shed corneocytes that vary between individual contributors. Recent research has suggested that optical properties such as autofluorescence at red wavelengths may be a potentially discriminating feature for epidermal cell populations in some touch mixture samples (Stanciu et al., 2016). In this study, we examined the consistency of such signatures using a different flow cytometry platform (BD Influx Cell Sorter) and set of contributors.

Methods

**Dataset 1. Influx touch epithelial samples**

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Source data files are organized into four different flow cytometry surveys, each involving a different set of donors, all of whom were sampled on the same day. File names are labeled with the anonymized sample ID number used for all experiments. Replicate measurements from the same cell solution are designated as ‘rep1’, ‘rep2’, and so forth. A table of analyzed samples (labeled by Donor ID) across each of the four experiments is provided.

Touch samples were collected from six volunteers using the following protocol which was approved by the VCU-IRB (#HM20000454; CR). Volunteers rubbed a sterile polypropylene conical tube (P/N 229421; Celltreat Scientific) for five minutes using their entire hand (i.e., palm and fingers). Cells were collected from the surface with six sterile pre-wetted swabs (P/N 22037924; Fisher Scientific) followed by two dry swabs. To elute the cells into solution, the swabs were manually stirred then vortexed for 15 seconds in 10 mL of ultrapure water (18.2 MΩ-cm). The entire solution was then passed through a 100 µm filter mesh prior to flow cytometry. Flow cytometry analysis of eluted cells was performed on the BD Influx Cell Sorter (Becton Dickinson) using the 488nm, 561nm, and 640nm lasers. Channel voltages were set as follows: Forward Scatter (FSC, 17.5V), Side Scatter (SSC, 16V) and Allophycocyanin (APC, 74.6V).

**Data availability**

*F1000Research*: Dataset 1. Influx touch epithelial samples, 10.5256/f1000research.8338.d116907 (Kwon et al., 2016).

**Author contributions**

CE conceived the study. CE, CS, and YK designed the experiments. CS and YK carried out the research. CE and KP prepared the first draft of the manuscript.

**Competing interests**

No competing interests were disclosed.

**Grant information**

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The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

**References**


The authors describe a dataset used for flow cytometric analysis of sloughed epithelial cells from a set of 6 individuals. It is not at all clear why these data are different from those reported in their copublished research note (http://f1000research.com/articles/5-180/v1) which provides an analysis of these data. The anonymized names of the individuals in the two papers are the same and they are both in .fcs format. The Data Note is larger than the data in the Research Note, presumably because it contains technical replicates. The differences are trivial. In fact, the Research Note would have greater validity if the Data Note dataset was incorporated into this document and discussed there.

The second point that should be addressed is how the statistics of the fluorescence distributions (FS, SS) for different samples from the same individual or from different individuals can be compared given that the intensities vary, presumably as a result of the differences in yield from each sample. The reliability of using these histograms for making comparisons between replicates or individuals for forensic or any other applications could be suspect (eg. sample R12) due to noise, broad distributions or other factors. I suggest that the authors determine and provide minimum threshold criteria for analysis of a sample or comparison with other samples.

**Competing Interests:** No competing interests were disclosed.

I have read this submission. I believe that I have an appropriate level of expertise to state that I do not consider it to be of an acceptable scientific standard, for reasons outlined above.
is correct that some of the same donors contributed samples to both studies, albeit collected on different days. Given observed differences in fluorescence profiles for some contributors from one day to the next, sampling from the same contributor over multiple days adds to our understanding of this phenomenon.

To further distinguish this Data Note from previous publication we have significantly revised the Data Note to include a more comprehensive set of samples and experimental conditions, investigating both optical properties and antibody binding capacities of ‘touch’ epithelial cell populations. This includes flow cytometry data for autofluorescence as well as antibody hybridizations with HLA and Cytokeratin probes. As the reviewer suggests, we are also in the process of incorporating aspects of this dataset into other manuscripts that discuss the biochemical and forensic applications of flow cytometry analysis of touch samples (e.g., pre-print available at http://biorxiv.org/content/early/2016/03/28/045948).

We also agree with the reviewer’s second point that there may be significant variability in the level of autofluorescence exhibited by touch samples. This in turn may be influenced by a variety of factors such as presence or absence of exogenous compounds, and/or intrinsic biological characteristics of the cells themselves. Understanding this variability, particularly for samples derived from the same contributor or for aged/degraded samples, is necessary to assess the potential utility of flow cytometry-based cell separation techniques such as FACS for downstream DNA profiling of separated cell populations derived from ‘touch’ biological mixtures.

We would note that although differences in cell yield change the number of cells (Y axis) fluorescing in the red portion of the spectrum at a given RFU value (X axis), it is nonetheless possible to develop a sense of how the average intensity of red autofluorescence exhibited by cells collected from different individuals, or from the same individual on different days, varies by comparing histograms, regardless of cell yield. We agree that an individual’s total cellular contribution to a biological mixture will ultimately be an important factor in whether that individual’s DNA profile can be successfully generated from a sorted mixture. However, any kind of minimum cell count threshold would be inextricably linked to the sensitivity/efficiency of downstream DNA typing methodologies used on sorted cell populations, and are beyond the scope of this Data Note.

**Competing Interests:** No competing interests were disclosed.

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The article suggests "touch samples" however the data set contains no data on forensic relevant touch samples. The six samples were from volunteers rubbing their entire hand. These are “fresh” cells and might not show the same flow characteristics as cells left behind on an object after a touch contact and
having “aged” on the object. The dataset provided is as it is not really relevant to the forensic field.

In addition I think that 6 samples is maybe a too limited number for this kind of studies.

**Competing Interests:** No competing interests were disclosed.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Author Response 29 Sep 2016

**Christopher Ehrhardt,** Virginia Commonwealth University, USA

We agree with the reviewer that there is an important distinction between ‘fresh’ biological samples and ones that are aged and/or degraded since the latter is more likely to be encountered in forensic caseworking samples. Uncompromised samples can nonetheless provide an important foundation for future studies that explicitly examine the effects of aging/degradation on optical signatures identified by the initial study. We also note that there are forensic scenarios where fresh touch samples may be collected and analyzed for an investigation. For example, a firearm may be discarded by a suspect and immediately collected during pursuit by law enforcement.

However, to increase the forensic relevance of this dataset the new version of the manuscript includes samples that were aged between 12 hours and seven days before collection, as well as samples that were collected from forensically relevant substrates such as replica firearms and knife handles.

For the updated version of the manuscript we have also expanded the dataset to include 33 different contributors.

**Competing Interests:** No competing interests were disclosed.