A comprehensive single cell data analysis of in lymphoblastoid cells reveals the role of Super-enhancers in maintaining EBV latency

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Abstract
We probed the lifecycle of EBV on a cell-by-cell basis using single cell RNA sequencing (scRNA-seq) data from nine publicly available lymphoblastoid cell lines (LCL). While the majority of LCLs comprised cells containing EBV in the latent phase, two other clusters of cells were clearly evident and were distinguished by distinct expression of host and viral genes. Notably, both were high expressors of EBV LMP1/BNLF2 and BZLF1 compared to another cluster that expressed neither gene. The two novel clusters differed from each other in their expression of EBV lytic genes, including glycoprotein gene GP350. The first cluster, comprising GP350\(^{-}\)LMP1\(^{hi}\) cells, expressed high levels of HIF1A and was transcriptionally regulated by HIF1-\(\alpha\). Treatment of LCLs with Pevonedistat, a drug that enhances HIF1-\(\alpha\) signaling, markedly induced this cluster. The second cluster, containing GP350\(^{+}\)LMP1\(^{hi}\) cells, expressed EBV lytic genes. Host genes that are controlled by super-enhancers (SEs), such as transcription factors MYC and IRF4, had the lowest expression in this cluster. Functionally, the expression of genes regulated by MYC and IRF4 in GP350\(^{-}\)LMP1\(^{hi}\) cells were lower compared to other cells. Indeed, induction of EBV lytic reactivation in EBV\(^{+}\) AKATA reduced the expression of these SE-regulated genes. Furthermore, CRISPR-mediated perturbation of the MYC or IRF4 SEs in LCLs induced the lytic EBV gene expression, suggesting that host SEs and/or SE target genes are required for maintenance of EBV latency. Collectively, our study revealed EBV associated heterogeneity among LCLs that may have functional consequence on host and viral biology.

Importance
Epstein-Barr virus (EBV) establishes a life-long latency program within host cells. As such, EBV immortalized lymphoblastoid cells (LCLs) often carry the latent EBV genome and only a small percentage of LCLs containing lytic EBV. However, the cellular programs that distinguish latent from lytic cells and the heterogeneity of cells in latent or lytic phases remains poorly explored. To explore these unknowns, we reanalyzed publicly available single cell RNA-seq data from nine LCLs. This approach permitted the simultaneous study of cells in both latent and lytic phases. We identified three cell populations with distinct lytic/latent activity and further characterized the transcriptomes of these cells. We also identified a new role of super-enhancers in regulating EBV lytic replication. Collectively, our studies revealed EBV associated heterogeneity among LCLs that contribute to EBV life cycle and biology.
**Introduction**

Epstein-Barr virus (EBV) is the first oncogenic human DNA virus discovered more than 50 years ago (1). EBV causes ~200,000 cases of diverse cancers every year (2), including lymphomas, nasopharyngeal carcinoma and gastric adenocarcinomas (3, 4). Most EBV infections occur early in life and are transmitted through saliva. EBV first infects oral epithelial cells and then B lymphocytes in the oral epithelium. EBV persists in memory B cells for life in a latent phase, so these cells express minimum EBV genes under host immune surveillance. However, when host immunity is impaired, for example by immunosuppressive treatment or HIV infection, EBV in infected B cells can enter type III latency where six EBV nuclear antigens (EBNAs), three latent membrane proteins, and a few noncoding RNAs and microRNAs, are expressed (3, 5). This can result in lymphoproliferative diseases or lymphomatous transformation (6). EBV in infected memory B cells can also enter a lytic phase to actively produce live virus. During EBV lytic replication, immediate early genes RTA and ZTA encoded by \textit{BRLF1} and \textit{BZLF1} genes, respectively, are first expressed. These are transcription factors (TFs) that turn on the expression of genes necessary for viral DNA replication and structural proteins, including viral membrane protein gp350, that binds to human B cell EBV receptor CD21, to assemble virions (7).

\textit{In vitro}, EBV infection of primary B lymphocytes leads to the establishment of lymphoblastoid cell lines (LCLs) (8). LCLs express EBV type III latency genes, the same genes seen in some EBV malignancies, including post-transplant lymphoproliferative disease and AIDS CNS lymphomas. Therefore, LCLs are an important model system to study EBV oncogenesis. Genetic studies have found that EBNA1, 2, LP, 3A, 3C and LMP1 are essential for EBV-mediated growth transformation (9, 10). EBNA1 tethers EBV episomes to host DNA (11-13). EBNA2 and LP are the major EBV transcription activators that activate expression of key oncogenes, including MYC (14, 15). EBNA3A and 3C repress expression of p16\textsubscript{INK4A} and p14\textsubscript{ARF}, to overcome senescence and BIM to avoid apoptosis (16-18). LMP1 activates NF-\kappaB to provide survival signals (19).

EBV infection significantly alters chromatin topology and function at EBV-interacting genomic loci of host cells (20). This alteration could be mediated by EBV-encoded transcription factors (21, 22) or via interaction between EBV episomes and the host genome (23), and may also depend on the EBV latency program (24, 25). Super-enhancers (SEs) are critical regions of mammalian genomes comprised of clusters of enhancers bound by arrays of transcription factors.
Viral transcription factors and host NF-κB subunits can form EBV SEs (22) with markedly high and broad histone 3 lysine 27 acetylation (H3K27ac) (27). These SEs are linked to many genes essential for LCL growth and survival, including MYC and IRF4, and their perturbation pauses LCL growth and causes cell death (28, 29). We have previously shown that EBV episomes physically interact with SE-containing genomic host loci in EBV-transformed lymphoblastoid cells (23). However, the consequences of these interactions and the effects of perturbations at these SE-containing loci on the EBV life cycle remains unexplored.

High throughput sequencing technologies can aid to dissect mechanisms underlying host-virus interactions (4, 23, 30, 31). However, the heterogenous nature of virally infected cells is an impediment to precisely probing the phases of virus life cycle and their effect on host genes in individual cells. Recent advances in single-cell transcriptomics have enabled successful resolution of tissue/cell heterogeneity in several species. Since these technologies agnostically capture both host and infecting viral sequences, they have also been utilized to explore host-virus interactions at a single cell level (32-34). Leveraging this feature here, we sought to identify the determinants of EBV latency in lymphoblastoid cells. Using single cell transcriptomics, we identified distinct clusters marked by differences in expression of GP350 and LMP1. Cells expressing high levels of LMP1, but not GP350, demonstrated high HIF1-α activity and could be induced by a HIF1-α stabilizer. Cells co-expressing GP350 and LMP1/BNLF2 had significantly reduced expression of SE-containing genes compared to cells containing EBV that was clearly in the latent phase (i.e., GP350- cells). Using proof-of-principle SE inactivation experiments, we found that host SEs are necessary for the maintenance of EBV latency. Collectively, our data not only highlight the heterogeneity among LCLs but also identifies common functional themes of the cells and their role in EBV associated biology.
Results

Single cell RNA-sequencing analyses resolve LCLs into three distinct populations

To better understand how EBV in infected cells spontaneously enter the lytic life cycle, we analyzed publicly available single cell RNA-sequencing (scRNA-seq) data from nine LCLs from several independent sources (see Methods) (33, 35-39). Briefly, we performed an unbiased integrative analysis across all these LCLs after regressing for potential batch effects, doublets and/or artifacts and known sources of heterogeneity, such as the stage of cell cycle using the Seurat platform (40) (Figs. S1a-c- see methods). Unsupervised clustering of all 46,205 cells according to expressions of both host and viral genes at three different resolutions yielded several clusters (Fig. S1d). Further examination of these clusters based on the expression of salient EBV genes, including GP350 and LMP1/BNLF2, and separation in UMAP space revealed that these clusters fall into three major groups according to the status of EBV gene expression, namely latent, early lytic and full lytic EBV cell clusters. The multiple clusters corresponding to EBV in the latent state were recently examined thoroughly (33). Since our focus was mainly on understanding the biology of EBV lytic life cycle, we combined all the latent cells into one cluster, resulting in three major clusters (Fig. 1a). These were evident in all LCL datasets examined (Figs. S1a-b). These clusters contained GP350\(^{-}\)LMP1\(^{lo}\), GP350\(^{-}\)LMP1\(^{hi}\) and GP350\(^{+}\)LMP1\(^{hi}\) cells, representing cells with EBV in the latent, early lytic and fully lytic states (Figs. 1a, S1e). Approximately 50-100 host genes were differentially expressed in each cluster compared to all other clusters (Figs. 1b, S1f and Table S1). Consistently, GP350\(^{+}\)LMP1\(^{hi}\) cells was the cluster expressing the most EBV genes, while GP350\(^{+}\)LMP1\(^{lo}\) cells represented the cluster showing the lowest expression of EBV genes (Figs. 1b-d).

GP350\(^{lo}\) cells comprised 94-98% of all LCLs across all the samples (Figs. 1a, S1a-b). They displayed minimal expression of LMP1/BNLF2 and minimal or no expression of EBV lytic genes, including GP350, BMRF1, BALF1 and BALF3 (Fig. 1c). Additionally, these cells expressed latency genes, including EBNA1 and EBNA2, indicating that this cluster mainly consisted of transformed cells that were in the EBV latent phase (Fig. 1d). This cluster was also the highest expressor of genes from immunoglobulin heavy or light chains, indicating the mature status of these transformed B cells (Figs. 1e, S1g). Consistently, nearly a quarter of cells in this cluster expressed high levels of PRDM1, indicating that these cells might have entered plasmacytic differentiation (41).
The two lytic clusters, $GP350^+LMP1^{hi}$ and $GP350^+LMP1^{hi}$ cells, respectively, each accounted for 1-5% of all LCLs (Figs. 1a, S1b). The salient viral feature of both clusters was the high expression of $LMP1/BNLF2$, a gene with well-established contribution to oncogenic human B-cell transformation (42) and $BZLF1$. $GP350^+LMP1^{hi}$ cells were the highest expressors of EBV lytic genes, including $GP350$, $BZLF1$ and $BMRF1$, while $GP350^-LMP1^{hi}$ cells express very few lytic genes (Fig. 1c). Remarkably, these two clusters had distinct expressions of host genes (Figs. 1e, S1f). Consistent with previous reports (33, 34), $GP350^+LMP1^{hi}$ cells highly expressed host $NFATC1$, $MIER2$, $SFN$ and $SGK1$ genes and were the highest expressors of host box-dependent myc-interacting protein 1 ($BIN1$). Conversely, $GP350^-LMP1^{hi}$ cells had the highest expression of host genes $HSPB1$, $ABCB10$, $MALAT1$ and $CD44$ (Fig. 1e).

To obtain insights into the functional state of cells in each cluster, we performed geneset enrichment analysis (GSEA), comparing the transcriptomes of cells in each cluster against those of cells from the other two clusters (Fig. 1f), and querying enrichment of all 50 hallmark genesets curated by the Molecular Signatures Database (MSigDB) (43). Genes differently regulated in $GP350^+LMP1^{lo}$ cells were significantly enriched in MYC targets, MTORC1 signaling and inflammatory response. Conversely, genes differently regulated in $GP350^-LMP1^{hi}$ cells were enriched in tumor necrosis factor alpha signaling, apoptosis and hypoxia (Fig. 1f). As expected, genes differently regulated in $GP350^-LMP1^{hi}$ cells were significantly depleted of genesets from most hallmark pathways, including MYC targets, MTORC1 signaling and interferon responses (Figs. 1f, S1h). This is consistent with the fact that fully lytic EBV reactivation pauses transcription of most host genes and pathways, which is evidenced by significantly reduced numbers of total host transcripts in lytic cells (Fig. S1i).

Collectively these analyses indicated that LCLs are predominantly comprised of three distinct cell populations characterized by differences in expression of both host and viral genes, notably cells containing EBV in the latent phase ($GP350^-LMP1^{lo}$), cells containing virus in the lytic phase ($GP350^-LMP1^{hi}$) and cells that were in between lytic and latent phases ($GP350^-LMP1^{hi}$). Furthermore, these data suggested that distinct functional states of individual LCL clusters may be related to expression of genes encoded by EBV and the host cell.
GP350–LMP1hi LCLs have a HIF1A-associated signature.

We next explored the transcriptional regulators of host gene expression. Our attention was drawn to HIF1A because GP350–LMP1hi cells were high expressors of several genes including HSPB1, MALAT1 and CD44 (Fig. 1e) that in other settings are known to be regulated by hypoxia or HIF1-α (44-46) and because our GSEA analysis had also indicated that the transcriptomes of these cells are highly enriched in the hypoxia gene set (Fig. 1f). HIF1-α is a critically important TF that is tightly regulated by oxygen tension and transactivates many genes essential for cellular responses and adaptation to hypoxia (47). To better characterize GP350–LMP1hi cells, we therefore quantified the mRNA expression of HIF1A, the gene that encodes HIF1-α, and its classical direct target PDL1 (48) in all clusters. HIF1A and PDL1 were both more highly expressed in GP350–LMP1hi cells compared to others (Fig. 2a). This was specifically evident for HIF1A as its expression levels were significantly higher in GP350–LMP1hi cells (Figs. 2a, S2a). To determine whether the changes in HIF1A expression could have any functional consequence, we next assessed the expression of HIF1-α target genes. We sourced a public list (49) of HIF1-α–induced (n=110) and HIF1-α–repressed (n=77) genes from MSigDB (50) and assessed the expression of these two sets in all 3 identified LCL clusters (Fig. 2b and Table S2). GP350–LMP1hi cells had the highest and lowest expressions among all clusters for HIF1-α–induced and HIF1-α–repressed genes, expressed as the module score (51), respectively (Fig. 2b). We confirmed these findings using two additional independent lists of HIF1-α–regulated genes (52) (Fig. S2b-c and Table S2).

We next predicted pharmaceutical agents that could induce the unique gene signatures of cells in the GP350–LMP1hi cells, using methods established previously by our group (32, 53). Among the topmost significant drugs predicted to enhance host gene expression pattern of GP350–LMP1hi cells was Pevonedistat (MLN4924) (Fig. 2c). Pevonedistat is a ubiquitin-activating enzyme E1 inhibitor that significantly stabilizes HIF1-α to potentiate its function (54). Because the HIF1-α pathway was one of the main features of GP350–LMP1hi cells, we hypothesized that enhancing HIF1-α signaling might preferentially induce this program. To test this hypothesis, we treated LCLs with Pevonedistat and measured HIF1A, LMP1, PDL1 and GP350. HIF1-α potentiation markedly induced mRNA expression of HIF1A, LMP1 and PDL1 (Fig. 2d), but not GP350 or BZLF1 (Fig. 2e). To further substantiate these observations at the
single cell level, and confirm expression of protein, we treated three different LCLs with Pevonedistat and performed flow cytometry. Pevonedistat reduced cell viability by nearly 30% (Figs. 2f, S2e). The frequency of LMP1\(^+\) cells was significantly increased (Figs. 2g-h, S2f-g) without increasing that of ZTA (Figs. 2i, S2h). The frequency of PDL1\(^+\) cells and PDL1 expression were also significantly increased upon treatment (Fig. 2j-k, S2i). We have also performed dose titration of Pevonedistat and have observed dose dependent increase of LMP1 and PD-L1, but not BZLF1, in gated live cells (Fig. S3), suggesting that Pevonedistat preferentially induce LMP1\(^+\) cells without increasing full lytic cycle.

**GP350\(^-\)LMP1\(^{lo}\) LCLs have distinct MYC-dependent transcriptional programs**

We next focused on transcriptional regulators of GP350\(^-\)LMP1\(^{lo}\) LCLs, the cluster containing EBV in the latent phase. MYC-regulated genes were among the top affected pathways when comparing transcriptomes of LCL clusters against each other (Fig. 1f) and box-dependent myc-interacting protein 1 (BIN1) was one of the top host genes distinguishing GP350\(^+\) from GP350\(^-\) cells (Fig. 1e). Moreover, MYC has been described as a key host factor repressing EBV lytic reactivation (55). Thus, we further examined the role of MYC in shaping the distinct LCL clusters. Because MYC is a transcription factor, we first determined the fraction of differently expressed genes in each cluster directly bound by MYC. For this, we sourced a publicly available ChIP-seq dataset for MYC in GM12878 (GSM822290, curated by ENCODE). Nearly 18-24% of genes differently expressed in each cluster were directly bound by MYC, with GP350\(^-\)LMP1\(^{hi}\) cluster having the most number of MYC targets (Fig. 3a and Table S2). This was significantly higher than what would be expected by chance because only ~10% of all human genes are bound by MYC in GM12878 (Fig. Sa).

The mRNA expression of MYC was significantly higher in GP350\(^-\)LMP1\(^{lo}\) LCLs than in either of the other two clusters (Fig. 3b). To determine whether MYC is biologically active, we looked for the signature of genes regulated by MYC. We curated a list of genes regulated by MYC in GM12878 from a publicly available dataset (55) (Table S2). Expression of MYC-induced genes was significantly higher (Fig. 3c, left panel) and MYC-repressed genes significantly lower (Fig. 3c, right panel) in GP350\(^-\)LMP1\(^{lo}\) cells than in the other two clusters. We also performed GSEA comparing transcriptomes of cells from each cluster against MYC targets curated by MSigDB (43). Consistent with our earlier observation (Fig. 1f), genes that
were more highly expressed in \(GP350^+\text{LMP1}^{lo}\) cells were highly enriched in MYC targets (Fig. 3d, left and right panels), while there was no significant difference between \(\text{LMP1}^{hi}\) clusters (Fig. 3d, middle panel). Collectively, these data indicated that MYC preferentially regulates a subset of genes that are differently expressed in \(GP350^+\text{LMP1}^{lo}\) LCLs.

**Super-enhancer-regulated genes are less highly expressed in \(GP350^+\text{LMP1}^{hi}\) LCLs**

EBV-infected cells periodically enter the lytic phase to produce progeny viruses but in EBV-immortalized lymphoblastoid cells EBV is mostly in the latent state. Earlier studies have shown that a small percentage of these cells are lytic (30). However, due to the technical challenges at the time, it was difficult to distinguish the cells in lytic phase from cells at latency phase in a mixed population. The recent development of scRNA-seq techniques allows us to capture the cells in lytic phase together with their transcriptome.

Nearly 10% of genes are regulated by multiple enhancers forming a complex architecture known as “super-enhancers” (SEs). SE-regulated genes are critically important for cell identity (27) and are associated with both Mendelian and polygenic diseases (56, 57) as well as cancers (58). Enhancer-promoter interactions are the cornerstones of mammalian gene regulation. We have previously shown that EBV episomes make reproducible contacts with the human genome at SE loci (23). To explore whether EBV disrupts modes of gene regulation in the three LCL subsets, we sourced a list of 257 annotated SE regulated genes from GM12878 (26) and determined whether these genes are differently expressed in the three identified LCL clusters. Unexpectedly, expression of SE-regulated genes, summarized as the module score, was significantly lower in \(GP350^+\text{LMP1}^{hi}\) cells, the cluster containing EBV in the lytic state, than in the other two subsets (Fig. 4a). We observed similar results when we used an independent curated set of 187 EBV-associated SEs (22) (Fig. S4a). Examples of such genes included \(\text{MYC}\), which contains one of the largest SEs in the genome (22), \(\text{IRF4, RUNX3, PAX5, IKZF3}\) and \(\text{DUSP22}\) (Fig. 4b). These findings suggested that lytic EBV is associated with disruption of expression of host genes regulated through SEs. To explore this possibility, we performed GSEA analysis comparing the transcriptomes of \(GP350^+\text{LMP1}^{+}\) cells to cells from the other two clusters for enrichment of all SE-regulated genes. This orthogonal approach also indicated that genes less highly expressed in \(GP350^+\text{LMP1}^{+}\) cells compared to the cells in the other two clusters were markedly enriched in SE-regulated genes (Fig. S4b).
We next assessed whether genes in these clusters were differently expressed when their associated SEs physically interact with EBV episomes. To this end, we divided SE-regulated genes into those that physically interact, or not, with EBV episomes and performed GSEA analysis comparing $GP350^+$ cells to the cells of other two $GP350^-$ subsets. Genes that were more highly expressed in $GP350^-$ cells were significantly enriched in SEs that interact with EBV episomes (Fig. 4c, left panel). This enrichment was less evident for SEs that do not interact with EBV episomes (Fig. 4c, right panel). To determine the functional consequences of differential expression of SE-regulated genes across LCL clusters, we focused on the transactivator IRF4 and transcription factor RUNX3 for which we could source their direct targets from ChIP-seq experiments and assess the expression of their targets. We noted that expression of both IRF4- and RUNX3-bound genes, summarized as the module score, was significantly lower in $GP350^+LMP1^{hi}$ cells (Figs. 4d, S4c), in which expression of both these TFs was also the lowest (Fig. 4b).

Since $GP350^+LMP1^{hi}$ cells represented the cluster in which lytic reactivation of EBV was apparent (Fig. 1e), we tested whether EBV reactivation affects the expression of SE containing genes. To this end, we treated EBV$^+$ AKATA cells with either anti-IgG or carrier. Anti-IgG is a potent inducer of EBV lytic reactivation in these cells (59). After stimulation, we measured mRNA and/or protein expression of EBV lytic markers and the host SE-regulated gene MYC (Figs. 4e-h). As expected, anti-IgG induced strong expression of the EBV lytic markers RTA, ZTA and BMRF1 (Fig. 4e). In contrast, the expression of both MYC and IKZF3 were significantly repressed following anti-IgG-treatment of cells (Fig. 4f). This effect was specifically a predicate of EBV-reactivation since anti-IgG did not repress MYC or IKZF3 expression in EBV$^-$ AKATA cells (Fig. 4g). These observations were further confirmed by immunoblots of ZTA, BMRF1 and MYC proteins (Fig. 4h). Consistently, a recent study has shown that depletion of MYC reactivates the EBV lytic cycle (55). To test whether depletion of IRF4 can similarly reactivate EBV lytic cycle, we reanalyzed RNA-seq from GM12878 LCLs that were subjected to IRF4 deletion via the CRISPR/Cas9 system (29). In this setting, depletion of IRF4 induced multiple EBV lytic genes, including $GP350$, $RTA$, $ZTA$ and $BMRF1$ (Fig. S4d). Consistently, a recent study has found that IRF4 knockdown in LCLs induces EBV lytic reactivation in LCLs and lytically infected cells have increased NFATc1 and decreased IRF4 expression (34). Collectively, our data suggest that SE-regulated genes are less highly expressed...
in \( GP350^{+}LMP1^{hi} \) LCLs, which show evidence of lytic EBV reactivation, and that experimental induction of EBV lytic cycle also represses expression of these genes.

**Disruption of super-enhancers in LCLs induces EBV lytic reactivation.**

The reciprocal relationship between expression of SE-regulated genes in \( GP350^{+}LMP1^{hi} \) LCLs and EBV reactivation suggested the possibility that SEs may be necessary for maintenance of EBV in the latent phase. To test this possibility, we initially selected SEs near \( MYC \) and \( IRF4/DUSP22 \) and performed CRISPR-mediated knockout or inactivation and then measured the expression of EBV lytic markers. For these experiments, appropriate guide RNAs were situated within the SEs at sites of maximal H3K27ac signal, a marker of active regions of the genome, especially promoters and enhancers. The selected sites were bound by one or more viral transcription factors (e.g. EBNA2, LP, 3A and 3C) and/or host NF-\( \kappa \)B family members (e.g. RelA, RelB, cRel, p50 and p52) and interacted within a topologically associated domain that contained the SE (Figs. 5a, 5c). Dual sgRNAs targeting both sides of \( MYC \) SE (~525 kb upstream) successfully deleted \( MYC \) SE from the genome (Fig. S5a), which led to a reduction of \( MYC \) transcription and upregulation of EBV lytic genes, namely ZTA, RTA, BGLF5 and BMRF1 expression (Fig. 5b). Similarly, inactivation of \( IRF4 \) SE by CRISPR-dcas9 tethered with a repressor consisting of KRAB and the transcription repression domain of MeCP2 successfully reduced \( IRF4 \) SE activity (Fig. S5b). This led to decrease in \( IRF4/DUSP22 \) mRNA expression and a significant increase of EBV lytic gene expression two days post inactivation (Fig. 5d). Deletion of both \( MYC \) and \( IRF4 \) genes have been previously shown to induce EBV lytic phase. Unexpectedly, however, when we measured the expression of EBV lytic genes at earlier time points, we observed that EBV lytic genes were significantly induced prior to decrease of \( IRF4 \) (Fig. S5c). This suggests the possibility that SE might be also necessary for the maintenance of EBV latency. To further explore this possibility, we selected another SE in the same topological associated domain of MIR155HG (Fig. 5e), using the same criteria as above and performed CRISPR-mediated inactivation. The disruption of this SE did not significantly reduce the expression of \( MIR155HG \); however, it significantly increased the expression of EBV lytic genes (Fig. 5f), namely ZTA, RTA, BGLF5 and BMRF1. CRISPRi disruption of RUNX3 SE also had similar activity (Fig. 5g, h). Collectively these data indicate that deletion of select host SEs leads...
to lytic reactivation of EBV and, by extension, that host SEs, or their target genes, are necessary for maintenance of EBV in the latent phase.

Discussion

LCLs have been instrumental for genetic and functional studies of human diseases over the past several decades (60). We and others have previously analyzed large numbers of LCL bulk RNA-seq data and found that EBV lytic gene expression correlates with cellular cancer-associated pathways, such as interferon-alpha, WNT and B cell receptor signaling (4, 30). However, these data were generated from bulk populations of cells, which biases insights towards those occurring in the largest sub-populations. While the majority of LCLs contain EBV in the latent phase of its life cycle, a small fraction (<5%) demonstrate spontaneous EBV reactivation, indicating that LCLs as a whole are heterogeneous. Important aspects of LCL heterogeneity have recently been explored using single cell RNA-sequencing (33). This analysis focused on heterogeneity within and across LCLs with respect to immunoglobulin isotypes, which further associated with pathways involving activation and differentiation of B cells. We have taken an integrative approach to combine the same data with several more datasets that are generated across different conditions and eliminate batch and technical effects. This integrative analysis provides a consistent representation of the data for downstream analyses and thus has the potential to uncover previously undetected biology. Specifically, we found LCLs to have higher heterogeneity in relation to the EBV status than previously appreciated. Specifically, we identified three prominent clusters that were marked by the expression of the EBV genes GP350 and LMP1/BNLF2. Of note, the EBV genome has extensive numbers of overlapping genes such as LMP1 and BNLF2a/b, making the quantification of their mRNA expression more challenging (61). This challenge could be further exacerbated by the 3’ mRNA capture bias in some of the current single cell technologies. Nevertheless, we showed that these clusters have distinct transcriptional programs and identified MYC and HIF1-α as transcriptional regulators of gene expression. LCLs in the GP350+ cluster expressed SE-regulated genes at significantly lower levels compared to cells in the other two clusters. Physical interactions between SE-containing loci and EBV episomes marked genes in GP350+ LCLs that were particularly lowly expressed. Indeed, in proof-of-principle experiments we found that experimental lytic reactivation of EBV disrupted expression of SE-regulated genes and, conversely, that disruption of SEs induced EBV
lytic reactivation. For IRF4 or MIR155HG associated SEs, lytic reactivation after SE disruption occurred prior to IRF4 downregulation, suggesting that these SEs themselves might be necessary for the maintenance of EBV latency. However, further studies are needed to fully discern this observation.

In the largest subset of LCLs, annotated as GP350–LMP1lo, EBV was clearly in the latent phase. This cluster showed a host gene signature enriched in MYC-regulated targets. As an oncogene, MYC is exquisitely carefully regulated by an archetypal SE. MYC itself binds to the EBV genome origin of lytic replication and suppresses DNA looping to the promoter of the lytic cycle initiator gene BZLF1 (55). MYC depletion reactivates the lytic cycle in different cells (55). Consistent with this, when we deleted the MYC SE, MYC expression was decreased and EBV lytic genes were induced, supporting the role of MYC as a repressor of EBV lytic activation. Thus, it appears that both the MYC gene and its associated SE have a role in maintenance of EBV latency.

Another GP350 negative cluster, characterized by high expression of LMP1/BNLF2, was the highest expressor of several host genes including HSPBI, MALAT1 and CD44 that are known features of cancer stem cells (62-64) and escape from apoptosis (65, 66). Interestingly, LMP1 alone induces CSC features in nasopharyngeal cell lines (67). However, such characteristics have not previously been ascribed to LCLs and warrant further investigation. LMP1 is a known oncogene and expressed in most EBV associated cancers (68) and has been previously associated with synthesis of HIF1-α protein and its DNA binding activity (69). Here, we also found that GP350–LMP1hi LCLs have a prominent HIF1-α signature and could be preferentially induced by Pevonedistat. These cells also expressed higher frequencies of PDL1, which were markedly increased upon Pevonedistat treatment. Interestingly, a recent study has found an association between numbers of PDL1 expressing B cells and the development of AIDS related non-Hodgkin lymphoma (70). Such PDL1 expressing B cells have previously been described to suppress effector function of immune cells (71). Thus, the identification of these cells might play an important role in understanding the oncogenesis and may suggest that drugs that stabilize HIF1-α might inadvertently induce LMP1 in diseases associated with EBV type III latency programs such as AIDS-associated B cell lymphoma, post-transplant lymphoproliferative disorder and diffuse large B cell lymphoma.
B cell differentiation into plasma cell has been linked to EBV lytic replication (72, 73). Specifically, PRDM1, a known driver of B cell differentiation into plasma cells (74), promotes EBV lytic replication by activating the transcription from immediate early gene promoters of ZTA and RTA (75). A recent single cell RNA-seq analysis of LCLs have revealed a positive correlation between specific immunoglobulin isotype and cell differentiation markers (33). However, these immunoglobin genes were not specifically characterized in lytic cells. We have found that the mRNA expression of PRDM1 and a range of immunoglobulin genes in \textit{GP350+LMP1}^\text{hi} cells was lower than in latent cells, which contrasts with previous reports about the role of PRDM1 in EBV lytic reactivation (Fig. S1e). It is possible that PRDM1 is important for initiation, but not maintenance, of the lytic cycle. Another possibility is that the transcription factor activity, but not the overall expression level, of PRDM1 is important for lytic replication. Further study is clearly required to delineate this relationship.

In summary, we performed integrative analysis of publicly available single cell RNA-seq data from different LCLs to help resolve their heterogeneity. We identified a novel cluster of cells that are between lytic and latent stage, marked by LMP1 and controlled by HIF1a. We also found that the mRNA expression of super-enhancer target genes is inversely correlated with lytic status of the cells and consistently CRISPR perturbation of super-enhancers increased the expression of EBV lytic genes. Our studies revealed EBV associated heterogeneity among LCLs that contribute to EBV life cycle and biology.
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Competing interests: The authors have no competing interests to declare.
Methods and Materials

Cell culture
LCL-358 (catalog no. 1038-3754NV17, Astarte Biologies), GTEX-UPJH-0001-SM-3YRE9, GM12878, AKATA EBV positive and AKATA EBV negative cells were cultured in RPMI1640 (VWR-0105-0500) media supplemented with 10% fetal calf serum (Gibco or Hyclone), 100 unit/mL streptomycin and 100 mg/mL penicillin (Gibco or Life Technologies). HEK293T cells purchased from ATCC were cultured in Dulbecco modified Eagle medium supplemented with 10% fetal calf serum (Gibco), 100 unit/mL streptomycin and 100 mg/mL penicillin. All the cells were maintained at 37°C in a 5% CO₂ humidified chamber. Cells were routinely confirmed to be mycoplasma negative according to PCR Mycoplasma Detection Kit (ABM Inc.) and were used at low passage (<10) number but were not independently authenticated.

CRISPRi repression
Plasmid dCAS9-KRAB-MeCP2 (#110821) purchased from Addgene was packaged with lentiviruses and used to transduce LCLs for 2 days followed by selection with 5ug/ml blasticidin for another 5 days. The expression of dCAS9-KRAB-MeCP2 was verified by western blot. sgRNAs targeting genomic sites of interest were designed with online software Benchling (www.benchling.com). sgRNAs were annealed and cloned into Lenti Guide-Puro vector according to previously published protocol (76). LentiGuide-Puro containing sgRNAs were packaged into lentiviruses and were used to transduce LCLs stably expressing dCAS9-KRAB-MeCP2. Cells were selected with 3ug/ml puromycin for 3 days and then allowed to grow for another 2 days. The list of sgRNAs are provided in Table S3

qRT-PCR
Cells were harvested and washed once with cold PBS. Total mRNAs were extracted using PureLink RNA mini kit (Life Technologies) or Direct-zol RNA extraction kit with DNase I treatment (Zymo Research) following manufacturer’s instructions. mRNAs were then reverse transcript into cDNA with iScript™ Reverse Transcription Supermix (Bio-rad) or OneScript Plus cDNA Synthesis SuperMix (ABM Inc.). cDNAs were amplified on an CFX96 Touch real-time PCR detection system (Bio-Rad) and SYBG Green (Thermo Fisher) was used to detect cDNA amplification. All experiments were performed in triplicates in total reaction volumes of 15 μL using BrightGreen 2X qPCR MasterMix-No Dye (ABM Inc.). A housekeeping gene was used to normalize gene expression. RNA relative expression was calculated using the 2 ΔΔCT method. The value for the cells transduced with non-targeting sgRNA was set to 1. The list of all qPCR probes is provided in Table S3.

ChIP-qPCR
LCLs stably expressing dCAS9-KRAB-MeCP2 were transduced with lentiviruses expressing sgRNAs. Two days after transduction cells were selected with 3ug/ml puromycin for another 3 days. Cells were then collected and fixed with 1% formaldehyde. The cells were lysed and sonicated with bioruptor (Diagenode). Sonicated chromatin was diluted and precleared with protein A beads followed by incubation with 4ug H3K27ac (Abcam, #ab4729) or control antibodies with rotating at 4°C overnight. The next day, Protein A/salmon DNA beads (Millipore, #16-157) were used to capture protein–DNA complexes. After precipitation, beads
were washed with low salt wash buffer (1% Triton-X-100, 0.1% SDS, 2mM EDTA (pH8.0), 150mM NaCl, 20mM Tris-HCl (pH 8.0)) once, high salt wash buffer (1% TritonX-100, 0.1% SDS, 2mM EDTA (pH8.0), 500mM NaCl, 20mM Tris-HCl (pH 8.0)) twice, LiCl wash buffer (0.25M LiCl, 1% NP-40, 1% NaDOC, 1mM EDTA, 10mM Tris-HCl (pH 8.0)) once, and TE buffer (1mM EDTA, 10mM Tris-HCl (pH 8.0)) once. Each wash was performed by gently spinning down beads at 300g for one minute and re-suspend beads with 1ml wash buffer followed by shaking at 4°C for 5 minutes. DNA and protein complexes were eluted with elution buffer (1% SDS, 100mM NaHCO3). Protein–DNA complexes were reverse cross-linked with proteinase K (Thermo Fisher, #EO0491). DNA was purified by using QIAquick Spin columns (Qiagen, #28104). qPCR was used to quantify the DNA from ChIP assay and normalize it to the percent of input DNA.

**Induction of EBV**

AKATA EBV positive and negative cells were treated with IgG (Agilent, # A042301-2) at a final concentration of 0.5% followed by incubation at 37°C, 5% CO2 for 6 hrs. Cells were then centrifuged and re-suspended with fresh RPMI1640 supplemented with 10% FBS and continue culture for another 48 hrs. mRNAs were extracted by using PureLink RNA mini kit (Life Technologies), qRT-PCR was used to detect EBV lytic gene expression. To induce LMP1 expression, LCLs were treated with 100 nM of NEDD8 inhibitor - MLN4924 (Pevonedistat) (A gift from Dr. Liu) or DMSO control and were collected at indicated time points for qRT-PCR and/or Flow cytometry.

**Dual CRISPR mediated DNA deletion**

Dual gRNAs were designed with webtools from benchling (www.benchling.com) and were cloned into pLentiGuide-Puro (Addgene Plasmid #52963) using the Multiplex gRNA kit (System Biosciences) according to the manufacturer protocol. The success of gRNAs insertion was verified by sequencing with U6 promoter primer. HEK293T cells were used to package lentiviruses by co-transfecting viral packaging plasmids pCMV-VSV-G (Addgene #8454), psPAX2 (Addgene #12260) and the pLentiGuide-Puro vector containing the target sgRNAs. 18 hrs after transfection, media were changed to fresh RPMI media containing 30% of FBS. 24 and 48hrs later, supernatant containing lentivirus was collected and filtered with a 0.45 micron filter. LCLs in which Cas9 was stably expressed were transduced with the filtered lentivirus (Day 0) for 2 days, and then selected with 3ug/mL Puromycin for 3 days. On Day 5, genomic DNA was extracted using the DNeasy Blood & Tissue kit (Qiagen) and RNA was extracted with the PureLink RNA Mini kit (Ambion). Genomic deletions were verified by PCR using the PrimeSTAR polymerase (Clonetech). qRT-PCR was done using the Power SYBR Green RNA-to-CT 1-Step Kit (Applied Biosystems).

**Single-cell RNA sequencing analysis**

10x Genomics Cell Ranger 6.0.2 count (77) was used to align the raw sequencing reads to a customized human (GRCh38) and EBV (NC 007605, obtained from (61)) hybrid reference genome to generate barcode and UMI counts. Seurat (v4) (78) was applied for the downstream analysis and visualization of the data as following: Genes that were expressed in less than 3 cells were discarded. Cells with >20% of their unique molecular identifiers (UMIs) mapping to
mitochondrial genes or cells with <250 detected genes were discarded. Only cells with >80% log10 (Genes per UMI) were retained. Cell cycle score for each cell was calculated by Seurat function CellCycleScoring using human cell cycle genes. SCTransform was then used to normalize the dataset using default parameters while regressing out mitochondrial genes and cell cycle scores (S and G2M) and identify variable genes. Doublets were removed by R package DoubletFinder (v2.0) (79). Then, the datasets were integrated based on “anchors” identified among datasets (nfeatures = 2000, normalization.method = “SCT”) prior to perform linear dimensional reduction by Principal Component Analysis (PCA). The top 50 PCs were included in a Uniform Manifold Approximation and Projection (UMAP) dimensionality reduction. Clusters were identified on a shared nearest neighbor (SNN) graph the top 50 PCs with the Louvain algorithm using three resolutions (i.e., 0.1, 0.3, 0.5). The clusters corresponding to latent EBV life cycle were combined as one cluster for the downstream analyses. Differential gene expression was determined by “FindMarkers” function on SCT normalized expression values with the default Wilcox Rank Sum test either as one versus rest or as a direct comparison with default parameters except logfc.threshold = 0. The cell annotation was based on the EBV genes and top differentially expressed genes. Gene list module scores were calculated with Seurat function AddModuleScore (51). This calculates the average scaled expression levels of each gene list, subtracted by the expression of control feature sets (n=100). All the displayed expression values on violin plots, feature plots and dot plots are SCT normalized expression values. The IRF4 bound genes are sourced from the ChIP-Atlas “Target Genes” database (80) with options: “hg38” as the genome and “+/− 5kb” as distance from TSS. Target genes with binding score not less than 500 in GM12878 cells are selected. All genesets used in this study are provided in Table S2.

Gene set enrichment analysis (GSEA)
GSEA was performed using pre-ranked mode and “No Collapse” options. The pre-ranked gene lists were ranked by the SCT normalized expression fold-change between comparison groups. EBV-contacted and EBV-non-contacted genesets are curated from our previous study (23) and provided in Table S2.

Statistical analysis and data visualization
Statistical analyses were performed using GraphPad PRISM 9 (La Jolla, CA, USA) with the method detailed in the legend.

Flow cytometry
All stained/fixed samples were acquired on Attune NxT Flow Cytometer (Thermo Fisher Scientific) with necessary internal controls to help assign gates. Fluorescence from multiple antibodies were compensated using AbC Total Compensation beads (Thermo Fisher Scientific, catalog no. A10497). In all experiments, cells were collected and stained with fixable viability dye eFluor780 (1:2000 dilution Life Technologies, catalog no. 65-0865-14;) followed by surface staining for PDL1 (CD274; clone 29E.2A3; BioLegend catalog no. 329714; 1:60 dilution) as per the manufacturer's instructions. Cells were then fixed with 4% methanol-free formaldehyde (Thermo Fisher Scientific, catalog no. 28908) followed by intracellular staining for BZLF1 (Santa Cruz Biotechnology, catalog no. sc-53904; 1:60 dilution) and LMP1 (clone LMPO24; Novus Biologicals, catalog no. NBP2-50383; 1:60 dilution) using FoxP3/Transcription factor
staining buffer set (eBioscience, catalog no. 5523) as per manufacturer’s instructions. Data were analyzed using FlowJo and cumulated using GraphPad PRISM software.

Data sources and availability
The single cell RNA-seq data are sourced from GSE126321 for GM12878 and GM18502; GSE111912 for GM12891; GSE158275 for LCL777B958, LCL777M81 and LCL461B958; GSE162528 for LCL; and GSE121926 for GM22648 and GM22649. The ChIA-PET in GM12878 is from GSE127053. The ChIP-seq data are from EBNA2: GSE29498; EBNA1L: GSE49338; EBNA3A: GSM1429820; EBNA3C: GSE52632; NF-kB: GSE55105 and H3K27Ac: GSM733771.


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Figure Legends

**Figure 1.** Single cell RNA-sequencing analyses resolve LCLs into three distinct populations. (a) Integrated UMAP showing 3 major cell types w.r.t. EBV status in nine LCLs used in this study. (b) Numbers of differentially expressed genes (FC>1.5 and adjusted p-val < 0.05) in indicated cluster compared to other clusters. (c-d) mRNA expression of EBV genes across all clusters shown as dot plot (c) or projected on the UMAP (d). (e) mRNA expression of top 10 human host cell defining genes across all clusters. (f) Significantly enriched hallmark pathways by GSEA comparing transcriptomes of cells in indicated cluster with all other cells. The positive and negative enrichment scores indicate activation and inactivation of the indicated pathway in each cell cluster, respectively. Only pathways that are enriched (FDR<5%) in at least one of the clusters are shown.

**Figure 2.** *GP350–LMP1*hi LCLs have a HIF1A-associated signature. (a) mRNA expression of *HIF1A* or *CD274* genes across all clusters as dot plot. (b) Module score of HIF1A induced genes (left panel) or HIF1A-repressed genes (right panel). HIF1A induced and repressed genes are sourced from MSigDB (M1308). **** p<0.0001 by two-tailed Wilcoxon rank-sum test. (c) Enrichr based drugs predicted (out of 906 total drugs) to counteract genes induced in *GP350–LMP1*hi LCLs compared to other cells, ordered by adjusted p-value. Drugs are sourced from Enrichr library “Drug_Perturbations_from_GEO_down”. (d-e) mRNA expression of indicated host (d) or EBV (e) genes in LCLs treated with 100nM DMSO or Pevonedistat. UBC was used as a housekeeping gene. (f-k) flow cytometry on BLCL-358 treated with 100nM DMSO or Pevonedistat for 72hr. Plots showing cell viability and LMP1, BZLF1 or PDL1 expression in LCLs treated with DMSO or Pevonedistat. Shown are cumulative %viability plots (f), representative flow cytometry plots (g) and cumulative data showing %LMP1+ (h) and %BZLF1+ (i) in gated live LCLs. (j-k) Representative PDL1 expression as mean fluorescent intensity (MFI) or cumulative %PDL1+ in gated live LCLs. Data in (d-k) are from n=3 or n = 4 independent experiments; gating strategy is shown in Fig. S2d. * p<0.05; ** p<0.01; *** p<0.001; **** p<0.0001 by two-tailed paired ratio t-test.

**Figure 3.** *GP350–LMP1*lo LCLs have distinct MYC-dependent transcriptional regulation. (a) fraction of differentially expressed genes (Fig. 1b) from each indicated cluster or all human genes (n=35541) that are bound by MYC. MYC bound genes (n=3534) in GM12878 are obtained from (81). *** p<0.001 by fisher exact test. (b) MYC mRNA expression across all clusters. **** p<0.0001 by two-tailed Wilcoxon rank-sum test. (c) Module score of MYC induced genes (left panel) or MYC-repressed genes (right panel). **** p<0.0001 by two-tailed Wilcoxon rank-sum test. MYC induced or repressed genes are sourced from (55). (d) GSEA plots comparing transcriptomes of indicated clusters for the enrichment in hallmark of MYC targets. NES: Normalized enrichment scores. FDR: False Discovery Rate.

**Figure 4.** Super-enhancer-regulated genes are less highly expressed in *GP350*+ LCLs. (a) Module score of SE containing genes in GM12878. SEs and their annotation are sourced from (26). (b) mRNA expression of select SE containing genes across all cell types. (c) GSEA plots comparing transcriptomes of *GP350*+ and *GP350*− cells for enrichment in genes neighboring SEs with (left panel) or without (right panel) EBV contacts. NES: Normalized enrichment scores. (d) Module score of n=500 top IRF4 (left panel) and RUNX3 (right panel) bound genes. ****
p<0.0001 by two-tailed Wilcoxon rank-sum test. (e-f) mRNA expression of indicated EBV (e) or host (f) genes in EBV⁺-AKATA cells with or without anti-IgG (1:200) treatment after 48 hours. (g) Control EBV⁻-AKATA cells are included when measuring host genes. Data are from n=3 independent experiments. * p<0.05; ** p<0.01; *** p<0.001 by two-tailed unpaired t-test. (h) Western blots of lysates of AKATA cells treated with or without anti-IgG after 48 hours. Shown are representative (n=2) images of ZTA (BZLF1), BMRF1 and MYC with GAPDH as loading control.

**Figure 5. Disruption of super-enhancers in LCLs induces EBV lytic reactivation. (a,c,e,g)** Genome browser tracks showing EBV transcription factors EBNA2, EBNALP, EBNA3A and EBNA 3C, host transcription factors RELA, RELB, c-REL, p50 and p52 and H3K27ac at the MYC (a), IRF4/DUSP22 (c), MIR155HG (e) and RUNX3 (g) loci. The CRISPR cleavage site or inactivation site is highlighted with vertical blue box. The expected affected target genes are highlighted by vertical red box. (b,d,f,h) mRNA expression of select host and EBV genes after CRISPR mediated knockout (b) or inactivation (d,f,h) of indicated site (bottom black triangle in a,c, e or g) in GM12878 cells. Data are from n=3 independent experiments. * p<0.05, ** p<0.01, *** p<0.001 by two-tailed unpaired t-test.
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