Human CD4+ T Lymphocytes Consistently Respond to the Latent Epstein-Barr Virus Nuclear Antigen EBNA1

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Abstract

The Epstein-Barr virus (EBV)-encoded nuclear antigen EBNA1 is critical for the persistence of the viral episome in replicating EBV-transformed human B cells. Therefore, all EBV-induced tumors express this foreign antigen. However, EBNA1 is invisible to CD8+ cytotoxic T lymphocytes because its Gly/Ala repeat domain prevents proteasome-dependent processing for presentation on major histocompatibility complex (MHC) class I. We now describe that CD4+ T cells from healthy adults are primed to EBNA1. In fact, among latent EBV antigens that stimulate CD4+ T cells, EBNA1 is preferentially recognized. We present evidence that the CD4+ response may provide a protective role, including interferon-γ secretion and direct cytolysis after encounter of transformed B lymphocyte cell lines (B-LCLs). Dendritic cells (DCs) process EBNA1 from purified protein and from MHC class II–mismatched, EBNA1-expressing cells including B-LCLs. In contrast, B-LCLs and Burkitt's lymphoma lines likely present EBNA1 after endogenous processing, as their capacity to cross-present from exogenous sources is weak or undetectable. By limiting dilution, there is a tight correlation between the capacity of CD4+ T cell lines to recognize autologous B-LCL–expressing EBNA1 and DCs that have captured EBNA1. Therefore, CD4+ T cells can respond to the EBNA1 protein that is crucial for EBV persistence. We suggest that this immune response is initiated in vivo by DCs that present EBV-infected B cells, and that EBNA1-specific CD4+ T cell immunity be enhanced to prevent and treat EBV-associated malignancies.

Key words: Epstein-Barr virus • Epstein-Barr virus nuclear antigen 1 • CD4+ T cell • cross-presentation • dendritic cells

Introduction

EBV is a human gamma herpesvirus with a tropism for B lymphocytes (1). More than 95% of the adult population carries EBV as a lifelong asymptomatic infection. Nevertheless, EBV has strong growth-transforming capacities (2). Each of its three latency programs gives rise to specific tumors originating from B cells or other cell types. As exemplified by EBV-transformed B cells (B lymphocyte cell lines [B-LCLs])1 or lymphoproliferative disease, the latency III phenotype is characterized by the expression of nine gene products: six EBV nuclear antigens (EBNA1, EBNA2, EBNA3A, EBNA3B, EBNA3C, and EBNALP) and three latent membrane proteins (LMP1, LMP2A, and LMP2B) (3). In latency II, EBV-associated malignancies like Hodgkin's lymphoma, nasopharyngeal carcinoma, T cell lymphoma, gastric carcinoma, and uterine leiomyosarcoma, three specific EBV genes, EBNA1, LMP1, and LMP2, are

1Abbreviations used in this paper: B-LCL, B lymphocyte cell line; DC, dendritic cell; EBNA, EBV nuclear antigen; ELISPOT, enzyme-linked immunospot; GA, Gly/Ala; HS, human serum; LMP, latent membrane protein; MOI, multiplicity of infection; MIP, matrix protein; NS1, nonstructural protein 1; SFC, spot-forming cell; SLAM, signalling lymphocyte activation molecule; vv, vaccinia virus

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maintained (3). Burkitt’s lymphoma exemplifies the EBV latency I phenotype. Only the EBNA1 protein is expressed in these transformed B cells. 

EBNA1, LMP1, and LMP2 are probably critical for tumorigenesis, inducing cell proliferation as well as resistance to apoptosis (4). EBNA1 binds as a dimer to the viral origin of replication and ensures episomal replication during B cell growth (5–8). The C O O H-terminal part of LMP1 can act as a direct oncogene (9–11) by mimicking CD40-mediated B cell activation (12–15). Thus, LMP1 engages signal proteins for the TNF receptor family (16, 17) and janus kinase 3 (18), ultimately leading to nuclear factor κB (17) and AP-1 (Jun/Jun or Jun/Fos) (19) induction. In addition, LMP1 expression protects against apoptosis by induction of bcl-2 (20, 21). Instead, LMP2 mimics B cell receptor signaling, constitutively engaging syk and lyn, protein tyrosine kinases (22). In Burkitt’s lymphoma, only EBNA1 seems to be required, as transformation is achieved by additional mechanisms probably involving c-myc uncoupling through chromosomal translocation (2). However, the increased incidence of B cell neoplasia in EBNA1 transgenic mice argues for an involvement of this EBV product in transformation even in latency I malignancies (23).

How then is transformation to latency I and II malignancies avoided in most carriers of EBV? Immunity to EBNA1 a priori could provide resistance to transformed cells, but it has proven difficult to detect specific T cell responses to this essential protein for EBV persistence. In fact, EBNA1 blocks its own processing for MHC class I presentation (24). This has been attributed to a deficit in proteasomal processing, caused by the NH₂-terminal Gly/Ala (GA) repeat domain (25, 26). A similar GA stretch prevents IkBα degradation by the proteasome (27). Other EBV latency gene products are the focus of a strong MHC class I–restricted CTL response, especially EBNA3A, EBNA3B, and EBNA3C (28). However, EBNA3 proteins are not expressed in most of the EBV–associated tumors mentioned above, and instead are expressed in transformed cultured lines (B-LCLs) and lymphoproliferative syndromes in immunosuppressed patients. CD8+ CTL responses to tumor-associated LMP1 (29) and LMP2 (30) proteins have only occasionally been detected.

It has been repeatedly apparent that the development (31–33) and maintenance (34–37) of effective CD8+ CTLs are dependent on CD4+ T cell help. Recognition of EBV products by CD4+ T cells has not been investigated in the same detail as the CD8+ T cell response (38). Only two EBV–specific CD4+ T cell clones have been described before (39, 54). While the EBNA2-specific CD4+ T cell clone recognized HLA-DQ-matched B-LCLs, the EBNA1-specific CD4+ T cell clone only killed targets after exogenous loading with recombinant EBNA1 protein. Therefore, it was suggested that EBNA1 prevents its own endogenous presentation onto MHC class II, and that CD4+ T cell recognition of EBNA1 does not mediate protective immunity against EBV–associated malignancies.

Dendritic cells (DCs) are potent APCs for CD4+ and CD8+ T cell immunity (40, 41). Therefore, we used DCs to search for CD4+ T cell responses to individual latent EBV products. For EBNA1, we delivered the antigen as recombinant protein (42–45), in recombinant vaccinia virus (vv) constructs, and by coculturing with B-LCLs. We have uncovered a strong CD4+ T cell response to EBNA1, as monitored by T cell activation and proliferation, IFN-γ secretion, and CTL activity. Paralleling the above results with one EBNA1-specific CD4+ T cell clone, we demonstrate that the EBNA1-specific CD4 responses—generated routinely from adult blood samples—recognize HLA-DR–matched B-LCLs and therefore could provide resistance to EBV infection and EBV–associated malignancy.

Materials and Methods

Cell Lines. The EBV–transformed B cell lines LR M (HLA-A2, -B44, -DRB1*0401, -DQA1*03, -DQB1*0301, and -DP4) (46), LG2 (HLA-DRB1*0101, -DQA1*0101, -DQB1*0501, -DP1*0101, and -DPB1*0201) (47), and newly generated B-LCL and the Burkitt’s lymphoma lines Ramos, EBV+, and Daudi, reverted to latency III (obtained from American Type Culture Collection), were cultured in RPMI 1640/10% FCS/glutamine/gentamicin. LCL-JT (HLA-DRB1*0301 and -DRB1*1301), LCL-BM (HLA-A1, -A3, -B7, -B8, -Cw6, -Cw7, -DR4, -DRw14, -DRw52, -DRw53, and -DQw3), LCL-DC (HLA-A2, -A24, -B38, -B46, -Cw1, -CD1502, -DRB1*0901, -DRB4*, -DRB5*0101, -DQB1*0502, and -DQB1*0303), LCL-BC (HLA-DRB1*0401, -DRB1*0701, DRB4*01, -DQB1*0302, and -DQB1*0201), and buffy coat–derived B-LCLs were generated by culturing PBMCs of healthy donors with supernatants of the marmoset cell line B95.8 in RPMI 1640/20% FCS/glutamin/gentamicin/1 μg/ml cyclosporin A. The rabbit RK13 and monkey BSC40 kidney cell lines were grown in DM EM/15% FCS/glutamin/gentamicin.

DC and PBMC Preparations. Leukocyte concentrates (buffy coats) from the New York Blood Center, as well as whole blood from lab donors, served as sources of PBMCs isolated by density centrifugation on Ficoll–Paque (Amersham Pharmacia Biotech). CD2+ PBMCs were separated by rosetting with neuraminidase-treated (Calbiochem) sheep RBCs (Colorado Serum Company) followed by red cell lysis with 1.66% ammonium chloride. Where indicated, CD8+ or CD4+ T cells were depleted with Leu2a or OKT8 (for CD8) or H P2/6 (for CD4) antibodies, followed by incubation with sheep anti–mouse IgG Dynabeads and a magnetic particle concentrator, M CP-1 (Dynal). Positive selection for CD4+ PBMCs was performed using anti-CD4 Microbeads, M S* / R S* columns, and MiniMACS separator (Miltenyi Biotech). DCs were generated from CD2+ PBMCs as described (48). 106 CD2+ PBMCs/ml were plated in 6-well plates with RPMI 1640/1% single donor plasma/glutamine/gentamicin. 100 μl medium was replaced at days 2, 4, and 6. Recombinant human (rh)IL-4 and rhGM-CSF were added to a final concentration of 1,000 U/ml at days 0, 2, 4, and 6. On day 7, the floating immature DCs were transferred to new plates at 3×10⁶ cells/ml and half of the medium was replaced by monocyte-conditioned medium to mature the DCs for 2 d. DCs and T cells were used fresh or after cryopreservation.

Vaccinia Virus Stock Generation and Infection of DCs. Recombinant vv were expanded in rabbit RK13 and titrated on monkey BSC40 kidney cells. Mature DCs were infected at a multiplicity of infection (MOI) of 2 for 1 h at 37°C and washed three times. The efficiency of infection was checked after 6–12 h by FACS®
as described using intracellular staining of a vaccinia early protein of 29 kD with the VV1-6B6 antibody (49).

Table I. Percentages of Blasting CD 4⁺ T Lymphocytes upon Stimulation with DCs Infected with Recombinant V accinia-EBV Viruses

| Donor no. | vvTK⁻ | vvEBNA1 | vvEBNA3A | vvEBNA3B | vvEBNA3C | vvLM P1 | vvLM P2A | vvBM LF1 | Influenza |
|-----------|-------|---------|----------|----------|----------|---------|---------|----------|-----------|
| 1         | 4*    | 17 (2)  | 22 (8)   | 11 (3)   | N        | 23 (3)  | 8 (4)   | 3 (3)    | 17 (6)    |
| 2         | 2     | 15 (3)  | 6 (2)    | 17 (8)   | 8 (3)    | 8 (2)   | 3 (2)   | 0 (0)    | 12 (3)    |
| 3         | 0     | 11 (6)  | 0 (0)    | 19 (8)   | 0 (0)    | 30 (18)| 0 (0)   | 0 (0)    | 28 (10)   |
| 4         | 1     | 11 (6)  | 1 (1)    | 12 (7)   | 2 (1)    | 14 (10)| 1 (0)   | 1 (0)    | 35 (6)    |
| 5         | 1     | 15 (4)  | 1 (1)    | 4 (2)    | 3 (1)    | 6 (1)  | 1 (1)   | 0 (1)    | 13 (10)   |
| 6         | 2     | 8 (3)   | N        | 2 (2)    | 1 (2)    | 5 (1)  | 3 (3)   | N        | 15 (7)    |
| 7         | 3     | 11 (6)  | 2 (2)    | 2 (2)    | 1 (1)    | 3 (2)  | 3 (1)   | 3 (3)    | 34 (11)   |
| 8         | 1     | 12 (4)  | 0 (0)    | 10 (2)   | 2 (1)    | 8 (9)  | 2 (2)   | 9 (8)    | 7 (3)     |
| 9         | 2     | 8 (0)   | 1 (0)    | 3 (0)    | 4 (2)    | 18 (5)| 5 (0)   | 4 (4)    | 2 (1)     |
| 10        | 5     | 18 (3)  | 5 (3)    | 3 (1)    | 17 (3)   | 8 (2)  | 4 (1)   | 5 (2)    | 25 (3)    |

*Values in parentheses reflect reactivity using vvTK⁻-infected DCs as control targets in the ELISPOT assay.

Table II. Number of IFN-γ-producing CD 4⁺ T Lymphocytes upon Stimulation with DCs Infected with Recombinant V accinia-EBV Viruses

| Donor no. | vvTK⁻ | vvEBNA1 | vvEBNA3A | vvEBNA3B | vvEBNA3C | vvLM P1 | vvLM P2A | vvBM LF1 | Influenza |
|-----------|-------|---------|----------|----------|----------|---------|---------|----------|-----------|
| 5         | 7 ± 1*| 23 ± 1  | 5 ± 4    | 2 ± 1    | 6 ± 2    | 4 ± 1   | 10 ± 4  | 1 ± 1    | 79 ± 3    |
|           | (7 ± 4†)| (7 ± 4) | (5 ± 1)  | (5 ± 1)  | (2 ± 1)  | (6 ± 1) | (2 ± 1) | (9 ± 1)  |           |
| 6         | 3 ± 1 | 88 ± 8  | N        | 5 ± 1    | 3 ± 1    | 4 ± 2   | 6 ± 3   | N        | 83 ± 3    |
|           | (37 ± 1)| (4 ± 2)| (2 ± 2)  | (3 ± 1)  | (2 ± 2)  | (2 ± 2) | (27 ± 6)|           |           |
| 7         | 1 ± 1 | 16 ± 1  | 4 ± 2    | 2 ± 1    | 2 ± 0    | 2 ± 2   | 6 ± 4   | 4 ± 2    | 77 ± 2    |
|           | (4 ± 1)| (2 ± 2)| (3 ± 0)  | (3 ± 0)  | (2 ± 1)  | (1 ± 0) | (5 ± 1) | (5 ± 1)  |           |
| 8         | 3 ± 1 | 36 ± 1  | 2 ± 1    | 25 ± 4   | 1 ± 1    | 64 ± 2  | 6 ± 2   | 18 ± 1   | 83 ± 1    |
|           | (6 ± 2)| (1 ± 0)| (1 ± 1)  | (2 ± 2)  | (5 ± 2)  | (2 ± 1) | (2 ± 1) | (3 ± 1)  |           |
| 9         | 2 ± 1 | 17 ± 1  | 2 ± 1    | 5 ± 2    | 8 ± 4    | 25 ± 5  | 10 ± 3  | 6 ± 2    | 15 ± 4    |
|           | (1 ± 1)| (2 ± 2)| (2 ± 2)  | (2 ± 1)  | (1 ± 1)  | (3 ± 1) | (2 ± 1) | (4 ± 1)  |           |
| 10        | 5 ± 1 | 48 ± 3  | 15 ± 2   | 13 ± 3   | 108 ± 3  | 38 ± 1  | 14 ± 1  | 5 ± 2    | 225 ± 5   |
|           | (15 ± 3)| (6 ± 4)| (4 ± 3)  | (9 ± 5)  | (10 ± 3) | (7 ± 2) | (3 ± 1) | (8 ± 4)  |           |

*Values in parentheses reflect reactivity using vvTK⁻-infected DCs as control targets in the ELISPOT assay.

as described using intracellular staining of a vaccinia early protein of 29 kD with the VV1-6B6 antibody (49).
the restimulation to a final concentration of 10 U/ml (Lympho-
cult). Microwell cultures were tested in split well 51Cr-release assays
against autologous DCs infected with vvEBNA1ΔGA or vvTK-, autologous B-LCLs, or LCL721.221. As <30% of the wells de-
developed CTLs or IFN-γ-secreting cells, it is >90% probable that
the responding wells represent clones (50). The BC cell line was
generated separating EBNA1-specific CD4+ T cells after stimula-
tion with vvEBNA1ΔGA-infected DCs using the IFN-γ secretion
assay according to the manufacturer’s instructions (Miltenyi
Biotech).

FACS Analysis of Stimulated CD4+ T Cell Populations and
PBMCs. Mature DCs were infected with recombinant vv at an
MOI of 2, or with influenza virus (PR8, Puerto Rico/8/34;
Spafas, Inc.) at an MOI of 0.5 for 1 h at 37°C in RPMI 1640 H.S.
DCs were washed twice, and 3 x 105 were added to 106 CD8-
CD2+ PBMCs in 96-well plates for 7 d. The cultures were restim-
ulated with irradiated (3,000 rads) 105 PBMCs and 3 x 105 DCs
per well and incubated for an additional 7 d. At day 14, cultures
were stained for 30 min on ice with 1 µg Simulset CD4-FITC/
CD8-PE (Becton Dickinson) and analyzed on a FACSscan™
(Becton Dickinson). CD56 antibody staining (BD PharMingen)
was used PE–goat anti–mouse IgG antibody (Biosource Interna-
tional) as secondary, PBMCs were typed for HLA-DR4 using HLA-
DR4 antibody (Accurate) as primary and FITC–goat anti–mouse
 IgG antibody (Biosource International) as secondary.

Enzyme-linked Immunospot Assay for IFN-γ-secreting Cells.
Enzyme-linked immunospot (ELISPOT) assays were performed
as described previously (51). MAHA S45 plates (Millipore) were
coated with anti–IFN-γ antibody 1-D1K (Mabtech). Afterwards, preassem-
lated EBNA1-specific IFN-γ-secreting CD4+ T cells (“blasts”) were stimulated and restimulated with vvEBNA1 (24, 25). Rponses were as-
essed by the presence of enlarged CD4+ T cells (“blasts”) after 2-wk-long stimulations with DCs. In the first week, one of a panel of vvEBV recombinants was used to stimu-
late the CD4+ T cells. Then the cultures were divided in two and restimulated for a second week with the original recombinant vv or with vvTK- as control. We looked for
blastogenesis specific to the EBV recombinant that stimu-
lated the CD4+ cultures in the first week.

All 10 donors showed strong responses to vvEBNA1 (Tables I and II, and Fig. 1, A and B). The response to the

\begin{figure}
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\caption{EBNA1 is recognized by CD4+ T cells from healthy EBV carriers. Blast formation (large forward scatter [FSC], x-axis) by CD4+ T cells (CD4-FITC, y-axis) was monitored by flow cytometry. Cultures of CD2-CD8+ T cells were stimulated with autologous DCs infected with vv constructs. (A) Culture stimulated with vvEBNA1ΔGA-infected DCs and restimulated with vvTK- infected DCs. (B) Blasting of a culture stimulated with vvEBNA1ΔGA-infected DCs. (C) Culture stimulated with vvEBNA3A-infected DCs and restimulated with vvEBNA1ΔGA-infected DCs. (D) T cells were stimulated and restimulated with vvEBNA3A-infected DCs. (E) Culture stimulated and restimulated with vvTK- infected DCs to evaluate the background of vaccinia stimulation. (F) CD2-CD8+ T cells responding to influenza virus-infected DCs as positive control. All cultures were prepared from the same donor (no. 5 in Tables I and II). Percentage of blasts was indicated. 1°, first stimulation; 2°, restimulation.}
\end{figure}
negative control (vvTK) were weak (Fig. 1 E) in all but one donor, excluded from the Tables. All donors responded to influenza-infected DCs as a positive control (Fig. 1 F). CD4 T cell responses by the 10 donors to the other vvEBV constructs were detected less consistently: EBNA3B (5/10), EBNA3A (1/10), EBNA3C (1/10), and LMP1 (6/10) (Table I). To ensure that all the recombinant vv infected a comparable proportion of the mature DCs, the intracellular expression of the 29-kD vaccinia early protein was measured by FACS®. Reproducibly, 40–60% of DCs were infected with the different vv (data not shown). The reliability of the CD4+ recognition of EBNA1 was also evident in an ELISPOT assay for IFN-γ secretion, where EBNA1 was the EBV latency gene most frequently recognized (Table II).

We regard these CD4+ T cell responses to EBNA1 to reflect priming by EBV infection of the blood donors in vivo, as we did not see blastogenesis in 2 wk if we stimulated neonatal T cells from cord blood specimens with EBNA1 (Fig. 2). The fetal CD4+ T cells performed similarly to adult CD4+ T cells in MLR proliferation assays, and the fetal DCs were likewise capable of eliciting strong MLR proliferation of adult CD4+ T cells (Fig. 2 A). However, none of the fetal samples were able to recognize EBV or influenza products in IFN-γ ELISPOT assays (Fig. 2). CD4+ T cells from adult controls recognized vvEBNA1ΔGA, vvEBNA3B, vvLMP1, and influenza-infected autologous DCs in the same assay (Fig. 2). CD4+ T cells recognize EBNA1 in an MHC class II–restricted fashion. We verified that our donors showed HLA class II diversity, as only two expressed HLA-DR4 (data not shown). To establish that MHC class II products were presenting EBNA1, we generated T cell lines, initially, from an HLA-DR4+ donor. We then assessed reactivity of the lines with DCs infected with recombinant vv expressing EBNA1 or pulsed with soluble EBNA1 protein. One line, CM 171198, was derived from CD8-CD2- PBMCs stimulated alternatively with autologous DCs infected with vvEBNA1ΔGA or the DR4-matched B-LCL LR M. The vvEBNA1 construct was deleted of the GA repeat that blocks MHC class I presentation and also reduces expression of EBNA1. Another line, CM 110199, was stimulated with DCs charged during their final maturation with recombinant EBNA1 protein expressed either in E. coli (EBNA1) or in a baculovirus/insect cell system (EBNA1). After 1 mo of culture, both lines were predominantly CD4+ T cells, 90% in CM 171198 and 76% in CM 110199, with CD56+ NK cells being the main contaminant (not shown).

The CD4+ T cell lines recognized DCs that were infected with vvEBNA1ΔGA or exposed to recombinant EBNA1 (Fig. 3, A and B). Reactivity could be measured as IFN-γ secretion (ELISPOT assays, Fig. 3 A) or by proliferation (Fig. 3 B). The T cell responses were blocked by an anti–HLA-DR antibody, L243, but not anti–HLA class I antibody, B-H9 (Fig. 3 A). In addition to DCs charged with EBNA1, the CM 171198 cell line recognized EBV-transformed B-LCLs without further addition of antigen (Fig. 3, A and B, bottom). In B-LCLs only full-length EBNA1 is expressed at detectable levels (24). This implies that full-length EBNA1, as expressed endogenously by B-LCLs, can be processed on MHC class II molecules for presentation.
CD4+ T cell recognition. The B-LCLs had to be matched at the DR4 allele to trigger T cell function. Thus, DR4+ B-LCLs (LRM and LCL-BM) induced proliferation, but DR4− cells (LG2 and LCL-DC) did not (Fig. 3, A and B, bottom).

EBNA1-specific CD4+ T Cells Kill B-LCLs. To determine if EBNA1 was an antigen for CD4+ CTLs, we stimulated CD8−CD2+ PBMCs from an HLA-DR4+ donor, JT, with irradiated autologous B-LCLs (expressing various known latent EBV antigens [3]) for 2 wk. In parallel, the B-LCLs were used to stimulate bulk CD2+ and CD4−CD2+ T cells. The content of the stimulated T cell populations was determined by FACSCalibur®. CD8-depleted responders were enriched for CD4+ cells, CD4-depleted responders were enriched for CD8+ cells, and the bulk T cells had a CD4+/CD8+ ratio of 1:2 (see Fig. 5, top). All contained ~25% CD56+ NK cells (not shown).

31Cr-release assays showed that the three populations of stimulated T cells killed autologous B-LCLs, with less recognition of the T2 cell line (Fig. 4, bottom left). As expected, the killing of autologous B-LCLs was completely blocked by L243 anti–HLA-DR antibody when CD4+ enriched populations, but not CD8+ enriched cultures, were tested (Fig. 4, bottom left). However, the CD8+ enriched cultures repeatedly developed stronger cytolytic activity than CD4+ enriched cultures. Killing by the bulk T cells was partially inhibited by the L243 antibody, but the block was ≤50% in our three experiments.

EBNA1-specificCTL function was also assessed using DC targets that had been infected with vvEBNA1ΔGA or recombinant eEBNA1 and bEBNA1 proteins (Fig. 4). CD4+ enriched T cells lyzed EBNA1-pulsed DCs. In contrast, CD8+ enriched cultures and bulk T cells were able to kill LCLs, but not the EBNA1-pulsed DCs (Fig. 4, bottom right).

To begin to determine if individual CD4+ T cells could lyse DCs pulsed with EBNA1, as well as B-LCLs expressing EBNA1 endogenously, we used autologous B-LCLs to isolate CD4+ CTLs by limiting dilution from cryopreserved T cells. The DCs, T cells, and autologous B-LCLs were derived from leukocyte concentrates. All 11 limiting dilution sublines that killed DCs in an EBNA1-dependent fashion also killed autologous B-LCLs, and we did not find any clone that killed the DCs and not the B-LCLs (Fig. 5). This indicates a tight correlation between recognition of epitopes expressed by B-LCLs and recognition of DCs that have been pulsed with EBNA1. As we were studying cells obtained under limiting dilution conditions, with <30% recognizing B-LCLs, it is >90% likely that individual clones were responsible for killing DCs and B-LCLs, although formal cloning experiments will be required (50).

R recognition of DCs infected with the vvTK− control vector or LCL721.221, an HLA class I−negative NK target, was poor in all 11 sublines (Fig. 5). Therefore, CD4+ T cells can lyse autologous B-LCLs, and one target very likely is EBNA1.

EBNA1 Is Recognized on B-LCLs after Endogenous Processing. As we were observing EBNA1-specific CD4+ cells that make Th1 cytokines and exert cytolytic activity upon encountering transformed B-LCLs it was important to establish that this new and potentially protective mechanism would operate on EBNA1 that was processed endogenously by transformed B cells. The alternative would be that in our cultures some cells were dying and were being reprocessed via an exogenous or endocytic pathway in a fraction of LCLs. We are not aware of inhibitors that efficiently and selectively block processing of EBNA1 from an endogenous or exogenous route.

Therefore, we first compared the capacity of DCs (as a positive control) and the EBV− Burkitt's lymphoma cell line, Ramos, to present EBNA1 through an exogenous pathway, either rEBNA1 protein or EBNA1 expressed by
allogeneic B-LCLs (Fig. 6 A). Because of MHC class II mismatching, the allogeneic B-LCLs could not directly present EBNA1 to T cell lines that had been selected for IFN-γ secretion upon stimulation with vvEBNA1ΔGA-infected autologous DCs. The Ramos cell line as well as the autologous DCs could present vvEBNA1ΔGA to an EBNA1-specific CD4+ T cell line from a donor matched at HLA-DR7 to Ramos (Fig. 6 A). In contrast, only the DCs, and not the Ramos Burkitt's lymphoma cells, presented exogenous rEBNA1 and EBNA1 from allogeneic LCLs (having ~20% trypan blue-positive or dead cells; Fig. 6 A).

DCs also presented EBNA1 from additional exogenous sources. In Fig. 6 B, DCs presented EBNA1 from four different allogeneic DC preparations infected with vvEBNA1ΔGA, presumably because infection with vv is cytotoxic for some of the infected DCs (49). As vv-infected DCs cannot produce virus particles (49, 53), coinfection could not be responsible for the observed EBNA1 transfer from one DC to another. A fifth semiallogeneic DC preparation presented vvEBNA1ΔGA directly to the T cell line (Fig. 6 B).

Although there is already evidence in the literature that Burkitt's lymphoma cells are comparably efficient to B-LCLs in presenting soluble antigen on MHC class II (54), we wanted to show that both types of lymphoma also share the inability to cross-present antigen from cocultured cells. Therefore, we infected mismatched DCs and EBV latency III type Daudi cells with recombinant vv expressing influenza N1 or M1, and then looked for cross-presentation of these influenza products by allogeneic B-LCLs or DCs, to CD4+ T cells of an influenza-reactive donor. Again, the DCs could present antigen from the allogeneic vvN1- or vvM1-infected cells, but the B-LCLs could not (Fig. 7). As a positive control, we showed that autologous B-LCLs could present N1 (Fig. 7 A) and M1 (Fig. 7 B) to CD4+ T cells when directly infected with vvN1 or vvM1, respectively. Coculturing B-LCLs with allogeneic infected B-LCLs for 2 d apparently did not transfer sufficient antigen amounts to the matched B-LCLs to trigger IFN-γ secretion by CD4+ T cells. In contrast, N1- or M1-expressing allogeneic B-LCLs could be efficiently processed by DCs and presented to CD4+ T cells.

This series of experiments indicates that DCs efficiently cross-present on MHC class II antigens (EBNA1, N1, and M1) from allogeneic DCs or B-LCLs, but that transformed B cells (B-LCLs or Burkitt's lymphoma cells) are weak or inactive in this respect. The findings are consistent with prior publications showing that B-LCLs can present soluble proteins that bind to their Ig receptor, but not by a nonspecific exogenous route (55). Therefore, CD4 T cell recognition of EBNA1 in B-LCLs very likely represents recognition of the endogenously processed protein that is critical for EBV-induced transformation.

Discussion

EBNA1-specific CD4+ T Cell Immunity Exists in All Healthy Adults. In an effort to define the CD4+ T cell repertoire for EBV latency gene products, we have uncovered a consistent CD4+ T cell response to EBNA1 presented by DCs and by allogeneic B-LCLs. The EBNA1-specific CD4+ T cells proliferate, secrete IFN-γ, kill targets, and can be readily propagated as EBNA1-specific, MHC class II-restricted lines. Other latency antigens (LM P1, LM P2, EBNA3A, EBNA3B, and EBNA3C; Tables I and II) can be recognized by CD4+ T cells, but less consistently. The CD4+ T cells described here are found in
bulk cultures and kill B-LCLs without further addition of EBNA1. The results are in marked contrast to the prior literature that has described only a single EBNA1-specific CD4+ T cell clone, and it only killed B-LCL targets upon supplementation with exogenous EBNA1 (39). The consistent detection of EBNA1 immunity in our studies could reflect the use of CD4+ T cells as responders, and DCs as APCs.

The consistent recognition of EBNA1 by CD4+ T cells is strikingly different from CD8+ T cells. CD8+ T cells, specific for EBNA1-derived MHC class I epitopes, can be found in blood, but they do not recognize autologous or HLA-matched B-LCLs (24). This indicates that the physiological levels of EBNA1 are not sufficiently processed onto MHC class I, nor do B-LCLs cross-present EBNA1 when B-LCLs die in culture. The block of endogenous processing of EBNA1 is well understood, i.e., the Gly/Ala repeat prevents efficient proteasomal degradation and therefore epitope generation for MHC class I presentation (25, 26).

In light of the efficient cross-presentation of antigens on
MHCI by DCs (56), it is likely that the observed EBNA1-specific CD8+ T cells are primed by DCs that process B-LCLs onto MHCI class I. However, because of the Gly/Ala repeat, such CD8+ T cells would not see EBNA1 expressed by EBV-infected cells.

Evidence that endogenous processing and presentation of Physiological Epitope Levels on B-LCLs are sufficient to elicit EBNA1-specific CD4+ T cell immunity. The EBNA1-specific CD4+ T cells described in this study recognize autologous B-LCLs by proliferation, IFN-γ secretion, and cytotoxicity. As these responses are MHCI class II restricted, and because EBNA1 is expressed endogenously in B-LCLs, two routes of processing can be envisioned. Either EBNA1 is processed directly by the EBV-infected cell itself (endogenous pathway), or dying EBNA1-expressing B-LCLs are endocytosed and presented (indirect or exogenous or cross-presentation pathway). This does not necessarily mean that the processing machinery differs, but that in the first endogenous case the antigen is processed in the cell that synthesizes EBNA1, making it a more reliable target for protective immunity in humans. From our experiments, exogenous processing is an unlikely explanation for presentation of EBNA1 on B-LCLs, as antigen transfer from HLA-mismatched cells was not observed (Fig. 6). B-LCLs were previously shown to be poor APCs for exogenous proteins (55), 100–300 times less efficient than DCs. Similarly we found that the Ramos Burkitt’s lymphoma line inefficiently processed rEBNA1 and cross-presented EBNA1 from cocultured B-LCLs to CD4+ effector cells (Fig. 6 A). Cross-presentation of influenza NS1 and MP by B-LCLs from vvNS1- or vvMP-infected DCs or EBV latency III type Daudi cells was also inefficient (Fig. 7). In contrast, DCs cross-presented antigen from B-LCLs or vvEBNA1ΔGAG-infected DCs, and processed rEBNA1 (Fig. 6, A and B).

This was not restricted to EBNA1, as DCs also presented N31 and MP from cocultured vvNS1- and vvMP-infected DCs and Daudi cells (Fig. 7). As a positive control for the competence of the lymphoma cells as APCs, we showed that direct infection of Ramos or B-LCL cells with recombinant EBNA1 or influenza vv led to CD4+ T cell responses. This implies that in healthy carriers, the DCs that are known initiators of immune responses (41) cross-present B-LCL-derived EBNA1 for priming of specific CD4+ T cells. Once activated, these CD4+ T cells may attack B cells that process EBNA1 endogenously and contribute to EBV-specific immunity.

The Gly/Ala Repeat Domain Does Not Influence MHC Class II Processing of EBNA1. Our experiments confirm that EBNA1 can be processed onto MHCI class II irrespective of the presence or absence of its Gly/Ala repeat. This domain inhibits proteasome-dependent processing for MHCI class I (25, 26). B-LCLs, the only reliable source of full-length EBNA1 (24), readily present EBNA1 to specific CD4+ T cells (Figs. 2–4). Moreover, B-LCLs with full-length EBNA1 can be cross-presented by DCs (Fig. 5 B). Therefore, the proposal that EBNA1 prevents its endogenous MHCI class II presentation (39) cannot be supported by our findings.

Evidence that CD4+ T cell immunity could contribute to the control of EBV. EBNA1-specific CD4+ T cells could provide direct resistance to EBV-transformed cells, through their cytokines and lytic function or by sustaining the CD8+ CTL response to other lymphoma-related EBV products such as LMP1 and LMP2. A good deal of circumstantial evidence for CD4+ T cell protection against gamma herpesviruses in vivo exists. (a) CTLs to EBV in the cottontop tamarin Sanguinis oedipus (57) are to a large extent MHCI class II restricted (58). MHCI class I-restricted, EBV-specific CTLs have yet to be found in the New World monkey, and this species lacks classical MHCI class I (although it does express homologues of nonclassical class I genes like HLA-G and HLA-F; 59). (b) Gamma herpesvi-
isolated CD4+ T cell recognition hierarchy. Previously, the specificities of only two bulk T cell preparations place EBNA1 at the top of the recognition hierarchy. Impaired CD4+ T cell responses are thought to be responsible for the EBV-induced mononucleosis seen in X-linked lymphoproliferative disease patients who have a mutation or deletion in signaling lymphocyte activation molecule (SLAM)-associated protein (SAP), an inhibitor of the T cell costimulatory molecule SLAM or CDw150 (63). This lack in SLAM function especially affects Th1 immunity, as SLAM engagement mediates increased IFN-γ secretion (64, 65). Thus, EBNA1-specific Th1 cells are probably disabled in X-linked lymphoproliferative patients.

Our findings uncover an immune response that changes current thinking on immune surveillance against EBV. The data on CD4+ T cell responses to EBV latent antigens in bulk T cell preparations place EBNA1 at the top of the recognition hierarchy. Previously, the specificities of only two isolated CD4+ T cell clones have been described (39, 54). Since only the EBNA2-specific CD4+ T cell clone recognized B-CLLS, the significance for CD4+ T cell recognition was questionable, particularly in the context of latency I and II programs of EBV transformation. However, the new data reveal a hierarchy of CD4+ T cell recognition that differs significantly from CD8+ T cell responses to EBV (28). EBNA3A, EBNA3B, and EBNA3C antigens dominate for CD8+ T cell recognition, but play a subdominant role for CD4+ T cells. In contrast, EBNA1, believed to be invisible to the immune system because of a block in its MHC class I presentation (24), is the main target of MHC class II-restricted CD4+ T cell responses. As we find these responses in all tested healthy adults, we suggest that EBNA1-specific CD4+ T cells provide resistance to the development of Burkitt's lymphomas, Hodgkin's, and other EBV-associated malignancies. Likewise, the new data suggest that EBNA1 be tested as an antigen to prevent and treat such malignancies.

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