



**S6 Fig. Phylogenetic pol tree with diverse CRF02\_AG Reference strains and patient consensus sequences.** Representative pol consensus sequences (con) were generated for each analyzed NGS patient time point from 6501 (red) and 6564 (blue). Patient consensus sequences were aligned with diverse CRF02\_AG Reference sequences downloaded from the Los Alamos Database (grey). They include representatives of all major CRF02\_AG branches and CRF02\_AG strains with highest similarity to the patients' con sequences according to HIV Blast. Neighbor joining phylogenetic trees were generated using MEGA and FigTree software