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From single-cell to cell-pool transcriptomes: Stochasticity in gene expression and RNA splicing

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Single-cell RNA-seq mammalian transcriptome studies are at an early stage in uncovering cell-to-cell variation in gene expression, transcript processing and editing, and regulatory module activity. Despite great progress recently, substantial challenges remain, including discriminating biological variation from technical noise. Here we apply the SMART-seq single-cell RNA-seq protocol to study the reference lymphoblastoid cell line GM12878. By using spike-in quantification standards, we estimate the absolute number of RNA molecules per cell for each gene and find significant variation in total mRNA content: between 50,000 and 300,000 transcripts per cell. We directly measure technical stochasticity by a pool/split design and find that there are significant differences in expression between individual cells, over and above technical variation. Specific gene coexpression modules were preferentially expressed in subsets of individual cells, including one enriched for mRNA processing and splicing factors. We assess cell-to-cell variation in alternative splicing and allelic bias and report evidence of significant differences in splice site usage that exceed splice variation in the pool/split comparison. Finally, we show that transcriptomes from small pools of 30–100 cells approach the information content and reproducibility of contemporary RNA-seq from large amounts of input material. Together, our results define an experimental and computational path forward for analyzing gene expression in rare cell types and cell states.

Gene expression levels can differ widely between superficially similar cells. One source of variation is stochastic transcriptional “bursting” (Elowitz et al. 2002; Ozbudak et al. 2002; Blake et al. 2003; Raser and O’Shea 2005; Kaufmann and van Oudenaarden 2007). These studies mainly used fluorescent protein fusion genes to monitor the expression of one or a few genes. They revealed dynamic fluctuations through time that are seen as “salt-and-pepper” variation across a cell population at any given time. In addition to this bursting behavior, individual cells are expected to display controlled and coordinated differences in the expression of genes engaged in dynamic physiologic processes, such as cell cycle phase progression, paracrine or autocrine signaling response, or stress response. Beyond such already appreciated heterogeneity lie currently unknown cell-to-cell differences with biological implications for defining cell states, metabolic function, and, in complex tissues, cell identity.

Measuring RNA transcripts in single cells is now done in multiple ways, and similar conclusions about variability are emerging from the higher sensitivity methods. For individual genes, single molecule RNA fluorescence in situ hybridization (SM-RNA FISH) is highly informative (Femino et al. 1998; Raj et al. 2008), and multiplexed versions now enable multiple genes to be measured in parallel (Lubeck and Cai 2012). In principle, an advantage of SM-RNA FISH is the ability to accurately count the absolute number of transcripts in a cell. A second and older approach is multiplexed single-cell RT-qPCR (Cornelison and Wold 1997), which has now been advanced to increasingly high-throughput formats (White et al. 2011; Sanchez-Freire et al. 2012, Livak et al. 2013). It produces semiquantitative relative comparisons between individual cells. However, neither SM-RNA FISH nor the current forms of multiplex RT-qPCR cover the entire transcriptome or have the single-nucleotide resolution needed to study fine-structure features of gene expression such as allele specificity, RNA editing, and alternative splicing.

To address these and other limitations, elegant methods have recently been developed for performing RNA-seq with very small amounts of RNA, down to the level of individual cells. These are broadly referred to as “single-cell RNA-seq” (Tang et al. 2009, 2010, 2011; Ozsolak et al. 2010; Islam et al. 2011; Brouilette et al. 2012; Cann et al. 2012; Hashimshony et al. 2012; Pan et al. 2012; Qiu et al. 2012; Ramsköld et al. 2012). Despite these significant advances, there are substantial shortcomings in these methods, and a robust method for comprehensive and accurate measurement of the transcriptome of a single cell is not yet available.

A particular challenge for single-cell methods is the efficiency and uniformity with which each mRNA is copied into cDNA and ultimately represented in the library. This challenge intersects in crucial ways with transcriptome structure. Specifically, thousands of genes are expressed in the range of 1 to 30 mRNA copies per cell, including many essential mRNAs (for example, key transcription factors) (Zenkusen et al. 2008). Even lower transcript levels, averaging <1 mRNA per cell on the population level, are now being reliably detected by RNA-seq. This raises questions whether very rare RNAs represent background biological noise, or alternatively, are functional in only a small fraction of cells. Single-cell RNA-seq has the potential to address these issues,
but their resolution depends on how faithfully and efficiently RNAs are captured and represented in sequencing libraries (referred to throughout as the “single-molecule capture efficiency,” \(p_{\text{smc}}\)). In addition, the uniformity of transcript coverage in early single-cell RNA-seq protocols has typically been heavily biased toward the 3’ end, which affects both gene expression estimates and the ability to analyze alternative splicing, RNA editing, and allelic bias.

A second major use for single-cell RNA-seq is the transcriptomic characterization of rare cells. The human body consists of hundreds of distinct cell types, plus large numbers of neuronal and transient developmental cell types. Many of these are numerically minor components of complex tissues, making them inaccessible to standard methods relying on large RNA inputs. Isolation of single cells based on the cell surface markers or using micromanipulation coupled with single-cell RNA-seq could fill this gap in complex multicellular organisms. However, the feasibility of this approach also depends on the experimental robustness of single-cell RNA-seq protocols. Alternatively, single-cell resolution may not be absolutely required for this purpose, and small pools of cells may be sufficient to characterize rare cell-type transcriptomes. An open unresolved question is how small such pools can be to adequately meet that goal.

In this study, we address the issues highlighted above. We used the SMART-seq protocol (Ramsköld et al. 2012) to measure the transcriptome of single cells and small cell pools from the GM12878 lymphoblastoid cell line. This line is derived from the NA12878 individual, for which a fully sequenced genome with completely phased heterozygous single nucleotide polymorphisms (SNPs) and indels is available (The 1000 Genomes Project Consortium 2012). GM12878 cells have also been the subject of an extensive functional genomic characterization by the ENCODE Consortium (The ENCODE Project Consortium 2011, 2012) and have been used in prior population-level studies of allele-biased gene expression and transcription factor occupancy (Rozowsky et al. 2011; Reddy et al. 2012).

Using spike-in quantification standards of known abundance (Mortazavi et al. 2008), we derive estimates for the absolute number of transcript copies for each gene in each cell and directly measure the average value of \(p_{\text{smc}}\) - “Pool/split” experiments (consisting of pooling RNA from multiple single cells, splitting the pool into the same number of separate reactions and building libraries from them) allowed us to measure the extent of and control for technical variation. We find that the \(p_{\text{smc}}\) value is quite low: \(\approx 0.1\). An analysis framework accounting for technical stochasticity is described and used to assess variability in gene expression, allelic bias, and alternative splicing between single cells. Distinct from prior studies, our approach allowed us to parse findings into those that are just as likely to be due to technical origins and those that are more likely to be of biological interest.

We report evidence of significant variability in the total number of mRNA molecules per cell, and identify biologically coherent modules of coexpressed genes specifically expressed in individual cells or groups of cells. These include expected variation associated with cell cycle phases, and an unexpected module enriched for mRNA processing and splicing genes. We observe evidence of higher levels of autosomal allelic exclusion on the single-cell level, potentially associated with transcription bursts; however, it is at present difficult to confidently distinguish from technical variability. In contrast, we find much stronger evidence for widespread major splice site usage switches between individual cells. Finally, our analysis of similarly constructed small cell pools (30–100 cells) reveals a high robustness and reproducibility, approaching that of bulk RNA measurements. This presents a reliable path forward toward the future comprehensive transcriptomic characterization of rare cell types.

Results

In silico examination of major variables affecting informativeness of single-cell and small cell-pool RNA-seq

We began this study with two goals: first, to study gene expression heterogeneity in GM12878 cells on the single-cell level, and second, to determine the minimal optimal size of a cell pool that is informative of the characteristics of the larger cell population, with the goal of applying that approach to rare cell types in future studies. How well these goals are achieved depends on several parameters affecting biological and technical stochasticity and detection sensitivity, the values of which were unknown. To understand their influence, we carried out a simulation of single-cell and cell-pool transcriptomes (see Supplemental Methods for details) by varying the following parameters:

1. Single-molecule capture efficiency \(p_{\text{smc}}\). In contrast to bulk RNA-seq libraries, an individual cell contains a very limited total number of mRNA molecules. Individual genes can be present in single-digit transcript numbers. If only a fraction of mRNAs are successfully represented in a library, a technical stochasticity component is introduced. Depending on its magnitude, data interpretability can be significantly affected due to false negatives and a distortion of relative gene abundance estimates. The \(p_{\text{smc}}\) parameter is the probability that any given original RNA molecule is captured in the final library. We examined the effect on expression quantification of \(p_{\text{smc}}\) ranging from 0.01 to 1.

2. Total number of mRNA molecules per cell. The impact of low \(p_{\text{smc}}\) on expression measurements will be more severe if fewer mRNA molecules are present in a cell. The average total number of mRNA molecules in a single cell is not known for most cell types, but it is expected to vary with cell size, metabolic status, and even cell cycle phase. This means that single-cell expression measurements in some cell types are likely to be more robust to technical noise than in others. We varied the total number of mRNAs from 50,000 to 1,000,000 (while keeping the number of genes expressed constant).

3. Frequency of expression of individual genes in single cells. From prior studies we expect that some genes will be expressed in all or most cells, while others will be expressed in only a subset of cells. Genes detected at lower levels in bulk RNA-seq are the most obvious candidates to be expressed in a subset of cells in a population, although we do not know what fraction of low-abundance RNAs behave in such a way. This is particularly relevant to cell pools: a gene expressed at 50 copies per cell but only in 10% of cells would still be stochastically represented in a pool of 10 cells even if \(p_{\text{smc}}\) is high. In the absence of reliable data on this, we modeled the probability of expression in a given single cell with a distribution centered around very high values for genes highly expressed in bulk RNA-seq measurements, and progressively lower values with decreasing expression levels (details in Supplemental Methods).

The simulation results are summarized in Figure 1, A–C and Supplemental Figures 1–25. As expected, low \(p_{\text{smc}}\) has a profoundly negative impact on gene expression quantification accuracy and reliability, leading to frequent false negatives (Fig. 1A; Supplemental
Figure 1. (Legend on next page)
Stochasticity in gene expression and RNA splicing

Fig. 1), and to poor estimates of expression levels. For example, in a single cell with 100,000 mRNAs, $p_{\text{true}} = 0.1$ results in only 40% of genes expressed at 100 FPKM receiving FPKMs within 20% of the true values (Supplemental Fig. 1C), but this fraction rises to nearly 100% if $p_{\text{true}} = 0.8$ (Supplemental Fig. 1G). The quantification of relative expression levels is similarly affected, with only the most highly expressed genes being consistently well-quantified relative to each other at low $p_{\text{true}}$ (Supplemental Figs. 12–25).

In contrast, our simulation results indicate that cell pools are much more robust to technical noise, with 90% of genes expressed at 10 FPKM receiving FPKM estimates within 20% of their true value (Supplemental Fig. 1C) at $p_{\text{true}} = 0.1$ in a pool of 100 cells. They also represent the expression profiles of the general population reasonably well (Supplemental Fig. 1), even at low $p_{\text{true}}$, starting from a size of ~30 cells (10-cell pools seem not to be sufficient to achieve this). Finally, as expected, the larger the number of total mRNA molecules per cell, the greater is the buffer against technical noise, resulting in more robust quantification (Supplemental Figs. 2–11).

Transcriptome measurements of individual single cells and companion pool/splits

The simulation results informed our experimental design, which aimed to gain a firm grasp on technical stochasticity in two ways (Fig. 1D). First, we generated single-cell RNA-seq libraries and in parallel carried out “pool/split” experiments. In a pool/split, multiple cells are pooled and lysed together, then split into the same number of reactions, from which libraries are built. Variation between these libraries should be purely technical (with stochastic splitting possibly playing a role at the low end). Variation observed at similar levels in both single cells and pool/splits cannot be confidently attributed to biological differences, although the stringency of this criterion may cause some true biological variation to be obscured. However, variation above the pool/split level can be identified and ascribed to biological sources with high confidence.

We generated single-cell RNA-seq libraries from 15 single GM12878 cells and from two pairs of 10-cell pool/split experiments. We also sequenced replicates of pools of multiple cells (10, 30, and 100 cells), as well as 100-pg and 10-ng samples of bulk RNA (corresponding to ~10 and ~1000 cells), to assess the stability of measurements as a function of the amount of starting material.

We used the SMART-seq protocol (Supplemental Fig. 12; Ramskold et al. 2012) to generate our libraries. A detailed description of the protocol, as we implemented it, is presented in Supplemental Methods. We obtained nearly uniform full-length transcript coverage (Fig. 1E; Supplemental Fig. 29). Uniformity of coverage, which depends on the intactness of RNAs and the successful copying of full-length molecules, is highly desirable for several reasons. First, RNA-seq data quantification using the RPKM/FPKM metric (Mortazavi et al. 2008; Trapnell et al. 2010) makes an implicit assumption of full coverage. Second, it enables the analysis of alternative splicing and allelic bias, as read coverage of 5′-proximal splice sites and heterozygous positions is ensured.

We added spike-in quantification standards of known abundance (in absolute number of RNA copies) (Supplemental Table 2) at the very beginning of cDNA synthesis. This allows us to, first, estimate $p_{\text{true}}$ and second, derive gene expression estimates in absolute numbers of copies per cell. The latter is important because while FPKM is useful for comparing expression levels within a library, it can only be used to compare directly across different libraries when the total amount of RNA in each starting sample is roughly the same (Anders and Huber 2010). This assumption is usually only mildly violated when working with bulk samples, but when single cells are compared, it becomes significantly more problematic as the variation in the total amount of RNA in each cell is expected to be much larger.

Figures 1 and 2 summarize the technical characterization of the SMART-seq protocol applied to GM12878 cells. In addition to the mostly complete coverage along transcript length, sequencing libraries were also highly enriched for exonic sequences (Supplemental Fig. 28), indicating a high efficiency of enrichment for polyadenylated molecules.

Gene detection in single cells versus pools of varied sizes

We compared single-cell and pool/split libraries, as well as cell pools, with bulk RNA samples from GM12878 cells (Fig. 1F). In bulk RNA libraries, we detect about 12,000 genes expressed at more than 0.1 FPKM. A lower number of genes, between 4000 and 5000, is detected in both single-cell and pool/split libraries. These differences between single cells and bulk libraries are due mostly to genes expressed at low levels. Genes expressed at more than 100 FPKM in 10-ng bulk RNA samples are detected in almost all libraries, while only ~30% of genes expressed at ~10 FPKM and 10% of genes expressed at ~1 FPKM were detected in any given single-cell library (Fig. 1G). Notably, the number of genes detected in both 100-cell and 30-cell pools was similar to that detected in the 10-ng libraries (~11,000). In contrast, in the 10-cell pools and 100-pg libraries, lower numbers of genes were detected, between

Figure 1. Simulated and measured transcriptome profiles from individual cells and small cell pools. (A) Number of detected genes in simulated data sets as a function of the number of cells pooled and the single molecule capture efficiency ($p_{\text{true}}$) (assuming 100,000 mRNA molecules per cell). See Supplemental Figure 1 for full details. (B, C) Accuracy of gene expression estimation as a function of the number of cells pooled and the single molecule capture efficiency; $p_{\text{true}} = 0.1$ in B and $p_{\text{true}} = 0.8$ in C. 100,000 mRNA molecules per cell assumed. Shown is the fraction of genes at the indicated expression levels in FPKM, whose estimated expression level in FPKM in simulated libraries was within 20% of their true value, after modeling the stochasticity due to the single-molecule capture efficiency of the library-building protocol. See the Methods section and Supplemental Figures 2–11 for full details. Note that the simulation is intended to illuminate the relative effects of the various parameters studied, and the absolute numbers of genes should not be directly compared to the real-life data shown in Fig. 1G. (D) Experimental design. Single cells are combined with spike-in quantification standards and SMART-seq libraries are generated. In parallel, multiple single cells are pooled together and combined with spikes, then lysed and split into the same number of reactions and converted into SMART-seq libraries. Libraries are then sequenced, data processed computationally, and estimates for the absolute number of copies per cell are derived based on the spikes. Variation in pool/split experiments is due to technical stochasticity, while variation in single-cell libraries is a combination of biological variation and technical noise. (E) Uniformity of transcript coverage. Shown is the average coverage along the length of an mRNA for single cells and pool/split experiments. Only mRNAs longer than 1 kb from genes with a single annotated isoform in the RefSeq annotation set were included. See Supplemental Figure 29 for more details. (F) Number of detected protein-coding genes for libraries built from 10 ng and 100 pg of poly(A)+ RNA, pools of 100, 30, and 10 cells, representative pool/split experiments (individually and summed across all libraries), and representative single cells (individually and summed across all libraries). (G) Fraction of genes from 100-ng bulk poly(A)+ RNA libraries that were detected in pools of 100, 30, or 10 cells, 100 pg of poly(A)+ RNA, pools/split experiments, and single cells. FPKM is shown on the x-axis.
Figure 2. (Legend on next page)
6000 and 7000. This is consistent with simulation results suggesting that 30 cells is the lower limit of cell number at which the transcriptome library complexity begins to approach that of the larger cell population. This is corroborated by the correlation between the expression levels of replicate measurements (Fig. 2A; Supplemental Fig. 50). In contrast, a sizable population of genes present at high levels in one replicate and at very low levels or completely absent in the other appears in 10-cell pools (Fig. 2B) and especially in pool/split libraries (Fig. 2C). Finally, union sets of genes detected in all individual cell libraries and in all pool/split libraries was ~10,000, which was in the range seen for 30- to 100-cell pools.

Pool/splits measure technical variation and reveal biological variation among single cells

The observed variations in gene expression levels and detection can be explained as a combination of some genes not being expressed in each and every cell and low $p_{\text{true}}$ resulting in large numbers of false negatives. We calculated the average $p_{\text{true}}$ across all libraries based on the detection of spike-ins (details in Methods). This number is in our estimates: ~0.1. We also estimate that for GM12878 single cells, one transcript copy corresponds to ~10 FPKM on average. This agrees well with the observation that detection of genes becomes unstable below ~100 FPKM (Fig. 2B,C), which is also consistent with previous observations (Ramsköld et al. 2012).

We next compared expression measurements in single-cell and pool/split libraries. Hierarchical clustering of each group is shown in Figure 2, D and E (with two independent biological replicate pool/split experiments shown in Fig. 2E). The distances between the expression profiles within the same pool/split experiment were significantly smaller than those for individual single cells (branch lengths in Fig. 2D,E), and average correlations between single cells were, accordingly, lower than those between libraries from the same pool/split (Fig. 2F,G; Supplemental Fig. 32). A notable feature of the data is small clusters of genes present at high levels in only one library. These are more prominent in single cells than in pool/splits, yet they are clearly present in all samples. In single cells, this is due to a mixture of stochastic capture effects and real biological variation. In pool/splits, stochastic capture is the predominant source. It is important to note that given the low $p_{\text{true}}$ it is difficult to determine the cause of variation for any given gene. Nevertheless, the major conclusion at the transcriptome level is that there are biological differences between single cells because the technical stochasticity in pool/splits is significantly less than variation across single cells.

Estimating absolute transcript levels in single cells

Absolute transcript counts are the biologically relevant values ideally obtained from a single-cell gene expression profiling experiment because, as discussed above, FPKM is a poor metric for comparing gene expression levels between individual cells if the total amount of RNA varies substantially. We derive transcript number estimates for each gene based on the FPKM values of spike-ins. We observed good agreement between the input number of spike-in RNA copies and the corresponding FPKM values in the final libraries (Supplemental Figs. 30, 31).

We use the transcripts-per-cell estimates for all subsequent analyses. Previous studies have reported that genes can be separated into two distinct groups based on their expression levels—one group expressed at high (>1 FPKM) levels and one at very low (<=1 FPKM) levels (Hebenstreit et al. 2011). We examined the distribution of estimated copies per cell in single cells and in pool/splits (Fig. 3A). We find that in individual cells, most protein-coding genes are expressed at levels between 1 and ~50 copies per cell. The distribution suggests a roughly equal number of genes at each level except for a larger group of transcripts with fractional transcript-per-cell values. Obviously, single-cell determinations are constrained in a way that population level measurements cannot be: One transcript per cell is the minimum nonzero value possible. The lower values likely represent a combination of mapping artifacts (due to high sequence homology of paralogs) and RNAs that were present at low levels to begin with and then poorly represented in the final library (due, for example, to the fragmentation of a single original RNA molecule resulting in artificially low FPKMs as a result of coverage only at the 3’ end). The distribution of estimated copies in pool/split libraries exhibited a more linear decrease in the number of more highly expressed genes, consistent with averaging of variation between cells.

We also examined the distribution of the expression levels of long noncoding RNAs (lncRNAs) (Guttman et al. 2009). Consistent with previous observations (Ramsköld et al. 2009; Guttman et al. 2010; Djebali et al. 2012), lncRNAs have generally much lower expression levels compared to protein-coding genes (Fig. 3B).

We were also able to directly assess the total number of mRNAs present in each cell (Fig. 3C,D). Based on the average mass of RNA in each cell (derived from bulk RNA samples from a known number of cells) and the average length of mRNAs in the human genome, we estimated that each GM12878 cell contains, on average, ~80,000 mRNAs. However, we observed striking cell-to-cell differences in the total transcript number of single cells, with some cells expressing <50,000 mRNAs and others almost 300,000. In contrast, pool/split experiments exhibited remarkable uniformity (between 50,000 and 100,000 transcripts) and agree well with prior expectations. It is therefore unlikely that the observed cell-to-cell variability is due to technical noise.

Because transcriptional regulators play a crucial role in defining the gene expression state of cells, we examined the expression of several well-known general transcription factors as well as major regulators of B-cell differentiation (Fig. 3E). Remarkably, except for IRF4, which was usually expressed at several dozen copies, most factors were detected at <10 copies per cell, and were often not detected at all. We stress that this does not mean that they are not expressed. Given the 10% $p_{\text{true}}$ of the protocol, these...
Figure 3. (Legend on next page)
observations are consistent with simple technical failure to detect them. It is also possible that there are no mRNA copies in some cells at the moment of harvest, especially if they are infrequently transcribed. Extending these observations to other functional groups, we assessed proteins involved in translation (as a major group of genes with housekeeping functions) (Fig. 3F), splicing regulators (Fig. 3G), and all transcription factors (Fig. 3H). The median number of copies per cell was ~100 for translation proteins, ~10 for splicing regulators, and strikingly, only ~3 for transcription factors. Beyond their biological interest, these large expression differences between functional gene categories mean that quantification is inherently less robust and less informative for some biological functions than it is for others.

Identification of modules of coexpressed genes

Cell-to-cell gene expression variability may occur on the level of individual genes, but it can also occur in a coordinated fashion. A well-studied example is cell cycle phase-specific gene expression. In an asynchronous culture, groups of genes expressed highly at specific times during the cell cycle should be present in a fraction of cells that is roughly proportional to the time cells spend in each identified phase. Population data do not, however, predict that most cells will be in a “pure” phase state nor that they will express phase-class genes at peak levels.

To test whether we are able to identify cell cycle-associated variation, and to search for any novel functional modules, we carried out weighted gene coexpression network analysis (WCGNA) (Zhang and Horvath 2005) using the copies per cell estimates for single cells and removing genes that were highly variant in pool/split libraries in order to minimize technical noise (see Methods; Supplemental Figs. 33, 34). We identified 19 coexpression modules containing ≥10 genes each (Supplemental Fig. 35). The expression patterns of these modules were mostly well-differentiated among single cells and were absent from pool/split libraries (Fig. 4B; Supplemental Fig. 34).

We then determined the Gene Ontology (GO) category enrichment of each module. The largest module (module 1) was highly enriched for GO categories relating to housekeeping and anabolic gene functions (Table 1; Supplemental Table 3). This included some enrichment for the G1- and S-phase GO terms, and also contained most genes that are generally highly expressed (Fig. 4A). Module 6 was enriched for genes involved in the M phase of the cell cycle. A single cell from the sample of 15 showed strong coordinated expression of genes from the M-phase GO categories enriched in this module. Transcripts from these M-phase genes were not similarly coordinated in other individual cells or in pool/split samples. We measured the fraction of unsynchronized GM12878 cells in the G0 + G1, S, and M phases of the cell cycle using flow cytometry (Fig. 4B). About 14% of cells were in M phase, and the probability of capturing exactly one such cell out of 15 is 0.25; that is, these observations are consistent with this cell being in the peak of M phase.

A more surprising observation was that the second largest module (module 2) was enriched for genes involved in splicing and mRNA processing. It is driven by an individual cell and two additional cells with a somewhat similar expression profile. The signature cell, however, was not an outlier when splice site usage patterns were compared between individual cells (data not shown). A simple interpretation of these observations is a general up-regulation of splice and mRNA processing factors in that cell that does not result in a distinctive alternative splicing program.

Module 3 was enriched for metabolic cofactor and iron-sulfur cluster binding proteins, including proteins involved in mitochondrial respiratory chains. This is an intriguing observation, as module 3 was mostly driven by the two cells exhibiting the highest total number of mRNA molecules per cell (Fig. 3C; fourth and fifth columns in clustergram in Fig. 4A), consistent with a generally elevated metabolic state.

We also carried out a mirrored WCGNA analysis in which the pool/splits were treated as single cells and vice versa. We did not observe significant GO enrichment beyond a few trivial terms in the largest modules (Supplemental Fig. 54; Supplemental Table 4). This is in contrast to the much more specific GO enrichment seen in single cells.

In addition to the coexpression analysis, we also examined the relationship between the expression variability of genes and various genomic data about their promoters, including long-range chromatin interactions, DNA methylation status, histone marks, transcription start site sequence elements, and CpG islands. No robust explanatory correlation was evident (Supplemental Figs. 46–50), and we expect that data with less technical stochasticity will be needed to illuminate relationships of this kind.

Allele-biased expression at the single-cell level

Allele-specific gene expression (either monoallelic or highly biased toward a monoallelic allele) has been previously reported to be widespread (Gimelbrant et al. 2007; Zhang and Borevitz 2009; McManus et al. 2010; Pickrell et al. 2010; Rozowsky et al. 2011; Reddy et al. 2012). An intriguing phenomenon observed for hundreds of genes in clonal lymphoblastoid cell lines (Gimelbrant et al. 2007; Chess 2012) is the random monoallelic expression of autosomal genes. However, those studies were conducted on large pools of cells, producing a snapshot of average allelic bias in the population, and leaving open the possibility that monoallelic expression is even more widespread on the single-cell level.

GM12878 cells are a good system for addressing this issue, as the fully phased heterozygous NA12878 genome sequence is available (The 1000 Genomes Project Consortium 2012). We aligned
RNA-seq reads in an allele-specific manner to the heterozygous GM12878 transcriptome and calculated allelic bias for each gene as the fraction of reads mapping to the maternal allele. As detailed in the Methods and Supplemental Methods, we applied very stringent criteria for determining statistically significant allele-biased expression events based on the absolute transcript number estimates and taking into account the challenges presented by the nature of single-cell RNA-seq data. We observed good reproducibility of allelic bias profiles in 10-ng bulk RNA libraries (Supplemental Fig. 37A), with most genes being expressed from both alleles (Supplemental Fig. 37D). Allelic bias was also highly reproducible in 30-cell and 100-cell pools (Supplemental Fig. 51). In contrast, allelic bias profiles of single cells correlated poorly with each other, and a large fraction of genes were apparently monoallelically expressed from different alleles in different cells (Supplemental Fig. 37B). The majority of highly expressed genes (≥100 copies per cell) exhibited biallelic expression, while most genes at low expression levels were measured as monoallelically expressed (Supplemental Fig. 37F). We then compared allelic bias variability for individual genes across individual single cells, focusing only on cells in which statistically significant allelic bias was observed, and observed frequent “switching” between the two alleles (Supplemental Figs. 37G, 38A).

These observations can be explained as a combination of biological and technical factors. First, it has been previously reported that allelic bias at the population level is more common among genes expressed at low levels (Gimelbrant et al. 2007, Reddy et al. 2012). A second explanation is the phenomenon of “transcriptional bursting” (Raj and van Oudenaarden 2008; Dar et al. 2012). A single transcription burst produces several mRNA molecules from a single allele. If all mRNAs from a gene in a given cell at a given moment are the product of one or a linked series of such bursts, all
copies would originate from one allele. Finally, stochastic effects due to the low single-molecule capture efficiency of the protocol undoubtedly play a role. The fewer founder molecules are captured, the more likely it is that they come from only one allele. To help parse these sources of variation, we performed the same analyses on pool/split libraries and observed a broadly similar (although always lower) fraction of genes passing all significance tests for allelic bias (Supplemental Figs. 37C,E, 38). The quantitative trend within the pool/fraction of genes passing all significance tests for allelic bias (Supplemental Fig. 40), suggesting that most novel splices are indeed only present in a small fraction of cells.

We asked how often multiple alternative splice sites are used per cell thresholds applied (Supplemental Fig. 40A). While this result could be greatly affected by \( p_{\text{unc}} \) issues, it was independent of the read and estimated transcript copies per cell thresholds used (Supplemental Fig. 40). We also examined the distribution of unannotated splices across individual single cells and found that the majority were detected in only a single cell, with \(<10\% \) found in two cells, and very few in three or more cells (Fig. 5B). This number is certainly an underestimate, given the low \( p_{\text{unc}} \). About 35\% of novel junctions connected two annotated exons (Fig. 5A); most of these represent novel exon skipping events. In another 60\%, the unannotated donor or acceptor site was internal to the gene. These were concentrated close to already annotated splice sites (Supplemental Fig. 40B,C). In particular, novel acceptor sites peaked at +3 and −3 position downstream from annotated sites representing mostly instances of NAGNAG tandem acceptor sites (Hiller et al. 2004; Bradley et al. 2012). Novel 5′ donor sites were fewer in number and peaked at +4 and −4 positions relative to annotated donor sites, thus shifting the coding frame of the transcript. This is a phenomenon we have previously also observed in bulk RNA-seq data (observations of the present study’s authors), the significance of which is at present not clear. The proportions observed were independent of the read coverage and estimated number of copies per cell thresholds applied (Supplemental Fig. 40A).

We quantified alternative splicing using the intron-centric splice inclusion \( \psi \) score approach (Pervouchine et al. 2013). Details of our mapping and analysis pipeline are described in the Supplemental Methods. For reasons given there, we adopted a conservative approach and only analyzed novel splice junctions for which at least one of the donor or acceptor sites has already been annotated in GENCODE v13 (Harrow et al. 2012), thus avoiding library-building artifacts.

We detected between 200 and 2000 novel splice junctions satisfying these criteria in each individual cell (Supplemental Fig. 43). This number is certainly an underestimate, given the low \( p_{\text{unc}} \). About 35\% of novel junctions connected two annotated exons (Fig. 5A); most of these represent novel exon skipping events. In another 60\%, the unannotated donor or acceptor site was internal to the gene. These were concentrated close to already annotated splice sites (Supplemental Fig. 40B,C). In particular, novel acceptor sites peaked at +3 and −3 position downstream from annotated sites representing mostly instances of NAGNAG tandem acceptor sites (Hiller et al. 2004; Bradley et al. 2012). Novel 5′ donor sites were fewer in number and peaked at +4 and −4 positions relative to annotated donor sites, thus shifting the coding frame of the transcript. This is a phenomenon we have previously also observed in bulk RNA-seq data (observations of the present study’s authors), the significance of which is at present not clear. The proportions observed were independent of the read coverage and estimated number of copies per cell thresholds applied (Supplemental Fig. 40A). We also examined the distribution of unannotated splices across individual single cells and found that the majority were detected in only a single cell, with \(<10\% \) found in two cells, and very few in three or more cells (Fig. 5B). While this result could be greatly affected by \( p_{\text{unc}} \) issues, it was independent of the read and estimated transcript copies per cell thresholds used (Supplemental Fig. 40). We also examined the distribution of unannotated splices across individual single cells and found that the majority were detected in only a single cell, with \(<10\% \) found in two cells, and very few in three or more cells (Fig. 5B). While this result could be greatly affected by \( p_{\text{unc}} \) issues, it was independent of the read and estimated transcript copies per cell thresholds used (Supplemental Fig. 40).

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Figure 5. (Legend on next page)
a significant proportion of novel splices had $\psi$ scores of 1 in single cells. This was true, however, only for genes expressed at lower levels ($\leq 50$ copies), where psmc artifacts are a likely cause. In contrast, in highly expressed genes, no novel junctions received a dominant ($\geq 0.5$) $\psi$ score. However, the scores were still consistently higher than what is observed for novel splices in bulk RNA samples.

Finally, we evaluated the consistency of splice site usage between individual cells. We applied a statistical framework similar to the one used to analyze allelic bias and derived a list of dominant splice junctions in each cell, taking into account the estimated absolute number of copies and the stochastic capture effects. We asked how often the dominant splice site changes between different cells. We found 282 such genes in single cells, suggesting the phenomenon may be widespread. The genes involved were enriched for ribosomal and translation proteins, and also, intriguingly, for proteins involved in RNA splicing and processing (Supplemental Table 6). We tested this single-cell variation against pool/split experiments, in which we found very few genes with different dominant splice sites across libraries. (Fig. 5E; Supplemental Fig. 44). This argues that much alternative splicing variation is in fact due to biological differences between cells, and is in agreement with the bimodality of splicing in individual mouse immune cells reported recently (Shalek et al. 2013).

Discussion

Two major goals for single-cell RNA-seq are to obtain high-resolution transcriptomes for rare cell types or states and to measure the differences in RNA expression and processing between individual cells. Here, we showed that the first goal can be achieved by studying 30- to 100-cell pool samples even in the absence of perfect capture of each and every individual RNA molecule. Our conclusion is consistent with independent 80-cell measurements (Ramskold et al. 2012). The pools reproduce the expression profiles (Supplemental Fig. 53) and allelic-bias patterns (Supplemental Fig. 51) of the larger population, and similar numbers of genes and splice junctions are detected in them (Supplemental Figs. 52, 53). The approach is applicable to cells collected by laser-capture (to be supplemented for future studies). We therefore consider this a provisional result in need of further investigation with improved experimental protocols. The observed frequency of major splice switching in single cells is a stronger effect, and based on comparison with pool/split experiments, it is unlikely to be the sole result of technical stochasticity. It has also been independently reported in a different system (Shalek et al. 2013).

Transcriptional bursting suggests an attractive biological explanation for these observations. If a gene is expressed in a series of infrequent (relative to the half life of its mRNAs) such bursts, at any given time the population of mRNAs in the cell is likely to originate from only one allele. Such bursting could also be the source of cell-to-cell variation in alternative splicing. It is possible that the same set of factors influencing splice-site choice maintain physical association with the gene during a transcriptional burst, leading to a particular splicing pattern being highly favored locally in space and time, even if factors supporting a different splice choice are present within the same nucleus. Alternatively, isoform choice could be driven by temporal switching of factors and would operate regardless of bursting behavior. These are testable alternatives for future studies.

Many specific biological processes, especially regulatory ones, involve genes whose transcript levels are in the range highly affected by technical variation, as shown by our survey of transcription factors. While measurements with current methods can give some important clues about coherent biological variation, especially when large numbers of individual cells are assayed, our results argue that considerable improvement in the single-

Figure 5. Alternative splicing at the single-cell level. (A) Classification of new junctions connecting known splice sites. (B) Frequency of detection of novel splice junctions. Novel junctions for which neither the donor nor acceptor site has been annotated were excluded for reasons described in the main text in both A and B. A threshold of 10 estimated copies and a coverage of 10 reads was applied, but results are essentially the same, independent of the thresholds used (Supplemental Fig. 40A). (C) Distribution of $\psi$ scores in bulk RNA samples for annotated and novel splice junctions. A threshold of 15 reads combined for all splice junctions in which a donor or acceptor site participates was applied. Note that for each $\psi_0$ score there is at least one matching $\psi_0 \leq 0.1$ – $\psi_0$ score corresponding to the other alternative junction; in some cases, more than two alternative donor or acceptor sites exist; thus the relative height of the $0 \leq \psi \leq 0.1$ bar. (D, upper and lower). Distribution of $\psi$ scores for annotated splice junctions at two different detection thresholds in single-cell libraries (see Supplemental Fig. 41 for more detail). (E, upper and lower) Distribution of $\psi$ scores for novel splice junctions at two different detection thresholds in single-cell libraries (see Supplemental Fig. 42 for more detail). (F) Frequency of major splice site usage switches between individual cells (F) and individual libraries in a pool/split experiment (G). Note the strong support for major splice site use switching across the collection of single cells.
molecule capture efficiency would profoundly advance the field. Based on our simulations and results from pool/split experiments, we estimate that an increase in \( p_{\text{muc}} \) from 0.1 to 0.5 would be a major leap forward, while \( p_{\text{muc}} \geq 0.8 \) would provide sufficient reliability for virtually any biological use. We anticipate that this empirical and analytical framework will be useful for evaluating future improvements in protocols, such as the recently described SMART-seq2 protocol (Picelli et al. 2013).

Finally, we found that the amount of mRNA per cell is highly variable between individual cells. Beyond biological interest, these differences in mRNA number are important for analysis pipelines. RPKM-type metrics are not reliable when there are large differences in total RNA per cell (Lin et al. 2012; Lovén et al. 2012). At present, the direct relationship between the absolute number of mRNA copies per cell and the number of sequencing reads in a library is lost due to the fragmentation of amplified cDNA molecules that is a common feature of available protocols, resulting in multiple distinct but overlapping sequencing fragments for each founder RNA molecule. mRNA copy number can be estimated back from FPKMs with the help of spike-in sequences. However, this is far from ideal, as it depends on the accuracy of quantification of the spike-ins and assumes the absence of systemic differences between spike-in RNAs and endogenous RNAs. If these assumptions are wrong, we expect a systematic error in the calculated number of mRNAs per cell, although the more interesting and important differences between individual cells versus pool/splits would remain. The above considerations make it very clear that a future ideal single-cell RNA-seq assay would combine a very high single-molecule capture efficiency with an amplification-free, and preferably also reverse transcription-free, direct RNA sequencing method to achieve direct counting of intact transcripts. Emerging sequencing technologies (Branton et al. 2008; Schadt et al. 2010) already hold promise for such radical improvements.

Methods

Cell growth and single-cell RNA-seq library construction

Individual GM12878 cells grown according to standard ENCODE protocols were picked with a glass micropipette, deposited into lysis buffer, and frozen. Cells were later lysed in reaction buffer, and single-cell SMART cDNA was generated following the SMART-seq protocol (Ramsköld et al. 2012) with the following modifications: (1) Carrier yeast tRNA was added in the lysis buffer to reduce handling losses and help maintain the integrity of the mRNA; (2) spikes of known copy number were introduced; and (3) the PCR cycle number was empirically titrated to accommodate the relatively small GM12878 cells. The SMART cDNA was tagedmented using Illumina/Nextera reagents as described in Gertz et al. (2012). A detailed description of experimental protocols is provided in the Supplemental Methods.

Sequence alignment and gene expression quantification

Reads were aligned against a combined Bowtie index of the human genome and spike-in sequences using TopHat (Trapnell et al. 2009, 2012). Gene expression was quantified using Cufflinks (Trapnell et al. 2013). Gene expression was measured by accounting for the fact that the number of all transcripts is not known but only the number of complete failures; a detailed description of the procedure is provided in the Supplemental Methods). The average \( p_{\text{muc}} \) for all spikes for which libraries with \( 0 \) FPKMs were observed was used, which is \( -0.01 \).

Analysis of allele-biased expression

We used the diploid (May 2011 release) NA12878 genome containing phased SNPs and indels based on the NCBI build 36 (hg18) version of the human genome (downloaded from http://sw. gersteinlab.org/NA12878_diploid/). Heterozygous transcriptomes containing two copies of each transcript were built, and reads were aligned using Bowtie (Langmead et al. 2009) (version 0.12.7) with zero mismatches allowed. Identical reads were collapsed, and reads were assigned to an allele if they mapped only to one of the alleles of a gene. Allele-biased expression was assessed by accounting for all of the following: (1) significance of allelic bias on the level of reads; (2) significance of allelic bias on the level of estimated copies per cell for each allele (derived from the total number of estimated copies for the gene); this is necessary, as a common feature of all current single-cell protocols is the production of multiple overlapping fragments from each original molecule; and (3) the possibility that the observed allelic bias is due to differential stochastic capture of the two alleles. A detailed description of the procedure is provided in the Supplemental Methods.

Alternative splicing analysis

We carried out alternative splicing analysis using the 5’ and 3’ splicing inclusion scores described by Pervouchine et al. (2013), and applying the same statistical procedure we used to assess allelic expression bias to determine statistically significant splice variant exclusion. A detailed description of the splicing analysis procedure is provided in the Supplemental Methods.

Gene expression clustering and weighted correlation network analysis

We used the WGCNA R package (Langfelder and Horvath 2008) to carry out the weighted correlation network analysis. Gene Ontology enrichment in modules was assessed using FuncAssociate2.0 (Berriz et al. 2009). Gene expression clustering was carried out using Cluster 3.0 (de Hoon et al. 2004) and visualized using TreeView (Saldanha 2004).

Data access

BAM files containing aligned and unaligned sequencing reads have been submitted to the NCBI Gene Expression Omnibus (GEO; http://www.ncbi.nlm.nih.gov/geo/) under accession number GSE44618.

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