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# Impact of transmitted HIV-1 drug resistance on the efficacy of first-line antiretroviral therapy with two nucleos(t)ide reverse transcriptase inhibitors plus an integrase inhibitor or a protease inhibitor.

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Keywords:	transmitted drug resistance, HIV-1, first-line antiretroviral therapy, integrase inhibitors, protease inhibitors



Title

2	Impact of transmitted HIV-1 drug resistance on the efficacy of first-line antiretroviral
3	therapy with two nucleos(t)ide reverse transcriptase inhibitors plus an integrase
4	inhibitor or a protease inhibitor.
5	
6	Running head: transmitted drug resistance and efficacy of first-line antiretroviral
7	therapy
8	
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- 34 **Key words**: transmitted drug resistance, HIV-1, first-line antiretroviral therapy,
- 35 integrase inhibitors, protease inhibitors

36 37

Abstract

38

## 39 Objectives

- The aim of this study was to examine the impact of transmitted drug resistance (TDR)
- 41 on response to first-line regimens with integrase strand transfer inhibitors (INSTI) or
- 42 boosted protease inhibitors (bPI).
- 43 Methods
- 44 From an Italian observational database (ARCA) we selected HIV-1 infected drug-naïve
- 45 patients starting 2 nucleoside reverse transcriptase inhibitors (NRTI) and either an
- 46 INSTI or a bPI, with available pre-ART resistance genotype. The endpoint was
- 47 virological failure (VF: plasma HIV-1 RNA >200 copies/ml after week 24,). WHO
- 48 surveillance drug resistance mutations and the Stanford algorithm were used to classify
- 49 patients into three resistance categories: no TDR (A), TDR but fully-active ART
- prescribed (B), TDR and at least low-level resistance to one or more prescribed drug
- 51 (C).
- 52 Results
- We included 1,365 patients with a median follow-up of 96-weeks (IQR 54-110): 1,205
- 54 (88.3%) starting bPI and 160 (11.7%) INSTI. Prevalence of TDR was 6.1%, 12.5%,
- 55 0.5% and 0% for NRTI, NNRTI, bPI, and INSTI, respectively.
- 56 Cumulative Kaplan-Meier estimates for VF at 48-weeks were 11% (10.1-11.9) for the
- 57 bPI- and 7.7% (5.4-10) for the INSTI-group.
- 58 In the INSTI-group, cumulative estimates for VF at 48-weeks were 6% (4-8) in
- 59 resistance category A, 5% (1-10) in B and 50% (30-70) in C (p<0.001). Resistance
- 60 category C (versus A, adjusted hazard ratio, aHR 12.6, 3.2-49.8, p<0.001) and nadir
- 61 CD4 (+100 cells/ $\mu$ L, aHR 0.6, 0.4-0.9, p=0.03) predicted VF. In the bPI-group, VF
- rates were not influenced by baseline resistance.
- 63 Conclusions
- 64 Our data support the need of NRTI-resistance genotyping in patients starting an INSTI-
- 65 based first-line ART.

66

### Introduction

Transmission of drug resistant HIV-1 is a well-known phenomenon detected in around 8% of newly diagnosed individuals in Europe, with significant differences depending on viral subtype, geographic area, risk group and migration timeline. Transmitted drug resistance (TDR) is increasing in Southern and Eastern Africa, particularly to the antiretroviral class of non-nucleoside reverse transcriptase inhibitors (NNRTI), a cornerstone of recommended first-line antiretroviral therapy (ART) in these countries. TDR may significantly influence the outcome of ART, therefore drug resistance testing is recommended for the choice of the first-line regimen in resource-rich countries.

The risk of virological failure was increased in patients harboring pre-treatment drug resistance to at least one of the prescribed drugs in NNRTI-based regimens, as compared with individuals without pre-treatment drug resistance, but not in patients with pre-treatment drug resistance and fully active ART.<sup>12</sup>

 International panels currently recommend first-line ART regimens including integrase strand transfer inhibitors (INSTI) or boosted protease inhibitors (bPI), because of their efficacy and tolerability. However, in the absence of resistance testing, some authors suggest to use bPI due to their higher genetic barrier compared with INSTI. Indeed, the influence of TDR on the efficacy of INSTI-based first-line regimens has not yet been established, due to the exclusion of individuals carrying TDR from clinical trials and the sparse data from observational cohorts. 5,6,13-16

The aim of this study was to examine the impact of TDR on response to first-line regimens in naïve patients starting INSTI-based 3-drug antiretroviral therapy. As a reference, we also analyzed the impact of TDR on the efficacy of boosted PI-based regimens.

#### Methods

Protease, reverse transcriptase and integrase genotype sequences from treatment-naïve HIV-1 infected adults starting a first-line therapy including 2 nucleoside or nucleotide

102	reverse transcriptase inhibitors (NRTI) plus 1 INSTI or 2 NRTI plus 1 bPI from January
103	2008 to June 2016 were selected from the Antiviral Response Cohort Analysis (ARCA),
104	an Italian multicenter virological and clinical database [http://www.dbarca.net],
105	including cases with at least 1 plasma HIV-1 RNA value after 24 weeks of follow up.
106	The database was approved by the local Ethics Committees and written informed
107	consent was obtained from all patients before participation. The study was performed in
108	accordance with the ethical guidelines of the Declaration of Helsinki (7th revision).
109	
110	Plasma genotypic resistance was determined by Sanger's population sequencing using
111	commercially available or homebrew systems. TDR was defined as the detection of at
112	least one mutation among those included in the WHO-recommended surveillance drug
113	resistance mutation (SDRM) list for NRTI, NNRTI, bPI <sup>17</sup> and those included in the
114	Stanford HIVdb SDRM Worksheet for INSTI
115	[https://hivdb.stanford.edu/pages/SDRM.worksheet.INI.html].  The  Stanford  HIVdb
116	algorithm (version 8.4, https://hivdb.stanford.edu) was used to classify patients into
117	three resistance categories: <sup>4</sup> absence of TDR mutations (resistance category A),
118	presence of TDR mutations but use of a fully-active ART regimen (B), or presence of
119	TDR mutations and at least low-level resistance to at least one prescribed drug (C).
120	HIV-1 subtyping was available as automatically performed by BLAST upon sequence
121	upload and further analyzed by phylogenetic analysis in case of <95% homology to the
122	pure subtype reference panel.
123	
124	The primary outcome was virological failure, defined as a plasma HIV-1 RNA $>$ 200
125	copies/mL after week 24, ignoring treatment changes. Survival analysis, using Kaplan-
126	Meier curves, was employed to estimate the probability of virological failure. Predictors
127	of virological failure were investigated using Cox regression models. All analyses were
128	performed using SPSS (version 22, IBM, Armonk, NY).
129	
130	Results
131	
132	A total of 1,365 patients were included, 1,205 (88.3%) treated with 2 NRTI plus 1 bPI
133	and 160 (11.7%) treated with 2 NRTI plus 1 INSTI. Baseline patients' characteristics
134	are shown in table 1. The main differences between the two treatment groups were a
135	higher baseline plasma HIV-1 RNA and lower baseline and nadir CD4 cells counts in

the bPI group. Patients in the INSTI group were cared more frequently in Southern Italy 136 137 and started therapy more recently. The most frequently prescribed INSTI was 138 raltegravir (RAL) (39%),followed by dolutegravir (DTG) (35%) and 139 elvitegravir/cobicistat (EVG/c) (26%). The most frequently used bPI was 140 lopinavir/ritonavir (LPV/r) (41%), followed by atazanavir/ritonavir (ATV/r) (30%) and 141 darunavir/ritonavir (DRV/r) (28%). 142 The overall prevalence of any TDR mutation was 18.4%, without differences between 143 144 groups. NRTI, NNRTI, PI and INSTI resistance mutations were detected in 83 (6.1%), 145 171 (12.5%), 35 (2.6%) and 0 (0.0%) patients, respectively. While there was a similar 146 prevalence of NRTI TDR in the two treatment groups, NNRTI TDR was more frequent 147 in the bPI group (13.1% versus 8.1% in the INSTI group, p=0.043), whereas PI TDR 148 was less frequent in the PI group (2.1% versus 6.3%, p=0.05). 149 150 During a median follow-up time of 96 weeks (IQR 54-110) virological failure occurred 151 in 195 individuals in the PI-group and in 11 in the INSTI-group, with an estimated 152 cumulative probability at 48 weeks of 11% (CI 95% 10.1-11.9) and 7.7% (CI 95% 5.4-153 10), respectively (p=0.01 by log-rank test). 154

In the INSTI group, resistance category C showed a significantly higher estimated 155 156 probability of 48-week virological failure (50%, 95% CI 30-70) versus A (6%, 95% CI 157 4-8) and B (5%, 1-10) (p<0.001). By contrast, in the bPI group the estimated probability 158 of virological failure at 48 weeks was similar in three categories: category A 11% (95%) 159 CI 10-12), B 12% (95% CI 10-14) and C 9% (95% CI 5-13) (p=0.390) (Fig.1). In the 160 INSTI group, but not in the PI group, resistance category C (versus A, adjusted hazard ration, aHR 12.6, 3.2-49.8, p<0.001) and nadir CD4 (+100 cells/μL higher, aHR 0.6, 161 0.4-0.9, p=0.03) independently predicted virological failure. In the PI group, in a 162 163 multivariable model adjusting for gender, nationality, TDF/FTC use, viral subtype, type of bPI and TDR to NRTI, independent predictors of virological failure were AZT/3TC 164 use (aHR 2.3, CI 95% 1.4-3.9, p=0.002), calendar year (per 1 year more recent, aHR 165 0.9, CI 95% 0.8-0.9, p=0.04) and LPV/r use (versus DRV/r, aHR 1.4, CI 95% 1.0-2.0, 166 p=0.03). 167

Eleven patients, mostly (9/11) harboring viral subtype B, experienced virological failure in the INSTI group: 8 were on treatment with RAL, 2 with DTG and 1 with EVG/c. At failure, plasma HIV-1 RNA ranged between 210 and 213,200 copies/mL and higher values were detected in patients with lower baseline CD4 counts. Three of the 11 failing INSTI carried TDR to NRTI: 2 M41L and 1 M184V, while none carried resistance to INSTI. Seven patients changed antiretroviral therapy after virological failure, with 6 patients switching to a bPI-based regimen. Among those that continued the previous regimen, 3 reached virological re-suppression at the subsequent visit and 1 was lost to follow up.

#### Discussion

 The key finding of this study is the impact of pre-treatment HIV-1 drug resistance on the risk of virological failure in patients initiating ART with 2 NRTI plus INSTI. Despite the small number of cases, the magnitude of this effect was very relevant, with a more than 10-fold higher adjusted hazard of virological failure as compared to patients without TDR. To our knowledge, this is the first report showing a significant impact of TDR to NRTI on the activity of first-line regimens with 2 NRTI plus INSTI, the current standard of care of first-line ART. Indeed, previous observational studies on the influence of TDR did not include INSTI-based regimens and clinical trials with INSTI excluded patients with TDR. 1,2,5,13,18,19 Interestingly, in the same group, TDR not affecting the activity of the prescribed drugs did not show any impact on virological efficacy. This finding is reassuring, suggesting that even in the presence of TDR, INSTI-based first-line regimens are effective when fully active accompanying drugs are selected based on the resistance test result.

The overall prevalence of TDR in this cohort was 18.4%, higher than usually reported in European cohorts, and was primarily driven by NNRTI resistance. This could be explained by the fact that, detection of TDRmay have advised to use high-genetic barrier bPI therapy, resulting in an overestimate of TDR in the case file. In addition, bPI-based regimens were preferentially prescribed to more challenging patients, such as those with higher baseline viral load and lower CD4 counts, possibly explaining at least in part the higher virological efficacy of INSTI-based as compared with PI-based regimens observed here. The relatively long period of observation (2008-2016) may

203	also explain some imbalance observed between the two treatment groups reflecting drug
204	availability over time. In the PI-group, AZT/3TC and LPV/r use were associated to
205	more frequent virological failure, suggesting a crucial role of their lower tolerability and
206	efficacy.
207	
208	The main limitations of this study are the retrospective nature, the small number of
209	patients treated with INSTIs and the relatively limited sample size in the INSTI
210	treatment groups, which did not allow a sufficient power to detect differences among
211	drugs with different genetic barrier. Future analyses including a larger and balanced
212	INSTI group are necessary to confirm our findings and clarify whether NRTI TDR has
213	a different impact on virological efficacy using different types of INSTI.
214	
215	In conclusion, our findings support the need of pre-treatment drug resistance testing to
216	NRTI in order to optimize antiretroviral therapy in patients starting first-line INSTI-
217	based regimens.
218	
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220	
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230	
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References

247

- 1 Hofstra LM, Sauvageot N, Albert J et al. Transmission of HIV Drug Resistance and
- the Predicted Effect on Current First-Line Regimens in Europe. Clin Infect Dis 2016;
- 250 62:655-663.

251

- 252 2 Colafigli M, Torti C, Trecarichi EM et al. Evolution of transmitted HIV-1 drug
- resistance in HIV-1-infected patients in Italy from 2000 to 2010. Clin Microbiol Infect
- 254 2012; 18:E299-E304.

255

- 3 Hauser A, Hofmann A, Hanke K et al. National molecular surveillance of recently
- acquired HIV infections in Germany, 2013 to 2014. Euro Surveill 2017; 22:pii=30436.

258

- 4 Gupta RK, Gregson J, Parkin N et al. HIV-1 drug resistance before initiation or re-
- 260 initiation of first-line antiretroviral therapy in low-income and middle-income countries:
- a systematic review and meta-regression analysis. Lancet Infect Dis 2017 Dec 5, pii:
- 262 S1473-3099(17)30702-8.

263

- 5 Wittkop L, Gunthard HF, de Wolf F et al. Effect of transmitted drug resistance on
- virological and immunological response to initial combination antiretroviral therapy for
- 266 HIV (EuroCoord-CHIAN joint project): a European multicohort study. Lancet Infect
- 267 Dis 2011; 11:363-371.

269 6 Di Biagio A, Rusconi S, Marzocchetti A et al. The role of baseline HIV-1 RNA, drug 270 resistance, and regimen type as determinants of response to first-line antiretroviral 271 therapy. J Med Virol 2014; 86: 1648-1655. 272 273 7 Kantor R, Smeaton L, Vardhanabhuti S et al. Pretreatment HIV drug Resistance and 274 HIV-1 Subtype C are independently associated with virological failure: results from the 275 multinational PEARLS (ACTG A5175) Clinical Trial. Clin Infect Dis 2015; 60: 1541-276 1549. 277 278 8 Linee guida Italiane sull'utilizzo dei farmaci antiretrovirali e sulla gestione HIV-1. 279 diagnostico-clinica delle persone con infezione da 280 http://www.simit.org/medias/1047-lg-hiv-2016-c17pubblicazioni2545allegato.pdf 281 9 Guidelines for the use of Antiretroviral Agents in HIV-1-Infected Adults and 282 283 Adolescentes. https://aidsinfo.nih.gov/guidelines on 9/9/2017 284 285 10 Guidelines, version 9.0. European AIDS Clinical Society (EACS), October 286 2017.http://www.eacsociety.org/guidelines/eacs-guidelines/eacs-guidelines.html. 287 288 11 Antiretroviral drugs for treatment and prevention of HIV infection in adults 2016 289 Recommendations of the International Antiviral-Society USA panel. 290 https://jamanetwork.com/journals/jama/fullarticle/2533073. 291 292 12 Hamers RL, Schuurman R, Sigaloff KC, et al. Effect of pretreatment HIV-1 drug 293 resistance on immunological, virological, and drug-resistance outcomes of first-line 294 antiretroviral treatment in sub-Saharan Africa: a multicentre cohort study. Lancet Infect 295 Dis 2012; 12:307-317.

296

13 Clotet B, Feinberg J, van Lunzen J et al. Once-daily dolutegravir versus darunavir plus ritonavir in antiretroviral-naive adults with HIV infection (FLAMINGO): 48 week results from the randomised open-label phase 3b study. *Lancet* 2014; 383:2222-2231.

300

301 14 Lennox Jl, Landovitz RJ, Ribaudo HJ et al. Efficacy and tolerability of 3 302 nonnucleoside reverse transcriptase inhibitor-sparing antiretroviral regimens for

303	treatment-naive volunteers infected with HIV-1: a randomized-controlled, equivalence
304	trial. Ann Intern Med 2014; 161:461-471.
305	
306	15 Socias ME, Nosova E, Kerr T et al. Patterns of transmitted drug resistance and
307	virological response to first-line antiretroviral treatment among HIV-positive people
308	who use illicit drugs in a Canadian setting. Clin Infect Dis 2017 May 6. Doi:
309	10.1093/cid/cix428 [Epub ahead of print].
310	
311	16 Zu Knyphausen F, Scheufele R, Kucherer C et al. First line treatment response in
312	patients with transmitted HIV drug resistance and well defined time point of HIV
313	infection: updates results from the German HIV-1 seroconverter study. PLoS One 2014;
314	9:e95956.
315	
316	17 Bennett DE, Camacho RJ, Otelea D et al. Drug resistance mutations for surveillance
317	of transmitted HIV-1 drug-resistance: 2009 update. PLoS One 2009; 4(3):e4724.
318	
319	18 Eron JJ Jr, Rockstroh JK, Reynes J, et al. Raltegravir once daily or twice daily in
320	previously untreated patients with HIV-1: a randomised, active-controlled, phase 3 non-
321	inferiority trial. Lancet Infect Dis 2011 Dec;11(12):907-15.
322	
323	19 Squires K, Kityo C, Hodder S, et al. Integrase inhibitor versus protease inhibitor
324	based regimen for HIV-1 infected women (WAVES): a randomised, controlled, double-
325	blind, phase 3 study. Lancet HIV 2016 Sep;3(9):e410-e420.
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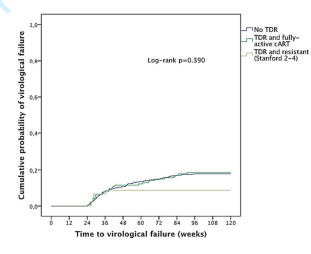
Fig.1 Kaplan–Meier curves showing the impact of the different pre-treatment HIV-1 drug resistance category on the virological outcome of first-line regimens based on 2 NRTI plus either a boosted PI (a) or an integrase inhibitor (b).

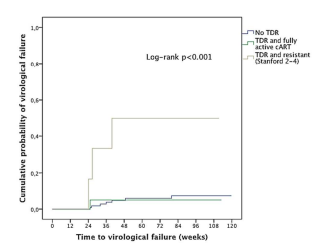
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a. PI-group

b. INSTI-group

Table 1. Baseline characteristics of the patient population.

	Overall	bPI group	INSTI group	P-value*
Characteristics	N=1,365	N= 1,205	N= 160	
Male, n (%)	1006/1355 (73.7)	880/1196 (73.0)	126/159 (78.8)	0.30
Age (year), median (IQR)	40 (33-48)	40 (33-48)	40 (30- 48)	0.60
Italian born, n (%)	982/1365 (71.9)	882/1205 (73.2)	100/160 (62.5)	0.05
Risk factor, n (%):				< 0.001
Heterosexual contacts	444 (32.5)	405 (33.6)	39 (24.4)	
MSM	272 (19.9)	245 (20.3)	27 (16.8)	
Injection drug users	109 (8.0)	101 (8.4)	8 (5.0)	
Other/Unknown	540 (39.6)	454 (37.7)	86 (53.8)	
Geographical area, n (%):				< 0.001
Northern Italy	584 (42.8)	540 (44.8)	44 (27.5)	
Central Italy	522 (38.2)	465 (38.6)	57 (35.6)	
Southern Italy and Islands	259 (19.0)	200 (16.6)	59 (36.9)	0.001
Calendar year of treatment start, median (IQR)	2011 (2009-2013)	2011 (2009-2012)	2015 (2014-2016)	< 0.001
Fime from HIV diagnosis (years), median (IQR)	0.3 (0.1-2.5)	0.3 (0.1-2.5)	0.5 (0.2-2.9)	0.60
Baseline plasma HIV-1 RNA (log <sub>10</sub> copies/mL), median (IQR)	4.9 (4.3-5.4)	4.9 (4.4-5.4)	4.7 (4-5.2)	< 0.001
Baseline CD4 cell count (cells/mm <sup>3</sup> ), median (IQR)	258.5 (103-383)	240 (96-364)	380 (198-557)	< 0.001
CD4 nadir cell count (cells/mm <sup>3</sup> ), median (IQR)	230 (95-346)	222 (89-334)	323 (167-496)	< 0.001
Subtype, n (%):				
В	944 (69.2)	839 (69.6)	105 (65.6)	0.30
non B	421 (30.8)	366 (30.4)	55 (34.4)	
Backbone, n (%):				
TDF/FTC	1011 (74.1)	895 (74.3)	116 (72.5)	0.63
ABC/3TC	240 (17.6)	198 (16.4)	42 (26.2)	0.002
AZT/3TC	102 (7.5)	102 (8.5)	0 (0)	< 0.001
other	12 (0.9)	10 (0.8)	2 (1.3)	0.59
Anchor drug DRV/r		339 (28.1)		

	LPV/r		499 (41.4)		
	ATV/r		367 (30.5)		
	RAL		207 (20.0)	63 (39.4)	
	EVG			41 (25.6)	
	DTG			56 (35.0)	
Patients with transmitted drug resistance, n (%):					
	Any class	251 (18.4)	222 (18.4)	29 (18.1)	0.514
	NRTI	83 (6.1)	74 (6.1)	9 (5.6)	0.484
	NNRTI	171 (12.5)	158 (13.1)	13 (8.1)	0.043
	PI	35 (2.6)	25 (2.1)	10 (6.3)	0.05
	INSTI	0 (0.0)	0 (0.0)	0 (0.0)	

./ritonavir; DTG, dolutegrav.
reverse transcriptase inhibitor; bPI, booss. ABC/3TC, abacavir/lamivudine; ATV/r, atazanavir/ritonavir; AZT/3TC, zidovudine/lamivudine; DRV/r, cps/mL, copies/mL; darunavir/ritonavir; DTG, dolutegravir; EVG, elvitegravir; INSTI, Integrase strand transfer inhibitors; LPV/r, lopinavir/ritonavir; MSM, man who have sex with man; NRTI, nucleoside reverse transcriptase inhibitor; NNRTI, non-nucleoside reverse transcriptase inhibitor; bPI, boosted protease inhibitor; RAL, raltegravir; TDF/FTC, tenofovir/emtricitabine.