

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