

Poster presentation

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## PI6-23. Antigen processing influences HIV-specific cytotoxic T lymphocyte immunodominance

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### Background

Cytotoxic T cells (CTL) play a key role in limiting human immunodeficiency virus (HIV)-1 replication. However, although the cellular immune response in HIV-infected individuals can potentially target multiple virus epitopes, the same few are repeatedly recognized. Here we investigated the factors determining observed CTL response hierarchies in Gag p17 and p24.

### Methods

We used constitutive and immuno-proteasomal digestion assays, transporter associated with antigen processing (TAP) binding assays, endoplasmic reticulum aminopeptidase (ERAAP) trimming assays, HLA binding assays, T cell cloning and ELISpot assays to evaluate the contribution of each of these factors to final epitope presentation and recognition. Key findings were further examined using structural analyses.

### Results

We show that CTL-immunodominance in regions of HIV-1 p17- and p24-Gag correlates with epitope abundance, which is influenced strongly by proteasomal digestion profiles, TAP-affinity and ERAAP-mediated trimming, and moderately by HLA affinity. Structural and functional

analyses demonstrate that proteasomal cleavage-preferences modulate the number and length of epitope-containing peptides, thereby affecting T cell response avidity and clonality. Cleavage patterns were affected by both flanking and intra-epitope CTL-escape mutations.

### Conclusion

Our analyses show that antigen processing shape CTL-response hierarchies, that viral evolution modify cleavage patterns, and suggest strategies for in vitro vaccine optimization.