

1 **c-Myc Represses Transcription of the Epstein-Barr Virus Latent Membrane Protein 1**
2 **Early After Primary B Cell Infection**

3 Running Title: c-Myc represses EBV LMP1 transcription (38/54 Characters)

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13 **ABSTRACT**

14 Recent evidence has shown that the EBV oncogene LMP1 is not expressed at high levels
15 early after EBV-infection of primary B cells, despite its being essential for the long-term outgrowth
16 of immortalized lymphoblastoid cell lines (LCLs). In this study, we found that expression of LMP1
17 increased fifty-fold between seven days post infection and the LCL state. Metabolic labeling of
18 nascently transcribed mRNA indicated this was primarily a transcription-mediated event. EBNA2,
19 the key viral transcription factor regulating LMP1, and CTCF, an important chromatin insulator,
20 were recruited to the LMP1 locus similarly early and late after infection. However, the activating
21 histone H3K9Ac mark was enriched at the LMP1 promoter in LCLs relative to early-infected B
22 cells. We found that high c-Myc activity in EBV-infected lymphoma cells as well as overexpression
23 of c-Myc in an LCL model system repressed LMP1 transcription. Finally, we found that chemical
24 inhibition of c-Myc expression both in LCLs and early after primary B-cell infection increased
25 LMP1 expression. These data support a model in which high levels of endogenous c-Myc activity
26 induced early after primary B-cell infection directly represses LMP1 transcription.

27 **IMPORTANCE**

28 EBV is a highly successful pathogen that latently infects greater than 90% of adults
29 worldwide and is also causally associated with a number of B-cell malignancies. EBV expresses
30 a set of viral oncoproteins and non-coding RNAs during the latent life cycle with the potential to
31 promote cancer. Critical among these is the viral latent membrane protein, LMP1. Prior work
32 suggests that LMP1 is essential for EBV to immortalize B cells, but our recent work indicates that
33 LMP1 is not produced at high levels during the first few weeks after infection. Here, we show that
34 the transcription of LMP1 can be negatively regulated by a host transcription factor, c-Myc.
35 Ultimately, understanding the regulation of EBV-encoded oncogenes will allow us to better treat
36 cancers that rely on these viral products for survival.

37 INTRODUCTION

38 Epstein-Barr virus (EBV) infection of primary human B cells leads to their immortalization,
39 or growth transformation (1). These immortalized cells, called lymphoblastoid cell lines (LCLs),
40 are latently infected by EBV and express a program, called latency III, consisting of nine viral
41 proteins and many non-coding RNAs. LCLs expressing Latency III genes are a model system for
42 EBV-associated malignancies such as Post-Transplant Lymphoproliferative Disease (PTLD) and
43 Diffuse Large B Cell Lymphoma (DLBCL) (1). Characteristic of Latency III is expression of the
44 essential viral oncogene latent membrane protein 1 (LMP1), which is a constitutively active TNF
45 receptor homologue that induces the host NF κ B and AP-1 signaling pathways to promote survival
46 (2-5). LMP1 is both necessary for EBV immortalization of primary human B cells and sufficient to
47 induce lymphomas when expressed in the murine B cell compartment (6, 7). The regions of the
48 LMP1 cytoplasmic tail that engage TNFR associated factors (TRAFs) and TNFR associated death
49 domain (TRADD) responsible for NF κ B activation are coincident with the regions required for
50 EBV-mediated immortalization (8, 9). LCLs are functionally addicted to the NF κ B signaling
51 induced by LMP1 as inhibition of this pathway results in apoptosis (3).

52 Studies of LMP1 transcriptional regulation have defined cis-acting elements and trans-
53 acting factors important in controlling its expression. In epithelial cells, LMP1 expression is driven
54 by a unique promoter found within the viral terminal repeats (10, 11). However, transcription of
55 LMP1 in Latency III expressing B cells is initiated at a bi-directional LMP1/LMP2B promoter (12).
56 The critical activator of this Latency III promoter is Epstein-Barr Nuclear Antigen 2 (EBNA2), which
57 binds to a so-called EBNA2-response element (E2RE) through the host factors RBP-J κ and PU.1
58 (13, 14). Other viral factors, such as EBNA3C, have also been shown to aid in the transcription
59 of LMP1 (15, 16). Additionally, a number of host factors have been shown to fine-tune LMP1
60 transcription, including ATF4 (17, 18), IRF7 (19), and NF κ B subunits themselves (20, 21). In fact,
61 host factors often lead to auto-regulatory feedback loops that maintain the levels of LMP1

62 expression, whether it is NF κ B in B cells (20) or STAT signaling on the terminal repeat promoter
63 in epithelial cells (22). Furthermore, the epigenetic state of the LMP1 promoter plays a key role in
64 transcription. Changes in cell type and latency state alter the prevalence of active and repressive
65 histone marks on the LMP1 promoter as well as CpG DNA methylation (23). In addition, viral
66 chromatin architecture and enhancer looping mediated by host factors such as CTCF and Rad21
67 play a direct role in the full activation of LMP1 (24).

68 LMP1 mRNA is also post-transcriptionally regulated to control protein output. While LMP1
69 is one of the most abundant viral transcripts detected in latently infected cells, it is transcribed at
70 a much lower rate implying a certain level of mRNA stability (25). This stability can be
71 counteracted by micro-RNA (miRNA) targeting of the long LMP1 3' untranslated region. LMP1
72 can be targeted by both EBV-encoded viral miRNAs such as miR-BART1-5p, miR-BART16-3p,
73 and miR-BART17-5p (26) as well as a number of host miRNAs (27). Of these host miRNAs,
74 inhibiting the c-Myc regulated miR17~92 miRNA cluster led to an upregulation of LMP1 protein
75 levels and slowed cell growth (27). This slowed growth could be explained in part by the known
76 ability of overexpressed LMP1 to be cytostatic (28-30).

77 The cellular oncogene c-Myc is overexpressed in a wide variety of cancers, including EBV-
78 associated Burkitt lymphoma where c-Myc is expressed at high levels due to a characteristic
79 chromosomal 8:14 translocation. Additionally, EBV induces c-Myc expression upon infection of B
80 cells by utilizing EBNA2 to co-opt native super-enhancer architecture upstream of c-Myc (31).
81 Upon *de novo* infection, c-Myc levels peak within the first week after infection, then wane, but
82 remain elevated throughout LCL outgrowth where c-Myc is crucial for the maintenance of the LCL
83 phenotype (32, 33). This conflicts with the NF κ B addiction observed in LCLs, as the c-Myc and
84 NF κ B signaling pathways have been shown to be directly incompatible (34).

85 We have previously shown that, despite being detectable early after infection of primary
86 B cells, LMP1 does not reach LCL levels of expression until greater than two weeks post infection
87 (35). These EBV-infected cells express EBNA2, proliferate, and show no signs of apoptosis even
88 when NF κ B is inhibited during the first weeks of infection (35, 36). Furthermore, the exact role of
89 LMP1 in the EBV life cycle has been called into question by recent work showing that LMP1 is
90 dispensable for tumor formation in a humanized mouse model (37, 38). In this work, we address
91 the question of how LMP1 is delayed in expression from early to late times after primary B-cell
92 infection. We investigate the nature of the mRNA change and the role of cellular factors in
93 temporal regulation of LMP1 expression.

94

95 RESULTS

96 **LMP1 transcription is robustly increased from early to late times after primary B cell**
97 **infection.** LMP1 mRNA and protein levels and, consequently, NF κ B targets are significantly lower
98 during the first two weeks following EBV infection of primary B cells relative to that found in
99 immortalized lymphoblastoid cell lines (LCLs) (35). Prior studies indicate that the LMP1 mRNA is
100 the most abundant latency transcript in LCLs despite being poorly transcribed (25). Therefore, to
101 determine the mechanism for low LMP1 mRNA levels at early times after infection, we assayed
102 the relative transcription rate and half-life using a 4-thiouridine (4sU) metabolic labeling approach
103 (**Fig. 1A** and (39)). This was performed by using Fluorescence Activated Cell Sorting (FACS) to
104 generate a pure population of proliferating (Celltrace Violet^o) EBV-infected B cells six days after
105 primary human Peripheral Blood Mononuclear Cell (PBMC) infection and allowing the cells to rest
106 overnight. Twenty-four hours later (Day 7 post infection), cells were pulsed with 4sU for exactly
107 one hour before total RNA was harvested. The process was repeated for LCLs that grew out from
108 matched PBMC donors five weeks post infection. As previously observed, the total LMP1 mRNA
109 level increased ~50-fold from day 7 post infection through LCL outgrowth (**Fig. 1B**). Comparing
110 the ratio of 4sU labeled nascent RNA and unlabeled decaying RNA (39), we found that the half-
111 life of the LMP1 mRNA increased two-fold between early and late times after infection from ~2
112 hours at 7 days post infection to ~4 hours in LCLs (**Fig. 1C**). Over the same time frame the relative
113 transcription rate of LMP1 mRNA increased nearly 25-fold (**Fig. 1D**). We also queried the EBNA2-
114 specific mRNA as well as Cp-derived EBNA transcripts and found only modest differences in their
115 overall expression, transcription rate, or stability through B cell outgrowth (**Fig. 1B-D**).

116 **The LMP1 locus is occupied by EBNA2 and CTCF similarly early and late after infection;**
117 **however lower levels of activated histones are detected early.** In B cells, the transcription of
118 LMP1 is regulated by EBNA2 (40). Prior work indicates that EBNA2-regulated genes, including c-
119 Myc and CD23, are induced to higher levels at day 7 post infection than in LCLs suggesting that

120 EBNA2 activity is not broadly suppressed during early infection (32). However, it remained a
121 possibility that LMP1 expression was low during early infection because of poor EBNA2
122 recruitment to the bi-directional LMP1/2B promoter. To address this, we performed Chromatin
123 Immuno-precipitation (ChIP) for EBNA2 at day 7 post infection and in LCLs. LMP1 and LMP2 are
124 transcribed from a complex and overlapping region of the EBV episome near the terminal repeats
125 (TR), diagrammed in **Fig. 2A**. We found that EBNA2 was recruited comparably to the LMP1
126 promoter and the viral C promoter both early after infection and in LCLs (**Fig. 2B**). A region in the
127 gene body of EBNA3C distal to both the LMP1p and the Cp was used as a negative control (**Fig.**
128 **2B**). Similarly, EBNA2 was recruited to the two major c-Myc upstream enhancer loci (31) as well
129 as the CD23 promoter comparably at 7 days post infection as in LCLs (**Fig. 2C**). Despite the lack
130 of change in EBNA2 occupancy at early and late times post infection, the activating histone mark
131 H3K9 Acetyl was enriched at the LMP1p in LCLs, correlating with the observed increased level
132 of transcription (**Fig. 2D**). A recent publication has implicated the chromatin architecture mediated
133 by CTCF at a single CTCF site downstream of LMP1 (here called the CTCF-Response Element)
134 to be important for H3K9 Acetylation and transcription at the LMP1 promoter (24). We assayed
135 for CTCF-occupancy at both this CTCF-RE as well as the EBV enhancer found in the EBER/OriP
136 region of the genome and found that CTCF occupancy did not change between day 7 post
137 infection cells and LCLs (**Fig. 2E**).

138 **c-Myc suppresses LMP1 transcription.** Despite EBNA2 recruitment to the LMP1 promoter,
139 reduced H3K9 acetylation suggested poor LMP1 transcription. Early after infection when LMP1
140 expression is lowest, c-Myc is strongly activated by EBNA2, and during the time frame that LMP1
141 expression increases the expression of c-Myc and its targets are subsequently attenuated (32).
142 Given this correlation, we sought to test the role of c-Myc in the regulation of LMP1 expression.
143 We first assayed LMP1 transcription using 4sU labeling in EBV-infected Burkitt lymphoma (BL)
144 cell lines that express very high levels of c-Myc relative to LCLs. We found that LMP1 mRNA and

145 protein levels were higher in LCLs than in EBV-infected BL41 cells (**Fig. 3A-B**). Consistent with
146 our hypothesis, the transcription of LMP1 was significantly lower in EBV-infected BL41 cells
147 relative to LCLs (**Fig. 3C**).

148 To directly assess the effect of c-Myc on LMP1 transcription in an LCL, we used the P493-
149 6 model of c-Myc and EBNA2 regulation (41, 42). In this system, EBNA2 (encoded endogenously
150 from the viral genome) is controlled post-translationally by Estrogen (β -Estradiol) due to its fusion
151 to a modified estrogen receptor, and heterologous c-Myc expression is provided *in trans* and
152 controlled transcriptionally through a tetracycline (tet-off) system. We verified that induction of
153 EBNA2 and inactivation of c-Myc (Estrogen+, Tetracycline+) led to an LCL-like phenotype with
154 high levels of LMP1 transcription (**Fig. 4A**). However, when removing EBNA2 activity (Estrogen-
155 , Tetracycline-) or inducing EBNA2 in the presence of high levels of c-Myc (Estrogen+,
156 Tetracycline-), we observed blunted endogenous LMP1 transcription relative to that induced in
157 the presence of low levels of c-Myc (**Fig. 4A**). These changes in transcription correlated with the
158 levels of total LMP1 mRNA and protein (**Fig. 4B-C**), while EBNA2-ER transcription from the
159 endogenous Cp was not affected by c-Myc overexpression (**Fig. 4B**). This controlled experiment
160 indicated that c-Myc overexpression is sufficient to suppress LMP1 transcription in an LCL and
161 suggested that c-Myc may be critical for suppression of LMP1 transcription early after primary B-
162 cell infection as well.

163 **c-Myc suppresses LMP1 expression early after primary B-cell infection.** To test the
164 hypothesis that c-Myc is responsible for LMP1 suppression early after infection, we chose to
165 target c-Myc pharmacologically given the intractability of reverse genetic experiments in primary
166 human B cells. The expression of c-Myc is exquisitely sensitivity to inhibition of bromodomain and
167 extra-terminal motif (BET) domain containing transcriptional activators including Brd4 (43, 44).
168 Therefore, we used two independent BET inhibitors, JQ1 and OTX015, to suppress c-Myc and
169 assay LMP1 transcription in EBV-infected cells. First, we treated LCLs with the BET inhibitors and

170 found that c-Myc mRNA levels decreased and LMP1 mRNA levels increased in a dose-dependent
171 manner (**Fig. 5A-B**). We next treated EBV-infected PBMCs 7 days post infection with BET
172 inhibitors. Consistently, we observed a dose-dependent decrease in c-Myc mRNA and increase
173 in LMP1 mRNA following both JQ1 and OTX015 treatment (**Fig. 5C-D**). Therefore, c-Myc activity
174 negatively correlates with LMP1 expression both early and late after infection.

175 **DISCUSSION**

176 In this study, we determined that transcription of the EBV major latent oncoprotein, LMP1,
177 is suppressed by c-Myc early after primary B cell infection. Transcription of LMP1 increased
178 twenty-five-fold from day 7 to LCL, while the mRNA half-life increased two-fold, leading to an
179 overall fifty-fold increase in LMP1 expression. This change correlated with an approximate two-
180 fold increase in the activating histone mark, H3K9Ac, at the LMP1 promoter, but the occupancy
181 of the major LMP1 trans-activator EBNA2, as well as CTCF-binding to the CTCF-RE, did not
182 change over the same time period. Furthermore, cellular c-Myc was shown to negatively regulate
183 LMP1 transcription and total LMP1 RNA levels in both the setting of highly expressed endogenous
184 c-Myc in a Burkitt lymphoma cell line as well as direct c-Myc overexpression in an LCL. Finally,
185 inactivating c-Myc expression with either of the BRD4-inhibiting compounds, JQ1 or OTX015, led
186 to a dose dependent increase in LMP1 expression both in LCLs and early after primary infection
187 of B cells.

188 The small but significant change in LMP1 half-life could be explained as a post-
189 transcriptional effect mediated by miRNAs that are known to target the LMP1 3'UTR. Previous
190 work has shown that the miR17~92 family of c-Myc-induced miRNAs target LMP1 and removal
191 of this miRNA family increased LMP1 protein production (27). Furthermore, our group has shown
192 that miR17~92 are expressed at the highest levels early after infection when LMP1 mRNA and
193 mRNA half-life is lowest (45). This low level "fine-tuning" of LMP1 expression might be critical in

194 the immortalized state when high levels of LMP1 can lead to cytostatic effects on cell growth (28,
195 30).

196 While the LMP1 promoter has been extensively studied in the immortalized LCL, no one
197 has previously characterized promoter occupancy early after infection. We have shown that early
198 after infection EBNA2 is recruited normally to its response element in the LMP1 promoter as well
199 as several canonical cellular targets including CD23 and c-Myc. This implies that there must be a
200 repressive element present in the LMP1p that supersedes the E2RE's function at early times after
201 infection. Such an element has been characterized in Burkitt lymphoma cell lines, and it is
202 responsive to members of the Myc family including MAD1 and MAX (46). Alternatively, c-Myc
203 activation could lead to decreased LMP1p occupancy and transcription through less direct
204 mechanisms. Previously c-Myc signaling and NF κ B signaling have been shown to activate
205 opposing and mutually exclusive growth programs (34). In this way, direct actions of c-Myc or
206 indirect actions of Myc-induced target genes might lead to less active NF κ B transcriptional
207 subunits that can no longer auto-regulate and act in a feed-forward manner on the LMP1p (20).

208 Many questions still remain regarding the functional relevance for EBV in delaying LMP1
209 expression. While still important for survival at late stages during infection, it has been shown that
210 EBV can protect B cells from apoptosis in the absence of NF κ B activity by activating the anti-
211 apoptotic cellular MCL-1 protein using EBNA3A (36). Additionally, our group has shown that
212 exogenously activating NF κ B during the early stage after infection when LMP1 is low increases
213 the transformation rate of the virus (35). However, if the ability of c-Myc to downregulate LMP1
214 expression was detrimental, one would assume that the virus would evolve to lack such
215 constraints.

216 Alternatively, we propose c-Myc repression of LMP1 may be beneficial in the life cycle of
217 EBV to maintain low levels of LMP1-induced NF κ B early after infection. It has been shown that c-

218 Myc overexpression suppresses recognition of EBV-infected cells by CD8+ T cells (47, 48). In
219 contrast, NF κ B activity is known to enhance MHC-mediated antigen presentation, making LCLs
220 excellent targets for CD8+ T cell killing (49). Thus, NF κ B activation in EBV-infected B cells in an
221 immune-competent host might ride a fine line between survival and immune recognition. Given
222 our data, we propose that early EBV-infected B cells with high c-Myc and low NF κ B activity
223 provide an ideal setting to escape CD8+ T-cell recognition. This is particularly important in sero-
224 positive individuals where reactivation and *de novo* naïve B-cell infection is thought to occur in
225 the setting of a robust CD8+ T-cell response. The re-seeding of the latency reservoir from these
226 naïve-infected cells into the memory B-cell compartment would then proceed with attenuated
227 MHC presentation of viral antigens. Therefore, c-Myc-suppressed LMP1/NF κ B activity together
228 with the recently described EBV miRNA-mediated attenuation of T-cell recognition and killing (50,
229 51) are likely the key elements of latent EBV-mediated immune evasion in the immune-competent
230 host.

231

232

233 **MATERIALS AND METHODS**

234 **Cell lines, culture conditions, and viruses**

235 Buffy coats were obtained from normal human donors through the Gulf Coast Regional
236 Blood Center (Houston, TX) and peripheral blood mononuclear cells (PBMCs) were isolated by
237 Ficoll Histopaque-1077 gradient (Sigma, H8889). B95-8 strain of Epstein-Barr virus was produced
238 from the B95-8 Z-HT cell line as previously described (52). Virus infections were performed in
239 bulk by adding 50 μ L of filtered B95-8 supernatant to 1×10^6 PBMCs.

240 Cell lines were cultured in RPMI 1640 media supplemented with 10-15% heat inactivated
241 fetal bovine serum (Corning), 2 mM L-Glutamine, 100 U/ml penicillin, 100 μ g/ml streptomycin
242 (Invitrogen), and 0.5 μ g/mL Cyclosporine A (Sigma). P493-6 cells (a kind gift of Dr. Georg
243 Bornkamm, Helmholtz Zentrum München) were cultured with 10% tetracycline-free FBS (Hyclone
244 SH30070), 1 μ M β -Estradiol, and 1 μ g/mL Tetracycline. All cells were cultured at 37°C in a
245 humidified incubator at 5% CO₂.

246 **Flow cytometry and sorting**

247 To track proliferation, cells were stained with CellTrace Violet (Invitrogen, C34557), a
248 fluorescent proliferation-tracking dye. Cells were first washed in FACS buffer (5% FBS in PBS),
249 stained with the appropriate antibody for 30min-1hr at 4°C in the dark, and then washed again
250 before being analyzed on a BD FACS Canto II.

251 Proliferating infected B cells were sorted to a pure population of CD19⁺/CellTraceViolet^{lo}
252 on a MoFlo Astrios Cell Sorter at the Duke Cancer Institute Flow Cytometry Shared Resource.
253 Mouse anti-human CD19 antibody (clone 33-6-6; gift from Tom Tedder, Duke University Medical
254 School) conjugated with either APC or PE was used as a surface B cell marker in flow cytometry.

255

256 **Nascent RNA extraction and profiling**

257 To assess both mRNA transcription and half-life cells were treated with 4-thiouridine (4sU,
258 Sigma) (200 μ M) for exactly one hour. Upon harvesting, total RNA was extracted via TRIzol
259 following the manufacturer's protocol (Life Technologies). 4sU-labelled nascent RNA was then
260 biotinylated using a highly efficient crosslinking reaction using EZ-Link Biotin-HPDP (Pierce)
261 dissolved in DMSO at a concentration of 1 mg/mL or MTSEA Biotin-XX (Biotium Cat# 90066) at
262 20 ng/ μ L dissolved in DMSO, and labeled RNA was separated from the total population using
263 streptavidin MyOne C1 Dynabeads (Invitrogen) as previously described (39). Subsequently, three
264 populations of RNA were reverse transcribed into cDNA using the High Capacity cDNA kit
265 (Applied Biosystems): total RNA (T), unlabeled RNA (U), and nascent RNA (N). Quantitative real-
266 time PCR was then performed on these three populations to discover total RNA abundance,
267 relative transcription rates, and mRNA half-life as previously described (53). In brief, the
268 abundance of mRNA was represented by the measurement of the total RNA sample (T). Relative
269 transcription was represented by the labeled fraction of mRNA (N). The decay rate (DR) was
270 calculated from measurements of nascent (N) and unlabeled (U) mRNA, as a function of N/U -
271 $\ln(1 - N/U)$. An apparent RNA half-life was calculated using the decay rate, $-t \times [\ln(2)/DR]$, where
272 t is the time of 4sU incorporation (1 h for the purposes of these experiments). Two assumptions
273 of this method are that transcription and stability are constant over the period of measurement.

274 **Chromatin Immunoprecipitation**

275 Chromatin Immunoprecipitation was performed using ChIP-IT High Sensitivity kit using the
276 manufactures directions (Active Motif). DNA was sonicated for 45 min with 30 second on/off
277 cycles on a Bioruptor (Diagenode). ChIP antibodies include EBNA2 (PE2; gift from Elliot Kieff),
278 H3K9ac (Active Motif cat # 51252 Clone 1B10) and CTCF (Active Motif cat #61311). Quantitative
279 real-time PCR primer sets include the C promoter (F: CCTAGGCCAGCCAGAGATAAT, R:

280 AGATAGCACTCGACGCACTG), LMP1 promoter (F: GGCCAAGTGCAACAGGAA, R:
281 GCAGATTACACTGCCGCTTC), c-Myc Enhancer 525 kb upstream (F:
282 CTAGTAGCAGGTGATGGGTTATG, R: CCTTTGGACCAGAAGAGGATG), c-Myc Enhancer 432
283 kb upstream (F: ACAGCCAGAGGTATTGGAAC, R: GGAAGGAACGAAACCCTAGAA), CD23
284 promoter (F: GATCGGCCATAGTGGTATGATT, R: CTCAGGTAAGAGAATTGGGTGAG), CTCF-
285 RE (F: CCACTAGGAACCCAAGATCAA, R: GCCCGCTTCTTCGTATATGT), and EBER/OriP (F:
286 GGGAAATGAGGGTTAGCATAGG, R: CAAGTCTACATCTCCTCAAGACAG. As a negative
287 control for the EBV genome EBNA3C (F: CAAGGTGCATTTACCCCACTG and R:
288 GGGCAGGTCCGTGAGAACT) was used. As negative controls from the human genome two
289 regions Neg1 (F: CCAATAACAGAAGCATTAAAATTCA, R: TTCAAGCACAGGCATACAGG) and
290 Neg2 (F: TCTCTGGGGAGATGGATTACA, R: CGTGAATCCTTTATTCTTGGA) were used.

291 **Gene Expression Analysis**

292 Total RNA was isolated from cells by using a Qiagen RNeasy kit and then reverse
293 transcribed to generate cDNA with the High Capacity cDNA kit (Applied Biosystems). Quantitative
294 PCR was performed by using SYBR green (Quanta Biosciences) in an Applied Biosystems Step
295 One Plus instrument. Primer sets include c-Myc (F: CTCCATGAGGAGACACCGC, R:
296 GAGCCTGCCTCTTTTCCACA), LMP1 (F: AATTTGCACGGACAGGCATT, R:
297 AAGGCCAAAAGCTGCCAGAT), EBNA2 (F: GCTTAGCCAGTAACCCAGCACT, R:
298 TGCTTAGAAGGTTGTTGGCATG), GAPDH (F: TGCACCACCAACTGCTTAGC, R:
299 GGCATGGACTGTGGTCATGAG) and SETDB1 (F: TCCATGGCATGGTGGAGCGG, R:
300 GAGAGGGTTCTTGCCCCGGT).

301 **Western blot**

302 Cells were pelleted and washed in PBS, and then lysed in 0.1% Triton-containing buffer
303 with Complete protease inhibitors. All protein lysates were run on NuPage 4–12% gradient gels

304 (LifeTechnology) and transferred to PVDF membrane (GE Healthcare). Membranes were blocked
305 in 5% milk in TBST and stained with primary antibody overnight at +4°C, followed by a wash and
306 staining with secondary HRP-conjugated antibody for 1 hour at room temperature. Antibodies
307 include LMP1 (S12; gift from Elliot Kieff, Harvard Medical School), c-Myc (Santa Cruz
308 Biotechnology, SC-764), and GAPDH (BioChain Institute, #Y3322).

309 **BRD4 Inhibition**

310 (+)-JQ1 and OTX015 were purchased from Selleckchem (cat. #S7110 and #S7360, respectively).

311 For LCLs:

312 LCLs were seeded at 3×10^5 /mL in the presence of either 0.1% DMSO, 100 or 500 nM JQ1 or
313 100 or 500 nM OTX015 for 24 hours. After 24 hours total mRNA was harvested using the Qiagen
314 RNeasy mini kit (ref #74104) according to manufacturer's instructions. 1 microgram of harvest
315 mRNA was reverse transcribed into cDNA using Applied Biosystems High-Capacity cDNA
316 reverse transcription kit (cat #4368814) according to manufacturer's instructions. RT-qPCR
317 analysis was performed using SYBR Green detection based system (Quanta Bio cat #95072-
318 05K) with 5 ng of cDNA used per reaction using an Applied Biosystems StepOne Plus Real Time
319 PCR System. All values were normalized to the endogenous loading control SETDB1. Relative
320 expression values were calculated using the $\Delta\Delta$ CT method.

321 For PBMCs:

322 Peripheral Blood Mononuclear Cells (PBMCs) were isolated from whole blood, infected with EBV
323 B95-8 and seeded at 1×10^6 /mL. Seven days post infection the cells were treated with either
324 0.1% DMSO, 100 or 500 nM JQ1 or 100 or 500 nM OTX015. After 24 hours, total mRNA was
325 harvested as previously described. 1 μ g of total RNA was reverse transcribed into cDNA as
326 previously described. RT-qPCR analysis was conducted as previously described.

327

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336

337 **FIGURE LEGENDS**

338 **Figure 1. 4sU metabolic pulsing reveals differences in LMP1 mRNA abundance, half-life**

339 **and transcription rate at Day 7 compared to LCL (A)** Schematic describing 4sU experiments.

340 Cells are pulsed for 1 hour with 4sU before RNA isolation by Trizol. 4sU labeled RNA is then

341 conjugated to biotin and isolated via magnetic streptavidin. Unbound RNA is saved as the “old”

342 fraction while bound RNA eluted from the beads becomes the “new” fraction. Isolated RNA is then

343 reverse transcribed for RT-qPCR analysis. (B) RT-qPCR analysis of total mRNA from 4sU

344 experiments for indicated genes at indicated times post infection. (C) RT-qPCR analysis of mRNA

345 half-life from 4sU experiments. Half-life is normalized to GAPDH. (D) Transcription rate analysis

346 of mRNA from 4sU experiments for indicated genes at indicated times post infection. All

347 transcription rates are relative to GAPDH. For all panels, Cp denotes EBV C promoter transcripts.

348 Each bar is representative of at least 3 independent donors. Error bars represent SEM. * denotes

349 $p < 0.05$, ** denotes $p < 0.01$ and *** denotes $p < 0.001$ by one-tailed pairwise student's t-test.

350 **Figure 2. Interrogation of the LMP1 locus by ChIP-PCR.** (A) Schematic of the LMP1 genomic

351 locus. Arrows with half arrow heads denote the location of primers used for ChIP-PCR in (B), (D)

352 and (E). (B) Sorted Day 7 proliferating primary EBV-infected B cells (black) or fully outgrown (>35

353 days) LCLs (blue) were harvested for chromatin and ChIP for EBNA2 was performed. LMP1p

354 denotes the LMP1 promoter, Cp denotes the C promoter and EBNA3C is an EBV genomic

355 negative control. All samples are normalized to the average of the two negative controls from the

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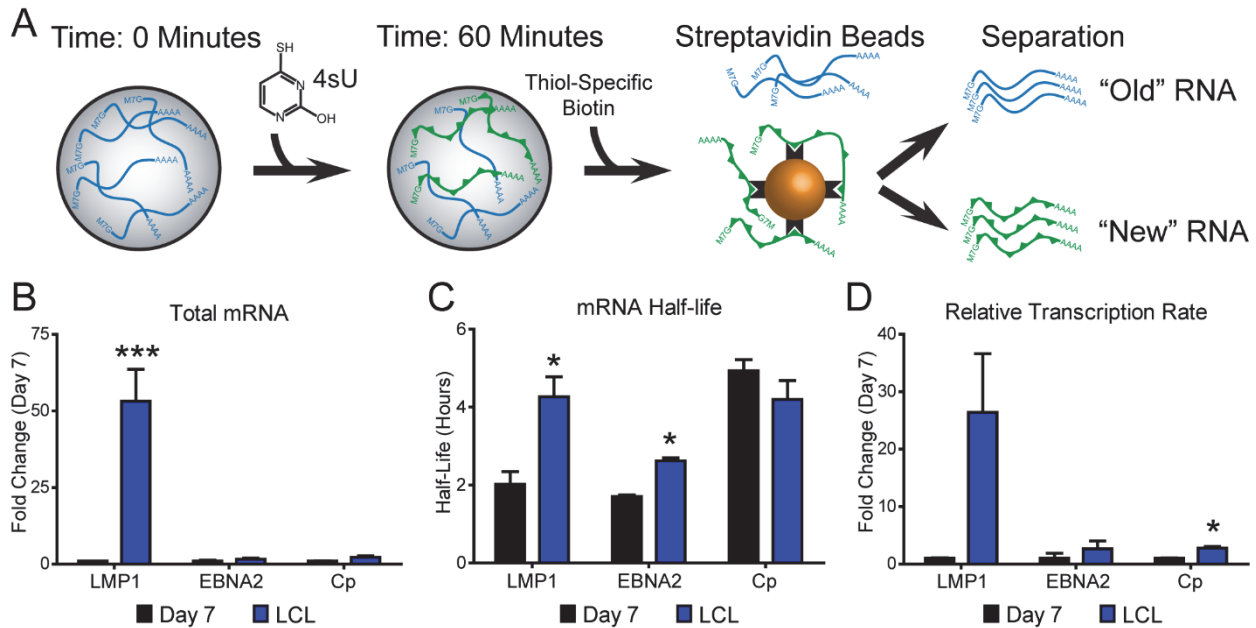
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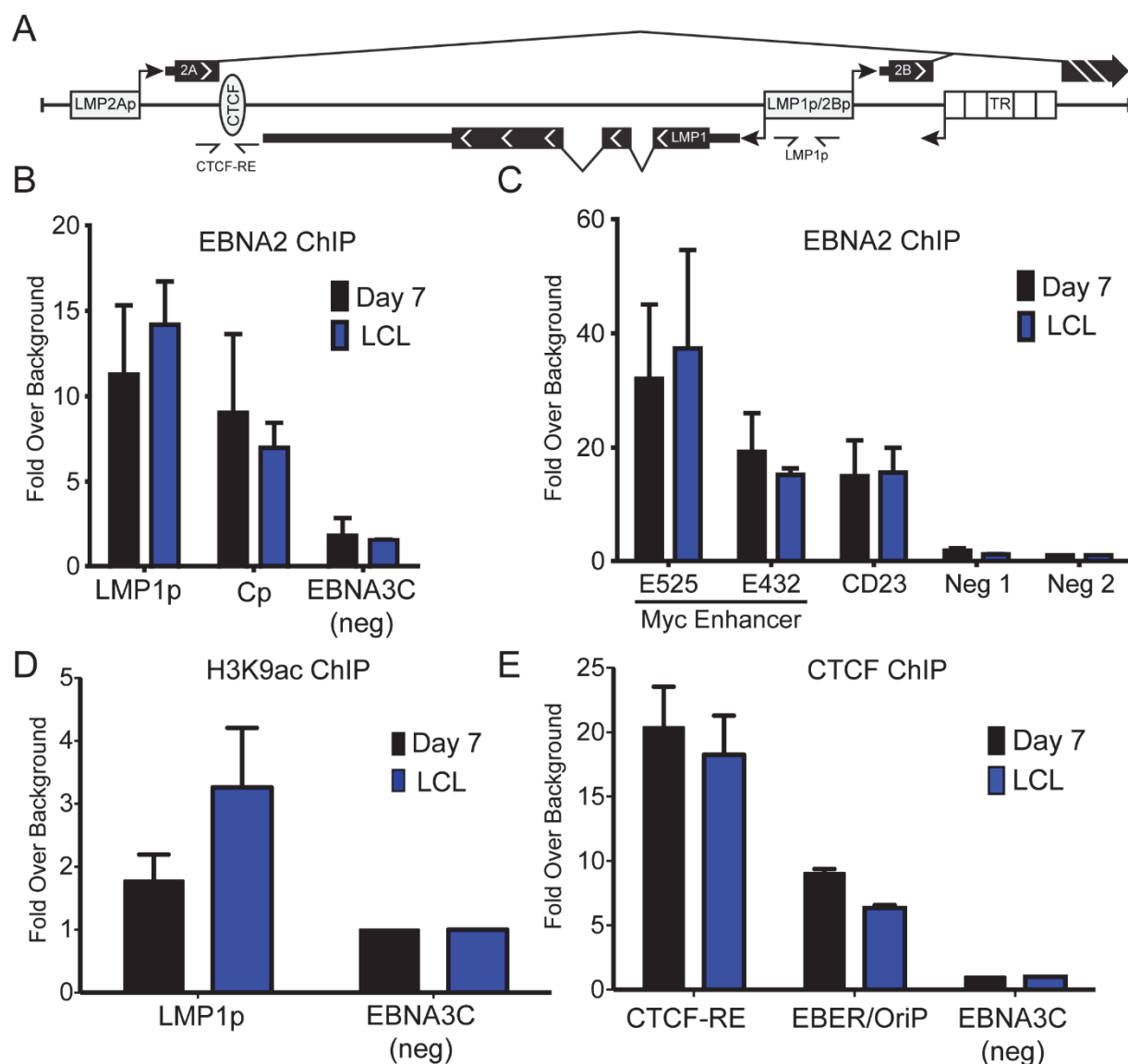
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569 **FIGURES and FIGURE LEGENDS**

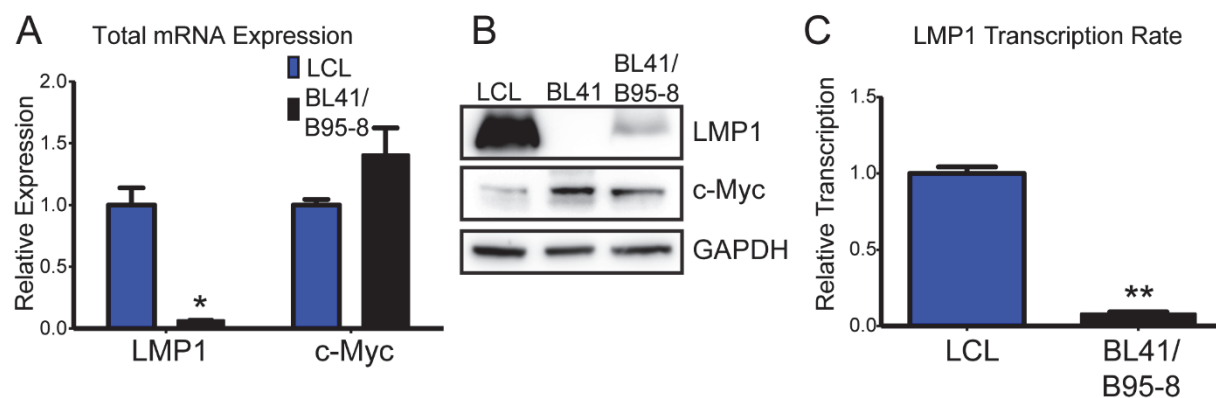


570
 571 **Figure 1. 4sU metabolic pulsing reveals differences in LMP1 mRNA abundance, half-life**
 572 **and transcription rate at Day 7 compared to LCL** (A) Schematic describing 4sU experiments.
 573 Cells are pulsed for 1 hour with 4sU before RNA isolation by Trizol. 4sU labeled RNA is then
 574 conjugated to biotin and isolated via magnetic streptavidin. Unbound RNA is saved as the “old”
 575 fraction while bound RNA eluted from the beads becomes the “new” fraction. Isolated RNA is then
 576 reverse transcribed for RT-qPCR analysis. (B) RT-qPCR analysis of total mRNA from 4sU
 577 experiments for indicated genes at indicated times post infection. (C) RT-qPCR analysis of mRNA
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 579 of mRNA from 4sU experiments for indicated genes at indicated times post infection. All
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 581 Each bar is representative of at least 3 independent donors. Error bars represent SEM. * denotes
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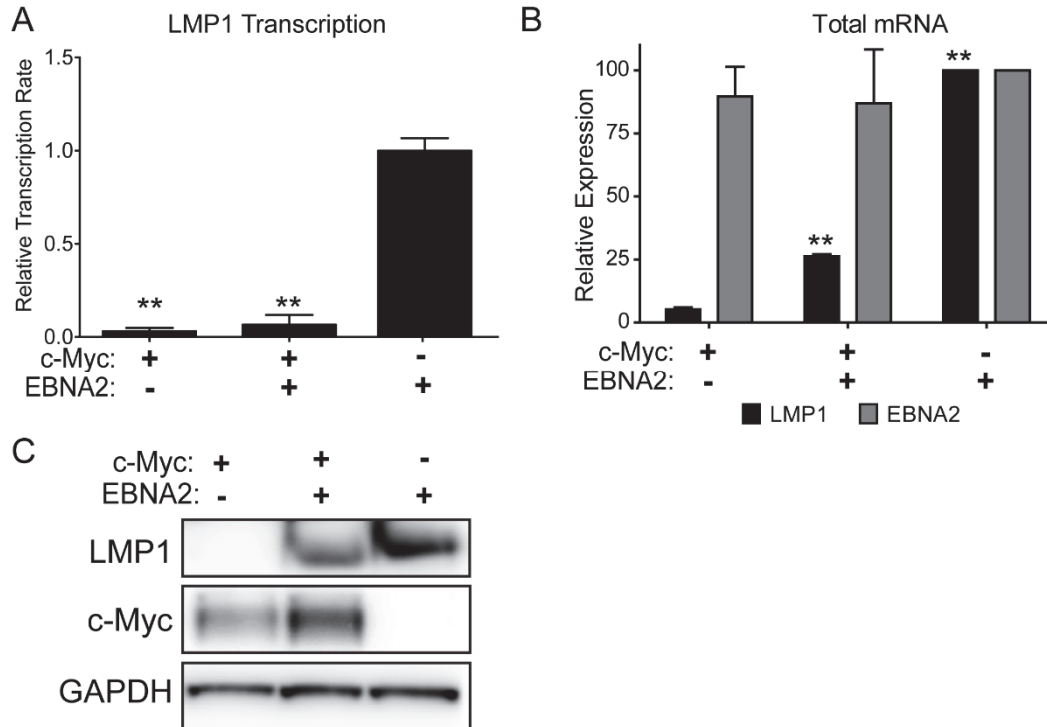


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 585 **Figure 2. Interrogation of the LMP1 locus by ChIP-PCR.** (A) Schematic of the LMP1 genomic
 586 locus. Arrows with half arrow heads denote the location of primers used for ChIP-PCR in (B), (D)
 587 and (E). (B) Sorted Day 7 proliferating primary EBV-infected B cells (black) or fully outgrown (>35
 588 days) LCLs (blue) were harvested for chromatin and ChIP for EBNA2 was performed. LMP1p
 589 denotes the LMP1 promoter, Cp denotes the C promoter and EBNA3C is an EBV genomic
 590 negative control. All samples are normalized to the average of the two negative controls from the
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 596 for CTCF. CTCF-RE denotes the CTCF binding site in the LMP1 locus (panel A), EBER/OriP
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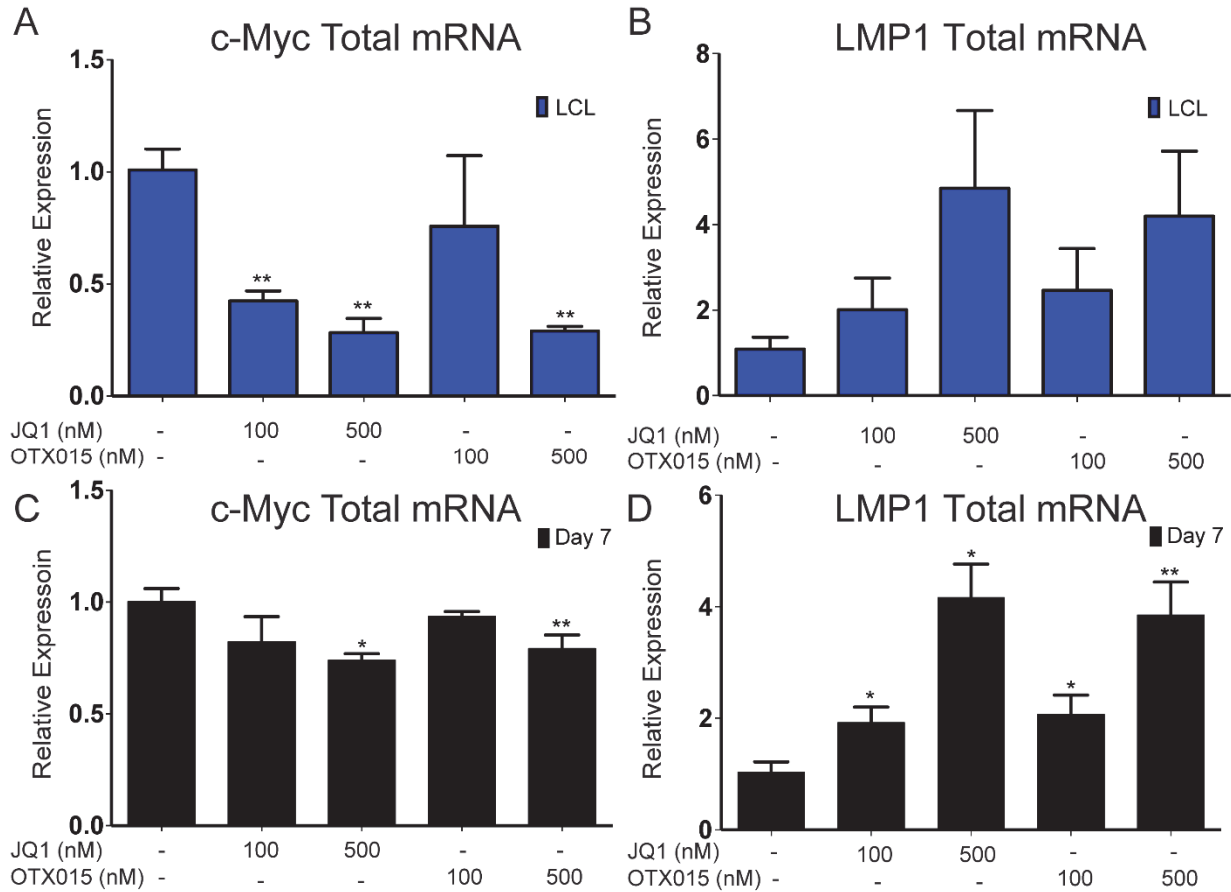
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619 **Figure 5. BET inhibition results in a dose dependent increase in LMP1 and simultaneous**
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