Epstein-Barr virus-positive gastric cancer involves enhancer activation through activating transcription factor 3

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Abstract
Epstein-Barr virus (EBV) is associated with particular forms of gastric cancer (GC). We previously showed that EBV infection into gastric epithelial cells induced aberrant DNA hypermethylation in promoter regions and silencing of tumor suppressor genes. We here undertook integrated analyses of transcriptome and epigenome alteration during EBV infection in gastric cells, to investigate activation of enhancer regions and related transcription factors (TFs) that could contribute to tumorigenesis. Formaldehyde-assisted isolation of regulatory elements (FAIRE) sequencing (-seq) data revealed 19 992 open chromatin regions in putative H3K4me1+ H3K4me3− enhancers in EBV-infected MKN7 cells (MKN7_EB), with 10 260 regions showing increase of H3K27ac. Motif analysis showed candidate TFs, eg activating transcription factor 3 (ATF3), to possibly bind to these activated enhancers. ATF3 was considerably upregulated in MKN7_EB due to EBV factors including EBV-determined nuclear antigen 1 (EBNA1), EBV-encoded RNA 1, and latent membrane protein 2A. Expression of mutant EBNA1 decreased copy number of the EBV genome, resulting in relative downregulation of ATF3 expression. Epstein-Barr virus was also infected into normal gastric epithelial cells, GES1, confirming upregulation of ATF3. Chromatin immunoprecipitation-seq analysis on ATF3 binding sites and RNA-seq analysis on ATF3 knocked-down MKN7_EB revealed 96 genes targeted by ATF3-activating enhancers, which are related with cancer hallmarks, eg evading growth suppressors. These 96 ATF3 target genes were significantly upregulated in MKN7_EB compared with MKN7 and significantly downregulated when ATF3 was knocked down in EBV-positive GC cells SNU719 and NCC24. Knockdown of ATF3 in EBV-infected MKN7, SNU719, and NCC24 cells all led to significant decrease of cellular growth through
INTRODUCTION

Gastric cancer is the second leading cause of cancer-related deaths in the world, accounting for approximately 783,000 deaths in 2018. The infectious agents, including the bacterium Helicobacter pylori and EBV, are major pathogens of GC. As a human oncogenic virus, EBV has been identified in various malignant diseases, including endemic Burkitt lymphoma, nasopharyngeal carcinoma, and 50% of Hodgkin’s disease, and 7%-16% of GC. Due to the particular molecular subtypes in GC, EBV infection status provided an independent classification that not only determined patient stratification, but revealed unique epigenetic phenotype and somatic genomic alterations.

A unique and characteristic feature of EBV is its capability to establish distinct latent gene expression patterns in vivo and in cultured cells (named latency type I, II, and III) in resting and proliferating cells. Epstein-Barr virus-positive GC, as well as Burkitt lymphoma, belong to latency I, in which the expression of viral latent genes is the most severely restricted to EBNA1, EBER, LMP2A, and BamHI A rightward transcripts. Oncogenic functions of these viral transcripts have been studied to elucidate the mechanisms of action of the viral proteins that contribute to tumorigenesis in EBV+ GC.

Although these functions of viral factors might partly contribute to tumorigenesis in EBV+ malignancies, other molecular aberrations, eg genomic and epigenomic aberrations, in host cells need to be elucidated. We have previously reported that EBV+ GC showed extensively high DNA methylation phenotype at gene promoter regions and abundant TSGs, eg p16, are repressed by the aberrant promoter hypermethylation. We also found that in vitro EBV infection in gastric epithelial cells introduces extensive DNA hypermethylation resembling the hypermethylation phenotype of clinical EBV+ GC tissue samples. Although this unique DNA methylation phenotype was reportedly confirmed in EBV+ GC through comprehensive analyses of GC by TCGA, epigenetic aberrations have been considered important for EBV+ GC tumorigenesis in addition to genetic alteration, eg PIK3CA mutation and PD-L1/2 overexpression.

Other than gene repression by aberrant DNA methylation, gene activation by aberrant enhancers is known as an epigenetic driver of many other types of cancer. It is also reported that mutations in enhancers or enhancer-binding TFs could cooperate with DNA methylation to aberrantly activate or repress neighboring genes and provoke cancer. In EBV+ GC, we preliminarily reported aberrant activation and repression at enhancer regions as well as promoter regions, however, the mechanisms to induce epigenetic activation during EBV infection in gastric epithelial cells is yet to be fully investigated.

To identify factors that can induce aberrant enhancer activation during EBV infection, we undertook comprehensive analyses of histone modification and open chromatin regions to predict their putative binding elements. We identified that ATSF3, which is activated by various environmental stresses and reportedly has dual functions as a TSG or an oncogene in several types of cancer, promoted proliferation in EBV+ gastric cells. This investigation could provide insights into the epigenomic activations in enhancer regions, and thus tumorigenesis, by EBV infection in gastric epithelial cells.

MATERIAL AND METHODS

2.1 Cell culture

The SNU719 EBV+ GC cell line was obtained from the Korean Cell Line Bank, and NCC24 from Creative Bioarray. The MKN7 EBV+ GC cell line (MKN7-WT), which shows a low DNA methylation epigenotype, was purchased from Riken BioResource Center Cell Bank.GES1 is a normal fetal gastric epithelial cell line immortalized with SV40 (Beijing Institute for Cancer Research). Cells were cultured in RPMI-1640 medium supplemented with 10% FBS and penicillin/streptomycin. MKN7 and GES1 cells were infected with recombinant EBV using the Akata system to establish EBV-infected MKN7 cells (MKN7,EB) as previously reported. The study design was approved by the institutional review board of Chiba University.

2.2 Chromatin immunoprecipitation-seq and FAIRE-seq analysis

Chromatin immunoprecipitation assays for histone modifications and FAIRE assays for open chromatin regions were carried out as previously reported. For ChIP assays for ATFS3 binding regions, cells were treated with 30 μM CDDP for 48 hours to induce ATFS3 expression. Approximately 1 × 10^7 cells were cross-linked with 1% formaldehyde.
for 10 minutes at room temperature and formaldehyde was quenched by addition of 2.5 M glycine to a final concentration of 0.125 M. Cross-linked chromatin was sonicated to a size of 0.2-1 kb using an ultrasonic disruptor (Branson Digital Sonifier). A total of 15 μg anti-ATF3 Ab (sc-188 X, rabbit mAb; Santa Cruz Biotechnology) and 20 μL Protein G Sepharose beads were mixed in IP dilution buffer and incubated for 6 hours at 4°C. After washing with IP dilution buffer, Ab-binding beads were added to the sonicated chromatin sample and incubated overnight at 4°C. Beads were washed and chromatin was eluted, followed by reversal of the cross-linking and DNA purification. Chromatin-immunoprecipitated DNA was dissolved in EB buffer (Qiagen). Libraries were constructed by using NEBNext ChIP-seq Library Prep Reagent Set for Illumina (NEB) according to the manufacturer’s instructions. The sequencing libraries were quantified by Bioanalyzer (Agilent) and sequenced at a concentration of 4 pM on an Illumina Hiseq (Illumina).

Sequenced reads in ChIP-seq and FAIRE-seq experiments were mapped to the UCSC human genome (hg19) using bowtie 2. Duplicated reads were removed with Picard tools. Peak calling and motif analyses were carried out by using HOMER software (http://homer.salk.edu/homer/index.html), which was also used to count reads for each signal. Peak annotation to the nearest genes was undertaken by using bedtools. Peak heatmaps were produced with the use of HOMER and TreeView for enrichment calculation and visualization.

2.3 | RNA sequencing

RNA was extracted by using the RNeasy Mini Kit (Qiagen) following the manufacturer’s protocol and treated with DNasel (Qiagen). Libraries for RNA-seq were prepared using the TruSeq Stranded mRNA Sample Prep Kit (Illumina), following the manufacturer’s protocol. Deep sequencing was carried out on the Illumina HiSeq 1500 or NextSeq 500 platform using the TruSeq Rapid SBS Kit (Illumina) in 50-base single-end mode according to the manufacturer’s protocol. Sequenced reads from the RNA-seq experiment were aligned by using HISAT2, and Cufflinks was used for transcript assembly. Gene expression levels were expressed as fragments per kilobase of exon per million mapped sequence reads.

2.4 | Cancer hallmark analysis

The ATF3 target genes identified were compared with GO categories related with cancer hallmarks. The GO categories used for the analysis of cancer hallmarks are shown in Table S1.

2.5 | Knockdown by shRNA

To knock down ATF3, double-stranded oligonucleotide DNA encoding shRNA against ATF3 was cloned into the pLKO.1 vector between EcoRI and AgeI sites, as previously described. Oligonucleotide sequences to ligate in the vector for construction of shATF3 and shNON are as follows, where 21-mer target sequences are underlined: shATF3, CCGG GCTGAAGCTTCTTTGGTTG. Viral packaging for shRNA retrovirus vectors was undertaken using 293T cells and FuGENE 6 (Promega), and medium containing the virus was collected 48 hours after transfection.

2.6 | Overexpression of latent EBV factors

Latent viral factors, BARF, EBER1, EBNA1, and LMP2A, were previously constructed in pcDNA3 vector. Each vector or mock vector was introduced in MKN7 cells using FuGene6, and gene expression was analyzed by RNA-seq as above.

2.7 | Overexpression of mutant EBNA1

Mutant ENBA1 to act as a dominant-negative inhibitor of EBNA1 was constructed according to the previous report. Mutant EBNA1 fused by mCherry fluorescent protein at the C-terminal region (mtEBNA1-mCherry) was inserted into the modified pcDNA5 vector. Mock vector with only mCherry inserted was also constructed and used as a control. Vectors with mtEBNA1-mCherry or mCherry were transfected in MKN7 EB cells by Lipofectamine 3000. On day 7, strongly red-colored cells caused by mCherry were sorted by a cell sorter HS800 (Sony). Decrease of EBV copy numbers was examined by qPCR.

2.8 | Real-time qPCR

Real-time qPCR was undertaken using SYBR Green and CFX96 Touch Real-Time PCR (Bio-Rad Laboratories). The quantity of genomic DNA in a sample was estimated by comparisons with standard samples that contained 10^7 to 10^6 gene copies, as previously described. The quantity of EBV genome was normalized to that of the HBB region in the human genome. Primers for the EBV genome are TCGTACATCT CCAGCCGACA CT and CATACAACCA CTGGCGATCC CC. Primers for HBB are CAGGGTGAGG TCTAAGTGAT GA and TTGAAGTCCA CCAGGCGACA CT and CATACAACCA CTGGCGATCC CC. Primers for HBB are CAGGGTGAGG TCTAAGTGAT GA and TTGAAGTCCA CCAGGCGACA CT and CATACAACCA CTGGCGATCC CC. Primers for HBB are CAGGGTGAGG TCTAAGTGAT GA and TTGAAGTCCA CCAGGCGACA CT and CATACAACCA CTGGCGATCC CC. Primers for HBB are CAGGGTGAGG TCTAAGTGAT GA and TTGAAGTCCA CCAGGCGACA CT and CATACAACCA CTGGCGATCC CC.
with standard samples that contained $10^3$ to $10^6$ gene copies.\textsuperscript{13} The expression level of \textit{ATF3} was normalized to that of \textit{B2M}. The PCR primers for \textit{ATF3} are GGATGTCCTC TGGCTGGAA TC and TCCTTTCATC TTCTCAGGG GC. Primers for \textit{B2M} are GATGATATG CCTGCCGTGT and CTGCTTACAT GTCTCATCC CA.

## 2.10 Western blot analysis

Cells were lysed in a lysis buffer containing 62.5 mM Tris-HCl (pH6.8), 2.3% SDS, 10% glycerol, 0.2% bromophenol blue for whole cell or 10 mM Tris-HCl (pH 7.4), 10 mM NaCl, 2.5 mM MgCl\textsubscript{2}, 1 mM EDTA, 1 mM EGTA, 0.5% NP40, and protease inhibitor cocktails (Roche) for cytoplasm and nuclear fraction. The following primary Abs were used: anti-\textit{ATF3} (sc-188 X; Santa Cruz Biotechnology for whole cell; or #33593, rabbit mAb; Cell Signaling Technology for cytoplasm and nuclear fraction), anti-\textit{Actin} (MA5-11889, mouse mAb; Invitrogen), anti-Lamin A (NA934V, rabbit polyclonal; GE Healthcare), and anti-\textit{α-tubulin} (sc-5286, mouse mAb; Santa Cruz Biotechnology). Protein-blotted membranes were incubated with Abs using Can Get Signal Immunoreaction Enhancer Solution (Toyobo) at 4°C overnight for the primary Abs, and at room temperature for 1 hour for secondary Abs, followed by visualization using the ECL prime system (GE Healthcare). The protein signals were detected using a Luminescent Image Analyzer LAS-3000 (Fujifilm).

## 2.11 Cell growth assay

Cells were seeded into 6 wells of 96-well plates at 2000 cells per well in 200 μL culture medium. After an appropriate period of culturing, WST-8 reagent (Dojindo) was added to each well, followed by incubation for 4 hours at 37°C in 5% CO\textsubscript{2}. Absorbance at 450 nm was measured using a SpectraMax PLUS 384 microplate reader (Molecular Devices).

## 2.12 Caspase 3/7 apoptosis staining

Analysis of apoptosis staining was undertaken using CellEvent Caspase-3/7 Green Detection Reagent (Caspase 3/7) (Invitrogen). The apoptotic cells were imaged using a BZ-X710 microscope (Keyence). Imaged data were processed and quantified using ImageJ.\textsuperscript{30}
2.13 | Statistical analysis

The P values for expression analysis were obtained using paired t test to evaluate differences between 2 groups, with P less than .05 considered statistically significant. Gene Ontology analysis was undertaken by using Metascape (http://metascape.org/gp/index.html#/main/step1). Gene set enrichment analysis was undertaken using GSEA software version 4.0.3 (http://software.broadinstitute.org/gsea/index.jsp).

3 | RESULTS

3.1 | Detection of activated enhancers in EBV-infected MKN7 cells

We undertook a FAIRE-seq analysis in MKN7_EB to detect open chromatin regions in gene promoters and enhancers. Promoter regions were defined by H3K4me3 peaks, and enhancer regions were defined by H3K4me1 peaks without H3K4me3 signal (Figure 1A,B). We identified a total of 52 850 FAIRE peaks in MKN7_EB, including 13 209 peaks (25%) in promoters and 19 992 peaks (38%) in enhancers. While the majority of FAIRE peaks were distributed to enhancer regions, 10 260 (51%) of those showed an increase of H3K27ac signals and thus were activated, 4523 (23%) showed no change in H3K27ac signals, and 5209 (26%) showed a decrease of H3K27ac signals and thus were inactivated (Figure 1C).

Using the top 3000 regions showing higher H3K27ac signals among 10 260 activated enhancers, motif analysis was used to search for candidate TFs binding to these regions (Figure 2A). The top 5 significant motifs (P values ranging 1 × 10^{-55} to 1 × 10^{-819}) were detected, and candidate TFs were identified to match those significant motifs. Expression levels of genes near the motifs were analyzed by RNA-seq and compared between MKN7_WT and MKN7_EB (Figure 2B). Significant upregulation of these neighboring genes was observed in MKN7_EB, with ATF3, TEAD2, and RUNX3 showing the most significant changes.

**FIGURE 2** Detection of activating transcription factor 3 (ATF3) as a candidate TF. A, Motif analysis for activated enhancer regions. Among 10 260 activated enhancer regions, the top 3000 regions with highly expressed H3K27ac signal were extracted and subjected to de novo motif analysis. Top 5 significant motifs and candidate TFs, ATF3, JUN, JUNB, FOSL2, and BATF, are shown. B, Expression of genes neighboring the top 5 motifs. Neighboring genes were significantly upregulated in Epstein-Barr virus (EBV)-infected MKN7 cells (MKN7_EB). C, Relative expression levels of candidate TFs in MKN7_EB. While ATF3, TEAD2, and RUNX3 were clearly upregulated in MKN7_EB, other candidate TFs were not. D, Expression of genes in EBV-infected GES1 cells. ATF3 was confirmed to be upregulated after EBV infection, whereas TEAD2 and RUNX3 were not. E, ATF3 protein level in cytoplasm and nucleus. Western blot showed ATF3 localized in nucleus, and expressed at higher levels in MKN7_EB cells. Open and closed boxes indicate relative ratio of the intensity of ATF3 to that of lamin A in the nucleus in MKN7_WT and MKN7_EB cells, respectively (n = 3). Values are shown as the mean ± SD.
genes was confirmed in MKN7_EB, suggesting that the activated enhancers could function in upregulation of the nearby genes. Among the candidate TFs, ATF3, TEAD2, and RUNX3 were upregulated more than 2-fold in MKN7_EB in mRNA levels (Figure 2C). When EBV was infected in GES1 normal gastric epithelial cells, upregulation of ATF3 was confirmed, but TEAD2 and RUNX3 were not upregulated (Figure 2D). We therefore focused on ATF3, and upregulation of ATF3 in MKN7_EB was detected in not only mRNA levels but also protein levels (Figure 2E).

To gain insight into the viral factors that might upregulate ATF3 expression in host cells, viral factors including BARF, EBER1, EBNA1, and LMP2A, which are known to be expressed in latent infection of EBV, were overexpressed in MKN7 cells, and RNA-seq was carried out. Activating transcription factor 3 was found to be upregulated the most by EBNA1 overexpression, and relative upregulation by more than 1.5-fold was also detected by EBER1 or LMP2A overexpression (Figure 3A). Previous ChIP-seq analysis for EBNA1 target regions showed that EBNA1 can bind to an enhancer region upstream of ATF3. When mutant EBNA1 was overexpressed in MKN7_EB, the copy number of the EBV genome was confirmed to be decreased, resulting in a considerable decrease of ATF3 expression level (Figure 3C). These results indicate that ATF3 could be upregulated, at least partly, by expression of EBV factors, eg EBNA1.

3.2 Identification of possible ATF3 binding sites by ChIP-seq

To identify target regions of ATF3 genome-wide, ChIP-seq using anti-ATF3 Ab was used against ATF3 binding sites (Figure 4). As ATF3 overexpression using pcDNA5 expression vector in MKN7 led to more than 100-fold upregulation of ATF3 and cell death at 36 hours, ATF3 upregulation was induced in MKN7 cells by exposure to 30 μM CDDP instead. As ATF3 expression was increased in a time-dependent manner up to 48 hours (Figure 4A), ChIP-seq analysis of possible ATF3 binding sites was carried out using MKN7 cells at 48 hours after CDDP exposure (Figure 4B,C). Among a total of 3345 ATF3 peaks defined by ChIP-seq analysis, 1339 peaks were distributed to enhancer regions, and 590 of those showed increase of H3K27ac signals and thus were activated (Figure 4D). Expression levels of 659 genes neighboring ATF3 peaks on the 590 activated enhancers were significantly increased ($P = 1.5 \times 10^{-6}$).

3.3 Identification of ATF3 target genes

As the possible ATF3 binding sites might include regions targeted specifically in the condition of CDDP exposure, ATF3 was knocked down...
in MKN7\_EB cells by a shATF3 to screen genes targeted by ATF3 in EBV-infected cells. Repression of ATF3 was confirmed by western blot (Figure 6). RNA sequencing was used to analyze gene expression alteration. Gene set enrichment analysis showed that the 96 ATF3 target genes were significantly enriched in genes downregulated by knockdown of ATF3, in both NCC24 and SNU719 cells, compared with shNON control cells (Figure 6A-D). We analyzed the cellular growth of ATF3 knocked-down cells by WST-8 assay (Figure 6E). Significant repression of cellular growth was detected in MKN7\_EB, NCC24, and SNU719 cells when ATF3 was knocked down. Caspase 3/7 activity was analyzed to detect apoptotic cells, and apoptosis was observed significantly frequently in MKN7\_EB, NCC24, and SNU719 cells when ATF3 was knocked down (Figure 6F).

4 | DISCUSSION

Through integrated analyses on alterations of transcriptome, histone modification, and open chromatin status, we here identified ATF3 as a critical transcriptional activator after EBV infection in gastric cells. We undertook screening of putative TF binding to activated enhancer regions that are predicted by motif analysis and refined by
gene expression analysis. ATF3 was identified as a candidate TF that is upregulated by EBV infection and induces aberrant enhancer activation. Expression of viral factors might be the cause of upregulated ATF3 expression, at least partly, and the effect of ATF3 expression on cellular proliferation was confirmed, suggesting a tumorigenic role of aberrant enhancer activation by ATF3 upregulation.

ATF3 is known to play different roles depending on various interacting partners. It is activated by a variety of environmental stress signals and is associated with the pathogenesis of various diseases including cancer, cardiac hypertrophy, and infection. In cancer, reciprocal functions of ATF3 as either an oncogene or a TSG have been reported, depending on the condition of the cell. It reportedly promotes apoptosis of human cancer cells, such as T-cell lymphoma, multiple myeloma, blood, lung, prostate, and colon cancer cells, and overexpression of ATF3 reportedly promotes the metastasis of prostate and breast cancer. As an oncogene, TGF-β, p53, and Wnt/β-catenin signaling pathways are the known factors to activate ATF3, leading to the expression of downstream factors, eg MMP13, Snail, Slug, and Twist, which are key regulators of epithelial-mesenchymal transition and metastasis. ATF3 was also reported to promote Runx2 expression in metastatic breast cancer cells, aiding bone metastasis of tumor cells. In this study, ATF3 was found to be upregulated by EBV infection through expression of EBV factors, eg EBNA1, EBER1, and LMP2A, and downregulation of ATF3 was confirmed when the copy number of the EBV genome was decreased in MKN7_EB due to overexpression of mutant EBNA1. Knockdown experiments of ATF3 showed that ATF3 has an oncogenic role to promote proliferation in EBV-infected gastric cells. Downstream targets of ATF3 were also shown to correlate with hallmarks of cancer, which might contribute to EBV+ GC progression.
The sequence-specific DNA binding protein EBNA1 binds with high affinity to 3 well-characterized sites in the viral genome that are important for maintenance of the episomal viral genome, DNA replication, and viral gene regulation. Several papers reported that there exist numerous EBNA1 binding sites in the host genome and that EBNA1 binds to these sites to activate the expression of neighboring host genes. It has been reported that EBNA1 activates many other host genes directly, which can function in various pathways related to cancer proliferation, including several cytokine pathways (IL-18, IL-6, IL-12, and IL-2), the p38 MAPK and MAP3K1 network in B cells, the activator protein 1 (AP1) transcription factor pathway in nasopharyngeal carcinoma cells, and signal transducer and activator of transcription 1 (STAT1) in carcinoma and B-lymphoma cells, although ATF3 had not been identified as an EBNA1 downstream target. This study showed that EBNA1 might be an EBV factor to potentially induce upregulation of ATF3, and thus promote proliferation in GC cells; mechanisms to activate ATF3 expression, eg function analysis of EBNA1 binding site nearby ATF3 as an enhancer, should be further investigated.

Previous comprehensive analyses of GC by us or TCGA revealed that EBV+ GC shows a unique DNA hypermethylation phenotype that is the most extensive hypermethylation among all the human malignancies. Distinct mutations of genes, eg PIK3CA and ARID1A, are also observed in EBV+ GC. Most of these aberrations are related with repression of tumor suppressive functions in gastric epithelial cells, and viral proteins also mainly function to repress tumor suppressive functions, except EBERs that reportedly promote tumor progression through activation of growth factors. Recently, aberrant enhancer activation has been reported in many
types of cancer to promote oncogene activation and is reportedly caused by genomic aberrations such as amplification or translocation, or activation of oncogenic master regulators in various cancers. Whereas enhancer activation in EBV+ GC has been reported in only a few published works, we here found that ATF3 is up-regulated and contributes to enhancer activation in EBV-infected gastric cells.

In summary, we undertook integrated analyses on alterations of the transcriptome and epigenome during EBV infection in gastric epithelial cells and identified upregulation of ATF3 that could contribute to aberrant enhancer activation in EBV+ GC and enhance proliferation of gastric cells by EBV infection.

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CONFLICT OF INTEREST
Yuta Asakawa is hired by Otsuka Pharmaceutical Co., Ltd. The authors have no other conflicts of interest to disclose.

DATA AVAILABILITY STATEMENT
NGS data generated in this study have been deposited in Gene Expression Omnibus under accession # GSE141385 (GSM4202156 - GSM4202165, GSM4308257, GSM4308258). Previously deposited NGS data (GSE97837 and GSE97838) that were used in this study are also available in Gene Expression Omnibus. The authors declare that all other data are available within the article or associated supplementary information files, or available from the authors on request.

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REFERENCES
1. Ferlay J, Colombet M, Soerjomataram I, et al. Estimating the global cancer incidence and mortality in 2018: GLOBOCAN sources and methods. Int J Cancer. 2019;144:1941-1953.
2. Amieva M, Peek RM. Pathobiology of Helicobacter pylori-Induced Gastric Cancer. Gastroenterology. 2016;150:64-78.
3. Uemura N, Okamoto S, Yamamoto S, et al. Helicobacter pylori infection and the development of gastric cancer. N Engl J Med. 2001;345:784-789.
4. Shibata D, Weiss LM. Epstein-Barr virus-associated gastric adenocarcinoma. Am J Pathol. 1992;140:769-774.
5. Burke AP, Yen TS, Sheikita KM, Sobin LH. Lymphoepithelial carcinomas of the stomach with Epstein-Barr virus demonstrated by polymerase chain reaction. Mod Pathol. 1990;3:377-380.
6. Murphy G, Pfeiffer R, Camargo MC, Rabkin CS. Meta-analysis shows that prevalence of Epstein-Barr virus-positive gastric cancer differs based on sex and anatomic location. Gastroenterology. 2009;137:824-833.
7. Cohen JI, Fauci AS, Varmus H, Nabel GJ. Epstein-Barr virus: an important vaccine target for cancer prevention. Sci Transl Med. 2011;3:107fs107.
8. Cancer T, Atlas G, Bass AJ, et al. Comprehensive molecular characterization of gastric adenocarcinoma. Nature. 2014;513:202-209.
9. Baer R, Bankier AT, Biggin MD, et al. DNA sequence and expression of the B95–8 Epstein-Barr virus genome. Nature. 1984;310:207-211.
10. Young LS, Yap LF, Murray PG. Epstein-Barr virus: more than 50 years old and still providing surprises. Nat Rev Cancer. 2016;16:789-802.
11. Fukayama M, Hino R, Uozaki H. Epstein-Barr virus and gastric carcinoma: virus-host interactions leading to carcinoma. Cancer Sci. 2008;99:1726-1733.
12. Ikwiriri E, Eizuru Y, Tokunaga M, Takada K. Autocrine growth of Epstein-Barr virus-positive gastric carcinoma cells mediated by an Epstein-Barr virus-encoded small RNA. Cancer Res. 2003;63:7062.
13. Matsuoka K, Kandara A, Nagae G, et al. Classification of Epstein-Barr virus-positive gastric cancers by definition of DNA methylation epigenotypes. Cancer Res. 2011;71:7187-7197.
14. Matsuoka K, Funata S, Fukuyo M, et al. Epstein-Barr virus infection induces genome-wide de novo DNA methylation in non-neoplastic gastric epithelial cells. J Pathol. 2017;242:391-399.
15. Saito R, Abe H, Kunita A, Yamashita H, Seto Y, Fukayama M. Overexpression and gene amplification of PD-L1 in cancer cells and PD-L1+ immune cells in Epstein-Barr virus-associated gastric cancer: the prognostic implications. Mod Pathol. 2017;30:427-439.
16. Herz HM, Hu D, Shilatifard A. Enhancer malfunction in cancer. Mol Cell. 2014;53:859-866.
17. Bradner JE, Hnisz D, Young RA. Transcriptional addiction in cancer. Cell. 2016;168:629-643.
18. Aran D, Hellman A. DNA methylation of transcriptional enhancers and cancer predisposition. Cell. 2013;154:11-13.
19. Okabe A, Funata S, Matsuoka K, et al. Regulation of tumour related genes by dynamic epigenetic alteration at enhancer regions in gastric epithelial cells infected by Epstein-Barr virus. Sci Rep. 2017;7:7924-7924.
20. Funata S, Matsuoka K, Yamanaka R, et al. Histone modification alteration coordinated with acquisition of promoter DNA methylation during Epstein-Barr virus infection. Oncotarget. 2017;8:55265-55279.
21. Li X, Zang S, Cheng H, Li J, Huang A. Overexpression of activating transcription factor 3 exerts suppressive effects in HepG2 cells. Mol Med Rep. 2019;19:869-876.
22. Rohini M, Haritha Menon A, Selvamurugan N. Role of activating transcription factor 3 and its interacting proteins under physiological and pathological conditions. Int J Biol Macromol. 2018;120:310-317.
23. Ke Y, Ning T, Wang B. Establishment and characterization of a SV40 transformed human fetal gastric epithelial cell line-GES-1. Zhonghua Zhong Liu Za Zhi. 1994;16:7-10.
24. Imai S, Nishikawa JUN, Takada K. Cell-to-cell contact as an efficient mode of Epstein-Barr virus infection in diverse human epithelial cell lines. J Virol. 1998;72:4371-4378.
25. Kaneda A, Fujita T, Anai M, et al. Activation of Bmp2-Smad1 signal and its regulation by coordinated alteration of H3K27 trimethylation in Ras-induced senescence. PLoS Genet. 2011;7:e1002359.
26. Ke Y, Ning T, Wang B. Establishment and characterization of a SV40 transformed human fetal gastric epithelial cell line-GES-1. Zhonghua Zhong Liu Za Zhi. 1994;16:7-10.
29. Kirchmaier AL, Sugden B. Dominant-negative inhibitors of EBNA-1 of Epstein-Barr virus. *J Virol*. 1997;71:1766-1775.
30. Schneider CA, Rasband WS, Eliceiri KW. NIH Image to ImageJ: 25 years of image analysis. *Nat Methods*. 2012;9:671-675.
31. Tripathi S, Pohl MQ, Zhou Y, et al. Meta- and orthogonal integration of influenza "oMICs" data defines a role for UBR4 in virus budding. *Cell Host Microbe*. 2015;18:723-735.
32. Subramanian A, Tamayo P, Mootha VK, et al. Gene set enrichment analysis: a knowledge-based approach for interpreting genome-wide expression profiles. *Proc Natl Acad Sci USA*. 2005;102:15545-15550.
33. Tempera I, Leo AD, Kossenkov AV, et al. Identification of MEF2B, EBF1, and IL6R as direct gene targets of Epstein-Barr virus (EBV) nuclear antigen 1 critical for EBV-infected B-lymphocyte survival. *J Virol*. 2016;90:345-355.
34. Lu F, Wikramasinghe P, Norseen J, et al. Genome-wide analysis of host-chromosome binding sites for Epstein-Barr Virus Nuclear Antigen 1 (EBNA1). *Virol J*. 2010;7:1-17.
35. Zhao J, Li X, Guo M, Yu J, Yan C. The common stress responsive transcription factor ATF3 binds genomic sites enriched with p300 and H3K27ac for transcriptional regulation. *BMC Genomics*. 2016;17:1-14.
36. Chüeh AC, Tse JWT, Dickinson M, et al. ATF3 repression of BCL-X<sub>L</sub> determines apoptotic sensitivity to HDAC inhibitors across tumor types. *Clin Cancer Res*. 2017;23:5573-5584.
37. Wang Z, Xu D, Ding HF, et al. Loss of ATF3 promotes Akt activation and prostate cancer development in a Pten knockout mouse model. *Oncogene*. 2015;34:4975-4984.
38. Joo JH, Ueda E, Bortner CD, Yang XP, Liao G, Jetten AM. Farnesol activates the intrinsic pathway of apoptosis and the ATF4-ATF3-CHOP cascade of ER stress in human T lymphoblastic leukemia Molt4 cells. *Biochem Pharmacol*. 2015;97:256-268.
39. Jan YH, Tsai HY, Yang CJ, et al. Adenylate kinase-4 is a marker of poor clinical outcomes that promotes metastasis of lung cancer by downregulating the transcription factor ATF3. *Cancer Res*. 2012;72:5119-5129.
40. Wolford CC, McConoughey SJ, Jalganokar SP, et al. Transcription factor ATF3 links host adaptive response to breast cancer metastasis. *J Clin Invest*. 2013;123:2893-2906.
41. Bandyopadhyay S, Wang Y, Zhan R, et al. The tumor metastasis suppressor gene <em>Dr-1</em> down-regulates the expression of activating transcription factor 3 in prostate cancer. *Cancer Res*. 2006;66:11983-11990.
42. Yin X, Wolford CC, Chang YS, et al. ATF3, an adaptive-response gene, enhances TGF[beta] signaling and cancer-initiating cell features in breast cancer cells. *J Cell Sci*. 2010;123:3558-3565.
43. Gokulnath M, Partridge NC, Selvanmurugan N, Runx2, a target gene for activating transcription factor-3 in human breast cancer cells. *Tumour Biol*. 2015;36:1923-1931.
44. O’Neil JD, Owen TJ, Wood VHJ, et al. Epstein-Barr virus-encoded EBNA1 modulates the AP-1 transcription factor pathway in nasopharyngeal carcinoma cells and enhances angiogenesis in vitro. *J Gen Virol*. 2008;89:2833-2842.
45. AlQarni S, Al-Sheikh Y, Campbell D, et al. Lymphomas driven by Epstein-Barr virus nuclear antigen-1 (EBNA1) are dependant upon Mdm2. *Oncogene*. 2018;37:3998-4012.
46. Wood VHJ, O’Neil JD, Wei W, Stewart SE, Dawson CW, Young LS. Epstein-Barr virus-encoded EBNA1 regulates cellular gene transcription and modulates the STAT1 and TGFβ signaling pathways. *Oncogene*. 2007;26:4135-4147.

**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section.

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