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# Resistance. Is it still a problem? 

# OC 57 IMPACT OF RESISTANCE MUTATIONS ON VIROLOGICAL EFFICACY OF DTG-BASED MAINTENANCE TWODRUG REGIMENS: AN ARCA COHORT STUDY 

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Background: Two-drug regimens (2DR) are largely prescribed as maintenance therapy, nowadays mainly based on DTG. While many data have been reported about PI-based 2DR, the impact of resistance mutations and duration of virological suppression on DTG-based 2DR remains to be clarified. The aim of this study was to evaluate the impact of resistance mutations on virological outcome of DTG-based 2DR maintenance ART.
Material and methods: Virologically suppressed patients (pts) switching to DTG+3TC or DTG+RPV with prebaseline (time of switch=baseline, BL) resistance genotype (at least PR/RT) were selected from the ARCA database. Primary endpoint was virological failure (VF: an HIV-RNA, VL, $>200 \mathrm{cps} / \mathrm{mL}$ or 2 consecutive $>50$ $\mathrm{cps} / \mathrm{mL}$ ). The probability of VF was estimated by Kaplan-Meier analysis. Resistance to 2DR was defined as occurrence of at least Stanford HIVdb (v.8.5) low-level resistance (LLR) to at least one drug included in the current 2DR, based on cumulative genotype. CD4 changes were assessed using Student's t-test for paired samples. A secondary analysis comparing 2DR with DTG-based 3D regimens was also performed.
Results: A total of 318 2DR pts were analysed: 260 (82\%) switching to DTG+3TC, 58 (18\%) to DTG+RPV; 68\% were males, median age was 51 (44-56) years, 12 (6-23) years of HIV infection, 5 (3-8) years of virological suppression, nadir CD4 231 (121-329), 5 (3-9) previous ARV lines, $59 \%$ previously exposed to INSTI, $11 \%$ with resistance to current 2DR. The integrase sequence was available in $14 \%$ of patients, none harbouring resistance to DTG. 20 VF were observed, of whom 4 ( $3 / 17 \mathrm{VF}$ in DTG+3TC, $1 / 3$ in DTG+RPV) in patients with at least LLR at BL (M184V+K219Q; D67N+K70R+K219Q; D67N+K70R+T215Y+219Q; E138A), in a median FU of 1.3 years (IQR 0.6-2). The 2-year estimated probability of VF was $8.7 \%(95 \% \mathrm{CI} 4.4 ; 13) ; 8.6 \%(4.1 ; 13.1)$ in those without resistance and $9.7 \% ~(-4.4 ; 23.8)$ in those with resistance ( $\log$ rank: $\mathrm{p}=\mathrm{ns}$, figure 1). No factor was significantly associated with VF at multivariate analysis, but in pts with <6 years of virological suppression, BL resistance was associated with a higher probability of VF ( $p=0.003$ ). After 48 weeks, a statistically significant increase in CD4+ was detected ( +56 cells $/ \mathrm{mmc}, \mathrm{p}<0.001$ ), independently from baseline resistance. The 2-year estimated probability of VF in the reference 3DR group ( $n=564$ ) was not different from that for the 2DR group: $8.8 \%(5.9 ; 11.7)$ in the whole case file and $9.7 \%(6.6 ; 12.8)$ in the presence of baseline resistance. Longer time of virological suppression was the only factor associated with a lower risk of VF in the 3DR dataset.
Conclusions: DTG-based 2DRs show high virological efficacy, even in the context of predicted incomplete activity, at least within a short-term follow-up. A longer duration of virological suppression seems to decrease the impact of resistance on virological outcome, however further studies are warranted to confirm this hypothesis and possibly define a clinically useful threshold.

