The 9th Kumamoto AIDS Seminar: Abstract Submission Form

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Impact of V2 mutations for escape from a potent neutralizing anti-V3 monoclonal antibody during in vitro selection of a pr ※タイトルはボールド

Junji Shibata¹, Kazuhisa Yoshimura¹, Akiko Honda¹, Atsushi Koito¹, Toshio Murakami² and Shuzo Matsushita^{1*} ※著者はイタリック体、1st のみボールドで。

Division of Clinical Retrovirology and Infectious Diseases, Center for AIDS Research, Kumamoto University, Kumamoto¹ and The Chemo-Sero-Therapeutic Research Institute, Kyokushi, Kikuc, ※所属はイタリック体

KD-247, a humanized monoclonal antibody (MAb) to an epitope of gp120-V3-tip, has potent cross-neutralizing activity against subtype R primary HIV-1 isolates. To assess how KD-247 escape muta ※ フォントは Times, 12 ポイント variants by exposing bulked primary R5 virt ※図は挿入しないでください。 of relatively low concentrations of KD-247, viruses with two amino acid (aa) mutations (R166K/D167N) in V2 expanded, and under high KD-247 pressure, a V3-tip substitution (P313L) emerged in addition to the V2 mutations. On the other hand, a virus with a V2 175P mutation dominated during passaging in the absence of KD-247. By domain swapping analysis, we demonstrated that the V2 mutations and the P313L mutation in V3 contribute to partially and completely resistant phenotypes against KD-247, respectively. To identify the V2 mutation responsible for resistance to KD-247, we constructed pseudoviruses with single or double aa mutations in V2 and measured their sensitivity to neutralization. Interestingly, the neutralization phenotypes were switched, such that the 175th aa (Pro or Leu) located in the center of V2 was exchanged, indicating that the 175th aa has a key role in dramatically changing the Env oligomeric state on the membrane surface and affecting the neutralization phenotype against not only anti-V3 antibody but also rsCD4. These data suggest that HIV-1 can escape from anti-V3 antibody attack by changing the conformation of the functional envelope oligomer by acquiring mutations in the V2 region in environments with relatively low antibody concentrations.