Host-specific adaptation of HIV-1 subtype B in the Japanese population


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Running title: HLA-associated polymorphisms in Japanese

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Word count: Text 7,021 words, Abstract 249 words
The extent to which HIV-1 clade B strains exhibit population-specific adaptations to host HLA alleles remains incompletely known, in part due to incomplete characterization of HLA-associated HIV-1 polymorphisms (HLA-APs) in different global populations. Moreover, it remains unknown to what extent the same HLA alleles may drive significantly different escape pathways across populations. As the Japanese population exhibits distinctive HLA class I allele distributions, comparative analysis of HLA-APs between HIV-1 clade B-infected Japanese and non-Asian cohorts could shed light on these questions. However HLA-APs remain incompletely mapped in Japan. In a cohort of 430 treatment-naive Japanese with chronic HIV-1 clade B infection, we identified 284 HLA-APs in Gag, Pol and Nef using phylogenetically-corrected methods. The number of HLA-associated substitutions in Pol, notably those restricted by HLA-B*52:01, was weakly inversely correlated with plasma viral load (pVL), suggesting that the transmission and persistence of B*52:01-driven Pol mutations could modulate pVL. Differential selection of HLA-APs between HLA subtype members, including those differing only with respect to substitutions outside the peptide-binding groove, was observed, meriting further investigation as to their mechanisms of selection. Notably, two-thirds of HLA-AP identified in Japan had not been reported in previous studies of predominantly Caucasian cohorts, and were attributable to HLA alleles unique to, or enriched in, Japan. We also identified 71 cases where the same HLA allele drove significantly different escape pathways between Japan versus predominantly Caucasian cohorts. Our results underscore the distinct global evolution of HIV-1 clade B as a result of host population-specific cellular immune pressures.
Importance section

CTL escape mutations in HIV-1 are broadly predictable based on the HLA class I alleles expressed by the host. Because HLA allele distributions differ among worldwide populations, the pattern and diversity of HLA-associated escape mutations are likely to be somewhat distinct to each race and region. HLA-associated polymorphisms (HLA-APs) in HIV-1 have previously been identified at the population level in European, North American, Australian and African cohorts, however, large-scale analyses of HIV-1 clade B-specific HLA-APs in Asians are lacking. Differential intra-clade HIV-1 adaptation to global populations can be investigated via comparative analyses of HLA-associated polymorphisms across ethnic groups, but such studies are rare. Here, we identify HLA-APs in a large Japanese HIV-1 clade B cohort using phylogeneticaly-informed methods and observe that the majority of them had not been previously characterized in predominantly Caucasian populations. Results highlight HIV’s unique adaptation to cellular immune pressures imposed by different global populations.
HIV CTL escape occurs in a manner that is highly reproducible in context of the HLA class I alleles expressed by the host (1-8). By extension, HIV sequences circulating in a given host population will exhibit polymorphisms that reflect the HLA allele distribution of that population (9). Because HLA class I allele distributions differ among racial and ethnic groups worldwide (10), the pattern and diversity of HLA-associated escape mutations is also likely to be somewhat distinct to each race and region. Numerous population-based studies identifying HLA-associated polymorphisms (HLA-APs) have been conducted in European, North American, Australian, and African cohorts (2, 6, 8). However, comparably fewer have been undertaken in Asian cohorts, where HIV-1 prevalence is also substantial (11). Since Asian populations differ in their HLA allele distributions from the cohorts previously studied, it is important to identify and analyze HLA-APs to achieve a better understanding of HIV-1 pathogenesis in Asia and to inform future HIV vaccine design efforts targeted to these populations. The Japanese epidemic is somewhat unique in Asia. While clades A/E and C predominate in many Asian countries (12-14), the Japanese HIV-1 epidemic comprises 80% clade B infections (12). As such, the analysis of Japanese cohorts also provides the opportunity to undertake comparative analyses of HLA-APs between Asian and non-Asian populations infected with HIV clade B.

Previous studies have investigated differential HLA-driven HIV evolution across human populations. For example, a study of HLA-specific adaptations in HIV Pol in a Mexican cohort identified “unique” HLA-APs in this population that were not present in an international cohort from Canada, the USA and Australia, even though both cohorts harbored HIV clade B (15). Most of the unique Mexican HLA-APs were restricted by HLA alleles particular to this population (e.g. HLA-B*39) but that were underrepresented or absent in the international cohort (15). This study therefore illustrates population-specific HIV adaptation in its most intuitive manifestation: where distinctive HLA-associated
polymorphisms are observed in a population due to the presence (or comparatively higher
frequency) of an HLA allele in this population compared to another.

What remains unknown however, is the extent to which the same HLA allele may drive
divergent escape pathways in different human populations. Two critical features are
required to address this question. First, the identification of HLA-AP must be undertaken at
the HLA subtype level. This is because the majority (>60%) of HLA-associated
polymorphisms are best defined at the subtype-level (16) – even for closely related HLA
subtype members that present the same or similar peptide epitopes (16, 17, 18, 19).

Comparative studies undertaken at allele-level (two-digit) resolution cannot disentangle
whether population-specific HLA-AP are attributable to differential HLA subtype
distributions between cohorts, or whether they are “true” cases where the same HLA
subtype drives different escape pathways across populations. Indeed, a study investigating
>500 Americans with chronic HIV-1 clade B infection observed distinct patterns of
HLA-APs among White, Black, and Hispanic individuals that were likely attributable to the
differential distribution of closely-related HLA subtypes among these groups (18) rather
than true differential escape. The present study is therefore undertaken at subtype-level
resolution. Secondly, the identification of population-specific escape pathways driven by
the same HLA allele requires a method to do so. Here, we adapt phylogenetically-corrected
statistical methods originally developed to assess differential escape among related HLA
subtypes (17) and apply them to investigate differential escape across host populations.

The present study is divided into two parts, each with a specific major objective. Our
first objective was to identify and characterize HLA-AP in HIV-1 Gag, Pol, and Nef
proteins in a cohort of 430 chronically clade B-infected Japanese individuals using
phylogenetically-informed approaches (20), and to investigate their associations with
clinical parameters (CD4+ T cell count and plasma viral load). Importantly, HLA
genotyping (and thus HLA-AP identification) was undertaken at subtype-level resolution,
allowing us to analyze the effect of genetic differences among closely-related HLA
subtypes on the selection of HLA-APs in the Japanese cohort as part of this objective. Our second major objective was to perform a comparative analysis of HLA-AP identified in Japan to those identified in a large international (Canada/USA/Australia) cohort of antiretroviral-naïve, chronically clade B infected, predominantly Caucasian individuals. As expected, a substantial proportion of Japanese HLA-AP were restricted by alleles unique to (or highly enriched in) Japan compared to the non-Asian cohort. Notably we also observed numerous cases where the same HLA allele drove significantly different – sometimes opposing – escape pathways in these two populations. Our results highlight HIV’s unique adaptation to cellular immune pressures imposed by different global populations.
Materials and Methods

Ethics statement

This study was approved as a part of “the study of immunological and virological analysis in HIV-1 infection (#540)” by the ethics committee for epidemiology and general study in the faculty of life science in Kumamoto University and the National Center for Global Health and Medicine. All studied individuals were adults. Written informed consent was obtained from all studied individuals according to the Declaration of Helsinki.

Subjects

Four-hundred-thirty treatment-naïve Japanese individuals with chronic HIV-1 clade B infection were enrolled in the National Center for Global Health and Medicine (NCGHM) from 2008 to 2011. HLA alleles of these individuals were determined at the 4-digit level by a probe-based sequence-specific oligonucleotide (SSO) typing method (HLA Laboratory, Kyoto, Japan). The median CD4+ T cell count (CD4 count) and plasma viral loads (pVL) at the first visit to NCGHM were 321 cells/μl (IQR: 190 to 440 cells/μl) and 25,000 copies/ml (IQR: 6,800 to 98,000 copies/ml), respectively.

HLA-associated polymorphisms derived from the International HIV Adaptation Collaborative (IHAC) cohort, comprising 1,888 treatment-naïve individuals with chronic clade B infection from Canada, USA, and Western Australia (16), identified using identical methods, were used for comparison. The IHAC cohort comprises predominantly Caucasian individuals, with Asians making up less 5% of the total. The median CD4 count in IHAC was 260 cells/μl (IQR: 110 to 418 cells/μl).

RT-PCR and sequencing of plasma HIV RNA

HIV-1 viral RNA was extracted from plasma samples using either a QIAamp MinElute virus spin kit (Qiagen, Valencia, CA) or an EZ1 Virus Mini Kit v2.0 (Qiagen, Valencia, CA).
Reverse transcription was performed using random hexamers with the SuperScript III First-Strand Synthesis System for RT-PCR (Invitrogen, Carlsbad, CA). HIV-1 Gag, Pol, and Nef genes were amplified from cDNA by nested PCR using Taq DNA polymerase (Promega, Fitchburg, WI) and 10 primer pairs that were designed based on the clade B strain. For subjects with a viral load below 1000 copies/ml, RT-PCR was performed with region-specific primers using the SuperScript III One-Step RT-PCR System with Platinum Taq kit (Invitrogen, Carlsbad, CA). The 1st round PCR product was then used in the 2nd round PCR amplification using Taq DNA polymerase (Promega, Fitchburg, WI) and the 10 primer pairs. The 2nd round PCR product was purified by using the ExoSAP-IT reagent containing Exonuclease I and alkaline phosphatase (GE Healthcare, Buckinghamshire). Gag, Pol, and Nef sequences were determined by using the BigDye Terminator v3.1 cycle sequencing kit (Applied Biosystems, Carlsbad, CA) and an ABI 3500 genetic analyzer (Applied Biosystems, Carlsbad, CA). Sequencing reactions were performed in both 5’ and 3’ directions to yield a minimum of bidirectional coverage of all regions. Sequence data was then aligned by using SeqScape software (Applied Biosystems, Carlsbad, CA) based on the HXB2 reference sequence (K03455). Accession numbers are AB873205 to AB873601 (Gag), AB873908 to AB874270 (Pol), and AB873602 to AB873907 (Nef).

Identification of HLA-associated polymorphisms

HLA class I associated polymorphisms in HIV-1 (HLA-AP) can be identified in large, cross-sectional linked datasets of host (HLA) and HIV genotypes using statistical association approaches that identify viral polymorphisms significantly over- or under-represented in individuals harboring a specific HLA class I allele (1, 2, 4, 16-18, 21). HLA-APs that are over-represented in individuals harboring the relevant HLA are commonly referred to as “adapted” forms, while those under-represented in individuals harboring the relevant HLA are referred to as “nonadapted” forms (2, 18). As such, “nonadapted” and “adapted” forms can be conceptualized to represent the
“immunologically susceptible” and “escape mutant” forms, respectively, for the specific
HLA allele in question at that HIV codon position. Statistical association approaches for the
identification of HLA-AP also correct for the confounding influences of viral phylogeny,
HIV codon covariation and linkage disequilibrium between HLA class I alleles (2, 16, 17,
21).
Associations between HLA class I alleles and HIV-1 amino acid polymorphisms in the
Japanese and IHAC datasets were identified using a published phylogenetically-corrected
logistic regression model that corrects for HLA linkage disequilibrium, HIV phylogeny, and
HIV codon covariation as potential confounders (17, 20). Briefly, maximum likelihood
phylogenetic trees were constructed using Gag, Pol and Nef sequences (one tree per gene),
and a model of conditional adaptation was inferred for each observed amino acid at each
codon. Amino acids are assumed to evolve independently along the phylogeny, until the
tree tips (representing the present host). In each host, HLA-mediated selection and HIV
amino acid covariation are directly modeled using weighted logistic regression, in which
the individual’s HLA repertoire and covarying HIV amino acids are used as binary
predictors and the bias is determined by the possible transmitted sequences as inferred by
the phylogeny (17). To identify which factors (HLA and/or HIV covariation) contribute to
selection pressure, we employ a forward selection procedure where the most significant
association is iteratively added to the model, with p-values computed using the likelihood
ratio test. We performed post-hoc filtering of the resulting HLA-associated polymorphisms
list, restricting our output to instances in which at least 10 individuals carried the allele or
polymorphism and at least 10 individuals did not carry the allele or polymorphism.
Multiple tests were accounted for using q-values, the p-value analog of the false discovery
rate (FDR) (22). The FDR is the expected proportion of false positives among results
deemed significant at a given threshold; for example, at $q < 0.2$, we expect 20% of
identified associations to be false positives. In the analyses identifying HLA-associated
polymorphisms (HLA-AP), significance threshold of $q < 0.2$ was employed.
209  **Statistical analysis**

Correlations between the total number of HLA-associated substitutions in each individual and clinical parameters (pVL and CD4 count) were performed using Spearman’s correlation. To count the total number of HLA-associated substitutions within a given HIV-1 sequence, we first identified all HIV-1 sites within that sequence identified as being associated with any HLA allele. The specific residue at each site was counted as “HLA-associated” if it matched any HLA-associated adapted form, or any residue other than a nonadapted form identified at that position. The HLA alleles expressed by the individual were not considered (unless specifically stated) – rather, our goal was to enumerate the number of HLA-AP associated with any HLA allele in each viral sequence. In analyses where host HLA alleles were not considered, HIV sites harboring residues that simultaneously represented a nonadapted and Adapted form associated with different HLA alleles were excluded from consideration.

223 **Detection of differential escape between closely-related HLA alleles, and between cohorts.**

Two types of differential escape were investigated. First, we investigated differential escape between closely-related HLA class I alleles, defined here as (four-digit) HLA subtype members belonging to the same (two-digit) allele group, in the Japanese cohort. Specifically, seven HLA allele groups (A*02, A*26, B*15, B*40, C*03, C*08, and C*14) for which a minimum of two subtype members were represented in the Japanese cohort, were investigated. For example, the HLA-A*02 allele group featured subtypes A*02:01, A*02:06 and A*02:07, while the A*26 allele group featured subtypes A*26:01 and A*26:03. For each allele group, we took the union of all HLA-AP identified for all subtype members of the group. Then, in a pairwise manner, we compared their strengths of selection between all HLA subtype members using a previously-described
phylogenetically-corrected interaction test (17). In this analysis, thresholds of $p < 0.05$, $q < 0.2$ were used to define significance.

Second, we investigated differential HLA-driven escape pathways between Japanese and IHAC cohorts. As outlined in the introduction, HLA-AP identified in human populations will differ to some extent due to the presence (or enrichment) of certain HLA alleles in one population versus another. However in this analysis we were specifically interested in identifying cases where the same HLA allele drove significantly different escape pathways in the two cohorts. To do this, we took the union of all HLA-AP identified in Japan and IHAC cohorts, that were restricted by HLA subtypes observed a minimum of 10 times in both cohorts. We then compared the strength of selection of each HLA-AP in a pairwise manner, between cohorts. The statistical methods used investigate differential escape between Japanese vs. IHAC cohorts are similar to those used to investigate differential escape between HLA subtype members (17), with some modifications as follows. Briefly, a phylogenetically-corrected logistic regression model is constructed using a single HLA allele as a predictor. Using a likelihood ratio test, we then compare this model to a more expressive one that includes an additional interaction term that is 1 if the individual expresses the HLA allele and is in the IHAC cohort, or 0 otherwise. In this way, we can obtain a $p$-value testing the hypothesis that selection is the same in both cohorts (null hypothesis) or whether selection differs across cohorts (alternative hypothesis). In contrast to the HLA-AP analyses described thus far, the present one does not feature corrections for HLA linkage disequilibrium or HIV codon covariation – and therefore will yield odds ratios of association and $p$-values that differ slightly from the original cohort-specific values. In the inter-cohort differential escape analysis, significance was defined as $p < 0.01$, $q < 0.05$. 

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Results

Identification of HLA-associated polymorphisms in chronically HIV-1 clade B-infected Japanese individuals.

The first objective of our study was to identify and characterize HLA-AP in Japan, a unique population in terms of its HLA class I distribution and predominantly HIV clade B epidemic. Towards this end, we analyzed linked HIV/HLA genotypes from 430 antiretroviral therapy-naive Japanese individuals chronically infected with HIV-1 clade B. A total of 78 unique HLA class I alleles, defined at subtype-level (4-digit) resolution, were observed in our cohort (Fig. S1) at frequencies consistent with the published literature (23). Of these, 37 (including 9 HLA-A, 17 HLA-B, and 11 HLA-C alleles) were observed in at least 10 individuals, and thus were included in the statistical analysis of HLA-APs (see methods). Amplification and sequencing of HIV-1 Gag, Pol without the transframe (TF) protein, and Nef was successful for 397 (92.3%), 363 (84.4%), and 306 (71.2%) individuals, respectively. As described in the methods, HLA-APs within these three genes were identified using a phylogenetically corrected logistic regression model which corrects for the confounding effects of viral phylogeny, HIV-1 codon covariation and linkage disequilibrium between host HLA class I alleles (16, 17, 20). A false discovery rate (q-value) approach was employed to address multiple tests.

At a threshold of $q < 0.2$, a total of 284 HLA-APs, comprising 143 adapted and 141 non-adapted associations, were identified in Gag (N=94), Pol (N=86), and Nef (N=104 associations) (Fig. 1 and Table S1). HLA-APs were more frequently detected in Nef (occurring at 45 of 206 codons; 21.8%) compared to Gag (51 of 500 of codons; 10.2%) or Pol (51 of 947 codons; 5.1%). Although HLA class I allele frequencies in Japan are somewhat distinct globally, the distribution of HLA-AP across HIV-1 proteins was consistent with that reported in previous studies of other populations infected with clades B or C (1, 2, 6, 7, 16). Broken down by HLA locus, the numbers of HLA-A-, HLA-B-, and
Correlation between the total number of HLA-associated substitutions and clinical parameters in Japanese individuals.

We next wished to investigate the relationship between the presence of HLA-associated substitutions in each gene and patient HIV-1 plasma viral load (pVL) and CD4+ T cell count (CD4 count) in the Japanese cohort. As described in the methods, substitutions within a given HIV-1 sequence were counted as “HLA-associated” if they had been identified as being associated with any HLA class I allele in our study, regardless of the HLA alleles expressed by the patient. For example, Gag-9S is a HLA-B*15:01-associated nonadapted polymorphism (Fig. 1 and Table S1); as such, any amino acid other than “S” at codon 9 was counted as an HLA-associated substitution. Similarly, Gag-123G is an HLA-C*01:02-associated adapted polymorphism (but no specific nonadapted forms, restricted by C*01:02 or others, were identified at this position); as such, any sequence harboring “G” at codon 123 was counted as having an HLA-associated substitution at this site.

A weak yet statistically significant inverse correlation was observed between pVL and the total number of HLA-associated substitutions in Pol (Spearman’s R = −0.11; p = 0.04) (Fig. 2A). However, no such correlations were observed for Gag (Spearman’s R = −0.056, p = 0.3) or Nef (Spearman’s R = −0.029, p = 0.6) (Fig. 2A). Moreover, no significant correlations were observed between the total number of HLA-associated substitutions in any HIV protein, and CD4 count (Fig. 2A). Though the overall association is weak, results raise the intriguing hypothesis that selection of certain HLA-driven substitutions in Pol could modulate VL in the Japanese population.

We next wondered whether the observed correlation between Pol polymorphisms and
lower pVL could be attributed to polymorphisms restricted by HLA alleles that are protective in Japanese populations. HLA-B*67:01 and the HLA-B*52:01-HLA-C*12:02 haplotype are examples of such protective alleles (24). As such, we investigated whether they could play a role in the observed pVL correlation. No HLA-B*67:01-associated substitution was identified in Pol, whereas four HLA-B*52:01-associated and one HLA-C*12:02-associated substitutions were detected in this protein (Table S1). Exclusion of the single HLA-C*12:02-associated substitution from analysis did not affect the relationship between the number of HLA-associated substitutions in Pol and pVL (data not shown). In contrast, exclusion of the four HLA-B*52:01-associated Pol substitutions substantially weakened the overall relationship between the number of HLA-associated Pol substitutions and pVL (Spearman’s R = −0.057; p = 0.3) (Fig. 2B). Similarly, specific consideration of only HLA-B*52:01-associated Pol substitutions revealed a highly significant inverse correlation with pVL (Spearman’s R = −0.18; p = 0.0007) (Fig. 2C) that represented the strongest such relationship detected in Pol for common HLA alleles observed in our cohort (Fig. S2). We therefore reasoned that B*52:01-restricted substitutions were likely to be critical mediators of the observed pVL effect.

Finally, stratification of B*52:01-associated Pol substitutions by host B*52:01 expression revealed that the inverse correlation with pVL remained strongly detectable in HLA-B*52:01− individuals (Spearman’s R = −0.18; p = 0.003), but not in HLA-B*52:01+ individuals (Spearman’s R = 0.026; p = 0.8) (Fig. 2D). We interpret our observations to suggest that HLA-B*52:01-restricted Pol substitutions possess fitness costs that manifest themselves in terms of lower pVL upon transmission to, and persistence in, HLA-B*52:01− individuals. In contrast, no such pVL effects are detectable in B*52:01+ individuals, likely because the fitness costs of these substitutions are outweighed by the advantages conferred by immune escape.

Differential escape between HLA subtypes in Japanese individuals.
Our final goal in characterizing HLA-AP in Japan was to investigate the extent of differential escape between closely-related HLA subtypes. In particular, we hypothesized that HLA subtype members differing with respect to amino acids located within the peptide-binding groove of the HLA molecule may differ with respect to the nature (or binding affinity) of the specific HIV epitopes presented (25-28), and therefore that they may exhibit differential escape pathways. In contrast, HLA subtype members that differ with respect to amino acids located outside the peptide-binding groove may be more likely to present the same epitopes (29-31), and therefore will generally exhibit less evidence for differential escape between them. Of the 284 HLA-AP identified in our cohort, 128 were restricted by HLA allele groups (A*02, A*26, B*15, B*40, C*03, C*08, and C*14) containing two or more subtype members (Table S1). For five of these allele groups (A*02, A*26, B*15, B*40, and C*08), subtype members differed by substitutions within the peptide-binding groove (Fig. S3), supporting them as potential candidates for differential HLA-AP selection. In contrast, members of the C*03 and C*14 subtypes differed by substitutions outside the peptide-binding groove (Fig. S3), suggesting that their epitope repertoire (and thus escape pathways) would be more similar to one another.

We began by simply comparing HLA-AP identified in context of the different HLA subtypes. As expected, viral polymorphisms associated with HLA subtype members differing within their peptide-binding grooves appeared to be quite specific to each HLA subtype (Fig. S3A, S3B, S3C, S3D, and S3F). Surprisingly however, viral polymorphisms associated with HLA subtype members differing only with respect to amino acids located outside their peptide-binding grooves also appeared to be quite specific to each HLA subtype (Fig. S3E and S3G). For example, HLA-C*03:03 and C*03:04, which differ only by substitutions at position 91 that have no contact with the groove (29-31), were associated with a total of 11 HLA-APs, none of which appeared to be shared (Fig. S2E).

Similarly, HLA-C*14:02 and C*14:03, that differ only by a substitution at position 21 located outside of the floor of the peptide-binding groove (Fig. S2G), shared only 10 of the
24 HLA-APs identified between them. However, qualitative comparisons of HLA-AP meeting a specific significance threshold, such as those described above, are not statistically robust (since individual associations may fail to meet the threshold and thus not be detected, or variations in allele frequency may limit power to detect associations). Thus, to explicitly investigate whether the above examples represent statistically significant instances of differential escape between subtype members we applied a phylogenetically-corrected interaction test to compare their strengths of selection between subtypes (17). For each HLA allele group, we took the union of all HLA-AP identified for all subtype members, and compared their strength of selection between all subtype members in a pairwise manner. Representative examples of our results are shown in Fig. 3. For example, HLA-A*26:01 or A*26:03 differ with respect to substitutions at amino acids 74, 76 and 77, located within the peptide-binding groove of the HLA molecule (Fig. S3B). A total of 10 HLA-APs, located at 8 HIV codons, were originally identified as associated with either HLA-A*26:01 or A*26:03 (Fig. S3B). Although qualitatively, all 10 HLA-APs appear to be differentially selected by HLA-A*26:01 or A*26:03 (Fig. S3B), the phylogenetically-corrected interaction test revealed only 3 of them (located at Pol residues 276 and 551, and Nef residue 85) to be significantly differentially selected in terms of their natural logarithm of the odds ratios of association (p < 0.05, q < 0.2) (Fig. 3A). Surprisingly, significant differential escape was also observed between subtype members that differed only with respect to substitutions outside of their peptide-binding grooves: 3 of 9 (33.3%) sites restricted by HLA-C*03 allele group members and 5 of 14 (35.7%) sites restricted by C*14 allele group members similarly exhibited statistically significant evidence of differential selection (Fig. 3B and 3C).

To compare whether the extent of differential escape between HLA subtype members varied between HLA allele groups that differed with respect to substitutions within or outside the binding groove, we asked whether the extent of differential escape between
subtype members of the former group (comprising A*02, A*26, B*15, B*40, and C*08) differed compared to the latter group (comprising HLA-C*03 and C*14). Overall, we found no significant differences in the proportion of differential escape between them (34.8% for HLA-C*03/C*14 subtypes compared to 36.8% for subtypes of all other HLA alleles, \( p = 0.5 \) (Table S2). This intriguing result suggests that variations outside the HLA binding groove may contribute as much to differential escape as variations within the binding groove.

Comparison of HLA-APs between Japanese and non-Asian individuals chronically infected with HIV-1 clade B.

Our second objective was to investigate HLA-AP identified in Japan versus those previously identified in non-Asian cohorts infected with HIV clade B. The comparison cohort in this analysis was the International HIV Adaptation Collaborative (IHAC) cohort, comprising 1,888 antiretroviral-naïve individuals with chronic clade B infection in Canada, the USA, and Australia (of which <5% of cohort participants are Asian) (16). HLA-AP will differ to some extent between human populations due to the presence (or enrichment) of certain HLA alleles in one population versus another. Indeed, HLA allele frequencies differed markedly between the Japan and IHAC cohorts (Fig. S1). As such, we begin with a qualitative comparison of HLA-AP between them. We begin with a simple positional analysis. In the Japanese cohort, HLA-APs were observed at a total of 147 codon positions in Gag, Pol, and Nef (Fig. 4). Of these, 117 (79.6%) were also associated with at least one HLA allele in IHAC. In contrast, the remaining 30 positions (including 16, 7 and 7 in Gag, Pol, and Nef, respectively) that harbored HLA associations in Japan were not associated with any HLA alleles in IHAC (Fig. 4). That 30/147 (20.4%) of HIV codons exhibited evidence of HLA-driven selection in Japan but not IHAC already strongly suggests that HIV is evolving under population-specific selection pressures in Japan compared to other regions.
Next we compared HLA-AP over HIV position and specific HLA restriction. Of the 284 HLA-APs identified in Japan, 188 (66.2%) were not reported in IHAC. As expected, a substantial portion of these (46 of 188, 24.5%) were associated with 8 HLA subtypes (A*26:03, B*40:06, B*54:01, B*55:02, B*59:01, B*67:01, C*08:03, and C*14:03) common in Japan but essentially absent (<1% frequency) in IHAC. Others were likely attributable to alleles observed at much higher frequencies in Japan compared to IHAC: for example, an additional 27.1% were associated with HLA alleles present in both cohorts, but whose frequencies were at least fourfold higher in Japan compared to IHAC. Overall, results suggest that HLA-APs identified in Japan are quite distinctive, in large part reflecting the unique HLA allele distribution in the Japanese population.

We also wished to investigate the existence of differential HLA-associated escape pathways between the two populations, that are not attributable to HLA frequency differences between them — in other words, cases where the same HLA subtypes drive significantly different escape pathways in Japan versus IHAC cohorts. This required the application of statistical tests (see methods and below). Specifically, we first identified a list of N=551 HLA-AP in HIV Gag, Pol and Nef, which represented the union of all HLA-AP identified in either Japan or IHAC for which both the viral polymorphism and the restricting HLA allele were observed in a minimum of 10 individuals per cohort (not shown). The latter criteria were employed in order to achieve some minimal statistical power to compare strengths of individual associations between cohorts. It is important to emphasize that these criteria would by definition exclude HLA alleles (and/or viral polymorphisms) present in one cohort but essentially absent in the other (as we would have no power, and in fact no rationale, to test whether their strengths of selection were statistically significantly different between cohorts).

For each HLA-AP, we calculated its natural logarithm of the odds ratio (lnOR) of association in each cohort — a measure that can be interpreted as an estimate of the strength of selection exerted by the HLA allele on that particular HIV codon, in that cohort. We then
applied a phylogenetically-corrected interaction test (17) to assess whether these lnORs of selection were significantly different in the Japanese versus the IHAC cohorts. In these analyses, statistical significance was defined as $p < 0.01$ and $q < 0.05$. Overall, 71 (of 551, 12.8%) HLA-APs originally identified in either Japan or IHAC cohorts exhibited significantly different strengths of selection between the two populations (Figure 5 and Table S3). The HLA-B*44:03-associated 125H substitution in Nef serves as an example of how to interpret these data. The lnOR of this association is 1.73 in Japan (with a cohort-specific $p$-value of $3.26 \times 10^{-6}$) versus 0.42 for IHAC (with a cohort-specific $p$-value of 0.36). Both lnORs are positive, indicating that 125H is positively associated with B*44:03 in both cohorts, but the higher lnOR in Japan indicates that the strength of selection of Nef-125H by B*44:03 is greater in Japan compared to IHAC (indeed, the cohort-specific $p$-values reveal that this association is significant in Japan but not IHAC). Finally, the $p$- and $q$-values for the intercohort comparison ($p = 1.02 \times 10^{-6}$ and $q = 1.19 \times 10^{-4}$; Table S3) confirm that the strength of selection of Nef-125H by B*44:03 is significantly greater in Japan compared to IHAC. Importantly, this difference is not simply attributable to intercohort differences in B*44:03 frequency (which is comparable between populations; Fig S1).

In addition to the HLA-B*44:03-associated 125H polymorphism in Nef, we identified 21 other HLA-AP whose strengths of selection were significantly greater in Japan compared to IHAC, yielding a total of 22 (of 71; 31.0%) HLA-APs in this category. Conversely, 39 (of 71; 54.9%) differentially-selected HLA-AP exhibited strengths of selection that were greater in IHAC compared to Japan. The HLA-A*26:01-associated 889S substitution in Pol serves as an example. The lnOR of this association is $-0.18$ in Japan (with a cohort-specific $p$-value of 0.3) versus $-1.17$ for IHAC (with a cohort-specific $p$-value of $7.92 \times 10^{-5}$). Both lnORs are negative, indicating that 889S is negatively associated with A*26:01 in both cohorts, but the more negative value for IHAC indicates that this association is stronger in IHAC compared to Japan. Finally, the $p$- and $q$-values for
the intercohort comparison \(p = 1.15 \times 10^{-4}\) and \(q = 4.48 \times 10^{-3}\); Table S3) confirm that of the strength of the negative association between Pol-889S by A*26:01 is significantly greater in IHAC compared to Japan.

Strikingly, the remaining 10 (of 71; 14.1%) differentially-selected HLA-APs displayed diametrically opposed directions of selection between the cohorts (defined here as lnORs of association that were positive in one cohort but negative in the other, where the cohort-specific \(p\)-values were <0.05 in both cases) (Fig. 5). The HLA-B*44:03-associated 120F substitution in Nef serves as an example. The lnOR of this association is 1.44 in Japan (with a cohort-specific \(p\)-value of \(2.03 \times 10^{-4}\)), indicating that HLA-B*44:03 is significantly positively associated with 120F in Japan. In contrast, the lnOR of this association is \(-0.69\) in IHAC (with a cohort-specific \(p\)-value of \(9.50 \times 10^{-3}\)), indicating that HLA-B*44:03 is significantly negatively associated with 120F in Japan. The \(p\)- and \(q\)-values for the intercohort comparison \((p = 2.15 \times 10^{-8}\) and \(q = 3.75 \times 10^{-6}\); Table S3) confirm that the opposing direction of selection of Nef-120F by B*44:03 between Japanese and IHAC cohorts is a statistically significant observation.

Of interest, the 71 HLA-APs identified as being under significantly different selection between Japan and IHAC cohorts were differentially distributed across HLA loci and HIV proteins (Fig. 6A and 6B). Specifically, HLA-A-associated polymorphisms that were significantly differentially selected across cohorts were most abundant in Gag, followed by Pol and Nef, whereas differentially-selected HLA-B-associated and HLA-C-associated polymorphisms were most numerous in Nef, followed by Pol and Gag. Taken together, results support the existence of HLA class I alleles that drive significantly different HIV escape pathways in global populations infected with the same viral clade. The uneven distribution of the locations of these differentially-selected polymorphisms across HLA loci and HIV regions raise the intriguing hypothesis that Gag and Pol/Nef may differentially evolve under selection pressures dominated by HLA-A versus HLA-B/C allele-restricted immune responses, respectively.
The present study comprised two major objectives, both of which are novel in terms of populations studied and/or analytical methods used. First, we characterized HLA-AP in HIV-1 clade B Gag, Pol and Nef and their relationship with clinical parameters in a large Japanese cohort. Second, we compared HLA-AP in Japanese versus non-Asian populations infected with HIV clade B, to identify population-specific differences in their selection. In particular, we wished to identify HLA-AP that are unique to Japan by virtue of the distinctive HLA distribution in this population, as well as cases where the same HLA allele drives divergent escape pathways in Japan vs. non-Asian populations.

This study is the first to identify HLA-APs in HIV-1’s structural and functional genes in Japanese populations. Only one previous study investigated HLA-AP in HIV-1 clade B-infected Asians (11): this study comprised 231 Chinese individuals infected during a narrow-source outbreak, and identified 141 HLA-associated polymorphisms at two-digit resolution. Our study differs from the latter with respect to cohort size, HLA genetics of the host population, HLA typing resolution and epidemiology of the epidemic. Using phylogenetically-informed approaches, we identified 284 HLA-APs within HIV-1 Gag, Pol and Nef in our cohort, supporting a strong influence of population-specific, HLA-driven immune pressures in shaping HIV-1 evolution in Japan. In contrast to a previous study undertaken in a predominantly Caucasian population that observed approximately one-half of the total number of Gag HLA-APs to be located within or flanking reported CTL epitopes (3), the majority of HLA-APs identified in the present study were not located nearby reported CTL epitopes. This discrepancy may be due to the limited number of Asian-specific HLA-restricted CTL epitopes identified to date, underscoring the need for further epitope discovery in these populations.

This study revealed differential frequencies of HLA-APs across HIV genes in the Japanese population. Consistent with previous studies of HLA-AP in HIV clade B (2, 16,
HLA-APs were more frequently detected in Nef than in Gag and Pol. Also consistent with previous observations in Caucasian, African, Chinese, and Mexican populations (1, 6, 11, 15, 18), the number of HLA-B-associated polymorphisms in our cohort was higher than that of HLA-A- or HLA-C-associated ones, further supporting a dominant role of HLA-B in HIV evolution (32). An interesting feature of the Japanese population is that approximately 70% of individuals carry HLA-A*24:02 (23). Despite sufficient statistical power to detect HLA-A*24:02-associated polymorphisms in our cohort, we identified only 9 of these, 6 of which were located in epitopes identified by our group (33-35). A possible explanation for the relatively low number of A*24:02-associated polymorphisms in Japan is that they have accumulated over time in circulating sequences such that they are no longer significantly enriched among persons expressing HLA-A*24:02. Further analysis of mutations selected by HLA-A*24:02-restricted CTLs should clarify the mechanism whereby high frequency HLA alleles influence the formation of HIV-1 polymorphisms.

Protective HLA alleles such as HLA-B*57, B*58, and B*27, select Gag mutations affecting viral replication in Caucasians and Africans (36-41) that may also provide some clinical benefit if they are transmitted to hosts lacking these alleles (42, 43). HLA-B*57, B*58, and B*27 are not present at appreciable frequencies in Japan (23). It is therefore perhaps unsurprising that no correlations between HLA-associated substitutions in Gag and HIV clinical parameters were observed in our cohort. In contrast, we observed a weak but significant inverse correlation between the frequency of HLA-APs in Pol and plasma viral load, which appeared to be driven by polymorphisms selected by HLA-B*52:01, an allele identified as protective in Japan (24). Upon further stratification by HLA-B*52:01 expression, the inverse correlation between VL and the total number of B*52:01-associated Pol substitutions was maintained in HLA-B*52:01−, but not in HLA-B*52:01+ individuals. Taken together, these findings suggest that transmitted B*52:01-associated polymorphisms could reduce viral fitness in a dose-dependent manner, though further studies would be required to assess this. In addition, these substitutions were not located within or nearby
known B*52:01-restricted epitopes. Thus, further research would be required to identify these epitopes and elucidate their mechanisms of escape.

Many previous studies of HLA-APs were performed at 2-digit HLA resolution (1-4, 6). Here, we performed HLA genotyping at 4-digit resolution, which allowed us to investigate differential escape between closely-related HLA subtypes in the Japanese cohort. Nearly one half of the HLA-AP identified in Japan were restricted by HLA allele groups containing two or more subtype members (A*02, A*26, B*15, B*40, C*03, C*08, and C*14). For five of these groups (A*02, A*26, B*15, B*40, and C*08), subtype members differed by substitutions within the peptide-binding groove, while for the remaining two groups (HLA-C*03 and -C*14), subtype members differed by substitutions located outside the peptide binding groove. Reasoning that amino acid differences located within the peptide-binding groove could modulate the nature or presentation of CTL epitopes, we hypothesized that the former group would generally exhibit distinct HLA-AP between subtype members, while the latter would generally exhibit similar or identical HLA-AP. However, we were surprised to observe substantial evidence for differential HLA-AP selection between closely-related HLA subtypes regardless of whether they differed in sequence within or outside the peptide binding groove. Significantly differential HLA-AP selection was observed at 3 of 9 HLA-C*03 associated sites and 5 of 14 HLA-C*14 associated sites (Fig. 3), proportions that were not significantly lower than the frequency of differential selection between subtypes that differed in their peptide-binding groove. This observation raised several hypotheses. HLA polymorphic sites outside the peptide-binding groove may indirectly influence the binding groove conformation, thus altering HLA-peptide interactions and/or T cell recognition. Another possibility is selection by NK cells, as KIR may recognize sites outside the peptide-binding groove. Indeed, KIR3DL1 bind to the loop including position 91 of HLA-B*57:01 (44). However it is not clear whether KIR2DLs, which are receptors for HLA-C, can bind to the loop outside the peptide-binding groove of HLA-C molecules. A recent study showed that HLA-C antigens...
are expressed at different levels on the cell surface, even among HLA-C subtypes (45). This study also observed a strong positive correlation between HLA-C expression level and the strength of HLA-C-mediated selection pressure conferred on HIV. Differential expression levels of these HLA-C subtype members in Japanese populations thus provide another potential explanation for this observation, for future follow-up.

Our second objective was to investigate differential HLA-AP between Japanese and non-Asian cohorts infected with HIV clade B. Here, the IHAC cohort (comprising clade B-infected Canadians, Americans and Australians) was used as a comparison group (16). HLA-AP identified in human populations will differ to some extent due to population-specific HLA distributions, yielding population-specific HLA-AP driven by HLA alleles present in one population but not another (15). Indeed, two-thirds of the HLA-APs identified in Japan had not previously been identified due to the presence of the restricting HLA alleles in Japan, but its absence (or far lower prevalence) in IHAC.

What remains unknown however, is the extent to which the same HLA allele may drive significantly different escape pathways in different human populations. To this end, we applied novel phylogenetically-corrected statistical approaches to assess the extent to which HLA-AP identified in either Japan or IHAC, that were restricted by HLA alleles present in both populations, exhibited significantly different strengths of selection between them. Of the 551 HLA-AP investigated, 71 (12.9%) were significantly differentially selected in Japan versus IHAC at a stringent statistical threshold of \( q < 0.05 \). Of these 71, 31% exhibited significantly greater strengths of selection in Japan compared to IHAC whereas 55% exhibited greater strengths of selection in IHAC compared to Japan. Surprisingly, the remaining 14% displayed diametrically opposed selection pathways in the two cohorts (where an HIV polymorphism represented the “adapted” form associated with a given allele in one cohort, but the “nonadapted” form associated with the same allele in the other cohort). It is important to emphasize that these significantly different pathways of HLA-AP selection are not simply attributable to differences in HLA frequency between the cohorts.
We feel that these are intriguing observations that merit further study. Nevertheless we propose the following potential interpretations. Firstly, these differences could be explained by functional differences in HIV-1 specific T cells elicited between Japanese and Caucasian cohorts, possibly as a result of differences in host genetics (for example in the genes that encode the T-cell receptor and/or modulate their expression). Such differences may influence the structure of the T-cell receptor(s) and thus the quality, quantity and/or makeup of the HIV-1 specific T cell repertoire, thus influencing the specific escape mutations selected in context of peptide-bound HLA. Further analysis of HIV-1 specific T cells driving the selection of these mutants in both cohorts is therefore warranted. It is also important to note that the intercohort HLA-AP comparisons, unlike previous analyses, did not correct for HLA linkage disequilibrium (LD) or HIV codon covariation. Although both Japan and IHAC cohorts feature HIV clade B infections, intra-clade differences in the viral backbone could also influence differential escape via epistatic effects. In-depth analyses of intercohort differences in HIV codon covariation relationships are also therefore warranted. Intercohort differences in HLA linkage disequilibrium are another possible contributor. Finally, the differentially-selected HLA-AP between cohorts appeared to be unevenly distributed by HLA locus: while HLA-A associated polymorphisms exhibiting differential selection between cohorts were more abundant in Gag compared to other proteins, HLA-B and HLA-C associated polymorphisms exhibiting differential selection between cohorts tended to be more abundant in Nef. This suggests that inter-cohort differential HLA-APs across HIV proteins may be arising as a result of cellular immune pressures exerted by distinct HLA class I loci, though this also requires further study.

Nevertheless, the present study confirms of the existence of population-specific HIV-1 adaptations that are attributable to the unique HLA allele distributions of that population (15). We additionally provide evidence of population-specific HIV adaptation to HLA-restricted immune responses that cannot be explained by differential HLA frequencies alone: cases where the same HLA allele drives significantly different, sometimes opposing,
escape pathways in different host populations. Taken together, results support differential HIV-1 adaptation to human populations worldwide, that might be driven by multiple host and viral mechanisms.
Acknowledgements

The authors thank Dr. Madoka Koyanagi and Rie Maruyama for collection of samples from the patients, Mari Hasegawa and Sayaka Nagata for technical assistance, Sachiko Sakai for her secretarial assistance, and Kyle Cobarrubias for assistance in constructing Figure 1.

TC is a JSPS Research Fellow. AQL is the recipient of a Frederick Banting and Charles Best Masters award from the Canadian Institutes for Health Research (CIHR). ZLB is the recipient of a New Investigator Award from the CIHR and a Scholar Award from the Michael Smith Foundation for Health Research (MSFHR). This research was supported by the Global COE program “Global Education and Research Center Aiming at the control of AIDS,” launched as a project commissioned by the Ministry of Education, Science, Sports, and Culture, Japan; and by Grants-in Aid for AIDS Research from the Ministry of Health, Labour, and Welfare, Japan.

Disclosure

The authors have no financial conflicts of interest.
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FIG 1 Escape Map of HLA-associated polymorphisms for Gag, Pol, and Nef.

Escape maps indicate the locations, specific residues and HLA restrictions of HLA-associated polymorphisms (all \( q < 0.2 \)). The global HIV-1 clade B consensus amino acid sequence is used as a reference. Shaded vertical bars separate blocks of 10 amino acids. “adapted” amino acids (those significantly overrepresented in the presence of a given HLA allele) are red. “nonadapted” amino acids (those significantly underrepresented in the presence of a given HLA allele) are blue. Polymorphisms associated with the same HLA allele that occur in proximity to one another are grouped together in yellow boxes. A list of all HLA-associated polymorphisms is provided in Table S1.

FIG 2 Correlation between HLA-associated substitutions in Gag, Pol, and Nef and viral load or CD4 count.

The total number of HLA-associated substitutions in each subject’s Gag, Pol, and Nef sequence was counted (see methods). (A) Correlation between the number of HLA-associated substitutions in Gag, Pol or Nef and pVL or CD4 count, (B) Correlation between pVL and the number of HLA-associated substitutions in Pol, with HLA-B*52:01-associated substitutions excluded (C) Correlation between pVL and the number of HLA-B*52:01-associated substitutions in Pol (all patients). (D) Correlation between the number of HLA-B*52:01 associated substitutions in Pol in HLA-B*52:01-positive individuals (left panel) and HLA-B*52:01-negative individuals (right panel). Analyses were performed using Spearman’s correlation. Linear regression lines are included in the plots.

FIG 3 Polymorphic positions in HLA class I molecules and differential escape between pairs of HLA subtypes.
In each ribbon diagram depicting the HLA-peptide-binding groove, the locations of residues differing among subtype members of the (A) HLA-A*26, (B) HLA-C*03, and (C) HLA-C*14 allele groups are highlighted in red and labeled with their locations and amino acids. HLA-AP comparisons between subtype members are shown in the corresponding plot below. Horizontal bars represent the natural logarithm of the odds ratio (lnOR), with colors indicating the restricting allele. Infinite lnORs are set to values of ±4. Boldface type indicates HLA-AP whose strengths of selection are statistically significantly different between the two subtype members ($p < 0.05, q < 0.2$).

**FIG 4 Location of HLA-associated sites common to HIV-1 clade B-infected Japanese and Caucasian cohorts, and those unique to Japan.**

The locations of all HLA-APs in Gag (500 codons), Pol (1,003 codons), and Nef (206 codons) are illustrated. The residues in the Pol transframe protein (TF) were not analyzed in IHAC and are thus excluded (grey bar). Blue squares identify codons that harbored at least one HLA-AP in both Japanese and IHAC cohorts. Red squares indicate codons that harbored HLA-AP in Japan, but that were not associated with any HLA alleles in IHAC.

**FIG 5 HLA-AP displaying significantly different strengths of selection between Japanese and IHAC cohorts.**

A phylogenetically-corrected interaction test was used to compare the natural logarithm of the odds ratio (lnOR) of selection of HLA-APs in the Japanese cohort versus the IHAC cohort. Comparisons with a $p < 0.01$ and $q < 0.05$ are reported. Bars represent the lnOR. Infinite lnORs are set to values of ±4. Boldface type indicates HLA-AP that display diametrically opposed directions of selection between the cohorts (defined here as lnORs of association that were positive in one cohort but negative in the other, where the cohort-specific p-values were <0.05 in both cases). The complete list of all comparisons with $p < 0.05$ is available in Table S3 in the supplemental material.
FIG 6 HLA-AP identified as being under differential strength of selection in Japanese and IHAC cohorts.

At a threshold of $p < 0.01$, $q < 0.05$, a total of 71 HLA-APs were identified as being under significantly different strengths of selection in Japanese and IHAC cohorts. The restricting HLA alleles and their HIV-1 protein locations are shown in (A). The number of differentially-selected HLA-AP, broken down by HLA locus and HIV-1 protein, is shown in (B).
Figure 3

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