

Pregnancy and Neonatal Outcomes following Prenatal Exposure to Dolutegravir

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Disclosure slide

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Background

- Dolutegravir (DTG) is an integrase strand inhibitor approved for the treatment of HIV in adults and adolescents since 2013
- Marketed as a single agent as Tivicay[©] and as a fixed dose combination tablet as Triumeq[©] (DTG/abacavir/lamivudine)
- In animal and *ex vivo* human placenta perfusion studies, DTG was shown to cross the placenta
- However, there is minimal information on use and safety of DTG in pregnant women



• To assess maternal and fetal outcomes following DTG use during pregnancy in real-world European settings

Our objectives were:

- To describe the characteristics of pregnant women receiving DTGbased regimens
- To describe the frequency of adverse pregnancy and birth outcomes, by trimester of DTG-exposure

Study design

- Analysis of prospectively collected individual patient data (i.e. with ARV exposure data collected before outcome is known) in observational studies of pregnant women living with HIV and their infants in Europe
- Data collection through:
 - European Pregnancy and Paediatric HIV Cohort Collaboration (EPPICC)
 - NEAT-ID network
 - PANNA (Pharmacokinetics of newly developed ANtiretroviral agents in HIV-infected pregNAnt women) <u>www.pannastudy.com</u>
- Anonymised individual patient data collected from studies/sites using a data specification based on a modified HIV Data Exchange Protocol (www.hicdep.org)
- Data merger and analysis at UCL
- NB Presenting updated results since submission of abstract

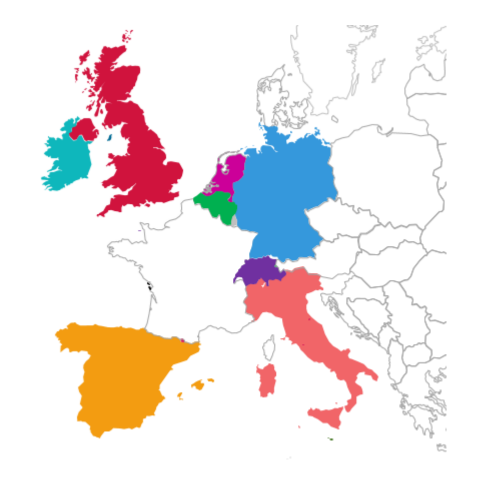
Definitions

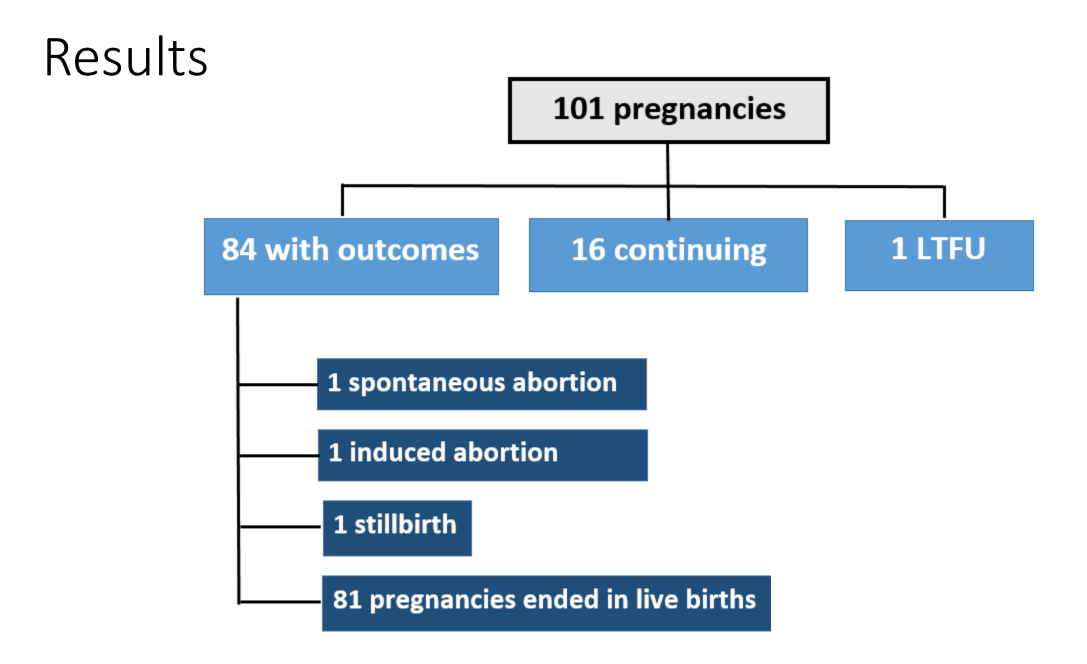
Pregnancy / birth outcome	Definition
Induced abortion	Voluntary termination of pregnancy before 22 weeks gestation
Spontaneous abortion	Death of a fetus or expulsion of the products of conception before 22 weeks gestation
Low birth weight	Birth weight of <2500 grams
Very low birth weight	Birth weight of <1500 grams
Small for gestational age	Based on sex-specific US standard ¹
Preterm birth	Birth of live infant at <37 weeks gestation
Stillbirth	Death of a fetus occurring at 22 weeks of gestation or more, or for situations in which the gestational age is unavailable, a fetus weighing at least 500 grams

1. Alexander GR, Himes JH, Kaufman RB, Mor J, Kogan M. A United States national reference for fetal growth. Obstetrics and Gynecology. 1996;87(2):163-8.

Results: geographic distribution

Country	Number of pregnancies
Belgium	2
Germany	19
Italy	5
Netherlands	2
Spain	9
Switzerland	3
UK & Ireland	61
TOTAL	101





Pregnancies: maternal characteristics

		N (%)	
Ethnicity	Black	71 (70%)	Timing of HIV
N=100	White	22 (22%)	diagnosis N=101
	Other	7 (7%)	History of AIDS
Region of origin	sub-Saharan Africa	62 (67%)	
N=93	Europe	22 (24%)	HCV status
	Other	9 (10%)	HBV status
Age at	<25 years	16 (16%)	N=91
conception N=101	25-34 years	46 (46%)	CD4 count (first
N-101	≥ 35 years	39 (39%)	in pregnancy)
Mode of HIV	Heterosexual	81 (86%)	
acquisition	Injecting drug use	3 (3%)	ART at conception
N=94	Vertical	9 (10%)	N=92
	Other	1 (1%)	

		N (%)
Timing of HIV	Pre-pregnancy	86 (85%)
diagnosis N=101	Antenatal	15 (15%)
History of AIDS N=89	Yes	10 (11%)
HCV status N=91	HCV seropositive	8 (9%)
HBV status N=91	HBsAg positive	4 (4%)
CD4 count (first in pregnancy)	≤350 cells/mm ³	38 (43%)
N=89	>350 cells/mm ³	51 (57%)
ART at conception N=92	Yes	55 (60%)

Pregnancies: earliest exposure to DTG

	T1 N (%)	T2 N (%)	T3 N (%)	Missing N (%)	Total
All pregnancies	58 (57.4)	24 (23.8)	18 (17.8)	1 (1.0)*	101 (100%)
Pregnancies ending in livebirths	42 (51.9)	21 (25.9)	17 (21.0)	1* (1.2)	81 (100%)
Stillbirth	0	1	0	0	1
Induced abortion~	1	0	0	0	1
Spontaneous abortion [#]	1	0	0	0	1

* Missing start date of DTG for 1 pregnancy ending in livebirth

~ Personal decision, no fetal abnormality

At 10 weeks gestation



• Among the 82 pregnancies ending in live birth or stillbirth

		Ν	(%)
Twin pregnancy N=82	No	80	(97.5)
	Yes*	2	(2.5)
Mode of delivery	Vaginal	45	(57.0)
N=79	Elective CS	21	(26.6)
	Emergency CS	9	(11.4)
	CS unspecified	4	(5.1)

* Twin pregnancies: 1 at 38 weeks, 1 at 33 weeks, livebirths

Gestational age

- Among 80 infants (79 pregnancies ending in singleton live birth and 1 pregnancy ending in stillbirth) 13.8% (11/80) infants were delivered preterm
- By earliest DTG exposure in pregnancy:

		Earliest DTG exposure in T1		Earliest DTG exposure in T2		Earliest DTG exposure in T3		Total	
Gestational	≥37 weeks	37/40	(92.5%)	16/22	(72.7%)	15/17	(88.2%)	68/79	(86.1%)
age N=79*	34-36 weeks	2/40	(5.0%)	5/22	(22.7%)	2/17	(11.8%)	9/79	(11.4%)
	<34 weeks	1/40	(2.6%)	1/22	(4.6%)	0/17	(0.0%)	2/79	(2.5%)

* 1 live-born infant excluded due to missing DTG start date (born at term)

Birthweight and small-for-gestational age

 Among 80 infants (79 pregnancies ending in singleton live birth and 1 pregnancy ending in stillbirth) 16.7% (13/78) had LBW and 18.7% (24/75) were SGA

By earliest DTG exposure in pregnancy:

		Earliest DTG exposure in T1		Earliest DTG exposure in T2		Earliest DTG exposure in T3		Total	
Birthweight	≥2500g	35/39	(89.7%)	15/21	(71.4%)	14/17	(82.3%)	64/77	(83.1%)
N=77~	1500-2499g	4/39	(10.3%)	6/21	(28.6%)	3/17	(17.7%)	13/77	(16.9%)
	<1500g	0/39		0/21		0/17		0/77	
SGA	No	34/39	(87.1%)	13/20	(65.0%)	14/16	(87.5%)	62/75	(81.3%)
N=75	Yes	5/39	(12.8%)	7/20	(35.0%)	2/16	(12.5%)	24/75	(18.7%)

~ 1 live-born infant excluded due to missing DTG start date (BW >2500g); 2 excluded due to missing BW (earliest DTG exposure in T1)

Congenital abnormalities

- Data available for 81 of 84 live-born / stillborn infants
 - Abnormalities were reported in 4 infants (4.9%, 95% CI 1.4, 12.2%)
 - No defect in the stillborn infant
- By earliest DTG exposure in pregnancy:

		Earliest DTG exposure in T1		Earliest DTG exposure in T2		Earliest DTG exposure in T3		Missing earliest DTG exposure
Infant has ≥1	No	39/42	(92.9%)	23/24	(95.8%)	14	(100%)	1
abnormalities N=81	Yes	3/42 (7.1%)		1/24 (4.2%)		0		0

Congenital abnormalities

• Data available for 81 of 84 live-born / stillborn infants

	Abnormality	Earliest DTG exposure	Infant sex	Maternal details	Other ARV exposures	Country
Infant 1	Patent Foramen Ovale, with small left-to-right interatrial shunt	From conception	Male	Black African, aged 38 at delivery	ЗТС, ABC	Italy
Infant 2	Bilateral hexadactyly, hands (father has the same defect)	Week 3	Male	White, aged 40 at delivery	3TC/ABC, FTC/TDF in T1	Italy
	Hypospadias					
Infant 3	Ankyloglossia (tongue-tie)	Week 12	Male	White, vertically infected, aged 31 at delivery	DRV/r, FTC/TDF, ATZ/r, RAL, TDF in T1	Italy
Infant 4	Hyperpigmentation on back	Week 14	Male	Black African, aged 34 at delivery	3TC, ABC	Switzerland

Conclusions

- This is the largest study to date of DTG use in pregnancy in Europe
- Nearly 60% of the included pregnancies had first trimester DTG exposure
- PTD (14%) and SGA (19%) rates were similar to those reported in UK¹:
 - PTD: 14% among women on ART at conception with CD4<350, 11% not on ART at conception irrespective of CD4
 - SGA: 20%
- Maternal characteristics of this "first wave" of DTG-exposed pregnancies differ from the larger population of pregnant women living with HIV
 - 10% vertically infected, 9% HCV co-infected
- Ongoing work to collect outcome data on the 16 continuing pregnancies
- These findings contribute to the evidence base on the real-world safety of DTG in pregnancy, but small numbers preclude firm conclusions
- Further prospective monitoring is required, particularly as DTG use expands

Acknowledgements

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- Italian Group on Surveillance of Antiretroviral Treatment in Pregnancy
- NENEXP Study (Catalonia)
- European Collaborative Study on HIV-infected pregnant women & their children
- Swiss Mother and Child HIV Cohort Study (MoCHiV)
- UK / Ireland National Study of HIV in Pregnancy and Childhood (NSHPC)
- PANNA Study
- NEAT-ID Network
- This study was supported by ViiV Healthcare









The European treatment network for HIV, hepatitis and global infectious diseases