Triple bypass: complicated paths to HIV escape

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Human immunodeficiency virus (HIV) type 1 is highly efficient at evading immune responses and persisting, ultimately causing fatal immunodeficiency in some patients. Mutation in the epitopes recognized by cytolytic CD8+ T cells (CTLs) is one such escape process. A new study now shows that one HIV−1 escape mutation may also result in impaired dendritic cell (DC) activity, possibly impairing later T cell responses to the same and other epitopes. The new data complete our understanding of the mechanisms by which the CTL response to an immunodominant gag epitope presented by human histocompatibility leukocyte antigen (HLA)−B27 is evaded. The complexity of the full escape helps to explain why patients with this HLA type progress to AIDS more slowly than average.

It is now well established that HIV can escape the immune responses that could otherwise control the infection, including both antibody and T cell responses. The latter has received much attention since it was first proposed in 1991 (1), and selection of escape mutations in several HIV epitopes targeted by CD8+ T cells has been carefully documented (2−5). Epitope escape usually occurs early in infection, as the rapidly replicating virus displays almost every possible mutation enabling it to evade any suppressive force. Although such escape mutations often undermine immune control, many also have a fitness cost for the virus (2, 3) so that escape is not always bad for the patient.

The first described and best-studied HIV CTL epitope comes from the p24 gag protein and spans amino acids 263−272 (KRWIILGLNK). This epitope is presented to CD8+ T cells by HLA B27 (2, 4−8). In patients with HLA B27, the CD8+ T cell response is dominated by T cells specific for this epitope, probably because these T cells are functionally diverse, bind antigen with relatively high avidity, and proliferate well (9). This response appears early in infection as the initial peak viremia declines (10), but unusually, virus escape does not occur until late in infection (8), probably because the escape requires more than one compensating mutation. In this issue, Lichterfeld et al. (p. 2813; reference 11) show that one of the mutations involved in this escape enhances HLA B27 binding to the inhibitory receptor immunoglobulin-like transcript (ILT) molecule 4 (also known as LIR2), which is expressed on monocytes and DCs (11).

Mechanics of a mutation

Peptides that bind to HLA B27 almost always have an arginine (R) at the second position, which fits snugly into the B pocket of the peptide-binding groove of the HLA molecule. The arginine at position 264 in the HIV−1 gag263−272 epitope is encoded by the amino acid codon AGA (or more rarely, AGG). Escape can occur when this arginine is mutated. The single mutations that have been recorded at this codon include AAA (lysine), ACA (threonine), and AGA or AGG (glycine). Although peptides with these substitutions may bind sufficiently to HLA B27 in in vitro assays, they are not recognized by T cells when expressed within the cell as viral or transfected sequences (8). These mutations thus effectively remove the epitope from the immune response. Given the high replication and mutation rates of HIV−1, one might expect the virus to escape rapidly and completely from the early T cells that recognize this epitope. Escape from this epitope, however, typically occurs only after several years, happening in at least 50% of HLA B27 patients who are studied for several years (3, 11, 12, and unpublished data). According to the Los Alamos HIV sequence database, these mutations are very rare in the general population of HIV-infected individuals, 95% of whom do not have HLA B27.

A possible explanation for the late escape was provided by Kelleher et al. (7), who showed that the critical arginine-to-lysine change at position 264 (R264K) only occurred when there had been a previous change of leucine-to-methionine at position six in the epitope (L268M). Unlike the arginine at position 2, the side chain at position 6 is not involved in HLA B27 binding but rather faces and interacts with the T cell receptor (Fig. 1). More recently, Schneidewind et al. (4) showed that there is nearly always a third amino acid change that accompanies the R264K and L268M mutations: an alanine-to-serine change at position 173 (A173S). This amino acid lies outside the epitope, but the three mutated amino acids cluster in the three-dimensional structure of p24, suggesting that they might compensate for each other. The R264K mutation imparts a fitness cost on the virus, as elegantly shown by Goulder et al. (2), who found that the virus rapidly mutated this sequence back to wild-type in HLA B27− babies that had been infected by B27− mothers carrying the mutated virus. This was directly tested in vitro by Schneidewind et al., who showed that an infectivity fitness cost for the R264K mutation was compensated for by the A173S mutation (4). This requirement for a triple mutation must be much rarer than a single mutation, and probably accounts for the delayed escape from this T cell response.
These observations imply a novel escape mechanism involving the inhibitory ILT4 receptor. This is the first demonstration that mutation in an HIV-1 CTL epitope can contribute to virus escape by influencing the interaction between HLA class I and a molecule other than the T cell receptor, although it is known that different epitope peptides (none yet known to escape) can influence the interaction between HLA molecules and other inhibitory receptors, such as the KIR family (12).

This is an important story for understanding HIV infection. The virus must be under considerable immune pressure to select this complex mosaic of mutations (Table I). Yet ultimately the virus evades immune control with the triple mutation and the virus load rises (8). Because this requires such a long period of time, other factors such as the loss of CD4 T cells must also contribute to the progression to AIDS. However, patients with B27 do progress to disease more slowly than the average HIV-infected individual, possibly because the B27-restricted T cells concentrate

**Table 1. Triple bypass**

| Mutation  | Timing       | Effect                              | Reference |
|-----------|--------------|-------------------------------------|-----------|
| L268M     | First        | Enhanced ILT4 binding              | (11)      |
| R264K (T/G) | Second/third | Abrogation of peptide binding to B27 | (7)       |
| S173A     | Second/third | Compensating effect                 | (4)       |

**Figure 1. Diagrammatic representation of the side chains of the HIV-1 gag peptide KRWILGLNK bound to HLA-B27, according to the structure determined by Stewart-Jones et al. (reference 13).** The α-1 helix is shown as a light blue dotted line. The floor of the peptide-binding groove below the peptide and α-2 helix are not depicted. Only the side chains of leucine 268 (L6) and asparagine 271 (N9) are exposed to the T cell receptor.
on a conserved part of the virus that requires this complicated and slow evasion process.

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