Host Ethnicity and Virus Genotype Shape the Hepatitis B Virus-Specific T-Cell Repertoire

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Repertoire composition, quantity, and qualitative functional ability are the parameters that define virus-specific T-cell responses and are linked with their potential to control infection. We took advantage of the segregation of different hepatitis B virus (HBV) genotypes in geographically and genetically distinct host populations to directly analyze the impact that host and virus variables exert on these virus-specific T-cell parameters. T-cell responses against the entire HBV proteome were analyzed in a total of 109 HBV-infected subjects of distinct ethnicities (47 of Chinese origin and 62 of Caucasian origin). We demonstrate that HBV-specific T-cell quantity is determined by the virological and clinical profiles of the patients, which outweigh any influence of race or viral diversity. In contrast, HBV-specific T-cell repertoires are divergent in the two ethnic groups, with T-cell epitopes frequently found in Caucasian patients seldom detected in Chinese patients. In conclusion, we provide a direct biological evaluation of the impact that host and virus variables exert on virus-specific T-cell responses. The discordance between HBV-specific CD8+ T-cell repertoires present in Caucasian and Chinese subjects shows the ability of HLA micropolymorphisms to diversify T-cell responses and has implications for the rational development of therapeutic and prophylactic vaccines for worldwide use.

Virus-specific CD8+ T cells recognize virus-encoded peptides associated with major histocompatibility complex (MHC) class I molecules displayed on the surfaces of the infected cells. Virally infected cells can produce thousands of potentially immunogenic peptides, but CD8+ T cells are usually directed against only a few peptides, and CD8+ T cells specific for different viral determinants can possess different antiviral activities (47). Information regarding virus-specific T-cell repertoires and the potential antiviral efficacies of CD8+ T cells with differing antigen specificities is essential to understand viral pathogenesis and develop vaccines. Such information is limited in the great majority of viral and bacterial infections due to cumbersome methods that are required for the detection and characterization of new MHC class I-restricted epitopes (47). In addition, the identification of the T-cell repertoire against viruses infecting different ethnic populations with distinct HLA class I alleles and haplotype frequencies is particularly complex because different ethnic groups are often infected by different viral strains, which are likely to have coevolved in these populations (11, 16, 29).

The influence that virus heterogeneity and the distinct HLA profiles of the infected subjects has on the repertoire and hierarchy of T-cell responses is difficult to predict. The existence of T-cell responses against conserved regions of different virus strains (13, 46) and the reported degeneracy in HLA-peptide binding, with identical peptides able to bind multiple HLA class I types (6, 14, 35, 37, 42), support the idea of a substantial overlap between the virus-specific T-cell repertoires of subjects of different ethnicities expressing closely related but distinct HLA class I molecules. On the other hand, viral heterogeneity might affect the generation of certain epitopes as strain-specific variations within the epitopes (3) or, in flanking regions, might impair their processing and presentation (23, 34). Even subtle differences in closely conserved HLA class I molecules (28, 37) may severely affect the presentation of specific epitopes (2, 40) or change their conformation (44) sufficiently so that individuals of different ethnicities may focus the response toward different T-cell epitopes.

Given the global distribution of hepatitis B virus (HBV), understanding the commonality or divergence of virus-specific T-cell responses present in HBV-infected patients with different ethnicities is necessary. HBV, a strictly hepatotropic virus, induces a functionally efficient, multispecific CD8+ T-cell response in subjects who resolve the infection, while HBV-specific CD8+ T cells are not present or functionally impaired in

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chronically infected patients (5). A comprehensive knowledge of HBV-specific CD8\(^+\) T-cell specificities is lacking, and with rare exceptions (7, 9, 43), CD8\(^+\) T-cell responses have been analyzed using preselected peptides able to bind to common HLA class I molecules (HLA-A2, -A3, -A24, -A11, and -B7) (19, 20, 26, 31, 32, 38, 43, 45). Attempts to define immunodominant regions in the HBV proteome were based on the use of HLA-A2-restricted epitopes (45) and on samples from HBV genotype A (HBV\(_{\text{genA}}\)) or HBV\(_{\text{genD}}\)-infected individuals of Caucasian decent. However, 75% of the population of chronically infected patients live in Asia (27), and Asian patients are infected, mostly at birth, by HBV\(_{\text{genC}}\) or HBV\(_{\text{genD}}\), which differ by nearly 8% in amino acid composition compared to genotypes A and D (21).

The HLA class I profiles of the two populations differ not only in the frequency of the major HLA class I alleles (i.e., HLA-A11 is present in 51.7% of Chinese and 14% of Caucasians; HLA-B40 is present in 31.5% of Chinese and 14.7% of Caucasians [28]) but are also characterized by substantial differences in allele subtypes. The HLA-A2 molecule, present in nearly 50% of both Caucasians and Chinese, is subdivided into HLA-A2 subtypes, which are differentially expressed in the two ethnic groups (22). More than 95% of HLA-A2\(^+\) Caucasians are HLA-A201\(^+\), whereas subtypes HLA-A0203, -A0206, and -A0207 are, respectively, present in 23%, 10%, and 45% of HLA-A2\(^+\) individuals of Chinese origin (22). Therefore, we performed the first direct comprehensive analysis of HBV-specific T-cell responses present in patients of different ethnicities (Chinese versus Caucasian) infected by different HBV genotypes (HBV\(_{\text{genA}}\) versus HBV\(_{\text{genD}}\)) to understand whether and, if so, to what degree host and virus variables influence the virus-specific T-cell response.

**MATERIALS AND METHODS**

**Patient selection.** A total of 76 HBV-infected patients of Chinese Han origin and 78 Caucasians of European descent were enrolled at the University of Infectious Diseases and Hepatology of the Azienda Ospedaliero-Universitaria of Parma, Italy; the University Hospital S. Orosola-Malpighi of Bologna, Italy; or the National University of Singapore Hospital. Twenty patients (8 Chinese and 12 Caucasians) had clinical, biochemical, and virological evidence of acute HBV infection (alanine aminotransferase [ALT] levels >20 times the upper limit of normal, detection of HBsAg and anti-HBe immunoglobulin M [IgM], and HBsAg clearance within 2 months from the clinical onset of hepatitis). A total of 64 HBV-infected patients of Chinese origin and 66 of Caucasian origin displayed HBsAg clearance within 2 months from the clinical onset of hepatitis. A total of 62 HBV genD-infected Caucasians were selected for HBV-specific T-cell analysis. Sixteen Caucasian HBV genA patients (5 Chinese and 11 Caucasian) were also excluded and 62 HBV genD-infected Caucasians were selected for further analysis. HLA-A2 subtypes of HLA-A2 patients (selected by a low-resolution genetic approach) were determined by high-resolution sequencing of the A2 locus (direct sequencing of the alpha 1 and alpha 2 chains).

This study was approved by the ethical committees of the Azienda Ospedaliero-Universitaria of Parma, the University Hospital of Bologna, and the National University of Singapore Hospital, and all subjects gave written informed consent.

**Virological assessment.** HBsAg, HBeAg, anti-HBs, anti-HBe, HBV IgG and IgM, anti-HBc, anti-HDV, anti-HCV, and anti-HIV were determined by commercial enzyme immunoassay kits (Abbott Labs, Abbott Park, IL; Ortho Clinical Diagnostics, Johnson & Johnson, Raritan, NJ; DiaSorin, Vercelli, Italy). Serum HBV DNA was quantified by PCR (Cobas Amplicor test; Roche Diagnostics, Basel, Switzerland). HBV genotyping was performed by restriction fragment length polymorphism analysis of a pre-S amplicon previously described by Lindt et al. (25).

**Isolation of PBMC and in vitro expansion of HBV-specific CD8\(^+\) cells.** Peripheral blood mononuclear cells (PBMC) were isolated from fresh heparinized blood by Ficoll-Paque density gradient centrifugation and resuspended in AIM-V medium (Invitrogen, Carlsbad, CA) with 2% pooled human AB serum. For the in vitro assays, the cells were used either directly ex vivo or after a 10-day antigen-specific in vitro stimulation. For the latter, 20% of the PBMC was first stimulated with 10 \(\mu\)g/ml of all the overlapping peptides from the respective HBV genotypes for 1 h at 37°C, then washed at 3.0 \(\times\) 10\(^6\) cells/ml before coculturing with the remaining PBMC in AIM-V medium with 2% pooled human AB serum supplemented with interleukin-2 (IL-2; R&D Systems, Abingdon, United Kingdom) (20 IU/ml) and seeded at 1 ml/well in 24-well plates. The immunological assays were performed on day 10 of the expansion.

**Synthetic peptides and antibodies.** Two panels of 313 15-mer peptides overlapping by 10 residues were used to test HBV-specific T-cell responses. The peptides covered the overall sequence of HBV\(_{\text{genA-D}}\) (GenBank accession number AF121241) and HBV\(_{\text{genA}}\) (GenBank accession number AF121243) and were purchased from Chiron Mimotopes (Victoria, Australia) or synthesized at the peptide synthesis facility of Massachusetts General Hospital using 9-fluorenylmethoxy carbonyl chemistry. The purity of the peptides was above 90%, and their composition was confirmed by mass spectrometry analysis. The designed peptides presented at least 95% similarity with those encoded by HBV\(_{\text{genA-D}}\) sequences from five Chinese and five Caucasian patients studied. The 15-mer core peptides were pooled in a 9 by 8 matrix containing right or left or both peptides/pool, respectively, using a concept similar to what was previously described for HIV (1). Envelope peptides were pooled in a 9 by 9 matrix containing nine peptides/pool, while polymerase peptides were pooled in a 14 by 12 matrix containing 12 or 14 peptides/pool, respectively. All peptides were first diluted to 40 mg/ml in dimethyl sulfoxide and then further diluted in RPMI medium at a working dilution (between 1 mg/ml and 1 ng/ml).

**Peptide- and HLA-defined HLA-A2-restricted cytolytic T lymphocyte (CTL) epitopes** were selected from five Chinese and five Caucasian patients studied. The 15-mer core and X peptides were pooled in a 9 by 8 matrix containing right or left or both peptides/pool, respectively, using a concept similar to what was previously described for HIV (1). Envelope peptides were pooled in a 9 by 9 matrix containing nine peptides/pool, while polymerase peptides were pooled in a 14 by 12 matrix containing 12 or 14 peptides/pool, respectively. All peptides were first diluted to 40 mg/ml in dimethyl sulfoxide and then further diluted in RPMI medium at a working dilution (between 1 mg/ml and 1 ng/ml).

**Hepatitis B virus (HBV) genotypes** were identified by high-resolution sequencing of the A2 locus and by mass spectrometry analysis. The designed peptides were purchased from GenScript (Piscataway, NJ). The peptide sequences were based on genotype-specific sequences of 24 GenBank entries (6 HBV\(_{\text{genA}}\), 8 HBV\(_{\text{genB}}\), 6 HBV\(_{\text{genC}}\), and 4 HBV\(_{\text{genD}}\)). Furthermore, a viral amino acid sequence analysis of the Core18-27 and EnvI35-41 regions of the Chinese and Caucasian HLA-A2\(^+\) patients studied confirmed the genotype-specific sequence of the infecting viral strain. Anti-CD8 (phycoerythrin [PE]-CY7), anti-CD3 (peridinin chlorophyll protein-Cy5.5), and anti-CD107a (fluorescein isothiocyanate) antibodies were purchased from Becton Dickinson Pharmingen (San Jose, CA). Anti-γ interferon (anti-IFN-γ; PE) was purchased from R&D Systems (Minneapolis, MN). Anti-granzyme B (clone H92, San Diego, CA) and 5\(\times\)10\(^5\) Plasmocin (InvivoGen, San Diego, CA) to prevent microbial contamination.

**IFN-γ ELISPOT assay.** IFN-γ, a TNF-α–like cytokine of entamoeba histolytica (ELISPOT) assays were performed as previously described (7) using a panel of 313 overlapping peptides covering the entire peptide sequence of HBV\(_{\text{genA-D}}\) or HBV\(_{\text{genA-D}}\) pooled in the described mixtures and used in patients infected with the respective HBV.
genotype. HBV-specific T-cell responses were analyzed in IFN-γ ELISPOT assays either ex vivo using fresh or frozen PBMC or after short-term peptide-specific polyclonal T-cell expansion (10 days). Briefly, 96-well plates (Multi-screen-HTS; Millipore, Billerica, MA) were coated overnight at 4°C as recommended by the manufacturer with 5 μg/ml capture mouse anti-human IFN-γ MAb (IDIK, Mabtech, Sweden). The plates were then washed five times with phosphate-buffered saline and blocked with AIM-V supplemented with 10% heat-inactivated fetal calf serum for 30 min at room temperature. A total of 1 × 10^5 PBMC or 5 × 10^4 cells from short-term polyclonal T-cell lines were seeded per well, in duplicate for each individual peptide mixture. The plates were incubated for 18 h at 37°C in the presence or absence of peptides (at a final concentration of 5 μM). After five washes with phosphate-buffered saline, 100 μl of 0.5 μg/ml biotinylated anti-human IFN-γ MAb (7B6-1; Mabtech, Sweden) was added; plates were incubated for 2 h at room temperature and thereafter washed five times, 100 μl of streptavidin-alkaline phosphatase (1:2,000 dilution) (Mabtech, Sweden) was added to each well for 1 h at room temperature, and plates were incubated in the dark. The plates were again washed five times, and 50 μl of alkaline phosphatase substrate (5-bromo-4-chloro-3-indolyl phosphate-nitro blue tetrazolium chloride [BCIP-NBT]; KPL, Gaithersburg, MD) was added. After 10 to 15 min, the colorimetric reaction was stopped by washing with distilled water. The plates were air-dried, and spots were counted using an automated ELISPOT reader (ImmunoSpot; CTL, Cleveland, OH). The number of IFN-γ-producing cells was expressed in spot-forming units (SFU) per 1 × 10^5 cells. The number of specific IFN-γ-secreting cells was calculated by subtracting the nonstimulated control value from the stimulated sample. Positive controls consisted of PBMC stimulated with staphylococcal enterotoxin B (SEB) or phytohemagglutinin. In the direct ex vivo assays, a well was considered positive when the number of SFU was more than 5 and at least three times above the mean of the unstimulated control wells (3 wells/patient). The responses to peptide mixtures were also analyzed directly ex vivo in nine healthy subjects, and only a single individual showed the presence of a positive response. The positivity criteria for ex vivo ELISPOT assays is less stringent, including wells that have SFU of more than 5 and at least two times above the mean of the unstimulated control wells. However, ICS was applied to every positive sample to reconfirm the response and to determine the T-cell subset (CD8 or CD4) responsible for IFN-γ production.

Image analysis. A series 3B ImmunoSpot image analyzer (Cellular Technology) specifically designed for the ELISPOT assay was used. The digitized images were analyzed for the presence of areas in which the color density exceeded the background by an amount set on the basis of the comparison of wells with peptide and wells without peptide. After background and noise subtraction, custom software is used to analyze spot morphology for circularity and density distribution to identify and separate touching and overlapping spots. Objects that meet these criteria are recognized as spots and counted. The measurement of the mean spot size is a built-in function of the software.

Statistical analysis. An unpaired t test was used in two instances: (i) to determine the significance of the difference in the mean percentage of positive mixtures between acute and chronic patients and (ii) to determine the significance of the difference in the ELISPOT assay-derived mean spot sizes between acute and chronic patients stimulated with HBV peptides or SEB/phytohemagglutinin. Differences with a P value less than 0.05 were considered statistically significant.

RESULTS

Comprehensive analysis of HBV-specific T-cell responses in HBVgenB- and HBVgenD-infected patients. To identify similarities/differences in the breadth and magnitude of HBV-specific T-cell responses between Caucasian and Chinese patients, HBV-specific IFN-γ-producing T cells were tested by direct ex vivo ELISPOT assays in 34 patients of Chinese origin (6 acute and 28 chronic HBVgenB infected) and in 37 Caucasian subjects (4 acute and 33 chronic HBVgenD infected) (the full clinical profiles of all patients tested are shown in Fig. S1 in the supplemental material). PBMC of Chinese patients were stimulated with the HBVgenB peptide panel, while Caucasian patients were stimulated with HBVgenD peptides. Consistent with previous data, obtained mainly for Caucasian patients (7, 9), HBV-specific T-cell responses were detected ex vivo only in patients with acute HBV infection (6/6 Chinese and 4/4 Caucasian subjects). Responses in chronic patients (18% [5/27] of Chinese and 15% [5/33] of all Caucasian individuals) were rarely observed ex vivo (Fig. 1a and b), indicating that clinical status was a stronger predictor of detectable responses than ethnicity or infecting genotype. In line with previous results (45), the envelope-specific T-cell response appears to be the only weak response detectable ex vivo in chronic HBV patients with a high HBV load (Fig. 1b).

HBV-specific T-cell responses were also compared after in vitro expansion in 17 HBVgenB-infected Chinese and 15 HBVgenD-infected Caucasian subjects. HBV-specific T-cell frequency was generally low directly ex vivo but became clearly detectable in all acute patients after in vitro expansion (Fig. 2a), while such expansion was defective in chronic patients irrespective of ethnicity and HBV genotype (Fig. 2b). The data for all patients are summarized in Fig. 2c, which provides a comparison of the numbers of peptide mixtures able to elicit ELISPOT assay responses in acute versus chronic HBV patients. We observed the same general pattern of efficient expansion and multispecific T-cell responses in acute patients compared to weak and narrower T-cell responses in chronic patients, irrespective of ethnicities and HBV genotypes. Furthermore, previous reports suggested that the magnitude of the HBV-specific T-cell response is inversely correlated with the serum HBV DNA level (7, 45); however, this relationship was not observed in our data, and there was no correlation between HBV-specific T-cell expansion and HBeAg/anti-HBe status, which is probably attributable to the limited sample size (data not shown).

Functional defects of HBV-specific T cells from chronic patients. The functional alteration of IFN-γ production by T cells has been reported in both Chinese (10) and Caucasian (7) chronic HBV patients. While such defects were shown to be HBV specific in Caucasians, a recent report suggested that T-cell defects present in Chinese chronic HBV patients were more pronounced and were caused by blockage of IFN-γ production (10). To determine if there was an impairment of IFN-γ production in chronic HBV patients, we examined the relative amount of IFN-γ secreted by HBV-specific and non-HBV-specific T cells from healthy and HBV-infected subjects. Spot sizes (18) obtained by direct ex vivo ELISPOT assays from a total of 48 Chinese chronic HBV patients (22 with HBV DNA loads of <10^6 and 26 with HBV DNA loads of >10^5) probed with 4 patients with acute HBV infection, and 9 healthy control subjects were measured. The average spot size of the non-HBV-specific T cells (SEB stimulated) was similar across all groups and was not influenced by HBV DNA load (Fig. 3a). Identical results were obtained with in vitro-expanded cells (Fig. 3b), confirming that non-HBV-specific T-cell function was not affected by HBV status. However, the average spot size of HBV-specific T cells was reduced nearly 50% in cells from chronic patients compared to individuals with resolved infection (P = 0.016) (Fig. 3b), demonstrating that the defect in IFN-γ production was selectively detectable in HBV-specific T-cell populations of chronic patients.

These results were validated by measuring the fluorescence intensity of IFN-γ, HBV-specific T cells. Representative results from one acute and one chronic patient are shown in Fig. 3c. The mean fluorescence intensity (MFI) of IFN-γ HBV-
specific cells detected in the acute and chronic patients differed significantly. IFN-γ+ HBV-specific cells from acute patients had a high MFI (MFI = 2,104), while IFN-γ+ HBV-specific T cells from chronic patients had a low MFI and were often difficult to distinguish from unstimulated cells. Taken together, these results show that Chinese patients with chronic hepatitis B, like Caucasians (7), seem to harbor a functional T-cell defect in IFN-γ production restricted to virus-specific cells. Ethnicity and HBV genotype influence the HBV-specific CD8+ T-cell repertoire of HBV-infected patients. The above results show that neither race nor HBV genotype significantly influences the general quantitative and qualitative profile of the HBV-specific T-cell response; however, these variables could still impact the diversity of the HBV-specific CD8+ T-cell repertoire. Therefore, we determined if ethnicity and HBV genotype can influence CD8+ T-cell responses against six HLA-A2-restricted epitopes that have previously been shown to be promiscuously presented by multiple HLA-A2 subtypes (Table 1) (6). High-resolution HLA-A2 typing was performed on the subjects to determine their HLA-A2 subtypes. All HLA-A2+ Caucasians displayed HLA-A0201, while HLA-A2+ Chinese patients displayed different HLA-A2 subtypes (A0201, n = 3; A0203, n = 5; A0206, n = 4; A0207, n = 8). Patient PBMC were stimulated for 10 days with the peptides corresponding to the sequence of the infecting HBV genotype, and the frequency of the HBV-specific CD8+ T-cells was analyzed by ICS for IFN-γ production.

Figure 4a and b show the results of the CD8+ T-cell responses against the Core18-27 and Env183-91 peptides, which differ by one amino acid between HBVgenB/C and HBVgenA/D and frequently stimulate virus-specific CD8+ T cells in HLA-A0201+ Caucasian HBV patients (4, 6, 30, 45). As expected, 13/16 HLA-A0201+ Caucasian patients responded to the Core18-27(V) epitope. In contrast, CD8+ T cells specific to Core18-27(I) were present in only 3/13 Chinese patients (Fig. 4A). Interestingly, none of the A201+ Chinese patients re-

FIG. 1. Ex vivo quantitative profile of HBV-specific T cells in Chinese and Caucasian HBV patients. (a) Overlapping peptide pools were used to stimulate PBMC directly ex vivo in the IFN-γ ELISPOT assay. Results from selected acute and chronic Chinese and Caucasian patients, segregated based on HBV DNA and ALT levels (shown above the graphs), are displayed. Each bar represents the response to an individual peptide mixture. (b) Mean ex vivo frequency of IFN-γ-producing cells in patients of different ethnicities. Each bar represents the response to the mixtures covering the indicated HBV protein, with error bars indicating the standard error.
sponded to HBV<sub>genB/C</sub> Core18-27(I), and specific responses were only detectable in Chinese patients expressing HLA-A*0207 (n/H11005 2) and HLA-A*0206 (n/H11005 1).

CD8<sup>+</sup> T-cell responses to the Env183-91 epitope were highly influenced by HLA-A2 micropolymorphisms: 14/16 HLA-A*0201<sup>+</sup> Caucasian HBV patients responded to the Env183-91(R) epitope from HBV<sub>genA/D</sub>, and 2/3 HLA-A*0201<sup>+</sup> Chinese patients (2/2 chronic patients) were capable of responding to the Env183-91(K) epitope from HBV<sub>genB/C</sub>. In contrast, all 11 Chinese patients expressing the A*0203, A*0206, or A*0207 HLA-A2 subtype were unable to respond to the Env183-91(K) peptide (Fig. 4b).

Analysis of CD8<sup>+</sup> T-cell responses against epitopes conserved between HBV<sub>genB/C</sub> and HBV<sub>genA/D</sub> (out of 24 full HBV genome entries in GenBank [see Materials and Methods and Table 1]) confirmed the impact of HLA-A2 subtypes on the HBV-specific T-cell repertoire. While patients expressing HLA-A*0201 were capable of responding to the Pol455-63, Env335-43, and Env348-57 epitopes (Fig. 4c), only sporadic promiscuous responses could be detected in the other HLA-A2 subtypes: one HLA-A*0203<sup>+</sup> patient (Env348-57), Responses to the Pol455-63 epitope, which was targeted by 6/8 HLA-A*0201<sup>+</sup> subjects, were not seen at all in HLA-A*0201-negative individuals (Fig. 4c). Conversely, responses to Env338-47, a peptide with only predicted HLA-A2-binding ability (30), were present in HLA-A*0206<sup>+</sup> and HLA-A*0207<sup>+</sup> acute subjects but undetectable in all HLA-A*0201<sup>+</sup> subjects tested.

**Hierarchy of HBV-specific CD8 T cells in HLA-A*0206<sup>+</sup> and HLA-A*0203<sup>+</sup> patients.** We then investigated whether the HBV-specific CD8<sup>+</sup> T-cell repertoire of Chinese patients expressing different HLA-A2 subtypes focused on specificities which differ from those previously defined in A*0201<sup>+</sup> Caucasian patients. To avoid the bias associated with focusing on previously identified epitopes, HBV-specific T-cell responses were assessed using 15-mer overlapping peptides covering the entire HBV proteome, followed by characterization of fine specificity and HLA restrictions of detectable responses. Given the extensive cell requirements for such comprehensive analyses, we were able to perform such thorough analysis in one HLA-A*0206<sup>+</sup> and one HLA-A*0203<sup>+</sup> patient with acute HBV infection. In the HLA-A*0206<sup>+</sup> acute HBV patient, three responses were found directly ex vivo, with a dominant response targeting a
region of the core protein (Fig. 5a). Detailed analysis of the phenotype, fine specificity, and HLA restriction of the responsive T cells revealed that this response recognized a novel HLA-A0206-restricted Core8-16 (EFGASVELL) epitope (data not shown). The two additional responses present at lower frequencies were directed against a second region in the core and one in the envelope which did not overlap with the previously known A2 epitopes (Fig. 5a).

FIG. 3. Quantification of IFN-γ production in acute and chronic Chinese patients. The mean spot size of the positive wells in the ELISPOT assay was used as an estimate of the quantity of IFN-γ produced by single cells after stimulation. (a) Direct ex vivo quantification of unicellular IFN-γ production in healthy, acute, and chronic Chinese patients after SEB stimulation. Each bar represents the mean spot size with error bars indicating the standard error of the mean. Chr, chronic. (b) Quantification of unicellular IFN-γ production in acute and chronic Chinese patients after 10 days of in vitro expansion. Only significant differences are shown (HBV-specific acute patients versus chronic patients [mean ± standard error of the mean], 10.9 ± 0.7 versus 6.9 ± 1.0 × 10⁻² mm²). (c) IFN-γ production by HBV-specific CD8 T cells determined by ICS. Representative acute and chronic HBV-infected Chinese patients are shown. The percentages of IFN-γ⁺ cells out of the CD8⁺ cells after gating on the CD3⁺ fraction of PBMC and the MFI of the double-positive population are indicated. Cells incubated without the HBV polymerase peptide pool were used as the unstimulated negative control in which the IFN-γ⁺ gate was set.

<table>
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<tr>
<th>Peptide</th>
<th>Amino acid sequence</th>
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*PBMC from HLA-A2⁺ Caucasian and Chinese acute or chronic patients (segregated by HBV DNA levels) were expanded in vitro for 10 days and tested against known HLA-A2-restricted HBV epitopes. The sequences of the Core18–27 and Env183–91 peptides differ by one amino acid (as indicated in bold) between the B and D genotypes.

b HLA-A2 supertype binding capacity.
Responses against “promiscuous” HLA-A2 epitopes were also analyzed but were not detected directly ex vivo (data not shown). However, after in vitro stimulation using the corresponding optimal HLA-A2 peptides, the A0206/H11001 acute patient demonstrated the presence of Core18-27(I) (HBV genB/C) and Env335-43 CD8+/H11001 responses, which were subdominant in terms of frequency compared to the Core8-16-specific CD8+/H11001 response (Fig. 5b).

The A2-restricted CD8+ T-cell repertoire in the HLA-A0203/H11001 patient differed significantly from that of the HLA-A0206/H11001 patient. Analysis of the HBV-specific CD8+ T-cell responses was performed only after in vitro expansion, and it revealed the presence of a HLA-A0203-restricted response to two HBV epitopes rarely recognized in Caucasian subjects (Env335-43 and Pol504-12; Fig. 5b). Responses against all other known HLA-A2-restricted HBV epitopes were undetectable, both directly ex vivo and after in vitro expansion (data not shown and Fig. 5b). These results confirm that the HLA-A2 polymorphisms in the Chinese subjects lead to a different focus of the HLA-A2-restricted responses compared to the response patterns seen in the HLA-A0201+ Caucasian population.

**Effect of HBV genotype and HLA-A2 subtypes on CTL recognition.** To further dissect how the combination of amino acid differences in HBV genotypes and HLA-A2 subtypes influence CD8+ T-cell response patterns, we assessed the recognition of genotype-specific epitope variants in the context of several HLA-A2 subtypes. Core18-27-specific CD8+ T-cell clones generated from an A201+ HBV-infected patient can efficiently recognize the Core18-27(I) peptide variant when presented by different A2 subtypes (Fig. 6). Peptide recognition was effective in the context of HLA-A0206-, HLA-A0207-, and HLA-A0201-positive EBV-transformed B-cell lines, which trigger CD8+ T-cell activation when pulsed with low (10 pM) concentrations of Core18-27(I) peptide. In contrast, only minimal CD8+ T-cell activation was observed after stimulation with peptide-loaded HLA-A0203+ target cells, even at the highest peptide concentration (100 nM) (Fig. 6).

**FIG. 4.** Induction of CD8+ T-cell response against known A2-restricted epitopes in HLA-A2+ Chinese and Caucasian patients. Bars represent the frequency of Core18-27 (a)- or Env183-91 (b)-specific CD8+ T cells in individual patients with the indicated HLA-A2 subtypes. PBMC of Chinese patients were expanded with HBVgenB/C Core18-27(I) or Env183-91(K) peptide, while PBMC of Caucasian patients were expanded with HBVgenA/D Core18-27(V) peptide or Env183-191(R) (all at 1 μM) for 10 days, before restimulating the lines with the corresponding stimulatory peptide and analyzing the frequency of the CD8+ cells producing IFN-γ with ICS. Sequencing of the Core18-27 region of the HBV infecting the chronic Chinese and Caucasian patients confirmed the presence of the Core18-27(I) and Core18-27(V) sequences in the Chinese and Caucasian patients, respectively. The hatched line demarcates the A0201-expressing subjects. (c) Frequency of CD8+ T-cell response against conserved HBV epitopes (genotype B = genotype D) Pol456-63, Env338-47, Env335-43, and Env348-57 in acute Chinese and Caucasian HBV patients.
We then analyzed, in parallel, the ability of HLA-A2 subtypes to present the two genotype-specific Core18-27 epitope variants as well as the stability of the peptide/HLA-A2 complex, a parameter associated with peptide immunogenicity (36). Antigen-presenting cells with defined HLA-A2 subtypes were pulsed with increasing concentrations of Core18-27(V) or Core18-27(I) peptides and then used to stimulate Core18-27(V)-specific CD8\(^+\) T-cell clones. Effector cells were added immediately or after a 12-h period. Both the Core18-27(V) as well as its 27(I) variant were presented effectively by HLA-A0201, -A0206, and -A0207 but not by HLA-A0203 if effector cells were added immediately. However, when the loaded peptides were allowed to dissociate from the antigen-presenting cells before the addition of the effector T cells (after a 12-h period), the Core18-27(I) variant was in all cases less stimulatory than the Core18-27(V) peptide. In the context of HLA-A0201, the genotype B Core18-27(I) peptide lost essentially all immunogenicity, whereas immunogenicity was retained when the Core18-27(I) peptide was presented by HLA-A0206. Not surprisingly, the little stimulatory efficiency of HLA-A203 targets was completely lost after the 12-h dissociation period (Fig. 7).

It is interesting to note that the peptide presentation data recapitulate the frequency of Core18-27-specific CD8\(^+\) T cells detected in HBV-infected patients. The hierarchy of Core18-27(I) presentation (A0201 > A0207 > A0206 > A0203) described by the presentation experiments in
Fig. 7 is in line with the presence of Core18-27(I)-specific CD8$^+$ T-cell responses found in some HLA-A0206$^+$ and -A0207$^+$ HBV patients and their absence in HLA-A0201$^+$ and -A0203$^+$ patients (Fig. 4a).

Finally, we tested whether HBV$^{\text{genB}}$-infected Asian individuals could recognize epitope variants from HBV$^{\text{genD}}$. T-cell lines specific for different HBV$^{\text{genB}}$ epitopes were generated from Chinese individuals with acute HBV$^{\text{genB}}$ infection and tested for recognition of epitope variants naturally present in genotypes D, A, and C. The single amino acid substitution within the Core8-16 peptide (S12T) did not affect Core8-16-specific CD8$^+$ T-cell recognition (Fig. 8). Similarly, despite the presence of three different amino acids, the peptide corresponding to the sequence Env370-79 of HBV$^{\text{genA,D}}$ was recognized, even though to a lesser degree, by the Env370-79 HBV$^{\text{genB}}$-specific HLA-A0203-restricted CD8$^+$ T-cell line. In contrast, the single amino acid difference present in the HBV$^{\text{genC}}$ (S373N) variant completely abolished CD8$^+$ T-cell recognition (Fig. 8). Whether such amino acid substitution (S373N) abolished the HLA-A203-binding ability of the peptide or instead altered the T-cell receptor recognition site will need to be properly analyzed.

DISCUSSION

In this study, we took advantage of the segregation of different HBV genotypes in geographically and genetically distinct host populations to directly analyze the impact that host and virus variables exert on the virus-specific T-cell response. Two important points emerged: (i) HBV-specific T-cell quantity is determined by the virological and clinical profiles of the patients, which outweight any influence of race or viral diversity, and (ii) HBV-specific T-cell repertoires are divergent in the two ethnic groups with T-cell epitopes frequently found in Caucasian patients and seldom detected in Chinese patients.

In Asia, chronic HBV infection is mainly derived from vertical infection at birth (39), whereas horizontal transmission (household/sexual contact) is the likely route of infection in Caucasians. In addition, infection by different HBV genotypes has also been suggested to influence clinical outcomes or disease severity (15). Furthermore, by inference from HCV infection, host ethnicity may impact clinical parameters and the antiviral CD4$^+$ T-cell response (41). To our knowledge, whether these epidemiological variables might alter the profile of the HBV-specific T-cell response has never been directly tested. Our results show that the epidemiological variables characteristic of the different ethnic populations do not impact the overall virus-specific T-cell quantity. Rather, the level of cellular immunity appears largely defined by the clinical status
of the tested patient. HBV-specific T cells were easily detected directly ex vivo in only acute patients, while HBV-specific T-cell responses were very weak in chronic patients. Although different HBV genotypes or transmission routes could trigger different pathogenic mechanisms resulting in viral persistence, our data indicate that the ability to control HBV, or conversely HBV persistence, results in highly similar quantitative profiles of HBV-specific T-cell responses, irrespective of ethnicity and HBV genotype.

We also confirmed, by analyzing IFN-γ production at a single-cell level, that HBV-specific T cells of Chinese chronic patients infected by HBV genB were defective for IFN-γ production. The IFN-γ defect was found exclusively in HBV-specific T cells while the general T-cell population was unaffected. While this finding is in contrast with recent data (10), which have implicated the apparent upregulation of PD-L1 on myeloid dendritic cells in the suppression of global T-cell function, the data confirm results obtained from anti-HBe + Caucasian patients, for which the impairment of IFN-γ production, restored by blocking PD-1/PD-L1, was restricted to HBV-specific T-cell responses, irrespective of ethnicity and HBV genotype.

HBV genotypes and the individuals’ genetic backgrounds instead play an important role in shaping the HBV-specific T-cell repertoire. We found that HLA-A2-restricted CD8 + T-cell epitopes, which frequently induced responses in HLA-A0201 Caucasian patients and possess “HLA class I supertype” binding capacity (6), are scarcely (Core18-27) or completely unable (Env183-91 and Pol455-65) to induce CD8 + T-cell responses in HLA-A0206 + , A0207 + , or A0203 + Chinese patients. Furthermore, a comprehensive analysis of the CD8 + T-cell repertoire performed in one HLA-A0206 + patient and one HLA-A0203 + HBV patient directly confirmed, for naturally infected subjects, how HLA micropolymorphisms (HLA-A0206/A0207 and -A0203 differ from HLA-A0201 by a single amino acid or three amino acids, respectively) and viral amino acid differences impact CD8 + T-cell responses. These two subjects preferentially focused T-cell responses on HLA-A2-restricted epitopes never (Core8-16 and Pol406-12) or scarcely (Env371–79) (33) detected in HLA-A0201 subjects, while responses to classical HLA-A2-restricted epitopes were not at all (HLA-A0203 + patient) or rarely (HLA-A0206 + patient) present.

The discordance of the HBV-specific CD8 + T-cell repertoires in the two ethnicities is not surprising if one considers the different mechanisms that might alter the immunogenicity of antigenic peptides. For example, the ability of a peptide epitope to induce CD8 + T cells can be influenced by its bound conformation without necessarily altering the HLA-binding ability. Work on EBV infection has shown that the conformational changes imposed by a single amino acid variation present in two closely related HLA class I molecules (HLA-B3501 and HLA-B3508) are sufficient to alter the ability of an identical peptide to induce a CTL response (44). This scenario might explain our inability to detect the HBV Pol455-63 epitope in HLA-A0203 + , A0206 + , and A0207 + patients, despite its conservation between different genotypes and its reported “promiscuous” ability to bind different HLA-A2 subtypes (6). Furthermore, the ability of HLA-A2 subtypes to preferentially present different sets of peptides (2, 40) is another important contributor to the distinct epitopes targeted in HBV patients expressing different HLA-A2 subtypes. In addition to the alterations imposed by the HLA-A2 polymorphisms, the presence of distinct HBV genotypes in different ethnicities can further increase the potential diversity in the pool of immunogenic peptides. HBV genotypes differ approximately 8% in their amino acid compositions (21), and so it is likely that HBV genotype amino acid variations located within or just outside HBV epitopes might modify epitope processing, presentation, or recognition (23, 34).

Our experiments prove that these amino acid variations can affect T-cell recognition of CD8-specific T-cell epitopes. In particular, we showed for the Core18-27 epitope that the presence of isoleucine instead of valine at position 27 reduced the stability of the peptide/HLA-A2 complex to various degrees in all the HLA-A2 subtypes tested. The Core18-27 isoleucine variation is characteristic of HBV genB/C and our findings help explain the generally poor immunogenicity of the Core18-27(I) epitope in HLA-A0201 + Chinese subjects. At the same time, the observation that the HBV genB/C Core18-27(I) peptide is more stably presented by HLA-A0206 and A0207 (only expressed by Chinese patients) than HLA-A0201 is puzzling. While it could be driven by a possible host-virus coevolution, its evolutionary significance is obscure, such coevolution would have led to the conservation of a highly avid epitope presented by some of the dominant HLA alleles. Whether this finding can be extrapolated to other epitopes and whether or not these responses are significantly involved in viral control clearly require broader analyses in larger cohorts and more diverse HBV backgrounds.

In conclusion, ethnic and viral differences do not alter the vigor of HBV-specific T-cell responses present in patients with HBV infection, but these variables shape distinct HBV-specific T-cell repertoires. The results from our findings show a limited contribution of HLA-A2 “promiscuous” HBV epitopes to the HBV-specific CD8 T-cell repertoire of ethnically different patients and question, at least in the context of the HLA-A2 allele, the ability of the supertype motifs to predict virus-specific CD8 + T-cell responses present in different ethnic groups. From our data, it seems clear that HBV-specific immune monitoring in Asian patients should not rely on the exclusive analysis of epitopes found in Caucasians or conform to algorithms that do not consider the clustering of different viral strains within different ethnic groups. Furthermore, the use of polyepitope vaccines (12), based on peptides selected by HLA class supermotifs or on responses found in Caucasian patients, might divert the response toward incorrect specificities, stimulating nonnatural responses which might be functionally incapable of recognizing the infectious virus, a situation that has already been reported in HIV infections (8, 24). However, it must also be considered that predicted epitopes that might not be naturally immunogenic can be advantageous in breaking immunological tolerance caused by a persistent virus infection, potentially expanding nontolerant, cross-reactive CD8 T cells able to recognize the infecting virus with therapeutic advantage (33). T-cell repertoires elicited by vaccines and their therapeutic effects in patients will have to be carefully analyzed in patients to resolve this conundrum and shed further light on the complex and fascinating diversity of the antiviral T-cell response in humans.
Degenerate immunogenicity of an HLA-A2-restricted hepatitis B virus nucleocapsid cytotoxic T-lymphocyte epitope that is also presented by HLA-B51. J. Virol. 75:3984–3987.


