A hyperactive transcriptional state marks genome reactivation at the mitosis–G1 transition

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During mitosis, RNA polymerase II (Pol II) and many transcription factors dissociate from chromatin, and transcription ceases globally. Transcription is known to restart in bulk by telophase, but whether de novo transcription at the mitosis–G1 transition is in any way distinct from later in interphase remains unknown. We tracked Pol II occupancy genome-wide in mammalian cells progressing from mitosis through late G1. Unexpectedly, during the earliest rounds of transcription at the mitosis–G1 transition, ∼50% of active genes and distal enhancers exhibit a spike in transcription, exceeding levels observed later in G1 phase. Enhancer–promoter chromatin contacts are depleted during mitosis and restored rapidly upon G1 entry but do not spike. Of the chromatin-associated features examined, histone H3 Lys27 acetylation levels at individual loci in mitosis best predict the mitosis–G1 transcriptional spike. Single-molecule RNA imaging supports that the mitosis–G1 transcriptional spike can constitute the maximum transcriptional activity per DNA copy throughout the cell division cycle. The transcriptional spike occurs heterogeneously and propagates to cell-to-cell differences in mature mRNA expression. Our results raise the possibility that passage through the mitosis–G1 transition might predispose cells to diverge in gene expression states.

[Keywords: chromatin; epigenetics; mitosis; transcription]

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Mitosis is accompanied by a dramatic interruption of nuclear processes. In metazoans, the nucleus is disassembled, and bulk RNA synthesis ceases (Prescott and Bender 1962). RNA polymerase II (Pol II) and other components of the eukaryotic transcriptional machinery dissociate from chromatin (Gottesfeld and Forbes 1997; Akoulitchev and Reinberg 1998; Prasanth et al. 2003) in part due to mitosis-specific post-translational modifications (Gottesfeld and Forbes 1997, Rizkallah et al. 2011). By late telophase, Pol II is known to re-enter the newborn nuclei in bulk and restore global RNA synthesis (Prasanth et al. 2003). However, we lack general principles of how individual genes reactivate transcription at the mitosis–G1 transition.

Many other interphase nuclear processes are also altered globally to varying extents during mitosis. Studies have described such alterations for the recruitment of transcriptional regulators (Raff et al. 1994; Martínez-Balbás et al. 1995; Dey et al. 2000; Christova and Oelgeschläger 2001; Kruhlak et al. 2001; Zaidi et al. 2003; Young et al. 2007, Yang et al. 2008, 2013; Blobel et al. 2009, 2012, Caravaca et al. 2013, Poleshko et al. 2013; Lake et al. 2014; Lodhi et al. 2014), deposition of histone variants and modifications (Kruhlak et al. 2001; Kelly et al. 2010, Varier et al. 2010, Wang and Higgins 2012), chromatin structure [Kuo et al. 1982, Michelotti et al. 1997, Kelly et al. 2010, Kadauke et al. 2012, Hsiung et al. 2014], long-range genome folding [Naumova et al. 2013, Dileep et al. 2015], lamina-associated genomic domains [Kind et al. 2013], and chromosome territories [Walter 2003]. Details related to the kinetics, order, and fidelity with which such structures and processes are regulated during mitosis is not well understood.
established during the mitosis–G1 transition are largely unknown, except for a few examples for factor localization (Prasanth et al. 2003; Poleshko et al. 2013), lamina-associated domains (Kind et al. 2013), and long-range chromosome interactions (Dileep et al. 2015).

Given these uncertainties in the gene regulatory milieu at the mitosis–G1 transition, might there be altered transcriptional output during this cell cycle phase? A microarray-based study identified ~200 mature mRNAs that fluctuate during early G1 in mammalian cells (Beyrouthy et al. 2008), but it is unknown to what extent changes in transcriptional activity versus post-transcriptional modulation are responsible for these fluctuations. Several studies have directly quantified transcriptional activity over time in cells transitioning from mitosis to interphase [Blobel et al. 2009; Dey et al. 2009; Muramoto et al. 2010; Zhao et al. 2011; Fukuoka et al. 2012; Kadauke et al. 2012; Caravaca et al. 2013] using RT-qPCR of primary transcripts of candidate genes [Blobel et al. 2009; Dey et al. 2009; Fukuoka et al. 2012; Kadauke et al. 2012; Caravaca et al. 2013], live-cell imaging of transcription of act-5 in *Dictyostelium* [Muramoto et al. 2010] and a multiplex reporter locus in a human cell line [Zhao et al. 2011], and microarray-based measurements of nascent transcripts [Fukuoka et al. 2012]. Several of these studies suggest or assume that transcriptional output early after mitosis starts off low and rises monotonically with G1 progression at varying kinetics [Blobel et al. 2009; Zhao et al. 2011; Fukuoka et al. 2012; Kadauke et al. 2012; Caravaca et al. 2013]. However, some genes show nonmonotonic changes in transcriptional output with cell cycle progression after mitosis, but no explanations for these observations have been proposed [Dey et al. 2009; Muramoto et al. 2010; Fukuoka et al. 2012; Caravaca et al. 2013]. It remains unclear which transcriptional pattern represents the general rule, as these previous approaches lacked genome-wide extraction of the most prominent patterns. Moreover, some of these studies are difficult to compare due to incongruities in their temporal coverage of transcriptional measurements and did not define a clear time frame for the occurrence of the first transcriptional cycle at the mitosis–G1 transition. Major questions remain unresolved. Genome-wide, when does de novo transcription upon reversal of mitotic silencing occur? Does the transcriptional program immediately after mitosis deviate significantly from later in interphase, and how might the mitosis–G1 transition influence the fidelity of transcriptional control?

To address these questions, we quantified transcriptional activity from mitosis through G1 phase using three independent methods: chromatin immunoprecipitation (ChIP) combined with high-throughput sequencing (ChIP-seq) of Pol II, RT-qPCR of primary transcripts, and simultaneous imaging of nascent and mature mRNA in single cells by single-molecule RNA fluorescence in situ hybridization (FISH). The temporal and genomic resolution of our strategy enabled visualization of the pioneering round of transcription at many genes upon reversal of mitotic silencing. We found that, during the earliest rounds of transcription, most active genes and intergenic enhancers are transcribed at a higher level than later in G1. This observation counters the prevailing assumption of generally lower initial transcriptional outputs immediately after reversal of mitotic silencing. Notably, the mitosis–G1 transcriptional spike does not scale with the frequency of enhancer–promoter chromatin contacts but is correlated with and preceded by higher levels of histone H3 Lys27 acetylation [H3K27ac] in mitosis. Single-molecule RNA FISH demonstrates that the early G1 transcriptional spike can constitute the maximum transcriptional activity in the entire cell cycle and propagate to cell-to-cell heterogeneity in mature mRNA levels. We discuss potential contributions of the mitosis–G1 spike in transcriptional compensation for changes in DNA copy number in the cell division cycle and as a source of gene expression heterogeneity.

**Results**

**Pol II ChIP-seq on synchronized and purified cell populations reveals the pioneering round of gene transcription at the mitosis–G1 transition**

We performed Pol II ChIP-seq during mitotic exit in murine erythroblast cells (G1E) that lack the hematopoietic transcription factor GATA1 [Weiss et al. 1997]. We used a well-characterized subline (G1E GATA1-ER) that expresses a GATA1-estrogen receptor (ER) fusion protein, enabling study of transcriptional control in the context of estradiol-inducible gene activation and repression [Weiss et al. 1997]. Tracking Pol II occupancy by ChIP-seq during brief cell cycle phases requires isolating a large number of cells specifically from the desired stages [Fig. 1A]. To accomplish this, we arrested G1E GATA1-ER cells [induced with estradiol for 13 h] in prometaphase by nocodazole treatment followed by release into nocodazole-free medium for 40–360 min. To minimize contamination with cells from undesired stages of the cell cycle, we purified cells from specific cell cycle phases at specified time points using a fluorescence-activated cell sorting (FACS) strategy [Fig. 1A]. This approach is based on a reporter [Kadauke et al. 2012] that consists of YFP fused to a mitotic degradation domain (MD), which confers degradation at the metaphase–anaphase transition [live-cell fluorescence microscopy in Supplemental Movie 1; Glotzer et al. 1991; Holloway et al. 1993]. The combination of synchronization coupled with FACS based on YFP-MD and DNA content enabled isolation of populations highly enriched for cells in prometaphase, between anaphase and cytokinesis, in early G1, and in late G1 [Fig. 1A]. One critical benefit of this strategy is that the G1 samples [sorted for 2N DNA content] are devoid of residual mitotic cells (4N) that might be delayed in their release from nocodazole arrest. Such contamination with transcriptionally silent mitotic cells would lead to an underestimate of the early G1 transcriptional activity in an ensemble assay.

We used these synchronized and sorted populations for ChIP-seq of total Pol II in three biological replicates. Examination of individual loci showed that the Pol II ChIP signal is eliminated in prometaphase [Fig. 1B], with
minimal residual signal attributable to contamination of this particular sample by $\sim$10% G2-phase cells [Supplemental Fig. S1], which are also 4N and high YFP-MD. This contamination of the 0-h sample does not affect the subsequent time points in our FACS purification strategy [Fig. 1A]. Our approach enabled the capture of the pioneering round of transcription, which is apparent as a synchronous wave of 5'-to-3' Pol II progression that initiates between anaphase and cytokinesis [4N and low YFP-MD] at 40 min after release [Fig. 1B]. This leading edge of Pol II ChIP signal represents a population-averaged position of the first polymerases to travel down a given gene over time, reaching the 3' end of genes at time points consistent with gene lengths, as shown for illustrative loci in Figure 1B. The partial progression of the Pol II leading edge can be seen for genes >50 kb at individual loci [Fig. 1B] and as a Pol II-binding profile averaged across all such genes [Supplemental Fig. S2]. Shorter genes appear to have already completed the first transcriptional cycle or the first several cycles sometime between the 40- and 60-min time points. Thus, the onset of transcriptional reactivation occurs within a narrow window between anaphase and cytokinesis (40–60 min after release from nocodazole arrest).
In addition to the progression of Pol II along the gene body, the amount of Pol II initiating transcription changes in gene-specific patterns over time. For example, at Chchd3, Zeb2, and Runx1, Pol II occupancy reaches maximum at the 60- to 90-min time points at the 5′ region of these genes followed by a decline through the remainder of G1 phase [Fig. 1B]. We refer specifically to this pattern of a sharp increase followed by sustained decrease as a “spike.” Importantly, at genes with this particular pattern, the increase in Pol II binding at the 5′ region propagates through the full gene length, visible as a spike in occupancy at the 3′ region with a time delay consistent with gene length. The downward-sloping part of the spike indicates that this spike in activity is diminished shortly after the completion of the initial transcriptional cycles. Because the spike in Pol II binding at the 3′ end follows that at the 5′ end in time (exemplified by Chchd3, Zeb2, and Runx1 in Fig. 1B), these Pol II-binding patterns reflect a spike in full-length gene transcription rather than just an increase of paused Pol II at the 5′ end. Not all genes display a transcriptional spike; for example, at Asb1, Pol II binding plateaus after ~90 min of release [Fig. 1B], whereas, at Mavs, Pol II binding rises continuously over a period of 360 min following release [Fig. 1B].

A spike in gene transcription at the mitosis–G1 transition is prevalent across the genome

To examine global distributions of Pol II occupancy over these time points, we measured Pol II occupancy at the 5′ regions of the 4309 nonoverlapping genes with above-background binding in at least one time point, as determined by a peak caller (Zhang et al. 2008). Globally, Pol II binding reaches substantial levels above background even prior to the completion of the first round of transcription for many genes at 60 min after release [Fig. 2A]. In terms of the rise in absolute Pol II binding prior to the 60-min time point, the onset of transcription occurs globally with minimal gene-to-gene differences in kinetics within the limits of our temporal resolution. Pol II binding at the 60- and 90-min time points overall overshoots that of the 360-min time point [Fig. 2A; individual replicates shown in Supplemental Fig. S5]. By 240 min, the distribution returns to roughly the same as 360 min [Fig. 2A]. Thus, contrary to prior expectations of transcriptional reactivation post-mitosis starting off with generally lower initial output, transient transcriptional hyperactivity is a widespread phenomenon associated with the earliest rounds of transcription upon reversal of mitotic silencing.

In general, comparisons of factor occupancy across ChIP-seq samples in the context of global changes in binding require that changes in normalized read counts accurately reflect absolute changes in binding. This important property holds true in our data due to the presence of a relatively large proportion of reads mapping to intergenic regions that represent nonspecific background [Supplemental Fig. S3]. This background serves as an internal calibration across sequencing libraries, enabling inferences of changes in Pol II occupancy on an absolute scale [Supplemental Fig. S3]. We also confirmed patterns observed by Pol II ChIP-seq at individual loci by Pol II ChIP-qPCR [Supplemental Fig. S4], further indicating that sequencing read counts reflect quantitation by qPCR.

While transcriptional hyperactivity upon reversal of mitotic silencing is a prevalent trend, individual genes can exhibit a variety of distinct temporal profiles of transcription, indicating a degree of gene specificity for such patterns [Fig. 1B]. To stratify Pol II-binding patterns at individual genes in an unbiased manner, we performed principal component analysis on Pol II binding at the 5′ region of genes at G1-phase time points (60–360 min). For this analysis, we first normalized Pol II binding at each time point by the sum of Pol II binding across all time points to remove gene-to-gene differences in transcriptional activity unrelated to cell cycle progression. The first principal component accounts for the most (47.2%) gene-to-gene variance and represents temporal shapes that fall along a continuum of early G1 spike versus late G1 up-regulation in Pol II binding [Fig. 2B]. The temporal shapes of individual genes, as defined by the projection onto the first principal component, is highly concordant across the three biological replicates [R = 0.8–0.9] [Supplemental Fig. S6]. Lower-ranking principal components are less clearly distinguishable from noise [Supplemental Fig. S6]. Ranking genes based on the degree of match to the first principal component (projection of each gene onto this principal component) reveals that ~50% of genes exhibit an early spike, 38% of genes exhibit a late plateau, and 12% of genes exhibit late up-regulation in Pol II binding, although these patterns are not discrete clusters [Fig. 2B]. Among the 50% of genes exhibiting some degree of match to the early spike pattern, the magnitude of the spike at the 90-min time point is, on average, 1.4-fold and can reach up to 4.3-fold—higher than the 360-min time point [Supplemental Fig. S7]. Here we refer to the early G1 spike as a trait defined quantitatively by the degree of match in the positive orientation of the first principal component, as shown in Figure 2B.

We found no association between the early G1 spike and the traveling ratio of Pol II, indicating that the occurrence of the early transcriptional spike does not involve a difference in the rate of Pol II promoter escape [Supplemental Fig. S8]. Pol II binding at the 3′ regions of genes often mirrors the temporal shape for the corresponding 5′ regions [Fig. 2B panels on right, and Fig. 2C top row], and very similar principal components were obtained from applying the analysis to Pol II binding at the 3′ region of genes [analysis not shown]. Thus, these temporal changes in Pol II binding reflect full-length gene transcription. Indeed, RT-qPCR of primary transcripts using primers flanking intron–exon junctions for a subset of genes demonstrate that the temporal patterns of Pol II binding at individual loci are well reflected at the level of RNA synthesis [Fig. 2C, bottom row].

The early G1 transcriptional spike pattern encompasses genes with functions general to many cell types (e.g., Tbp) [Fig. 2C] as well as genes involved in developmental regulation relevant to hematopoietic cells, such as Gata2, Myc, Kit, and Runx1 (Fig. 2B,C), with an enrichment for genes in p53 signaling pathways [Supplemental Fig. S9].
Figure 2. A spike in gene transcription is prevalent across the genome at the mitosis–G1 transition. (A) Replicate-averaged Pol II binding at the 5′ 2.5-kb region of 4309 genes active in at least one time point is plotted for each time point against the 360-min time point. Runx1 and Mavs, two genes with distinct temporal patterns shown in Figure 1B, are highlighted. Plots for individual replicates are shown in Supplemental Figure S5B. (B) For the same 4309 genes as in A, we performed principal component analysis on Pol II binding (RPKM), normalized by total Pol II binding for each gene over all time points, at the 5′ 2.5-kb region of each gene. For this analysis, only G1 time points (60, 90, 180, 240, and 360 min) were used. The temporal “shapes” of Pol II binding [eigenvector] of the first principal component is shown. Genes were ranked by their degree of match to (projection onto) the first principal component, and all of their gene-normalized RPKMs at the 5′ 2.5-kb regions were plotted in a heat map for all time points. Threshold for “early spike” is defined as the inflection of projection onto the first principal component from positive to negative. The threshold for separating “late plateau” and “late up-regulated” were chosen manually based on the appearance of the heat map. Four-hundred-thirty-four early spike genes and 432 late up-regulation genes meet the significance threshold ($P < 0.05$, determined by bootstrapping) for their projections onto the first principal component (Supplemental Fig. S5A). The positions of several genes in the heat map are shown at the right, together with their RPKMs for both the 5′ 2.5-kb and 3′ 2.5-kb regions. (C) Pol II ChIP-seq binding profiles (error bars denote SEM; $n = 3$) are shown together with quantification of primary transcripts by RT-qPCR using primers flanking the intron–exon junction (error bars denote SEM; $n = 5$–6). Pol II ChIP profiles from Runx1 and Mavs from Figure 1B are reproduced here for ease of comparison. In addition to these genes, profiles for Gata2, Kit, Hbb-b1, and Hba-a1 are shown in A for Pol II ChIP and in Figure 3D for primary transcript RT-qPCR.
The late up-regulation pattern enriches for gene ontology terms related to plasma membrane proteins [Supplemental Fig. S9] and, of relevance to erythroid biology, includes both $\alpha$-globin ($Hba-a1$) and $\beta$-globin ($Hbb-b1$) genes. Note that the late up-regulation category does not necessarily represent delayed transcriptional reactivation on an absolute level; rather, these include genes that tend to reach similarly high levels of Pol II binding at 60- to 90-min time points and then exhibit further sustained up-regulation through the remainder of $G1$, as exemplified by absolute Pol II binding for $Mavs$ [Fig. 1B] and $Hbb-b1$ [Fig. 2B].

To test whether the early $G1$ spike versus late $G1$ up-regulation patterns are cell type-specific, we performed nocodazole-mediated mitotic arrest-release in a murine embryonal carcinoma cell line [F9] [Alonso et al. 1991] and measured transcription by primary transcript RT-qPCR. Of 13 genes examined that are expressed in both $G1E$ GATA1-ER and F9 cells, 10 showed similar $G1$ transcriptional patterns in both cell types, indicating that the $G1$-phase modulation of transcription that we uncovered in $G1E$ GATA1-ER cells can be found across developmentally distinct murine cell types [Supplemental Fig. S10].

The mitosis–$G1$ transcriptional spike also occurs at intergenic enhancers, but restoration of enhancer–promoter chromatin contacts after mitosis does not spike Pol II. The late up-regulation pattern enriches for gene ontology terms related to plasma membrane proteins [Supplemental Fig. S9] and, of relevance to erythroid biology, includes both $\alpha$-globin ($Hba-a1$) and $\beta$-globin ($Hbb-b1$) genes. Note that the late up-regulation category does not necessarily represent delayed transcriptional reactivation on an absolute level; rather, these include genes that tend to reach similarly high levels of Pol II binding at 60- to 90-min time points and then exhibit further sustained up-regulation through the remainder of $G1$, as exemplified by absolute Pol II binding for $Mavs$ [Fig. 1B] and $Hbb-b1$ [Fig. 2B]. To test whether the early $G1$ spike versus late $G1$ up-regulation patterns are cell type-specific, we performed nocodazole-mediated mitotic arrest-release in a murine embryonal carcinoma cell line [F9] [Alonso et al. 1991] and measured transcription by primary transcript RT-qPCR. Of 13 genes examined that are expressed in both $G1E$ GATA1-ER and F9 cells, 10 showed similar $G1$ transcriptional patterns in both cell types, indicating that the $G1$-phase modulation of transcription that we uncovered in $G1E$ GATA1-ER cells can be found across developmentally distinct murine cell types [Supplemental Fig. S10].
Figure 3. Pol II binding at enhancers, but not enhancer–promoter contacts, also spikes at the mitosis–G1 transition. (A) For 809 intergenic enhancers, we performed principal component analysis on Pol II binding in the same fashion as that detailed for genes in Figure 2B. Shown are the results outlined in a fashion analogous to Figure 2B. For the enhancers that we highlight at the right of the heat map, we also show the raw RPKM of Pol II binding at the nearest gene. (B) Capture-C (a multiplexed derivative of chromosome conformation capture) analysis of chromatin contacts in a nocodazole arrest–release experiment using the Myc promoter as the anchor. The Y-axis of browser tracks are read counts of ligation products normalized to total number of ligation products in the library. We highlight a likely enhancer at the +211-kb region (resides within the transcribed region of noncoding RNA Pvt1, which is omitted in the graph for clarity) and a known enhancer at the +1.7-Mb region (Shi et al. 2013). (C, left) Quantification of Capture-C read densities for the Myc +211-kb and +1.7-Mb enhancers. The Y-axis denotes read counts reflecting ligation products normalized to total number of ligation products in the library. Error bars denote SEM, with $n = 3$ sequencing libraries encompassing two separate ligation libraries and three separate oligo captures. (Right) Pol II ChIP-seq read densities at the Myc gene (5′ 2.5 kb and 3′ 2.5 kb, duplicated from Fig. 2 for ease of comparison) and at the +211-kb and +1.7-Mb enhancers. (D) Quantification of enhancer–promoter contacts using Capture-C anchors at the promoters of early G1 transcriptional spike genes (Cd47 and Kit) and late G1 up-regulated genes (Hba-a1 and Hbb-b1). See Supplemental Figure S14 for browser tracks. These enhancers were described in prior studies: Cd47 −20 kb was described in Dogan et al. (2015), Kit −114 kb was described in Jing et al. (2008) and Lee et al. (2015), the Hba-a1 −34-kb R2 region was described in Hughes et al. (2005), and the Hbb-b1 +32.5-kb locus control region was described in Bender et al. (2000). The Y-axis is the normalized Capture-C contact frequency as described in C. Error bars and the number of replicates are as described in C, except for Hbb-b1, which represents $n = 2$ separate ligation libraries.
Likewise, to carry out H3K27ac ChIP-seq on mitotic cells, we applied a similar procedure using a more robust and cost-effective mitosis-specific antibody [MPM2] to achieve essentially 100% mitotic purity [Campbell et al. 2014]. The mitotic purity of these samples ensures that the data reflect properties of mitotic cells rather than contaminating interphase cells. H3K27ac levels in prometaphase strongly, but imperfectly, correlate with that in asynchronous cells at promoters \( R = 0.72 \) and intergenic enhancers \( R = 0.72 \), indicating a degree of locus specificity in the maintenance of H3K27ac in mitosis (Supplemental Fig. S15). To our knowledge, this is the first report of H3K27ac levels in mitosis measured by ChIP.

For each data set, we quantified read counts at promoters of active genes [4309 genes analyzed in Fig. 2] and intergenic enhancers [809 enhancers analyzed in Fig. 3]. In Figure 4, we show the Pearson correlation coefficient between the read counts for each feature with our measure of early spike versus late up-regulation G1 transcriptional patterns [Pol II ChIP-seq degree of match to the first principal component in Figs. 2, 3], analyzing promoters and intergenic enhancers separately. Among the features examined at promoters, the early G1 transcriptional spike is most strongly correlated with H3K27ac levels in mitosis, including both prometaphase \( R = 0.47 \) and anaphase–telophase \( R = 0.53 \) populations. By comparison, the correlation between the early G1 transcriptional spike and promoter H3K27ac levels in asynchronous cells is less positive \( R = 0.30 \), indicating that mitotic H3K27ac levels specifically provide some additional predictive power. Likewise, for intergenic enhancers, the H3K27ac level in mitosis \( R = 0.35 \) for prometaphase, \( R = 0.46 \) for anaphase–telophase is the most positively correlated with the early G1 transcriptional spike and is less positively correlated for the H3K27ac level in asynchronous cells \( R = 0.12 \) [Fig. 4].

In contrast, the mitotic or asynchronous levels of two features at promoters previously shown to have locus-specific degrees of persistence in mitosis—DNase sensitivity [Hsiung et al. 2014] and GATA1 occupancy [Kadauke et al. 2012]—showed no correlation with the early G1 transcriptional spike. However, this does not rule out that there might be a minimal level of mitotic DNase sensitivity required for the early G1 transcriptional spike because the promoters of virtually all transcriptionally active genes are at least somewhat DNase-sensitive in mitosis.

**Figure 4.** Higher levels of H3K27ac during mitosis predict the mitosis–G1 transcriptional spike at genes and intergenic enhancers. We examined the signal strength of chromatin-associated features in mitotic and/or asynchronous cells for their genome-wide correlation (Pearson R) with the early G1 transcriptional spike, defined in Figure 2B for genes and in Figure 3A for intergenic enhancers as the degree of match to the first principal component from Pol II ChIP-seq. Pol II ChIP-seq was generated in this study, DNase-seq was from Hsiung et al. [2014], GATA1 ChIP-seq was from Kadauke et al. [2012], H3K27ac ChIP-seq was generated in this study, and H3K4me1, H3K4me3, H3K9me3, H3K27me3 were from Wu et al. [2011]. The mitosis H3K27ac ChIP-seq sample consisted of ~100% mitotic purity obtained by FACS for MPM2 positivity as described in Campbell et al. [2014]. Error bars denote 95% confidence intervals.
The early G1 transcriptional spike is also weakly correlated with promoter levels of Pol II binding \((R = 0.26)\), suggesting a mild association with overall levels of transcriptional activity [Fig. 4]. Another active promoter modification, H3K4me3, is not predictive \((R = 0.09)\), whereas levels of repressive modifications H3K27me3 \((R = -0.23)\) and H3K9me \((R = -0.11)\) in asynchronous cells are weakly anti-correlated with the early G1 transcriptional spike [Fig. 4]. At intergenic enhancers, levels of DNase sensitivity, GATA1 occupancy, H3K4me1, H3K4me3, H3K27me3, and H3K9me3 in mitotic and/or asynchronous cells are all either weakly anti-correlated or uncorrelated with the early G1 transcriptional spike [Fig. 4].

We conclude that the H3K27ac level specifically during mitosis exceeds other indicators of active chromatin in its predictive power of the early G1 transcriptional spike at both genes and intergenic enhancers. Since this strongest predictor precedes the mitosis–G1 transcriptional spike, the temporality of the association is consistent with the possibility that mitotic H3K27ac may be involved in causing the mitosis–G1 transcriptional spike.

**The mitosis–G1 transcriptional spike propagates to cell-to-cell heterogeneity in mature mRNA expression**

Our findings thus far demonstrate a spike in transcriptional activity at the mitosis–G1 transition based on measurements of cell population average. Does this transcriptional spike occur in all cells in the population or only a subset of cells, thus potentially contributing to transcriptional heterogeneity among cells? Is the mitosis–G1 transcriptional spike buffered by post-transcriptional regulation, or does the transcriptional spike propagate to elevated mature mRNA levels? To address these questions, we used single-molecule RNA FISH to simultaneously quantify nascent and mature mRNA in single cells by three-dimensional (3D) microscopy [Femino 1998; Raj et al. 2008; Levesque and Raj 2013]. We imaged nascent and mature mRNAs in the same field by hybridizing fixed cells with spectrally distinguishable probes specific to introns or exons of a given gene. While the vast majority of exonic probe signals are from mature mRNAs, colocalized exonic and intronic probe signals are primary transcripts that mark active transcription sites in interphase cells [Fig. 5A]. In prometaphase, cells with condensed chromosomes show no detectable signal in the intronic channel due to mitotic transcriptional silencing, whereas stable mature mRNA molecules that presumably arose prior to mitosis are detectable [Fig. 5A]. Consistent with our Pol II ChIP-seq data, the earliest active transcription appears in cells between anaphase and cytokinesis (4N and low YFP-MD). This RNA synthesis occurs amid chromosomes that are still morphologically condensed, demonstrating that overt condensation does not prohibit gene transcription [Fig. 5A; Supplemental Fig. S16].

We focused on two genes that exhibit transcriptional spikes at the mitosis–G1 transition: Myc and Gata2 [Pol II ChIP patterns for both are in Fig. 2B]. Myc and Gata2 encode for transcription factors that regulate stemness and self-renewal and whose mature mRNA half-lives are relatively short (~15 min to 2 h for Myc [Dani et al. 1984; Watson 1988; Herrick and Ross 1994] and ~2.8 h for Gata2 [Sharova et al. 2009]). Expression of Myc and Gata2 is down-regulated upon exposure to estradiol for 13 h through transcriptional repression by GATA1-ER [Grass et al. 2003; Rylski et al. 2003], and, at this relatively low level of expression, the mature mRNA levels among single cells can vary by >100-fold [Fig. 5C; Supplemental Fig. S19]. The single-cell expression patterns of these genes allow evaluation of the degree to which mitosis–G1 transcriptional spiking of these genes can contribute to heterogeneity in mature mRNA levels.

Transcription visualized by single-cell imaging is known to occur intermittently, with intervals of active RNA synthesis interspersed with periods of inactivity [Golding et al. 2005; Chubb et al. 2006; Raj et al. 2006]. When cells are fixed and viewed as a static image of single-molecule RNA FISH, this pulsatile nature of transcription manifests as a mix of transcriptionally “on” and “off” cells [Fig. 5B; Supplemental Fig. S16]. Single-molecule RNA FISH shows that the early transcriptional spike that we observed as an average across cell populations [Fig. 2B] occurs by a spike in the fraction of cells actively transcribing at the mitosis–G1 transition for both Myc (45% for 4N cells at the 60-min point vs. 12.5% at the 240-min time point) [Fig. 5C left panel] and Gata2 (50% for 4N cells at the 60-min time point vs. 5% at the 240-min time point) [Fig. 5C right panel]. Of note, the zenith of the transcriptional spike coincides with the time point when chromosomes are still morphologically condensed [Fig. 5A,C]. In contrast, the intensity of transcription sites is relatively unchanged [Supplemental Fig. S18]. Thus, the mitosis–G1 spike in averaged transcriptional output mostly arises from an increase in the probability of the gene being in a transcriptionally “on” state and not from an increase in the number of nascent transcripts synthesized during each “on” period.

How does the spike in the probability of being in a transcriptionally “on” state at the mitosis–G1 transition manifest at the level of mature mRNA in single cells? To address this, we quantified the number of mature mRNA molecules in each cell. For both Gata2 and Myc, the transcriptionally “on” cells express higher levels of mature mRNA than the transcriptionally “off” cells across all time points [Fig. 5D], confirming that the intermittent nature of transcription for these genes contributes visibly to cell-to-cell variability in mature mRNA levels. Furthermore, among the transcriptionally “on” cells, the mature mRNA levels spike at the 90- to 120-min time points, subsequent to the spike in transcription for both Myc (1.9-fold increase in median mature mRNA concentration from 60 to 90 min in 2N cells) [Fig. 5D left panel] and Gata2 (1.5-fold increase in median mature mRNA) [Fig. 5D right panel]. Of note, this spike is not observed for the transcriptionally “off” cells in the corresponding time points. Additional biological replicates are shown in Supplemental Figure S17. In static images, the spike in mature mRNA specifically among transcriptionally “on” cells at the 90- to 120-min time points [Fig. 5D] must have arisen from transcriptional activity...
prior in time; thus, the spike in mature mRNA levels at these time points is enriched for the subset of cells that previously participated in the transcriptional spike at the 60-min time point. These results support that the mitosis–G1 transcriptional spike reflects an increased probability that individual cells are in a transcriptionally “on” state.
state and is substantial enough to produce a spike in mature mRNA levels to overcome any potential buffering by post-transcriptional regulation. While unable to directly offer mRNA expression trajectories of single cells over time, these data suggest that participation by individual cells in the transcriptional spike at the 60-min time point may predispose those cells to be in a transcriptionally “on” state in subsequent time points even when the overall fraction of the population transcribing has already declined.

Early spike and late-up-regulation G1 transcriptional patterns can be observed in the absence of cell cycle synchronization and can constitute the maximal transcriptional activity per DNA copy in the entire cell cycle

Our analyses thus far have relied on the use of cell cycle synchronization methods, as have previous studies of transcription at the mitosis–G1 transition (Blobel et al. 2009; Dey et al. 2009; Zhao et al. 2011; Fukuoka et al. 2012, Caravaca et al. 2013). While the resolution of cell cycle synchrony provided unambiguous visualization of the pioneering round of transcription upon reversal of mitotic silencing (Fig. 1B), the effects of synchronization on transcription are unknown and could potentially confound our gene expression observations. To avoid cell cycle synchronization completely, we sought to measure transcription in cells from different cell cycle stages in an asynchronous population by imaging. We used cell area as a proxy for cell cycle progression since cytokinesis, based on an empirically determined proportionality between the two variables (Supplemental Fig. S20). By tracking live cells through cell divisions using bright-field microscopy, we also demonstrated that newly divided cells in early G1 are enriched among the smallest cells (Supplemental Movie 1). Thus, combining RNA FISH with cell area provides a view of transcription with respect to approximate cell cycle progression since cytokinesis, enabling resolution within G1 phase that is difficult to achieve with other cell cycle markers. Satisfyingly, the transcriptional patterns for Gata2 and Kit [early G1 spike genes] and Hbb-b1 [a late G1 up-regulation gene] obtained by this method reflect that measured from approaches using synchronization [Fig. 6]. Furthermore, after normalizing for DNA copy number changes, the early G1 spike for Gata2 and Kit constitutes the highest transcriptional activity throughout the cell cycle for those genes, whereas the maximal activity for Hbb-b1 is near the late G1/S boundary [Fig. 6]. Thus, the G1-phase transcriptional patterns that we uncovered can be observed in naturally dividing cells in the absence of synchronization and can constitute periods of maximal activity in the entire cell cycle. We also note that, for both early spike genes, Gata2 and Kit, this contributes to a doubling of transcriptional activity per DNA copy when averaged across all of G1 relative to that in G2. In the Discussion, we explore the implications of this for gene dosage compensation for DNA copy changes during the cell division cycle.

Discussion

Our results uncover previously unknown genome-wide patterns of transcriptional modulation from mitosis through late G1, observing transcriptional spiking at the mitosis–G1 transition for approximately half of all active genes and intergenic enhancers [Fig. 7]. Prior to this work, an implicit expectation has been that early transcription at the mitosis–G1 transition resumes in a manner starting from initially low levels and then increases with varying kinetics to achieve maximal levels later in interphase. However, only some genes in prior studies appear to display these characteristics (Blobel et al. 2009; Zhao et al. 2011; Fukuoka et al. 2012; Kadauke et al. 2012; Caravaca et al. 2013), while others exhibit nonmonotonic patterns of transcription over time (Dey et al. 2009; Muramoto et al. 2010; Fukuoka et al. 2012; Caravaca et al. 2013).

Figure 6. The mitosis–G1 transcriptional spike can be observed in the absence of synchronization and can constitute the maximum transcriptional activity per DNA copy in the entire cell cycle. Asynchronously dividing cells were imaged by 3D wide-field microscopy after performing RNA FISH for two early spike genes [Gata2 and Kit] and a late-up-regulated gene [Hbb-b1] using intronic and exonic probes. Primary transcript content is shown in a dotted line in terms of absolute primary transcript equivalents per cell and in a solid line after normalization by DNA content as a moving mean across cell area manually determined from bright-field images. Using the proportionality between cell area and DAPI intensity [Supplemental Fig. S20], we roughly estimated the G1-, S-, and G2-phase boundaries demarcated by color. Solid horizontal lines indicate the average within the entire G1 or G2 compartment. Gata2 quantification is based on images of 5702 cells pooled across four biological replicates, Kit quantification is based on images of 640 cells, and Hbb-b1 quantification is based on images of 1339 cells. The gray shading around the moving mean denotes SEM within a sliding window of cell size.
Our present results demonstrate that the initial rounds of gene transcription upon reversal of mitotic silencing exhibit higher activities across the genome compared with late G1, providing an overall context for interpreting results from prior studies. In particular, Muramoto et al. (2010) showed a spike in transcriptional activity immediately after mitosis of the act-5 gene by live-cell imaging in Dictyostelium. In light of our findings that the mitosis–G1 transcriptional spike is shared by at least two developmentally distinct murine cell lines (Supplemental Fig. S10), these results together suggest that a spike in transcriptional activity after mitosis does not reflect a peculiarity of a specific gene or cell type but is a general phenomenon of the genome that can be observed in evolutionarily distant cell types. The strength of our observation is supported by genome-wide coverage, purity of cells from the relevant cell cycle stages, extraction of prominent transcriptional patterns by unsupervised pattern discovery, and evidence that the mitosis–G1 transcriptional spike propagates to heterogeneity in single-cell mature mRNA levels.

What might be the mechanistic underpinning of the mitosis–G1 transcriptional spike? An important consideration is the effect of mitosis on the bulk distribution of transcription regulators. For clarity of discussion, we use Pol II as an example to illustrate a likely and potentially generalizable biophysical process. Mitotic displacement of Pol II would be expected to increase the unbound fraction of Pol II that would subsequently be available to initiate transcription upon reversal of mitotic inhibition. Since transcription initiation is restricted to promoter and enhancer regions, this process would likely produce a transient increase in the ratio of effective enzyme concentration to the available DNA substrate. Upon reversal of mitotic silencing, such global shifts in factor distribution might predispose much of the genome to transcriptional spiking by mass action. Given that many general transcription factors have genomic occupancy distributions similar to Pol II during interphase and are likewise displaced from mitotic chromatin, the above scenario almost certainly applies to some factor that exists at a limiting concentration for transcriptional initiation. Such global changes in effective concentrations would be difficult to test for experimentally.

The degree of locus specificity observed for the mitosis–G1 transcriptional spike requires additional explanations. Figure 4 suggests that regions with high H3K27ac might be preferred by Pol II reinitiation, explaining their proclivity for transcriptional spiking. Numerous studies proposed or assumed that chromatin-associated molecular entities marking individual loci during mitosis would influence the subsequent reading of the genome by the transcriptional machinery at the mitosis–G1 transition. Hence, these entities are metaphorically alluded to as mitotic “bookmarks.” How is H3K27ac—now a candidate bookmark uncovered by genome-wide association—specified at individual loci during mitosis? Levels of histone acetylation in general are thought to result from the dynamic equilibrium of histone acetyltransferase versus histone deacetylase activities. Immunofluorescence microscopy previously showed that most acetylated histone H3 is globally reduced during mitosis (Kruhlak et al. 2001). This may reflect the outcome of bulk redistribution of both histone acetyltransferases—including p300 and CBP, known to be responsible for depositing H3K27ac (Tie et al. 2009)—and histone deacetylases away from chromatin in mitosis (Kruhlak et al. 2001). Thus, our observation of significant retention of H3K27ac by ChIP-
sequ at many loci during mitosis is likely an exception with respect to overall depletion of histone H3 acetylation and indicates that some level of histone acetyltransferase activity must remain and exert locus-specific effects during mitosis. How the activities of these enzymes are specified at individual loci during mitosis remains unexplored.

What might be the biological consequence of the mitosis–G1 transcriptional spike? It is unclear whether this phenomenon has been programmed to serve a biological function. Recent single-molecule RNA FISH studies of candidate genes in mammalian cells [Padovan-Merhar et al. 2015, Skinner et al. 2016] and genome-wide studies in Saccharomyces cerevisiae [Voichek et al. 2016] have shown that total transcriptional outputs for individual genes before and after DNA replication in S phase are equal. Our single-molecule RNA FISH measurements of per-copy gene transcription for the early G1 transcriptional spike genes Gata2 and Kit are consistent with these prior observations. Thus, on a per-DNA-copy level, transcriptional activity is twofold higher in G1 than in G2, an observation first proposed by Padovan-Merhar et al. (2015) as promoting transcriptional homeostasis in the face of increased DNA copy upon replication. This doubled transcriptional activity per DNA copy overall in G1 necessarily includes contributions from the mitosis–G1 transcriptional spike. Thus, at least a portion of the transcriptional compensation in G1 arises from an unknown mechanism that exerts the most effect at the mitosis–G1 transition rather than acting uniformly throughout G1. Such a model would not be mutually exclusive with, and could act in concert with, other potential mechanisms previously suggested to contribute to transcriptional gene dosage compensation, such as the dampening of transcriptional output upon nascent DNA synthesis in S phase [Padovan-Merhar et al. 2015, Voichek et al. 2016]. To what extent a dysregulation in gene dosage compensation at the transcriptional level might influence cellular function remains an intriguing open question.

Regardless of whether the mitosis–G1 transcriptional spike serves any particular biological function, our analysis of mature mRNA expression levels in transcriptionally “on” versus “off” cells [Fig. 5] suggests that the transcriptional spike does not occur uniformly across a cell population. The differences in mature mRNA levels between the transcriptionally “on” and “off” cells [Fig. 5] appear modest (1.5-fold for Myc and 1.9-fold for Gata2) in the context of the population’s overall >100-fold range in mature mRNA levels. However, our approach of imaging fixed cells cannot directly evaluate a cumulative effect size that might be extracted from observing the trajectories of mRNA production in live individual cells over multiple cell divisions. To illustrate this possibility, suppose that, upon division of a single cell, the early G1 transcriptional spike stochastically occurs in one daughter cell but not the other, perhaps, on average, resulting in a 1.9-fold difference of mature mRNA levels in those two cells. Such a difference, while moderate to begin with, might predispose one cell to a higher probability of subsequent higher expression levels. Such a scenario is consistent with, although not proven by, our indirect inferences from static images in Figure 5 and might be particularly applicable if the gene product is involved in an autoregulatory positive feedback loop. Repeated sampling of the mitosis–G1 transition over multiple cell divisions might account for at least part of the eventual substantial divergence in gene expression state among all of the progeny of the original founding cell. Consideration of the mitosis–G1 transition as a source of gene expression heterogeneity might pave the way for understanding why the probability of certain types of phenotypic transitions is modified by rapid proliferation [Smith et al. 2010] and passage through mitosis [Egli et al. 2008, Gani et al. 2011, Halley-Stott et al. 2014] or early G1 phase [Singh et al. 2013]. We envision that the mitosis–G1 transcriptional spike, on average, may promote gene expression homeostasis with respect to DNA dosage, yet its variable occurrence at the single-cell level may contribute to diversification of gene expression states.

Materials and methods

Cell culture, cell cycle synchronization, and cell sorting

G1E erythroblasts were previously derived through deletion of GATA1 in mouse embryonic stem cells followed by in vitro differentiation [Weiss et al. 1997]. We cultured a subline of G1E cells, G1E-ER4, in which GATA1-ER was retrovirally transduced [referred to in the text as G1E GATA1-ER], as described previously [Weiss et al. 1997]. We retrovirally transduced G1E-ER4 cells with the YFP-MD construct (Kadauke et al. 2012) and sorted for a pool of YFP-positive cells. Except where indicated in the text as uninduced, we induced cells to mature with 100 nM estradiol to activate GATA1-ER. During estradiol induction, cells were simultaneously treated with 200 ng/mL nocodazole for 7–13 h, washed once, and replated into fresh medium lacking nocodazole for varying times (40–360 min), ensuring that all samples were exposed to estradiol for the same duration of 13 h. Cells were fixed with 1% formaldehyde, stained with DAPI, and sorted on a BD FACSAria based on YFP-MD and DAPI signal. Sorting of MPM2-positive prometaphase populations and MPM2-negative interphase populations for H3K27ac ChIP-seq was carried out as described in Campbell et al. [2014].

F9 cells [Alonso et al. 1991] were cultured in plates precoated with 0.1% gelatin and grown in DMEM + 10% FBS. For mitotic arrest–release, cells were treated with 200 ng/mL nocodazole for 4 h, and a “shake-off” [gentle rinsing with medium] was performed to isolate mitotic cells followed by replating into fresh medium for varying durations of the release time course.

ChIP-seq

We performed ChIP-seq of a total of three biological replicates using N-20 antibody [Santa Cruz Biotechnology, catalog no. sc899] for the 0-, 60-, 90-, 180-, and 360-min time points; two biological replicates for the 240-min time point; and one replicate for the 40-min time point. Two replicates of input DNA at the corresponding time points were also sequenced. For ChIP-qPCR of the initiating form of Pol II, we used 8WG16 antibody [Covance, catalog no. MMS-126R]. H3K27ac antibody from ActiveMotif [catalog no. 39685] was used for H3K27ac ChIP-seq. To summarize briefly, cells fixed with 1% formaldehyde and subjected to lysis in detergents, sonication, and immunoprecipitation of chromatin. Following library construction through blunt end
Bioinformatic analysis of ChIP-seq data

To summarize briefly, reads were mapped to the mouse mm9 genome using Bowtie [Langmead et al. 2009]. Mapped reads were passed to MACS [Zhang et al. 2008] with a matched control (input) data set for peak calling and producing Bigwig files with reads shifted to account for fragment size. If the 5' or 3' 2.5-kb region of a gene overlapped at least one Pol II peak called by MACS in at least one sample (arrest-release and asynchronous samples with estradiol induction), then we deemed the gene active, arriving at 4309 nonoverlapping active genes defined by the single largest 3-kb and 330 base pairs (bp). See the Supplemental Material for details.

RT-qPCR of primary transcripts

We isolated RNA using TRIzol [Life Technologies] or TRIzol LS [Life Technologies]. Reverse transcriptase reaction was performed with iScript [Bio-Rad]. qPCR was performed with Power SYBR Green [Invitrogen]. All primer sequences are in the Supplemental Material. For primary transcript measurements, primers flank intron–exon junctions. Primary transcript quantifications are normalized to Gapdh mature mRNA. Results are similar when normalized to Hprt mature mRNA [data not shown].

Capture-C

After cell fixation with 1% formaldehyde for 10 min and cell sorting as described above, Capture-C was performed with a double-capture procedure [Davies et al. 2016]. Chromatin was digested using DpnII. We used biotin-labeled DNA oligos to pull down the target restriction fragments. Capture-C libraries were sequenced on Nextseq500 with paired-end sequencing (75 bp on each end). The raw reads were processed using published scripts [https://github.com/telenius/captureC/releases]. We wrote custom scripts to normalize data by the total number of reads representing fragments ligated to the anchor region in the library. Enhancer regions used for quantification were defined by assessment of Capture-C signal by eye together with consideration of prior literature and DNase sensitivity signals from Hsiung et al. (2014).

Single-molecule RNA FISH

We performed single-molecule RNA FISH as described previously [Feminó 1998; Raj et al. 2008; Raj and Tyagi 2010; Levesque and Raj 2013]. In brief, we fixed cells in 1.85% formaldehyde for 10 min at room temperature and stored them in 70% ethanol at 4°C until further processing. FISH probes consisted of oligonucleotides conjugated to fluorescent dyes as follows: Myc exons to Cy5, Gata2 exons to Cy3, Myc introns to Alexa594, and Gata2 introns to Alexa 594. Oligonucleotide sequences are in the Supplemental Material. Imaging was performed on a Nikon Ti-E inverted fluorescence microscope using a 100× plan-apo objective [numerical aperture of 1.43], a cooled CCD camera [Pixis 1024B from Princeton Instruments], and filter sets SP102v1 (Chroma), SP104v2 (Chroma), and 3100v2 (Chroma) for Cy5, Cy3 and DAPI, respectively. A custom filter (Omega) was used for Alexa 594. We took optical z-sections [typically 45] at intervals of 0.35 µm, spanning the vertical extent of cells, with 1 sec of exposure time for Cy3, Cy5, and Alexa 594 and 100 msec of exposure time for DAPI.

Data access

All raw and processed sequencing data will be deposited at Gene Expression Omnibus [GSE83293]. In addition, scripts that reproduce the majority of figures starting from processed data in tabular form [including RPKMs from ChIP-seq, spot counts and intensities from single-molecule RNA FISH, read counts from Capture-C, and Ct values from RT-qPCR] are provided and maintained in an online repository [https://cshiung@bitbucket.org/arjunrajlaboratory/hsiung_mitosisreactivation]. Pol II ChIP-seq tracks can be viewed at the genome browser hosted by Pennsylvania State University [http://main.genome-browser.bx.psu.edu/cgi-bin/hgTrackUi?hgsql=206746_6ux0ZSv9DeKChVjzcf2tRchNFS&c=chr5&g=meryYfpm].

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Repair and adaptor ligation using Illumina’s TruSeq ChIP sample preparation kit [Illumina, catalog no. IP-202-1012], size selection with SPRiSelect beads (Beckman Coulter, catalog no. B23318), and PCR amplification, libraries were multiplexed and sequenced on an Illumina HiSeq 2000. The mean fragment size was ~340 base pairs [bp]. See the Supplemental Material for details.

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by the DiGaetano family (to G.A.B. and P.E.); and National Institutes of Health 1RO1 HL119479 (to G.A.B.), C.C.-S.H., A.R., and G.A.B. planned the study. C.C.-S.H., C.R.B., P.H., A.J.S., C.A.K., C.F., K.S.J., and L.S. performed experiments. C.C.-S.H., C.R.B., P.G., P.E., P.H., and B.G. performed computational data processing and analyses. C.C.-S.H., R.C.H., A.R., and G.A.B. interpreted the results. C.C.-S.H., A.R., and G.A.B. wrote the manuscript with input from all authors.

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