All\(^2\): A tool for selecting mosaic mutations from comprehensive multi-cell comparisons

Vivekananda Sarangi\(^1\), Yeongjun Jang\(^1\), Milovan Suvakov\(^2\), Taejeong Bae\(^1\), Liana Fasching\(^1\), Shobana Sekar\(^1\), Livia Tomasini\(^2\), Jessica Mariani\(^2\), Flora M. Vaccarino\(^2,3\), Alexej Abyzov\(^1\)*

1 Department of Quantitative Health Sciences, Center for Individualized Medicine, Mayo Clinic, Rochester, Minnesota, United States of America, 2 Child Study Center, Yale University, New Haven, Connecticut, United States of America, 3 Department of Neuroscience, Yale University, New Haven, Connecticut, United States of America

* Abyzov.Alexej@mayo.edu

Abstract

Accurate discovery of somatic mutations in a cell is a challenge that partially lays in immaturity of dedicated analytical approaches. Approaches comparing a cell’s genome to a control bulk sample miss common mutations, while approaches to find such mutations from bulk suffer from low sensitivity. We developed a tool, All\(^2\), which enables accurate filtering of mutations in a cell without the need for data from bulk(s). It is based on pair-wise comparisons of all cells to each other where every call for base pair substitution and indel is classified as either a germline variant, mosaic mutation, or false positive. As All\(^2\) allows for considering dropped-out regions, it is applicable to whole genome and exome analysis of cloned and amplified cells. By applying the approach to a variety of available data, we showed that its application reduces false positives, enables sensitive discovery of high frequency mutations, and is indispensable for conducting high resolution cell lineage tracing.

Author summary

DNA make up in cells of a human is slightly different from one another because of cell specific mutations called mosaic mutations. Mosaic mutations can be introduced during early development in a fetus, during normal cell division throughout life or during an aggressive onset of cell division such as cancer. Thus, the extent of accumulation, time of acquiring and specific location of mosaic mutations in the genome are vital for the understanding of the biology of normal development as well as diseases that are caused by it. The ultimate way of discovering and analyzing mosaic mutations is by studying the genome of single cells. Our method, All\(^2\), uses a novel approach to compare the genome of single cells to one another to accurately recognize true mosaic mutations from natural variations and from noise by implementing a unique scoring method. We have shown that our method performs better than discovery of mutations from a bulk or from comparing cells to a bulk. We have applied the method to high resolution cell lineage tracing and demonstrated its superb performance for reconstructing individualized cell ancestry trees starting from the zygote.
**Funding:** This study was funded by the Foundation for the National Institutes of Health (NIH), grant number: R01MH100914, https://reporter.nih.gov/search/Ub0-VCBSJU6igHtJvKaVzw/projects, and grant number: U01MH106876 https://reporter.nih.gov/search/ZjYDqHtGaUeFFWwy3HrYw/projects to FV. AA received funding from the Division of Cancer Epidemiology and Genetics, National Cancer Institute, grant number U24CA220242, https://reporter.nih.gov/search/mJbqYiI0PUm r4Y9x0E72K w/projects FV and AA were both funded by the Simons Foundation, grant number 399558, https://www.simonsfoundation.org/ The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

**Competing interests:** The authors have declared that no competing interests exist.

**Introduction**

With advances in sequencing technologies, analysis of the genome of a single cell is gaining traction owing to its multiple applications. Applications include studying mutations in normal and cancer single cells, identification of driver mutations in cancer [1,2], tracing cell lineages in human development and identification of sub-clones in cancer, where mosaic mutations are used as barcodes to study cell lineages [1–4]. For detection of mosaic mutations in single cells, the most frequently used approach is to compare single cell genomes to that of a matched reference bulk. While this approach works well to find private mutations in a cell, it misses mutations that are present at higher frequency, and consequently present in multiple cells in the bulk. It also requires one to have bulk data which might not be always available. Here we present a tool called All^2 (pronounced ‘all square’) which detects mosaic mutations without the need for a reference bulk by relying on comprehensive cell-to-cell comparisons. By consolidating information from all comparisons, every call is categorized as either a germline variant, mosaic mutation or noise/false positive.

**Results**

**Concept**

All^2 is an easy-to-use tool which extends and implements an algorithm initially proposed in Bae et al. 2018 [5]. All^2 takes mutation calls from all pair-wise comparisons of N cells in the study and, for every non-redundant call, creates a NxN pairwise binary matrix corresponding to comparing different pairs of cells, where 1 corresponds to a call and 0 to no call. Patterns of values in the matrix are used to determine whether a call is a mosaic mutation, germline variant or false positive (Fig 1A–1D). In theory, the presence of these patterns should be sufficient to make the determination, however, real data has noise, smearing the patterns (S1 Fig).

For effective categorization, we developed a scoring system which reflects how likely it is for a call to be a mosaic mutation or a germline variant. The tool calculates two scores: a germline score and a mosaic score, each within a range between 0 and 1. A real mosaic variant could only be discovered when comparing a cell carrying the variant and a cell not carrying the same variant. The number of times a call for a variant shows up in the matrix is determined by

\[ n = f(1-f)N^2, \]

where \( n \) is the number of times a variant is seen in all comparison, \( f \) is fraction of cells carrying the variant, \( N \) is the total number of cells. By solving the above quadratic equation, we get two solutions:

\[ f_m \approx 0.5 - \sqrt{0.25 - n/N^2} \quad \text{and} \quad f_g \approx 0.5 + \sqrt{0.25 - n/N^2} \]

Since the mosaic mutations are typically present in a small fraction of cells in the bulk, and germline variants are present in (almost) all the cells, we conditionally call \( f_m \) as frequency of a mosaic mutation and \( f_g \) as frequency of a germline variant. Note, a germline variant can be lost or undetected in some cells, and that is why its cell frequency in a bulk may be below 1.

Since \( f_m = 1-f_g \) we can just use one equation, such as \( f = f_m \), where \( f \) is the fraction of cells with mosaic mutation or the fraction of cells without germline variant. Now, we can calculate the number of cells \( N' \) carrying the mosaic mutation or the number of cells not carrying germline variant as \( N' \approx fN \). In case of a true mosaic mutation, the corresponding calls are arranged
in rows in the matrix (Fig 1B), and would sum up to

$$n_m = \sum_{i=1}^{N'} nr_i,$$

where \(nr_i\) is the number of calls for the variant in a row corresponding to the \(i^{th}\) cell. From the data, the best estimate of \(n_m\) is the maximum from all possible subset of \(N'\) cells from \(N\) (S2 and S3 Figs). Similarly, for a germline variant, corresponding calls are arranged in columns (Fig 1C), and would sum up to

$$n_g = \sum_{i=1}^{N'} nc_i,$$

where \(nc_i\) is the number of calls for the variant in a column corresponding to \(i^{th}\) cell. And best estimate of \(n_g\) is the maximum from all possible subset of \(N'\) cells from \(N\) (S2 and S3 Figs). The mosaic and germline scores are then defined as

$$\text{Mosaic Score} = \frac{\max(n_m)}{n}$$

$$\text{Germline Score} = \frac{\max(n_g)}{n}$$

A call having a high mosaic score and low germline score is defined as a mosaic mutation. Similarly, a call with high germline score and low mosaic score is defined as a germline score. When a call has both high germline and mosaic scores, we define it as a high frequency mosaic mutation. Such mutations are likely present at a higher fraction of cells in a tissue. For
example, such mutations could occur during early development and be present in a high fraction of cells across tissues in the human body [6]. The distribution of mutations (as dots) on a plane with axes corresponding to the two scores can be used to divide the calls into mosaic mutations, germline variants, noise or false positive (low mosaic and low germline score) and high frequency mosaic mutations (high mosaic and high germline score) (Fig 1E).

Implementation and usage

Genomes of all pairs of cells need to be compared prior to using All\textsuperscript{2}. Variant calls can be made by a caller of choice (see Methods). All\textsuperscript{2} is written in python and has three commands: `score`, `call`, and `matrix`. The first command takes a manifest file with names of single cells, along with the VCF file containing calls (SNVs and INDELS) as rows. Case and control fields in the manifest file are used to define the directions of pairwise comparisons, where the case is compared to control. A user can optionally provide a BED file with the inclusion list of genomic regions where to apply the filtering. The output of this command is a file with mosaic and germline scores for each of the calls as well as density scatter plot of the scores showing distribution of calls based on their scores (S4A Fig). The second command relies on the output of the `score` command and annotates the calls as mosaic mutation, germline variant, noise, or high frequency mosaic mutations based on default or user specified score cut-offs. This command also annotates the density scatter plot (Fig 1E), provides a file with annotated calls for each cell, per category plots of call counts, per sample plots of call counts, VAF (variant allele frequency) plot, and mutation spectrum plots (S4B–S4D Fig). The third command plots a matrix of pairwise comparison for one or multiple calls. The plots also display calculated scores along with VAF for the call(s) in each cell (S5 Fig). Analogous to SNVs and indels, All\textsuperscript{2} is capable of filtering structural variant (SV) calls using commands `score_sv`, `call_sv` and `matrix_sv`. Two SV calls are considered the same if they have at least 50% reciprocal overlap. For this purpose, the tool supports VCF file as input, e.g., VCFs generated by the SV caller MANTA [7,8].

One implicit underlying assumption of the approach is that in each compared cell, the genome is covered/sequenced uniformly. This is true in case of the single cell cloning approach, however, single cell genome amplification may result in non-uniform coverage which, at the extreme, manifests in allelic dropouts [9]. To handle this, we have implemented a dedicated allele dropout analysis (ADA) mode, which considers allele dropout regions when calculating the scores, thereby reducing the noise (see below). The ADA mode can also be used for running All\textsuperscript{2} on exome data where the exome capture region can be specified per cell in the manifest file.

Application to reconstructing cell lineage tree

To demonstrate the uniqueness of All\textsuperscript{2} approach, we applied it to reconstruct post-zygotic cell divisions in a living individual. Analysis of developmental cell lineages is one of the central questions in developmental biology, resolving which can shed light on the etiology of developmental diseases. Unlike model organisms, lineage tracing in humans can only be done retrospectively using naturally occurring somatic variants that serve as permanent marks of the lineages. Mutations that occur during early development are present in a high fraction of cells across tissues in the human body, and their discovery is challenging for existing methods.

In the analyzed individual, we compared mosaic variant discovery using three approaches: 1) by analysis of bulk blood and saliva; 2) by pairwise comparison of 25 clonal iPSC lines (representing 25 fibroblast single cells) with the bulk blood; and 3) by comparing the clonal lines followed by application of All\textsuperscript{2}. To reconstruct the lineage tree, we selected mosaic variants
shared by clones or by multiple bulk tissues [6] (Fig 2). Analysis of bulks alone allowed discovering only high frequency mutations but not all. For example, mutations a, b, and c defining branches of the first zygotic cleavage (Fig 2B) could not be discovered because of resembling germline variants by frequency of occurrence in the bulks (i.e., in 80% to 90% of cells). Pairwise comparisons between clones and bulk tissues are powered to find mutations present in the analyzed cell and at low frequency (typically <1% VAF) in bulks but miss high frequency mutations. Remarkably, the All2 approach was able to call both high and low frequency mutations resulting in the most complete lineage tree—a tree that cannot be reconstructed even if we combine comparisons of clones relative to bulk tissues and analysis of bulks.

In this comparison we utilized data from bulk blood and saliva as these samples are easier and cost-effective, as compared to bulks of fibroblasts, to collect for an individual. Also, blood and saliva are made up of multiple early developmental cell lineages [6], while fibroblasts from a biopsy can be dominated by just a few lineages [10]. So, samples from fibroblasts can only
have a fraction of early mosaic mutations. Our additional analysis of 65 SNVs in the tree using high coverage data suggests that using any of the bulk samples (saliva, blood, or fibroblasts) does not outperform \( \text{ALL}^2 \) for lineage tree construction (S1 Table).

The advantage of calling mosaic mutations in bulk is that it allows discovering mutations with intermediate VAF (between 1% to 10%), which were not sampled by the 25 analyzed clones and consequently, not discovered by \( \text{ALL}^2 \). Increasing the number of analyzed clones will likely increase the overlap in discovered mutations between those two approaches but would also increase experimental cost. Thus, this observation suggests complementarity in analyzing clones/single cells and bulks for lineage reconstruction.

### Allele dropout mode for whole genome amplified single cells

Using clones as gold standard, we applied \( \text{ALL}^2 \) in Allele Dropout Analysis (ADA) mode (see Methods) to MDA amplified single cells, to demonstrate the effectiveness of this mode to filter out spurious calls originating from biases in the amplification process. MDA uses \( \varphi_{29} \) polymerase under isothermal condition, which results in an exponential DNA amplification. The exponential amplification leads to uneven coverage and over representation of one allele over the other (allele imbalance). In extreme, a locus can have only one allele amplified and germline variants on the unamplified locus will be lost. ADA mode is designed to address this issue. In ADA mode, \( \text{ALL}^2 \) takes a list of genomic regions (in BED format) where no allele dropout is observed (see Methods). Using this, for each call, \( \text{ALL}^2 \) excludes from the score calculation those cells where a call is not made, and the surrounding region has allele dropout. This exclusion may change the number of considered cells and pairwise comparisons, which eventually affects the mosaic and germline scores.

We called mosaic mutations in 11 clones (representing 11 brain progenitor cells) derived from a human fetal brain (specimen 316) [5], as well as in an MDA amplified single cell taken from one of the clones. Just adding the single MDA-amplified cell into the analysis, more than doubled the count of high frequency mosaic mutation (Fig 3A and 3B). Next, we applied a single cell specialized caller SCOUT [11]. We observed that it reduced the effect of MDA amplification artifacts; 47% reduction in high frequency calls and 33% reduction in call in the single

---

**Fig 3.** \( \text{ALL}^2 \) in ADA mode reduces false positive calls from allele dropout in MDA. (A) Score distribution when applying \( \text{ALL}^2 \) to 11 clones derived from single brain progenitor cells. There are 29 calls for high frequency (gray area) and 2548 calls for low frequency (green area) mosaic mutations. The ‘C’ points to mosaic calls in the clones). (B) Adding one MDA amplified cell to the analysis results in double the number of calls for high frequency mosaic mutations. Noise also increases. The ‘S’ points to the calls coming from the single cell. (C) Application of a specialized single cell caller SCOUT on the single cell partially mitigates issues with calling, i.e., reduces the noise and the number of mosaic calls. (D) Applying the ADA mode results in almost the same set of high frequency mosaic mutations. The mode also reduced calls for mosaic mutations in single cell without affecting calls in the clones. The color (and size) of the circles corresponds to the number of mutations sharing the same scores as depicted in the colorbar. The mosaic mutations are represented by the light green area, germline mutations are represented in light blue area, high frequency mosaic mutations are represented in the light gray area and noise is represented by the light red area.

https://doi.org/10.1371/journal.pcbi.1009487.g003
Further application of the ADA mode resulted in calling all but one (28 vs 29) high frequency mosaic calls and further 11% reduction in mosaic calls in the single cell (Fig 3D). Additionally, there was a 69% reduction in the germline variants after applying ADA. These variants, falsely called as homozygous reference due to allele dropout in the single cell, are effectively filtered by the ADA mode. Mutation counts per clone (Fig 3D) were also similar to those found when analyzing only clones (Fig 3A). This comparison yields evidence that even though the number of mutations called in the single cell is high, by applying ADA mode, we were able to reduce the number of likely false calls introduced by single cell amplification by half, without compromising the mutation calls from clones not affected by allele dropout. ADA mode is also effective in reducing false positive calls when using data for cells with low (i.e., highly non-uniform) amplification quality, however, that can come with the tradeoff of filtering likely true mosaic mutations (S6 Fig).

**Runtime**

Runtime depends on the number of cells in the study and the variant caller used (since some variant callers will output higher number of calls than others). For the first example with 25 iPSC lines (Fig 2B), application of All² using 8 GB memory on a 2.4 GHz dual-core processor took less than 15 minutes for the ‘score’ module and less than 10 minutes for the ‘call’ module to compute mutation annotation and plot the mutation count and VAF plots. For the second example with 11 clones and one single cell (Fig 3), application of All² in ADA mode took less than 90 minutes to complete the ‘score’ module and less than 20 minutes for the ‘call’ module. In this case, the runtime is longer because of the longer list of variant candidates from the single cell.

**Discussion**

We have developed and implemented All², which can discover mosaic SNVs, indels, and SVs from exhaustive cell-to-cell comparison of WGS data from single cells or clones. Our method is superior to using deep sequencing of bulk tissues and/or paired comparison of single cells versus bulk for detection of both low and high frequency mosaic mutations. A limitation relative to bulk method is that the mutations that are not sampled by the analyzed single cells cannot be discovered. This can be addressed by increasing the number of analyzed single cells. We have also applied All² for comprehensive reconstruction of a developmental lineage tree, showing that All² allows a vastly more comprehensive lineage discovery. Furthermore, the method is general and can be applied to any problem of lineage tracing that relies on the analysis of multiple cells, such as tracing cancer evolution.

We further demonstrate that All² facilitates removal of false positive calls (in ADA mode) from amplified single cells. Additionally, since ADA mode takes a BED file with inclusive regions as input, All² can be applied to the analyses of exome sequencing where a user can provide a file with target regions. The same mode can also be applied to exclude copy number altered regions when analyzing cancer cells. All² provides visualizations such as allele frequency distribution, mutation spectrum, mutation counts and score distribution plots to help guide the user to better understand their data as well as change parameter setting for calling mosaic mutations. The tool is open source and is freely available on GitHub: https://github.com/abyzovlab/All2.

**Methods**

**iPSC line generation**

The iPSC lines were derived from fibroblasts using the Epi5 Episomal iPSC Reprogramming Kit (Invitrogen catalog A15960) delivering the five reprogramming factors Oct4, Sox2, Klf4,
L-Myc, and Lin28. The iPSC lines were propagated using mTeSR1 media (Stem Cell Technologies) on 1X Matrigel-coated dishes (Matrigel). Genomic DNA was extracted at passage six, using QIamp DNA Minikit (Qiagen) following the manufacturer instructions.

Saliva collection and DNA extraction
Saliva DNA was collected and purified using the Oragene-Discover kit (DNA Genotek) following the manufacturer instructions. Saliva DNA was extracted using DNeasy Blood and Tissue kit (Qiagen) with the following modifications: 5 ml AL-buffer and 200 μl Proteinase K were added to saliva and incubated at 5600B0030C for 30 minutes. RNA was digested using 2003B0043l RNAse A (Qiagen) for 5 minutes and DNA was extracted using 4 extraction columns in parallel to optimize the yield.

Blood collection and DNA extraction
10–15 ml of blood was collected using BD Vacutainer ACD tubes. DNA was extracted using the Gentra Puregene Blood Kit (Qiagen) following standard manufacturer protocols.

Whole genome sequencing (WGS)
DNA extracted from iPSC lines were sequenced at 30X, while DNA extracted from saliva and blood was sequenced at 200X. All sequencing was conducted at BGI using with 2x100 bp paired reads. The sequencing library preparation was PCR-free.

Fetal brain tissue and MDA
Collection of fetal brain tissues for subject 316, derivation of clonal neurosphere lines and sequencing has been previously described [5]. Single cells from a clonal neurosphere line were manually picked using a micropipette under an inverted microscope. Whole genome amplification was performed by multiple displacement amplification (MDA) using the REPLI-g Single Cell Kit (QIAGEN) following the manufacturer recommendations. Genomic DNA was extracted using the DNeasy Blood & Tissue Kit (QIAGEN). Multiplex PCR for four arbitrary loci from different chromosomes was used to exclude single cells if less than four loci were amplified [12]. Five out of eight single cells (62.5%) passed the 4-locus multiplex PCR quality control and were selected for sequencing. Illumina Truseq DNA PCR-free libraries were prepared for the five cells and sequenced on a HiSeq X (2X150 bp) at 30X coverage.

Allele dropout analysis mode
We started with raw FASTQ files which were aligned to the GRCh37 human reference genome using BWA mem version 0.7.10 [13], the BAM files were then realigned and recalibrated using GATK 3.6 [14]. The clones and the single cells were compared to each other using Mutect2 [15], Strelka2 [16] and SCOUT [11]. For the clones, mutations called by both Mutect2 [15] and Strelka2 [16] with depth of 10 or more reads as well as a PASS value by both callers were used as input to All2. For the single cells, mutations called by Mutect2 [15], Strelka2 [16] and SCOUT [11] with a depth of greater than 10 reads and a PASS value from all callers were used. All2 was run four times with four different settings as depicted in Fig 2. Post All2, only mutations which had an allele frequency of 35% or more were considered, to further filter noise introduced during clone amplification, library preparation and sequencing. The allele dropout regions for single cells were calculated using CNVpytor [17], where the entire genome was divided into 5000 base pair bins. For each bin, a likelihood score was calculated using allele frequency of SNPs within the bin. Bins were marked as allele dropout if they satisfied both of the
following conditions: i) at least one heterozygous SNP in the bin had VAF smaller than 0.01 or larger than 0.99 ii) the maximum likelihood VAF within the bin deviated from 0.5 by more than 0.1. Additionally, we marked as dropout neighboring bins of the bin satisfying the above conditions. For each mutation call, only cells with no call (or calls with VAF < 50%) and marked as having a drop out at the corresponding locus were excluded from calculation of score by All$^2$ (S7 Fig). Single cell QC on the MDA amplified cells was performed using Scellector (9) and only one cell (cell5) passed QC (S8 Fig).

**Mutation calling for lineage analyses**

The FASTQ files were processed the same way as the clones above. Calls were made using an allele frequency cut-off of 35% to remove mutations introduced during culturing clones. Additionally, only INDELs shorter than 10bp (most confident calls) were used. Pairwise comparison between bulk data and the clones were done using consensus calls between Mutect2 and Strelka2. Mutations with a depth greater than 10 reads, at least 2 alternate supporting reads, and PASS values from both callers were used. For the allele frequency plots (Fig 2A and 2C), all mutations from All$^2$, bulk, and pairwise comparison were used. For details, including calling mosaic mutation from bulk tissue and lineage tree construction, please refer to the method section of Fasching et al [6].

**Supporting information**

- S1 Fig. Real data introduces noise/missing data that masks mutation type pattern. (PDF)
- S2 Fig. Example of calculating mosaic and germline scores for a mosaic variant. (PDF)
- S3 Fig. Example of calculating mosaic and germline scores for a germline variant. (PDF)
- S4 Fig. Plots generated by All$^2$ ‘call’ command. (PDF)
- S5 Fig. NxN pairwise binary matrices for an exemplar call. (PDF)
- S6 Fig. All$^2$ in ADA mode including 3 single cells (cell1, cell3, and cell5). (PDF)
- S7 Fig. Regions with allelic dropout for single cell. (PDF)
- S8 Fig. Allele frequency distribution of heterozygous germline variants in 5 MDA-amplified cells. (PDF)
- S1 Table. Fraction of mosaic mutations (SNVs) missed using different tissue types. (PDF)

**Author Contributions**

**Conceptualization:** Taejeong Bae, Flora M. Vaccarino, Alexej Abyzov.

**Data curation:** Yeongjun Jang, Liana Fasching, Livia Tomasini, Jessica Mariani.
Formal analysis: Vivekananda Sarangi.

Funding acquisition: Flora M. Vaccarino, Alexej Abyzov.

Investigation: Vivekananda Sarangi, Flora M. Vaccarino, Alexej Abyzov.

Methodology: Vivekananda Sarangi, Taejeong Bae, Shobana Sekar, Alexej Abyzov.

Resources: Yeongjun Jang, Milovan Suvakov, Taejeong Bae, Liana Fasching, Livia Tomasini, Jessica Mariani.

Software: Vivekananda Sarangi.

Supervision: Flora M. Vaccarino, Alexej Abyzov.

Visualization: Vivekananda Sarangi, Milovan Suvakov.

Writing – original draft: Vivekananda Sarangi.

Writing – review & editing: Vivekananda Sarangi, Shobana Sekar, Flora M. Vaccarino, Alexej Abyzov.

References


