

# GUIDELINES Version 11.1 October 2022

English

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# **Introduction to the EACS Guidelines 2022**

Welcome to the EACS Guidelines!

These Guidelines were developed by the European AIDS Clinical Society (EACS), a not-for-profit organisation, whose mission is to promote excellence in standards of care, research and education in HIV infection and related co-infections, and to actively engage in the formulation of public health policy, with the aim of reducing the HIV disease burden across Europe.

The EACS Guidelines were first published in 2005, and are currently available, online as a pdf and web-based version, and as a free App for iOS and Android devices. The Guidelines are no longer produced as a printed booklet, but continue to be translated into several different languages. The Guidelines undergo formal minor revisions annually and major revisions every second year. Interim updates may however also be provide at any time the panels consider it necessary.

The aim of the EACS Guidelines is to provide easily accessible and comprehensive recommendations to clinicians involved in all aspects of care. Unless mentioned otherwise, they always refer to the specific management of people living with HIV.

The EACS Guidelines cover a relatively large and diverse area geographically, with different national levels of access to care. As a natural consequence, the Guidelines aim to cover a relatively wide range of recommendations as opposed to the often more uniform national guidelines.

The 2022 version of the Guidelines includes updates of all existing sections. In order to emphasise person-centered language, the abbreviation PLWH referring to people living with HIV has been deleted throughout the guidelines. The most essential changes are listed in the Summary of changes from v11.0 to v11.1

Each respective section of the Guidelines is managed by a panel of experienced European HIV experts, with additional experts in other fields of expertise included where necessary. All recommendations are evidence-based whenever possible and based on expert opinions in the rare instances where adequate evidence is unavailable. The Guidelines do not provide formal grades of evidence, panels make decisions by consensus or by vote when necessary and we do not publish results of the votes or discrepancies if any occur

The EACS Guidelines panels are overseen by a Guidelines Chair who serves a three-year term and is elected from the Governing Board. Each panel is led by a Panel Chair, supported by a Vice-Chair and a Young Scientist. The Co-Chair will take over the role of Chair after the Chair's term expires. Panel membership is reviewed annually and rotation is overseen by the Panel Leads and Guidelines Chair according to a standard operating procedure. Operational matters of the EACS Guidelines are led by a Coordinator in the Medical Secretariat, supported by the EACS Secretariat.

Only the latest and key references used to produce the Guidelines are provided in a separate section, see References. A short summary of the key findings of highlighted references is included.

Please reference the EACS Guidelines as follows: EACS Guidelines version 11.1. October 2022.

Video links to the EACS online course on Management of HIV and Co-infections are provided throughout the Guidelines, see Video links.

The diagnosis and management of HIV infection and related co-infections, opportunistic diseases and co-morbidities across all ages continue to require a multidisciplinary effort for which we hope the 2022 version of the EACS Guidelines will provide you with an easily accessible overview.

All comments to the Guidelines are welcome and can be directed to quidelines@eacsociety.org

We wish to warmly thank all panelists, external experts, linguists, translators, the EACS Secretariat, the Sanford team and everyone else who helped to build up and to publish the EACS Guidelines for their dedicated work.

Enjoy!

Georg Behrens and Juan Ambrosioni

October 2022



# Summary of Changes from v11.0 to v11.1

The COVID-19 situation is rapidly changing, and evidence is constantly accumulating. Therefore, we refer to the regularly updated BHIVA, DAIG, EACS, GESIDA & Polish Scientific AIDS Society Statement on risk of COVID-19 for https://www.eacsociety.org/home/covid-19-and-hiv.html

### **ART** section

- Initial Combination Regimen for ART-naïve Adults, pages 13-14
  - ABC should not be used for same day start
  - Precision that DOR has not been compared to an INSTI and was shown to be non-inferior to EFV and DRV
  - Specification that EFV should be used at 400 or 600 mg qd and that if rifampicin-based regimen for tuberculosis is used 600 mg must be
  - · DRV/r should be used with caution in persons with a high CVD risk Switch strategies for virologically suppressed persons, page 16

  - EVG/c and unboosted ATV have been removed from alternatives
  - Intermittent therapy remains a not recommended strategy, with new wording about QUATUOR
- Virological failure, page 17
  - TDF and TAF can be used in association with 3TC or FTC if genotype shows only limited NRTI mutation(s)
  - EVG/c has been removed form alternatives
- Treatment of pregnant women living with HIV or women considering pregnancy, page 18
  - New wording about DTG and neural tube defects
  - Reference to EVG/c has been removed
- ART and TB co-infection, page 20
  - ATV/r and LPV/r have been removed from combinations to use with rifabutin

### **DDI** section

- The antiviral drugs molnupiravir, nirmatrelvir/ritonavir, sotrovimab and the immunosuppressant drug infliximab have been added to the COVID-19 drug interaction table
- The antimycobacterial drugs linezolid and pretomanid have been implemented in the anti-tuberculosis drug interaction table. Furthermore, a footnote has been added for EFV to indicate that EFV should be dosed at 600 mg qd in presence of rifampicin but can be dosed at 400 mg or 600 mg qd in absence of rifampicin
- All tables have been updated to include changes implemented in the HIV drug interaction website (University of Liverpool) in the past year
- A footnote was notably added to the contraceptive table to indicate a higher risk of sub-therapeutic intramuscular medroxyprogesterone concentrations at week 12 in women with higher BMI on EFV treatment. Dosing medroxyprogesterone every 8-10 weeks in women with a higher BMI on EFV and particularly on EFV plus rifampicin prevent this risk

### Co-morbidity section

- The impact of comorbid mental health disorders on adherence to opiate substitution therapy and the use of fixed dose combination with naloxone to reduce risk of overdose with buprenorphine have been included in the Opioid Addiction, Pharmacological Treatment section, page 58
- A new resource for information on cancer drug interactions has been added to the Cancer: Treatment monitoring section, page 60
- Updated guidance on HBV, pneumococcal and SARS-CoV-2 vaccina-
- Updated guidance on management of varices, page 80
- Updated guidance on nutrition of cirrhotic persons and management of hepatic encephalopathy, page 81
- Updated dietary advice for the management of non-alcoholic fatty liver disease (NAFLD), page 82
- Updated guidance on management of Hepatorenal Syndrome Acute Kidney Injury (HRS-AKI), page 83

### **Viral Hepatitis Co-infections section**

- Hepatitis D and E infection:
  - Hepatitis D Virus:
    - 6. Bulevirtide (2mg/d s.c) in combination with TDF/TAF is recommended in HDV-RNA positive persons with compensated liver disease and should be used where available. The optimal duration of treatment remains unclear. Treatment should be performed in centers with sufficient experience

### Opportunistic Infections and COVID-19 section

- COVID-19 section has been extensively modified according to the updated evidences from literature, see page 139-140
- TB treatment guidelines have been reformulated according to the recently published updates from WHO, pages 135-136
- Results from a large clinical trial on treatment of cryptococcal meningitis in resource-limited settings have been added in the comments section of cryptococcal meningitis induction therapy, page 128
- A comment on the results of a clinical trial investigating addition of miltefosine to amphotericin B for visceral leishmaniasis has been added, page 134
- Recommendations for toxicity monitoring in TMP-SMX therapy have been added, pages 126-127
- Diagnostic recommendations for HSV and VZV infections have been reformulated, pages 130
- Recommendation for secondary prophylaxis discontinuation in CMV retinitis has been reformulated, page 131
- Minor stylistic changes were made throughout the text

### **Paediatric HIV Treatment section**

- Relevant toxicities to paediatric/adolescent ART have been added in table 1, page 142
- Minor edition in the other sections

EACS Guidelines are available online at http://www.eacsociety.org and in the EACS Guidelines App

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# **Abbreviations**

Antiret	troviral drug (ARV) abbrevia	tions	
3ТС	lamivudine	NNRTI	non-nucleoside reverse
ABC	abacavir		transcriptase inhibitors
ATV	atazanavir	NVP	nevirapine
BIC	bictegravir	PI	protease inhibitors
CAB	cabotegravir	Pl/b	protease inhibitors
COBI	cobicistat		pharmacologically
	(used as booster=/c)		boosted with cobicistat
d4T	stavudine		or ritonavir
ddl	didanosine	PI/c	protease inhibitor
DOR	doravirine		pharmacologically
DRV	darunavir		boosted with
DTG	dolutegravir		cobicistat
EFV	efavirenz	PI/r	protease inhibitors
EVG	elvitegravir		pharmacologically
ENF	enfuvirtide (T20)		boosted with ritonavir
ETV	etravirine	RAL	raltegravir
FI	fusion inhibitor	RPV	rilpivirine
FPV	fosamprenavir	RTV	ritonavir (used as
FTC	emtricitabine		booster=/r)
FTR	fostemsavir	SQV	saquinavir
IDV	indinavir	TAF	tenofovir alafenamide
INSTI	integrase strand transfer	TDF	tenofovir disoproxil
	inhibitor		fumarate
LPV	lopinavir	TPV	tipranavir
MVC	maraviroc	ZDV	zidovudine
NRTI	nucleos(t)ide reverse transcriptase inhibitors	XTC	3TC or FTC
	transcriptase irinibitors		

Oth			

Other an	bieviations		
ACEi	angiotensin converting enzyme inhibitor	CSF CTC	cerebrospinal fluid computed tomography
AFP	alpha-foetoprotein		colonoscopy
ALP	alkaline phosphatase	CVD	cardiovascular disease
ALT	alanine aminotransferase	CXR	chest X-ray
aMDRD	abbreviated modification	DAA	direct acting antiviral drug
	of diet in renal disease	DDI	drug-drug interaction
	formula	DPP-4i	dipeptidyl peptidase 4
ARB	angiotensin receptor		inhibitor
	blocker	DRESS	drug rash with
ART	antiretroviral therapy	DIVEGO	eosinophilia and systemic
AST	aspartate		symptoms
A01	aminotransferase	DXA	dual energy X-ray
ASCVD	atherosclerotic	אא	absorptiometry
ASCAD	cardiovascular disease	ECG	electrocardiogram
В			O .
B	buprenorphine	eGFR	estimated glomerular filtration rate
bid	twice daily	-01 B	
BMD	bone mineral density	ESLD	end stage liver disease
BMI	body mass index	FBC	full blood count
BP	blood pressure	FH	familial
CABG	coronary artery bypass		hypercholesterolaemia
	grafting	FIT	faecal immunochemistry
CAPD	continuous ambulatory		test
	peritoneal dialysis	FRAX®	fracture risk assessment
CAD	coronary artery disease		tool
cART	combination antiretroviral	FRAT	falls risk assessment
	treatment		tool
CBT	cognitive behavioural	FS	frailty scale
	therapy	GAD-2	generalized anxiety
CCB	calcium channel blocker		disorder 2-item screening
CGA	comprehensive geriatric		tool
	assessment	GDR	genotypic drug resistance
CKD	chronic kidney disease		test
CKD-EPI	CKD epidemiology	GLP1RA	glucagon like peptide 1
	collaboration formula		receptor agonist
CMV	Cytomegalovirus	GT	genotype
CNS	central nervous system	HAV	Hepatitis A virus
COPD	chronic obstructive	HAD	HIV-associated dementia
001 D	pulmonary disease	HBV	Hepatitis B virus
COVID-	Coronavirus disease	HCC	hepatocellular carcinoma
19	2019	HCV	Hepatitis C virus
13	2013	1104	riepaulis C virus

HDL-c	
	HDL-cholesterol
HDV	Hepatitis D virus
HEV	Hepatitis E virus
HF	heart failure
HIVAN	HIV-associated
HIV-VL	nephropathy HIV viral load (HIV-RNA)
HMOD	hypertension-mediated
THIOD	organ disease
HPV	Human papillomavirus
HRS	hepatorenal syndrome
HSR	hypersensitivity reaction
HSV	Herpes simplex virus
ICS	Inhaled corticosteroid
IFG	Impaired Fasting Glucose
IFN IGRA	interferon interferon-gamma release
IGRA	assay
IGT	impaired glucose
	tolerance
IHD	ischaemic heart disease
im	intramuscular
IRIS	immune reconstitution
	inflammatory syndrome
iv	intravenous
IVDU LA	intravenous drug use long-acting
LA LABA	long-acting β2-agonist
LAMA	long-acting muscarinic
	antagonist
LDL-c	LDL-cholesterol
LGV	lymphogranuloma
	venereum
LOQ	limit of quantification
MDR-TB Mg	multidrug resistant TB magnesium
MND	mild neurocognitive
WII LD	disorder
MRI	magnetic resonance
	imaging
MSM	men who have sex with
мтот	men
MTCT	mother to child transmission
МТ	multitarget
sDNA	stool DNA
MRI	magnetic resonance
	imaging
MX	methylxanthines
N	norbuprenorphine
NAFLD	non-alcoholic fatty liver
	non-alcoholic fatty liver disease
NAFLD NASH	non-alcoholic fatty liver disease non-alcoholic
	non-alcoholic fatty liver disease
NASH	non-alcoholic fatty liver disease non-alcoholic steatohepatitis non-steroidal anti- inflammatory
NASH NSAID NP	non-alcoholic fatty liver disease non-alcoholic steatohepatitis non-steroidal anti- inflammatory neuropsychological
NASH NSAID NP Ols	non-alcoholic fatty liver disease non-alcoholic steatohepatitis non-steroidal anti- inflammatory neuropsychological opportunistic infections
NASH NSAID NP	non-alcoholic fatty liver disease non-alcoholic steatohepatitis non-steroidal anti-inflammatory neuropsychological opportunistic infections orthotopic liver
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NASH NSAID NP OIS OLTX PAP PCI PD4 PEP PrEP PEG-IFN PHI po PPD	non-alcoholic fatty liver disease non-alcoholic steatohepatitis non-steroidal anti-inflammatory neuropsychological opportunistic infections orthotopic liver transplantation papanicolaou test percutaneous coronary intervention phosphodiesterase 4 inhibitors post-exposure prophylaxis pre-exposure prophylaxis pre-exposure prophylaxis pegylated-interferon primary HIV infection per oral purified protein derivative
NASH NSAID NP OIS OLTX PAP PCI PD4 PEP PrEP PEG-IFN PHI po PPD PPI	non-alcoholic fatty liver disease non-alcoholic steatohepatitis non-steroidal anti-inflammatory neuropsychological opportunistic infections orthotopic liver transplantation papanicolaou test percutaneous coronary intervention phosphodiesterase 4 inhibitors post-exposure prophylaxis pre-exposure prophylaxis pre-exposure prophylaxis pegylated-interferon primary HIV infection per oral purified protein derivative proximal renal tubulopathy prostate specific antigen
NASH NSAID NP OIS OLTX PAP PCI PD4 PEP PEG-IFN PHI po PPD PPI PRT	non-alcoholic fatty liver disease non-alcoholic steatohepatitis non-steroidal anti-inflammatory neuropsychological opportunistic infections orthotopic liver transplantation papanicolaou test percutaneous coronary intervention phosphodiesterase 4 inhibitors post-exposure prophylaxis pre-exposure prophylaxis pre-exposure prophylaxis pegylated-interferon primary HIV infection per oral purified protein derivative proton pump inhibitor proximal renal tubulopathy prostate specific antigen proprotein convertase
NASH NSAID NP OIS OLTX PAP PCI PD4 PEP PEG-IFN PHI po PPD PPI PRT PSA PCSK9	non-alcoholic fatty liver disease non-alcoholic steatohepatitis non-steroidal anti-inflammatory neuropsychological opportunistic infections orthotopic liver transplantation papanicolaou test percutaneous coronary intervention phosphodiesterase 4 inhibitors post-exposure prophylaxis pre-exposure prophylaxis pre-exposure prophylaxis pegylated-interferon primary HIV infection per oral purified protein derivative proton pump inhibitor proximal renal tubulopathy prostate specific antigen proprotein convertase subtilisin/kexin type 9
NASH NSAID NP OIS OLTX PAP PCI PD4 PEP PEG-IFN PHI po PPD PPI PRT PSA PCSK9 PTH	non-alcoholic fatty liver disease non-alcoholic steatohepatitis non-steroidal anti-inflammatory neuropsychological opportunistic infections orthotopic liver transplantation papanicolaou test percutaneous coronary intervention phosphodiesterase 4 inhibitors post-exposure prophylaxis pre-exposure prophylaxis pre-exposure prophylaxis pegylated-interferon primary HIV infection per oral purified protein derivative proton pump inhibitor proximal renal tubulopathy prostate specific antigen proprotein convertase subtilisin/kexin type 9 parathyroid hormone
NASH NSAID NP OIS OLTX PAP PCI PD4 PEP PEG-IFN PHI po PPD PPI PRT PSA PCSK9	non-alcoholic fatty liver disease non-alcoholic steatohepatitis non-steroidal anti-inflammatory neuropsychological opportunistic infections orthotopic liver transplantation papanicolaou test percutaneous coronary intervention phosphodiesterase 4 inhibitors post-exposure prophylaxis pre-exposure prophylaxis pre-exposure prophylaxis pegylated-interferon primary HIV infection per oral purified protein derivative proton pump inhibitor proximal renal tubulopathy prostate specific antigen proprotein convertase subtilisin/kexin type 9

qid	four times daily
RAS	resistance-associated substitutions
RBV	ribavirin
RCT	randomized controlled trial
RIG	Rabies Immunoglobulin
SARS-	Severe Acute Respiratory
CoV-2	Syndrome Coronavirus-2
SABA	short-acting β2-agonist
SAMA	short-acting muscarinic
	antagonist
sc	subcutaneous
SCORE	systemic coronary risk
	estimation
SGLT-2i	sodium/glucose co-
	transporter 2 inhibitor
SOT	solid organ transplant
SPPB	short physical
	performance battery
SSRI	selective serotonin-
	reuptake inhibitor
STI	sexually transmitted
011	infection
SU SVR	sulfonylurea
SVK	sustained virological response
TBS	trabecular bone score
TC	total cholesterol
TDM	therapeutic drug
I DIVI	monitoring
TG	triglycerides
TIA	transient ischaemic attack
tid	three times daily
TMP-	trimethoprim-
SMX	sulfamethoxazole
TZD	thiazolidinediones
UA/C	urine albumin/creatinine
	ratio
UP/C	urine protein/creatinine
	ratio
US	ultrasound
VL	viral load (HIV-RNA)
VZV	varicella-zoster virus
WB	western blot
XDR-TB	extensively drug-
7	resistant TB
Zn	zinc

# Part I Assessment of Initial & Subsequent Visits

	Assessment	At HIV diagnosis	Prior to starting ART	Follow-up frequency	Comment	See page
HISTORY						
Medical	Complete medical history including:	+	+	First visit	On transfer of care repeat assessment	
	Family history (e.g. premature CVD, diabetes, hypertension, CKD)	+		First visit	Premature CVD: cardiovascular events in a first degree relative (male < 55, female < 65 years)	62, 63-64
	Concomitant medicines (i)	+	+	Every visit		
	Past and current co-morbidities	+	+	Every visit		
	Vaccination history	+		Annual	Measure antibody titres and offer vaccinations where indicated, see Vaccination	
Psychosocial	use, smoking, diet, exercise, drug use)  frequently	Adverse lifestyle habits should be addressed more frequently	61			
	Employment	+	+		Provide advice and support if needed	
	Social and welfare	+	+	Every visit	Provide counselling if needed	
	Psychological morbidity	+	+			
	Partner and children	+			Test partner and children if at risk	
Sexual and	Sexual history	+			Address issues concerning sexual dysfunction	91-95
Reproductive	Safe sex	+			Risk of sexual transmission should be addressed	
Health	Partner status and disclosure	+		6-12 months	Recommend starting ART in serodifferent couples	
	Conception issues	+	+			
	Hypogonadism	+	+	As indicated	Persons with complaints of sexual dysfunction	91, 94
	Menopause	+	+	Annual/ as indicated	Screen for perimenopause symptoms in women ≥ 40 years	91
HIV DISEASE						
Virology	Confirmation of HIV Ab pos	+			More frequent monitoring of HIV-VL at start of ART	12-14
virology	Plasma HIV-VL	+	+	3-6 months	Perform genotypic resistance test before starting ART if not previously tested or if at risk of super-infection	
	Genotypic resistance test and sub-type	+	+/-	At virological failure		
	R5 tropism (if available)		+/-		Screen if considering R5 antagonist in regimen	
Immunology	CD4 absolute count and %, CD4/CD8 ratio (optional: CD8 and %)	+	+	3-6 months	Annual CD4 count if stable on ART and CD4 count > 350 cells/µL (II) CD4/CD8 ratio is a stronger predictor of serious outcomes	12-14
	HLA-B*57:01 (if available)	+	+/-		Screen before starting ABC containing ART, if not previously tested, pages 12-13, 24	
CO-INFECTIONS						
STIs	Syphilis serology	+		Annual/ as indicated	Consider more frequent screening if at risk	15, 91
	STI screen	+		Annual/ as indicated	Screen if at risk and during pregnancy	
Viral Hepatitis	HAV screen	+			Screen if ongoing risk (e.g. MSM); vaccinate if non-immune	- 90, 115-
	HBV screen	+	+	As indicated	Annual screen if ongoing risk; vaccinate if non-immune. Use ART containing TDF or TAF in vaccine non-responders	117
	HCV screen	+			Further screen based on risk behaviour and local epidemiology.  Measure HCV-RNA if HCV Ab pos or if recently acquired infection suspected	
	HDV screen			As indicated	All Persons with positive HBs-Ag should also be screened for HDV co-infection	115, 122
	HEV screen			As indicated	Screen persons with symptoms consistent with acute hepatitis, unexplained flares of aminotransferases or elevated liver function tests, neuralgic amyotrophy, Guillain-Barré, encephalitis or proteinuria. Include anti-HEV IgG and IgM and NAAT for HEV-RNA in blood and if possible in stool	122



	Assessment	At HIV diagnosis	Prior to starting ART	Follow-up frequency	Comment	See page
Tuberculosis	CXR	+			Consider routine CXR in persons from high TB	20,
	PPD	+			prevalence populations.	135
	IGRA in selected high-risk	+		Re-screen if	Some national guidelines consider the ethnicity,	
	populations (if available)	T		exposure	CD4 count and ART usage to define indication for latent tuberculosis infection screening.  Use of PPD/IGRA depending on availability and	
					local standard of care. IGRA should, however, be tested before PPD if both are to be used, given the potential for a false positive IGRA after PPD. See Diagnosis and Treatment of TB	
Others	Varicella zoster virus serology	+			Offer vaccination where indicated	90
	Measles/Rubella serology	+			Offer vaccination where indicated	
	Toxoplasmosis serology	+				
	CMV serology	+				
	Cryptococcus antigen	+/-			Consider screening for cryptococcus antigen in serum in persons with CD4 count < 100 cells/µL	
	Leishmania serology	+/-			Screen according to travel history/origin	
	Tropical screen (e.g. Schistosoma serology)	+/-			Screen according to travel history/origin	
	Influenza virus	+		Annual	In all persons with HIV, see Vaccination	90
	Streptococcus pneumoniae	+			No recommendations available regarding the need for a booster dose, see Vaccination	90
	Human papilloma virus	+		As indicated	Vaccinate all persons with HIV with 3 doses be-	90
					tween ages 9 and 40. If HPV infection is established, efficacy of vaccine is questionable, see Vaccination	
	SARS-CoV-2				In a pandemic situation, vaccinate irrespective of CD4 count and HIV-VL according to national Guidelines	90
CO-MORBIDITIES	3					
Haematology	FBC	+	+	3-12 months		
0,	Haemoglobinopathies	+			Screen at risk persons	
	G6PD	+			Screen at risk persons	
Body Composition	Body-mass index	+	+	Annual		61
Cardiovascular Disease	Risk assessment (Framingham score (iii))	+	+	Annual	Should be performed in all men > 40 years and women > 50 years without CVD	62
	ECG	+	+/-	As indicated	Consider baseline ECG prior to starting ARVs associated with potential conduction problems	
Hypertension	Blood pressure	+	+	Annual		63-6
Lipids	TC, HDL-c, LDL-c, TG <sup>(iv)</sup>	+	+	Annual	Repeat in fasting state if used for medical intervention (i.e. $\geq$ 8 h without caloric intake)	69
Glucose	Serum glucose	+	+	Annual	Consider oral glucose tolerance test / HbA1c if fasting glucose levels of 5.7-6.9 mmol/L (100-125 mg/dL)	67-6
Pulmonary Disease	Respiratory symptoms and risk factors	+	+	Annual	If severe shortness of breath is reported with preserved spirometry, echocardiography may be performed to rule out heart failure and/or pulmonary hypertension	105
	Spirometry			As indicated	Spirometry should be performed in all symptomatic persons (xir)	
Liver Disease	Risk assessment <sup>(v)</sup>	+	+	Annual		79-8
	ALT/AST, ALP, Bilirubin	+	+	3-12 months	More frequent monitoring prior to starting and on treatment with hepatotoxic drugs	
	Staging of liver fibrosis			12 months	In HCV and/or HBV co-infected persons (e.g. FibroScan, serum fibrosis markers)	79-8
	Hepatic ultrasound			6 months	Persons with liver cirrhosis (Xiii)	79-8
Renal Disease	Risk assessment(vi)	+	+	Annual	More frequent monitoring if eGFR < 90 mL/min,	74-7
	eGFR (CKD-EPI)(vii)	+	+	3-12 months	CKD risk factors present <sup>(v)</sup> and/or prior to starting and on treatment with nephrotoxic drugs <sup>(x)</sup>	
	Urine dipstick analysis(viii)	+	+	Annual	Every 6 months if eGFR < 60 mL/min or rapid decline in eGFR, if proteinuria ≥ 1+ and/or eGFR < 60 mL/min perform UA/C or UP/C <sup>(viii)</sup>	
Bone Disease	Bone profile: calcium, PO <sub>4</sub> , ALP	+	+	6-12 months		71-7
	Risk assessment <sup>(x)</sup> (FRAX <sup>®(xi)</sup> in persons > 40 years)	+	+	2 years	Consider DXA in specific persons, see page 71 for details	
Vitamin D	25(OH) vitamin D	+		As indicated	Screen at risk persons	72



	Assessment	At HIV diagnosis	Prior to starting ART	Follow-up frequency	Comment	See page
Cognitive impairment	Screening questionnaire	+	+	As indicated	Screen all persons without highly confounding conditions. If abnormal or symptomatic, see algorithm page 104 for further assessment.	104
Anxiety	Questionnaire	±	±	As indicated	Screen at risk persons	100- 101
Depression	Questionnaire	+	+	As indicated	Screen at risk persons	96-97
Older persons	Polypharmacy review			Annual	Perform periodic medicines review	108- 109
	Frailty			Annual	Screen with Frail Scale, Walking Speed or short physical performance battery	110- 111
	Falls			Annual		112
Cancer	Mammography			1-3 years	Women 50-74 years	59
	Cervical PAP or liquid based cytology			1-3 years	women with HIV > 21 years	
	Rectal exam and anoscopy			1-3 years	MSM and persons with HPV-associated dysplasia. Evidence of benefit not known	
	Ultrasound and alpha-foe-toprotein			6 months	Controversial; persons with cirrhosis and persons with HBV co-infection at high risk of HCC <sup>(xiii)</sup>	
	Prostate cancer (PSA)			1-2 years	Men > 50 years with a life expectancy >10 years	
	Others			As indicated	Lung cancer and colorectal cancer screening according to local screening programmes	

If a person has been stable on ART for 6 months or more, with no other significant issues, clinicians could consider using alternative modalities such as email/phone/or other electronic means (Good practice point, GPP).

This form of consultation can have the same validity as a face-to-face consultation if properly instituted in a clinical protocol.

The European Union funded EmERGE project is currently looking at such interventions https://www.emergeproject.eu

- Review all concomitant medicines which may potentially interact with ARVs or increase co-morbidities, see
  - Drug-drug Interactions between Analgesics and ARVs
    Drug-drug Interactions between Anticoagulants/Antiplatelet Agents
    and ARVs
  - Drug-drug Interactions between Antidepressants and ARVs Drug-drug Interactions between Antihypertensives and ARVs
  - Drug-drug Interactions between Anti-malarial Drugs and ARVs Drug-drug Interactions between Anti-tuberculosis Drugs and ARVs
  - Drug-drug Interactions between Anxiolytics and ARVs
    Drug-drug Interactions between Bronchodilators (for COPD) and
  - Drug-drug Interactions between Contraceptives and ARVs
    Drug-drug Interactions between Corticosteroids and ARVs
    Drug-drug Interactions between COVID-19 Therapies and ARVs
    Drug-drug Interactions between Hormone Replacement Therapy
  - (HRT) and ARVs Drug-drug Intercations between Immunosuppressants (for SOT) and ARVs
  - Drug-drug Interactions between Pulmonary Antihypertensives and
  - Drug-drug Interactions between Viral Hepatitis Drugs and ARVs and http://www.hiv-druginteractions.org
- ii If stable on ART with undetectable HIV-VL and CD4 count > 350 cells/ μL, suggest annual CD4 count
- iii A risk equation developed from HIV populations is available, see https://www.chip.dk/Tools-Standards/Clinical-risk-scores.Of note, if an individual receives medicines to control dyslipidaemia and/or hypertension, the estimation should be interpreted with caution
- iv A calculator for LDL-cholesterol in cases where TG is not high can be found at https://www.mdcalc.com/ldl-calculated
- Risk factors for chronic liver disease include alcohol, viral hepatitis, obesity, diabetes, insulin resistance, hyperlipidaemia and hepatotoxic drugs.
- vi Risk factors for CKD: hypertension, diabetes, CVD, family history, black African ethnicity, viral hepatitis, low current CD4 count, smoking, older age, concomitant nephrotoxic drugs
- vii eGFR: use CKD-EPI formula based on serum creatinine, gender, age and ethnicity because eGFR quantification is validated > 60 mL/min. The abbreviated modification of diet in renal disease (aMDRD) or the Cockroft-Gault (CG) equation may be used as an alternative, see https://www.chip.dk/Tools-Standards/Clinical-risk-scores

- viii Some experts recommend UA/C (urinary albumin creatinine ratio) or UP/C (urinary protein creatinine ratio) as a screening test for proteinuria in all persons. UA/C predominantly detects glomerular disease. Use in persons with diabetes. UP/C detects total protein secondary to glomerular and tubular disease and can be used for screening for ARV toxicity, page 75
- ix Different models have been developed for calculating a 5-year CKD risk score while using different nephrotoxic ARVs, integrating HIV independent and HIV-related risk factors
- x Classic risk factors: older age, female gender, hypogonadism, family history of hip fracture, low BMI (≤ 19 kg/m²), vitamin D deficiency, smoking, physical inactivity, history of low impact fracture, alcohol excess (> 3 units/day), steroid exposure (minimum 5 mg for > 3 months)
- Xi WHO fracture risk assessment (FRAX®) tool: http://www.shef.ac.uk/
- xii Respiratory symptoms: shortness of breath, chronic cough and sputum. Risk factors: tobacco, occupation, in- and outdoor pollution and host factors including previous PCP or TB, recurrent pneumonia and Alpha-1 antitrypsin deficiency. A diagnosis of COPD should be considered in persons over the age of 35 years who have a risk factor (current or ex-smoker) and who present with exertional breathlessness, chronic cough, regular sputum production, frequent winter 'bronchitis' or wheeze
- kiii HCC screening is indicated in all cirrhotic HBV or HCV co-infected persons (even if HCV infection has been cured and HBV replication is medically suppressed) in a setting where treatment for HCC is available. Although the cost-effectiveness of HCC screening in persons with F3 fibrosis\* is uncertain, surveillance may be considered based on an individual risk assessment (https://easl.eu/publications/clinical-practice-guidelines/). In HBV-positive non-cirrhotics, HCC screening should follow current EASL guidelines. Risk factors for HCC in this population include family history of HCC, ethnicity (Asians, Africans), HDV and age > 45 years. EASL guidelines propose using the PAGE-B score in Caucasians to assess the HCC risk, however this score has not been validated in persons with HIV, see pages 59, 81 and 115
- \* See table on cut-off values of non-invasive tests for the detection of significant fibrosis and cirrhosis, page 121. The combination of blood biomarkers, the combination of liver stiffness measurement and blood tests or repeated assessments may improve accuracy, see EASL recommendations on treatment of Hepatitis C 2020 - EASL-The Home of Hepatology (free registration needed to get access)

# Part II ART

This section provides an overview of the important aspects of ART management. Recommendations are based on a range of evidence, in particular it is weighted towards randomised controlled clinical trials. Other data have been taken into account, including cohort studies, and where evidence is limited, the panel has reached a consensus around best clinical practice. The ART section is wide ranging and, with the recommendation to start therapy independently of CD4 count, there is an important section on readiness to start. Treatment recommendations are based on drugs licensed in Europe and range from initial therapy through to switching with or without virological failure. Two important areas of ART are highlighted: pregnancy and TB. Details on the use of PrEP, which is being rolled out across Europe, are also included.

# Assessing Readiness to Start and Maintain ART(1)

### Goal: to help persons start and/or maintain ART

Starting ART is recommended for all persons with HIV regardless of CD4 count to reduce the morbidity and mortality associated with HIV infection, and to prevent HIV transmission (START and TEMPRANO trials, HPTN 052, PARTNER Study). Evidence is accumulating that starting ART on the same day after establishing a diagnosis of HIV infection is feasible and acceptable for newly-diagnosed individuals. Nevertheless, assessment of the readiness to start ART is essential to allow to express the person's preference and not feel pressured to start ART immediately, unless clinically indicated

Given the need for lifelong treatment, successful ART requires a person's readiness to start and adhere to the regimen in a sustained manner. The trajectory from problem awareness to maintenance on ART can be divided into five stages. Knowing a person's stage, health care providers use appropriate techniques to assist them to start and maintain ART

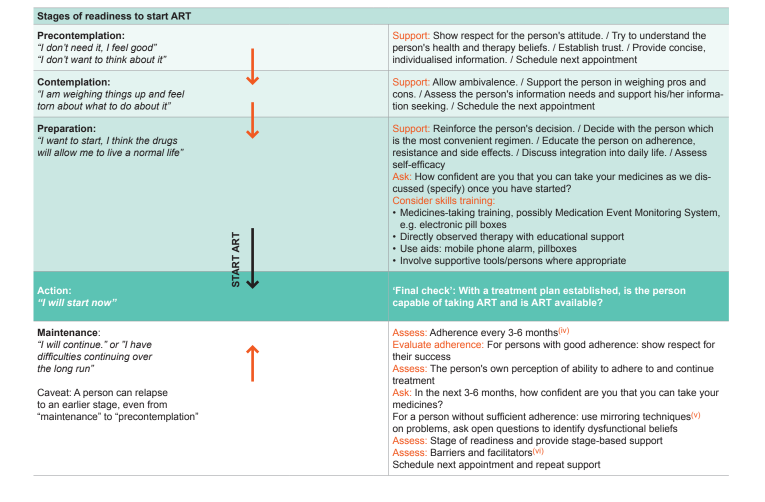
Identify the person's stage of readiness using WEMS<sup>(ii)</sup> techniques, and start discussion with an open question/invitation:

"I would like to talk about HIV medicines." <wait> "What do you think about them?"

Based on the person's response, identify his/her stage of readiness and intervene accordingly(iii)

Immediate (i.e. same day) start of ART should be considered, and especially in the following situations:

- In the setting of primary HIV infection, especially in case of clinical signs and symptoms of meningoencephalitis (within hours). In this situation, the clinician may start ART immediately after a positive screening HIV test and before obtaining confirmatory HIV test results such as a HIV-VL
- The wish to start ART immediately
- In a setting where loss-to-follow-up is more likely if ART is not started the same day



# Several barriers are known to influence ART decision making and adherence to $\ensuremath{\mathsf{ART}}$

Screen for and talk about problems and facilitators

Consider systematic assessment of:

- Depression(vii), see pages 96-97
- Cognitive problems<sup>(viii)</sup>, see page 104
- Harmful alcohol<sup>(ix)</sup> or recreational drug use, see page 58

Consider talking about:

- Social support and disclosure
- Health insurance and continuity of drug supply
- · Therapy-related factors

Recognise, discuss and reduce problems wherever possible in a multidisciplinary team approach

- i Algorithm adapted from Fehr et al.
- ii WEMS: Waiting (> 3 sec), Echoing, Mirroring, Summarising
- iii The person presenting in the clinic may be at different stages of readiness: precontemplation, contemplation or preparation. The first step is to assess the stage, and then to support/intervene accordingly. In the case of late presentation (CD4 count < 350 cells/µL), the initiation of ART should not be delayed. The person should be closely followed and optimally supported. Schedule the next appointment within a short time, i.e. 1-2 weeks
- iv Suggested adherence questions: "In the past 4 weeks, how often have you missed a dose of your HIV medicines: every day, more than once a week, once a week, once every 2 weeks, once a month, never?" /

- "Have you missed more than one dose in a row?"
- Mirroring: reflecting back on what a person has said or non-verbally demonstrated (e.g. anger or disappointment) WITHOUT introducing new material by asking questions or giving information
- vi Adherence to long-term therapies
- vii See Mental Health section, Depression: Screening and Diagnosis
  Meta-analysis shows a consistent relationship between depression and
  ART non-adherence that is not limited to those with clinical depression.
  Therefore, assessment and intervention aimed at reducing depressive
  symptom severity, even at subclinical level is important.
- viii See Algorithm for Diagnosis and Management of Cognitive Impairment in persons without Obvious Confounding Conditions
- ix FAST-alcohol use, ask: How often have you had 6 or more units if female, or 8 or more units if male, on a single occasion in the last year? Never = 0, Less than monthly = 1, Monthly = 2, Weekly = 3, Daily or almost daily = 4. Stop if the answer is 0 (Never). Ask more questions if the answer is 1, 2, 3 or 4.



# Recommendations for Initiation of ART in persons with Chronic Infection without Prior ART Exposure<sup>(1)</sup>

Recommendations take into account the level of evidence, the degree of progression of HIV disease and the presence of, or high risk for, developing various types of (co-morbid) conditions.

ART is recommended in all adult persons with HIV, irrespective of CD4 counts<sup>(1)</sup>

- ART is recommended irrespective of the CD4 count. In certain situations (i.e lower CD4 count or pregnancy), there is a greater urgency to start ART immediately
  - In persons with OIs, ART initiation may have to be deferred, see page 123, for ART initiation in the presence of specific OIs. For ART initiation in persons with TB, see page 20
  - A possible exception to immediate start of ART might be HIV controllers, persons with high CD4 counts and HIV-VL < 1000 copies/mL, although even in such persons ART initiation has been shown to increase CD4 count, decrease inflammation, lower the risk of clinical events and prevent HIV transmission</li>
  - Genotypic resistance testing is recommended prior to initiation of ART, ideally at the time of HIV diagnosis. Genotypic testing should not delay ART initiation (it may be re-adjusted after genotypic test results)
  - If ART needs to be initiated before genotypic testing results are available, it is recommended to select a first-line regimen with a high barrier to resistance, including a PI/b or second generation INSTI
  - Whether rapid, possibly same-day ART start is proposed to newly diagnosed persons or postponed until complementary assessments depends on the setting and medical circumstances, medical indications to start ART more urgently and risk of loss from care. To reduce loss to follow-up between diagnosis and ART initiation, structural barriers delaying the process should be addressed



# **Initial Combination Regimen for ART-naïve Adults**

Before selecting an ART regimen, it is critical to review:

- If a woman wishes to conceive or is pregnant: Treatment of Pregnant Women Living with HIV or Women Considering Pregnancy
- · If the person has an opportunistic infection: Initiation of ART regimen in persons with opportunistic infections
- If the person has **TB**: Antiretroviral regimens in TB/HIV co-infection
- If the person has potential treatment limiting comorbidities: Comorbidity section, dose adjustment for renal and liver impairment
- If the person is treated with other medications: Drug-drug interactions
- If the person has Swallowing Difficulties: Administration of ARVs in persons with swallowing difficulties
- If the person has acquired HIV while receiving PrEP: In this situation, change PrEP to a triple-drug ART regimen including a third drug with a high barrier to resistance (preferably DRV/b, DTG or BIC) plus two nucleoside analogues without interrupting antiretrovirals. The danger of acute seroconversion syndrome and higher infectiousness would be arguments for immediate switch to triple therapy. ART should be adjusted if more extensive resistance is demonstrated by genotypic resistance analysis
- Only drugs currently licensed for initiation of therapy by the EMA are included (in alphabetical order)
- Recommended regimens should be considered first and are preferable for most persons. Antiretroviral drugs in the Recommended category provide a
  combination of essential characteristics for an optimal treatment such as long-term efficacy, barrier to resistance, safety, tolerability and few drug-drug
  interactions. Alternative regimens should be considered if recommended regimens are not feasible
- An increasing number of generic HIV drugs are now available, and their use can lead to large cost savings. The use of generic forms of drugs included in recommended regimens should therefore be encouraged, even if single tablet regimens are not used, as recent studies have shown similar virologic outcomes in ART-naïve persons receiving either a single pill or two pills qd
- Tailoring antiretroviral regimens for each individual is essential in the presence of resistance
- For a wider review of possible drug-related adverse events, please see: Adverse Effects of ARVs and Drug Classes

Regimen	Main requirements	Additional guidance (see footnotes)
Recommended regimens		
2 NRTIs + INSTI		
ABC/3TC + DTG ABC/3TC/DTG	HLA-B*57:01 negative HBsAg negative	I (ABC: HLA-B*57:01, cardiovascular risk) II (Weight increase (DTG))
TAF/FTC/BIC		II (Weight increase (BIC, TAF))
TAF/FTC or TDF/XTC + DTG		<ul><li>(Weight increase (DTG, TAF))</li><li>(TDF: prodrug types. Renal and bone toxicity. TAF dosing)</li></ul>
TAF/FTC or TDF/XTC + RAL qd or bid		<ul> <li>(Weight increase (RAL, TAF))</li> <li>(TDF: prodrug types. Renal and bone toxicity. TAF dosing)</li> <li>(RAL: dosing)</li> </ul>
1 NRTI + INSTI		
XTC + DTG or 3TC/DTG	HBsAg negative HIV-VL < 500,000 copies/mL Not recommended after PrEP failure	<ul><li>(Weight increase (DTG))</li><li>(3TC/DTG not after PrEP failure)</li></ul>
2 NRTIs + NNRTI		
TAF/FTC or TDF/XTC + DOR or TDF/3TC/DOR		<ul> <li>(Weight increase (TAF))</li> <li>(TDF: prodrug types. Renal and bone toxicity. TAF dosing)</li> <li>(DOR: caveats, HIV-2)</li> </ul>
Alternative regimens		
2 NRTIs + NNRTI		
TAF/FTC or TDF/XTC + EFV or TDF/FTC/EFV	At bedtime or 2 hours before dinner	<ul> <li>(Weight increase (TAF)</li> <li>(TDF: prodrug types. Renal and bone toxicity. TAF dosing)</li> <li>(EFV: neuro-psychiatric adverse events. HIV-2 or HIV-1 group 0, dosing)</li> </ul>
TAF/FTC or TDF/XTC + RPV or TAF/FTC/RPV or TDF/FTC/RPV	CD4 count > 200 cells/µL HIV-VL < 100,000 copies/mL Not on gastric pH increasing agents With food	(Weight increase (TAF))   (TDF: prodrug types. Renal and bone toxicity.   TAF dosing)   VIII (RPV: HIV-2)
2 NRTIs + PI/r or PI/c		
TAF/FTC or TDF/XTC + DRV/c or DRV/r or TAF/FTC/DRV/c	With food	(Weight increase (TAF))   (TDF: prodrug types. Renal and bone toxicity.   TAF dosing)   X (DRV/r: cardiovascular risk)   X (Boosted regimens and drug-drug interactions)



### **Additional Guidance**

- ABC contraindicated if HLA-B\*57:01 positive, not to be used for same day start. Even if HLA-B\*57:01 negative, counselling on HSR risk still mandatory. ABC should be used with caution in persons with a high CVD risk (> 10%), page 62
- II Treatment with INSTIs or TAF may be associated with weight increase
- III In certain countries, TDF is labelled as 245 mg rather than 300 mg to reflect the amount of the prodrug (tenofovir disoproxil) rather than the fumarate salt (tenofovir disoproxil fumarate). There are available generic forms of TDF, which instead of fumarate use phosphate, maleate, and succinate salts. They can be used interchangeably

When available, combinations containing TDF can be replaced by the same combinations containing TAF. TAF is used at 10 mg when coadministered with drugs that inhibit P-gp, and at 25 mg when coadministered with drugs that do not inhibit P-gp

The decision whether to use TDF or TAF depends on individual characteristics as well as availability

If the ART regimen does not include a booster, TAF and TDF have a similar short-term risk of renal adverse events leading to discontinuation and bone fractures

TAF\*\*\* should be considered as a first choice\*\*\*\* over TDF in individuals with:

- established or high risk of CKD, see page 74;
- coadministration of medicines with nephrotoxic drugs or prior TDF toxicity, see page 75;
- osteoporosis / progressive osteopenia, high FRAX score or risk factors, see page 71;
- history of fragility fracture, see pages 71 and 73
- V RAL can be given as RAL 400 mg bid or RAL 1200 mg (two, 600 mg tablets) qd. Note: RAL qd should not be given in presence of an inducer (i.e. TB drugs, antiepileptics) or divalent cations (i.e. calcium, magnesium, iron), in which case RAL should be used bid
- V HIV infections occurring in the context of PrEP failure may be associated with resistance-associated mutations. 3TC/DTG may be used in in this context only if there is no documented resistance in genotypic test
- VI DOR is not active against HIV-2. DOR has not been compared to an INSTI and was shown to be non inferior to EFV and DRV. There is risk of resistance associated mutations in case of virological failure. Results of genotypic resistance test are necessary before starting DOR
- VII EFV: not to be given if history of suicide attempts or mental illness; 400 or 600mg daily should be used; if rifampicine based regimen for tuberculosis is used, 600 mg dosing must be used; not active against HIV-2 and HIV-1 group O strains
- VIII RPV is not active against HIV-2
- IX A single large study has shown increase in CVD risk with cumulative use of DRV/r, not confirmed in other studies. DRV/r should be used with caution in persons with a high risk of cardiovascular risk
- X Boosted regimens with RTV or COBI are at higher risk of drug-drug interactions, see Part III Drug-drug interactions
- \*\*\* There are limited data on use of TAF with eGFR < 10 mL/min
- \*\*\*\* Expert opinion pending clinical data

# **Primary HIV Infection (PHI)**

### Definition of PHI(i-iv)

- · High-risk exposure within previous 6 weeks, and
- · Detectable virus in plasma (p24 Ag and/or HIV-RNA) and/or
- Evolving anti-HIV antibody reactivity (negative or indeterminate to positive)
- · With or without clinical symptoms

### Classification of PHI(i-v)

- Acute infection: HIV detection (p24 Ag and/or HIV-RNA) in the absence of HIV antibody
- · Recent infection: HIV antibody detection; up to 6 months after infection
- Where available, Western Blot (WB) or Immunoblot patterns of reactivity can be used to stage the infection as follows:
  - Stage I: HIV-RNA positive only (average duration 5 days)
     HIV-VL levels are median 2,000 copies/mL (IQR 300-20,000 copies/mL),
     and are < 100 copies/mL in approximately 10% of cases.</li>
     Low HIV-VL levels should be interpreted with caution due to the risk of false positivity
  - Stage II: HIV-RNA and p24 Ag positive only (average duration 5.3 days)
     HIV-VL levels are usually > 10,000 copies/mL
  - Stage III: HIV-RNA, p24 Ag and anti-HIV antibody positive by immune assay, no specific WB bands (average duration 3.2 days)
  - Stage IV: as Stage III but indeterminate WB pattern (5.6 days)
  - Stage V: as Stage III, but reactive WB pattern lacking p31 reactivity (average duration 69.5 days)
  - Stage VI: as stage III but full WB reactivity including a p31 band (indefinite)

### Starting treatment

Treatment of PHI is recommended for all cases

The recommendation is based on:

- Improvement of clinical symptoms of PHI, when present, especially severe general symptoms and/or neurological disease
- Benefits of early therapy:
- virological: decrease of the HIV-VL set-point and size of the viral reservoir; reduction of viral genetic evolution
- immunological: decrease of immune activation and inflammation; preservation of immune function and integrity of lymphoid tissue; possibly neurological and gut protection; possibly enhancement of post-treatment control and response to future eradication strategies
- Usually short interval between identification of PHI and a CD4 count < 500 cells/µL</li>
- Potential benefits of treatment for the community: reduced risk of transmission. Most infections are transmitted by persons who are unaware of their HIV status
- Reduced anxiety and facilitated disclosure to contacts
   The person should be counselled on indications and benefits of starting treatment as soon as possible, despite absence of demonstrated improved long-term clinical benefits<sup>(v)</sup>
  - Once treatment is started, it should be continued. A subsequent interruption is not recommended

### **Treatment selection**

- The person should preferably be recruited into a clinical trial or studies investigating HIV curative strategies
- Any use of PrEP or PEP should be identified and taken into account when choosing the initial regimen
- A drug resistance test is recommended in all cases as soon as possible after diagnosis
- Therapy may have to start before the results of resistance testing become available. In such cases, preference should be given to starting a PI/b or an INSTI with high barrier to resistance (DTG or BIC), in order to increase the barrier to resistance of the overall regimen. More than three active drugs are not needed.

A potential advantage for selecting DTG or BIC is the faster VL suppression. The benefit of combining PI/b with INSTI has not been shown. A combination of TDF or TAF, FTC, and either DRV/b, DTG or BIC, should therefore be considered, and the regimen adjusted, if needed, once the resistance test becomes available and viral load suppression is achieved. Where such a regimen is not available, national epidemiological data on prevalence and patterns of transmitted drug resistance (where available and sufficiently representative) may assist with the treatment selection process.

### Other considerations

- All newly infected persons should undergo investigations to diagnose sexually transmitted infections (e.g. syphilis, gonorrhoea, chlamydia), HBV, HCV and HPV, pages 7-9. Antibody seroconversion can be delayed and tests to identify the viral RNA are required in order to identify a recent HCV infection, page 120
- All women living with HIV of reproductive age should have a pregnancy test
- All persons should be counselled about the high risk of transmission, preventive measures, and importance of notifying partners
- i HIV-1 RNA becomes detectable in plasma around day 11 after exposure, approximately 7 days before p24 Ag and 12 days before anti-HIV antibodies
- ii Everyone with detectable HIV-VL and negative or indeterminate serology must receive confirmation of anti-HIV antibody seroconversion in follow-up testing. The interval of testing (up to stage V) is one week
- iii Some centres may have access to sero-incidence markers (e.g., anti-body avidity testing) that identify an infection acquired within the previous 3-6 months. Assay reliability varies and results should be interpreted with caution when they are the sole indicators of a recent infection
- iv A small subset of persons can spontaneously control the infection without treatment (elite controllers)
- Post-treatment controllers. A small proportion of recently-infected persons have been able to spontaneously control HIV-infection following ART discontinuation, when ART was initiated during PHI

See online video lectures When to start ART Part 1, When to start ART Part 2, What ART to start Part 1 and What ART to start Part 2 from the EACS online course on Management of HIV and Co-infections

# **Switch Strategies for Virologically Suppressed Persons**

### **Definition of virologically suppressed**

Clinical trials exploring switching strategies have generally defined suppression as an HIV-VL < 50 copies/mL for at least 6 months

### Indications

- Documented toxicity caused by one or more of the antiretrovirals included in the regimen, see Adverse Effects of ARVs and Drug Classes
- 2. Prevention of long-term toxicity, see Adverse Effects of ARVs and Drug Classes. This may include person's concerns about safety
- Avoidance of drug-drug interactions, page 26. This includes ART switch when starting HCV treatment to avoid DDIs, see Drug-drug Interactions between Viral Hepatitis Drugs and ARVs
- Planned pregnancy or women wishing to conceive, see Treatment of Pregnant Women Living with HIV or Women Considering Pregnancy
- Ageing and/or comorbidity with a possible negative impact of drug(s) in current regimen, e.g. on CVD risk, metabolic parameters
- Simplification: to reduce pill burden, adjust food restrictions, improve adherence and reduce monitoring needs
- Protection from HBV infection or reactivation by including tenofovir in the regimen
- 8. **Regimen fortification**: Increasing the barrier to resistance of a regimen in order to prevent VF (e.g. in persons with reduced adherence)
- Cost reduction: switching to the generic form of their current regimen, if available

### **Principles**

Clinicians should always review possible adverse events or tolerability issues with current antiretroviral regimens. Just because the viremia is suppressed it should not be assumed that the person is well adapted and tolerating the current regimen

- 1. The objectives of treatment modification should be to eliminate or improve adverse events, facilitate adequate treatment of comorbid conditions, and improve quality of life. The primary concern when switching should be to sustain and not to jeopardize virological suppression. In persons without prior virological failures and no archived resistance, switching regimens entail a low risk of subsequent failure if clinicians select one of the recommended combinations for first-line therapy. The majority of clinical trials showing non-inferiority of the new regimen after the switch have actively excluded persons with prior virological failures and historical resistance
- The complete ARV history with HIV-VL, tolerability issue, cumulative genotypic resistance history and/or phases of viremia on previous regimens with the potential of resistance development should be evaluated prior to any drug switch
- Switches within the same drug class (i.e. TDF/FTC -> TAF/FTC, EFV -> DOR or RPV) are usually virologically safe if equal potency and in the absence of resistance
- Cross-class switches of single drugs with the same barrier to resistance (for example EFV to RAL) are usually virologically safe in the absence of resistance to the new compound
- 5. In case of prior virologic failures, with or without evidence of resistance, switches have to be planned especially carefully when they result in a lower barrier to resistance of the regimen. A PI/b may only be switched to an NNRTI, INSTIs RAL if full activity of the 2 NRTIs in the new regimen can be assumed based on resistance data, ARV history and HIV-VL results before switching (see 2.) Due to the higher barrier to resistance of DTG and BIC, it is currently unclear if a switch to DTG- or BIC-based regimens also requires full activity of 2 NRTIs in the combination
- 6. Before switching, remaining treatment options in case of potential virological failure of the new regimen should be taken into consideration. This requires knowledge about the resistance selection profile of the switch regimen. Especially, when reducing the number of drugs in a regimen or its barrier to resistance, the chances of composing a fully suppressive regimen after potential failure following switch should be considered

- 7. Proviral DNA genotyping may be useful in persons with multiple virological failures, unavailable resistance history or low-level viremia at the time of switch. Results ought to be taken cautiously as proviral DNA genotype may not detect previous resistance mutations and can also detect clinically irrelevant mutations. Therefore, routine proviral DNA genotyping is currently not recommended
- 8. When selecting a new regimen, clinicians should carefully review the possibility of new drug-drug interactions with antiretroviral and concomitant medication leading to suboptimal drug exposure or toxicity, as well as the lag time for hepatic enzyme induction or blockade following discontinuation of the offending drug. Examples are: increased TDF toxicity with a PI/b or an increase in metformin exposure with DTG
- If the switch implies discontinuing TDF and not starting TAF, clinicians should check the HBV and HBV vaccination status. TDF or TAF should not be discontinued in persons with chronic HBV
- Persons should be seen soon (e.g. 4 weeks) after treatment switches to check for maintenance of suppression and possible toxicity or tolerability issues of the new regimen
- If someone receives and tolerates a regimen that is no longer a preferred option, and none of the other reasons for change applies, there is no need to change. Example: persons tolerating EFV-containing regimens
- See online video lecture How to Change ART from the EACS online course Management of HIV and Co-infections

### **Dual therapies**

In persons with suppression of HIV-VL < 50 copies/mL for the past 6 months these dual therapy strategies should only be given if there is

- a) no historical resistance and
- b) HBV immunity or if non-immune concomitant HBV Vaccination

# Dual therapies supported by large randomized clinical trials or meta-analyses:

DTG + RPV

XTC + DTG

XTC + DRV/b

Long-acting CAB + RPV bi-monthly injections

In clinical trials, these strategies have not been associated with more virological rebounds than triple therapy. There were a few cases of resistance development on DTG + RPV and CAB + RPV

### Dual therapy options supported only by small trials:

These regimens should be indicated only in persons not eligible for other treatment combinations due to intolerance or resistance to other drugs

DRV/b + RPV

DRV/b + DTG

### Strategies not recommended

- a. Monotherapy with a PI/b
- b. Monotherapy with DTG
- c. Dual or triple NRTIs combinations
- d. Specific two-drug combination, i.e.1 NRTI + 1 NNRTI or 1 NRTI + 1 unboosted PI, 1 NRTI + RAL, MVC + RAL, PI/b + MVC, ATV/b + RAL
- e. Intermittent therapy, sequential or prolonged treatment interruptions. In one open-label randomized study, 4 consecutive days a week of triple therapy was non inferior to 7 days a week, at 48 weeks in the context of close monitoring and counseling with visits every 3 months

# **Virological Failure**

Definition	INCOMPLETE SUPPRESSION: HIV-VL > 50 copies/ mL at 6 months after starting therapy in a person not previously on ART. In persons with very high baseline HIV-VL (> 100,000 copies/mL), achieving viral suppression may take longer than 6 months  REBOUND: confirmed HIV-VL > 50 copies/mL in someone with previously undetectable HIV-VL
General measures	Review expected potency of the regimen, taking into account all available historical genotypes
	Evaluate adherence, tolerability, drug-drug interactions, drug-food interactions, psychosocial issues
	Perform resistance testing preferably on failing therapy (usually routinely available for HIV-VL levels > 200-500 copies/mL and in specialised laboratories for lower levels of viraemia) and obtain historical resistance testing for archived mutations
	Tropism testing if considering MVC
	Consider TDM
	Review ART history
	Identify treatment options, active and potentially active drugs/combinations
Management	If HIV-VL > 50 and < 200 copies/mL:
of virological	Check for adherence, reinforce adherence
failure (VF)	Check HIV-VL 1 to 2 months later(i)
	If genotype shows no resistance mutations <sup>®</sup> : maintain current ART if it contains INSTI with high barrier to resistance (BIC, DTG) or PI/b, otherwise monitor carefully
	If HIV-VL confirmed > 200 copies/mL:
	Therapeutic decision will depend on the resistance testing (genotype) results:
	If no resistance mutations found: check for adherence, reinforce adherence, perform TDM, discuss change to a different regimen
	If resistance mutations found: switch to a suppressive regimen based on drug and genotype history; multidisciplinary expert discussion advised in case of multiclass resistance
	Goal of new regimen: HIV-VL < 50 copies/mL within 6 months

### In case of demonstrated resistance mutations

### General recommendations:

Use at least 2 and preferably 3 active drugs in the new regimen (including active drugs from previously used classes) based on resistance mutations present in current and earlier genotypic analyses

- \* If genotype shows only limited NRTI mutation(s) e.g. M184V and/or 1-2 TAMs<sup>(iii)</sup>: new regimen can include 2 NRTIs (3TC or FTC plus TDF or TAF) and either 1 active PI/b (i.e. DRV/b) or BIC or DTG (RAL or NNRTI not recommended)
- \* If genotype shows multiclass resistance (i.e. ≥ 2 classes): new regimen will usually use
- at least 1 fully active PI/b (i.e. DRV/b) or 1 fully active 2<sup>nd</sup> generation INSTI (BIC, DTG)
- plus 1 or 2 drugs remaining fully active despite resistance to other drugs from the class (i.e. 1 or 2 NRTIs and/or DOR)
- and/or from a class not used previously i.e. INSTI, NNRTI, PI/b, assessed by genotypic testing
- \* When a 2-3 drugs active regimen cannot be constructed with NRTI, NNRTI, PI/b and INSTI, a drug with a new mechanism of action such as fostemsavir or ibalizumab can be added to obtain such a 2-3 drugs active regimen
- \* In any case monotherapy is not recommended. In such situations, consider access to experimental drug therapy through early access program or clinical trials (e.g lenacapavir)

If < 2 active drugs are available, discuss on case by case situation deferring change, except in persons with low CD4 count (< 100 cells/ $\mu$ L) or with high risk of clinical deterioration for whom the goal is the preservation of immune function through partial reduction of HIV-VL (> 1 log<sub>10</sub> copies/mL reduction) by recycling drugs

### Other considerations:

- Treatment interruption is not recommended
- Continuation of 3TC or FTC even if documented resistance mutation (M184V/I) might be beneficial

If many options are available, criteria of preferred choice include: simplicity of the regimen, toxicity risks evaluation, drug-drug interactions, and sparing of future salvage therapy

- In the absence of resistance and in persons fully adherent to treatment, consider non-suppressible viremia due to cellular proliferation
- Take into consideration that certain mutations can revert and/or disappear if there is no drug pressure
- iii Thymidine Analog Mutations (TAMs) are non-polymorphic mutations selected by the thymidine analogs ZDV and/or d4T. For more detailed information NRTI Resistance Notes, see HIV Drug Resistance Database https://hivdb.stanford.edu/ or French ANRS resistance web page www.hivfrenchresistance.org



# **Treatment of Pregnant Women Living with HIV or Women Considering Pregnancy**

Scenarios for pregnant women or women who wish to conceive

Women planning to be pregnant or becoming pregnant while already on ART	<ul> <li>Maintain ART: The main goal of ART during pregnancy is maintaining treatment efficacy, both for the women's benefit and HIV transmission risk.</li> <li>ART may be switched temporarily for the duration of pregnancy to the preferred combinations recommended for ART naïve pregnant women, see table 1</li> <li>The decision on switching ART should be individualized taking into account the person's history of treatment, adherence and tolerability, and weighed against potential risk coming from ART exposure or suboptimal pharmacokinetics in pregnancy</li> <li>If the purpose for switching is insufficient data about safety and efficacy in pregnancy, it should be explained to the pregnant woman and her decision/willingness to switch current regimen taken into account:</li> <li>Lower serum concentration was observed in persons on therapies boosted with COBI, DRV/r qd and RPV</li> <li>There is insufficient data in pregnancy for BIC, DOR, RAL qd, and dual regimens</li> <li>Pregnant women should be monitored monthly or bimonthly (depending on adherence and length of virological suppression) and as close as possible to the predicted delivery date. HIV-VL should be tested every two months of pregnancy and including 36 weeks of gestation</li> </ul>
2. Women becoming pregnant while treatment-naïve	Starting ART as soon as possible is highly recommended, see table 1
Women whose follow-up starts late in the second or in the third trimester	Start ART immediately (see table 1) and consider RAL or DTG as the preferred choice to obtain rapid HIV-VL decline and to ensure the HIV-VL is undetectable by the time of delivery
Women whose HIV-VL is not undetectable at third trimester	Perform resistance testing and consider changing to or adding INSTI (RAL or DTG) if not on this class to obtain rapid HIV-VL decline
<ol><li>Women whose HIV-VL is &gt; 50 copies/mL at week 34-36 of pregnancy</li></ol>	Elective cesarean section to be planned at week 38, see labour and breastfeeding
6. Women diagnosed with HIV in labour	See labour and breastfeeding
	<ul> <li>Elective cesarean section to be planned at week 38</li> <li>iv ZDV: During labour and delivery: 2 mg/kg loading dose followed by continuous iv infusion of 1 mg/kg/hour until delivery</li> <li>Scheduled cesarean delivery: start iv ZDV 3 hours before surgery</li> <li>Unscheduled cesarean delivery: consider administering loading dose then proceed to delivery</li> <li>2) Women diagnosed with HIV during labour:</li> <li>If possible, perform caesarean section</li> <li>iv ZDV: During labour and delivery: 2 mg/kg loading dose followed by continuous iv infusion of 1 mg/kg/hour until delivery. Consider administering loading dose then proceed to delivery</li> <li>PEP should be given to all newborns born to mothers living with HIV according to local guidelines.</li> <li>For antiretroviral therapy in children with HIV, See page 140</li> </ul>
8. Breastfeeding	<ul> <li>The topic of feeding intentions should be discussed with a pregnant woman as early as possible in pregnancy, together with providing education and support to the mother</li> <li>We advise against breastfeeding, as in high-income settings the optimal way to prevent mother-to-child transmission is to feed infants born to mothers living with HIV with formula milk</li> <li>To reduce the potential physical and emotional discomfort associated with breast engorgement, together with the risk of covert breastfeeding, women living with HIV should be given cabergoline to suppress lactation after delivery</li> <li>In situations where a woman chooses to breastfeed, we recommend input from an interdisciplinary team including adult HIV specialist, paediatrician and obstetrician/gynecologist</li> <li>We recommend monthly follow-up during the whole breastfeeding period with increased clinical and virological monitoring of both the mother and the infant. Measurement of drug concentrations in the milk could be done to inform clinical practice</li> <li>Maternal HIV-VL &gt; 50 copies/mL should result in cessation of breastfeeding, providing cabergoline and support from interdisciplinary team and a nursing specialist</li> <li>Immediate consulting by the interdisciplinary team should be provided in case of signs and symptoms of mastitis, infant mouth or gut infections</li> <li>Currently there is no evidence supporting PrEP recommendation for the infants who are breastfed</li> <li>After stopping the breastfeeding, the child should undergo routine diagnostics as recommended in HIV-exposed children</li> </ul>

### Table 1. Antiretroviral regimen for ART-naïve pregnant women

ART-naïve pregnant women should initiate treatment as soon as possible. The decision of ART regimen should be discussed with the person and individualized taking into account tolerability, possible adherence issues, as well weighed against potential risk coming from ART exposure or suboptimal pharmacokinetics in pregnancy.

Pregnant women starting ART should be monitored monthly or bimonthly (depending on adherence and length of virological suppression) and as close as possible to the predicted delivery date. HIV-VL should be tested every two months of pregnancy and including 36 weeks of gestation

Regimen	Main requirements	Additional guidance (see footnotes)
Recommended regimens		
2 NRTIs + INSTI (PREFERRED)		
ABC/3TC + DTG or ABC/3TC/DTG	DTG to be discussed with women considering to become pregnant or if to be used in first 6 weeks of pregnancy HLA-B*57:01 negative HBsAg negative	I (ABC: HLA-B*57:01, may delay starting ART) II (DTG in pregnancy: see footnote)
TDF/XTC or TAF/FTC + DTG	DTG to be discussed with women considering to become pregnant or if to be used in first 6 weeks of pregnancy. TAF/FTC not recommended in first 14 weeks of pregnancy	II (DTG in pregnancy: see footnote) III (Tenofovir salts) IV (TAF & pregnancy)
TDF/XTC or TAF/FTC + RAL 400 mg bid	TAF/FTC not recommended in first 14 weeks of pregnancy	III (Tenofovir salts)  IV (TAF & pregnancy)  V (RAL in pregnancy, bid dosing)
2 NRTIs + PI/r		
TDF/XTC or TAF/FTC + DRV/r 600 mg/100 mg bid	With food TAF/FTC not recommended in first 14 weeks of pregnancy	III (Tenofovir salts) IV (TAF & pregnancy) VI (DRV dosing) VII (COBI boosting)
Alternative regimens		
2 NRTIs + INSTI		
ABC/3TC + RAL 400 mg bid	HBsAg negative HLA-B*57:01 negative	I (ABC: HLA-B*57:01, may delay starting ART) V (RAL in pregnancy, bid dosing)
2 NRTIs + NNRTI		
ABC/3TC + EFV	HLA-B*57:01 negative HBsAg negative HIV-VL < 100,000 copies/mL At bedtime or 2 hours before dinner	I (ABC: HLA-B*57:01, may delay starting ART) VIII (EFV HIV-2 & group O)
TDF/XTC or TAF/FTC + EFV or TDF/FTC/EFV	At bedtime or 2 hours before dinner TAF/FTC not recommended in first 14 weeks of pregnancy	III (Tenofovir salts) IV (TAF & pregnancy) VIII (EFV HIV-2 & group O)
TDF/XTC or TAF/FTC + RPV or TDF/FTC/RPV or TAF/FTC/RPV	CD4 count > 200 cells/µL HIV-VL < 100,000 copies/mL Not on gastric pH increasing agents With food TAF/FTC not recommended in first 14 weeks of pregnancy	II (Tenofovir salts) IV (TAF & pregnancy) IX (RPV exposure during 2 <sup>nd</sup> and 3 <sup>rd</sup> trimester, HIV-2) X (Interactions)
2 NRTIs + PI/r		
ABC/3TC + DRV/r 600 mg/100 mg bid	HLA-B*57:01 negative HBsAg negative With food	I (ABC: HLA-B*57:01, may delay starting ART) VI (DRV dosing) VII (COBI boosting)

### Additional guidance

- ABC contraindicated if HLA-B\*57:01 positive. Even if HLA-B\*57:01 negative, counselling on HSR risk still mandatory. If testing for HLA-B\*57:01 results in delay of ART initiation, consider other recommended backbone
- II A minimal non-significant increase in neural tube defects was shown among women receiving DTG from conception compared with all other antiretroviral exposure
- III Some generic forms of TDF use phosphate, maleate, and succinate salts instead of fumarate. They may be used interchangeably. In certain countries, TDF is labelled as 245 mg rather than 300 mg to reflect the amount of the prodrug (tenofovir disoproxil) rather than the fumarate salt (tenofovir disoproxil fumarate)
- IV TAF/FTC not recommended in first 14 weeks of gestational age as IMPACT 2010/VESTED randomized study evaluating the safety and virologic efficacy of this combination recruited women only between 14-28 weeks of pregnancy
- V There were no reports of neural tube defects among 1991 prospective reports of RAL exposure in pregnancy, 456 of which were in the periconception period. No data on RAL 1200 mg qd: not recommended
- VI DRV/r 800/100 mg qd not recommended during pregnancy due to decreased levels. DRV/c is not recommended during pregnancy due to significant lower exposures of DRV and COBI in the second and third trimester of pregnancy
- VII Boosting with COBI is not recommended after the second trimester of pregnancy (insufficient drug levels)
- VIII EFV not active against HIV-2 and HIV-1 group O strains
- IX Lower RPV exposure during second and third trimesters; Consider monitoring VL more frequently. RPV is not active against HIV-2
- X Pregnant women are often prescribed anti-H2 or proton pump inhibitors for nausea. Careful review of concomitant medicines at each visit and providing pregnant women with information on potential interactions is recommended



# **ART in TB/HIV Co-infection**

### **Principles**

Persons with TB should be started on standard TB therapy with 2 months rifampicin/isoniazid/pyrazinamide/ethambutol followed by 4 months rifampicin/isoniazid (choice of drugs and length of treatment depends on drug susceptibility and site of disease), see Diagnosis and Treatment of TB in Persons with HIV

All persons with TB/HIV co-infection should start ART irrespective of CD4 count. Treatment supervision and adherence evaluation are very important. If the person is already on ART, check for potential DDIs and if these are significant, consider switching to one of the recommended regimens for TB/HIV co-infection

### Suggested timing of ART initiation in TB/HIV co-infection

ART should be started as soon as possible (within two weeks of initiating TB treatment) regardless of CD4 count

However, if TB meningitis signs and symptoms are present ART initiation may be delayed. See When to start ART in persons with Opportunistic Infections (OIs)

Be aware of IRIS reaction in persons starting ART at low CD4 count levels and with early initiation of ART. Prophylactic prednisone for 4 weeks at the time of ART initiation (prednisone 40 mg qd for 14 days, then 20 mg qd for 14 days) can prevent paradoxical TB-associated IRIS in persons with CD4 < 100 cells/ $\mu$ L receiving TB treatment

Corticosteroids should be considered for treatment of symptomatic IRIS, with dosages and duration tailored according to response

### Table 1. Antiretroviral regimens in TB/HIV co-infection

These recommendations are for persons initiating ART with susceptible Mycobacterium tuberculosis infection. When treating MDR-TB or XDR-TB, careful review of DDIs and potential toxicities is mandatory before initiating ART. For a wider review of potential DDIs of ART and TB treatment, see page 35

Regimen	Main requirements	Additional guidance (footnotes)
Recommended regimens with rif	ampicin	
2 NRTIs + NNRTI		
TDF/XTC + EFV or TDF/FTC/EFV	At bedtime or 2 hours before dinner	l (tenofovir salts) ll (EFV: suicidality. HIV2 or HIV-1 group 0)
ABC/3TC + EFV	HLA-B*57:01 negative HBsAg negative HIV-VL < 100,000 copies/mL At bedtime or 2 hours before dinner	III (ABC: HLA-B*57:01) II (EFV: suicidality. HIV-2 or HIV-1 group 0)
Alternative regimens with rifamp	icin	
2 NRTIs + INSTI		
TDF/XTC + DTG bid		I (tenofovir salts) IV (DTG: dosing)
TDF/XTC + RAL bid		l (tenofovir salts) V (RAL: dosing)
ABC/3TC + RAL bid	HBsAg negative HLA-B*57:01 negative	III (ABC: HLA-B*57:01) V (RAL: dosing)
Other combinations with rifabuti	n	
2 NRTIs + PI/r		
TDF/XTC + DRV/r	With food	VI (rifabutin dosing)
ABC/3TC + DRV/r	HLA-B*57:01 negative HBsAg negative HIV-VL < 100,000 copies/mL With food	III (ABC: HLA-B*57:01) VI (rifabutin dosing)

### Additional guidance

- I There are available generic forms of TDF, which instead of fumarate use phosphate, maleate, and succinate salts. They can be used interchangeably. In certain countries, TDF is labelled as 245 mg rather than 300 mg to reflect the amount of the prodrug (tenofovir disoproxil) rather than the fumarate salt (tenofovir disoproxil fumarate)
- II EFV: not to be given if history of suicide attempts or mental illness; not active against HIV-2 and HIV-1 group O strains
- III ABC contraindicated if HLA-B\*57:01 positive. Even if HLA-B\*57:01 negative, counselling on HSR risk still mandatory. ABC should be used with caution in persons with a high CVD risk (> 10%)
- IV DTG should be dosed 50 mg bid when given with rifampicin since rifampicin lowers DTG exposure. This dose adjustment should be maintained for 2 weeks after stopping rifampicin as the inducing effect persists after discontinuation of a strong inducer
- V RAL 400 or 800 mg bid. With RAL 400 bid a large phase 3 study showed non-inferiority at week 24 but failed to show non-inferiority at week 48 compared to EFV. With 800 mg bid only limited date from a phase 2 study with potential increases in liver toxicity
- VI For guidance on ARV and rifabutin dosing, see TB Drug Doses, DDI table on Anti-tuberculosis drugs and ARVs



### Non-rifamycin regimens

Tuberculosis can be treated with regimens that do not contain rifamycins. Their use should be contemplated only in persons with serious toxicity to rifamycins where desensitisation has failed, or in persons with rifamycin-resistant isolates. Although non-rifamycin regimens have fewer drug-drug interactions, such regimens are inferior to a rifampicin-based regimen for fully drug-sensitive TB treatment

Poorer outcomes have also been seen in cases where rifampicin is used for the initial two months before the regimen is switched to isoniazid and ethambutol in the continuation phase

In countries where neither DTG nor rifabutin are available, or there is no possibility to use RAL or EFV, following combinations could also represent a short-term alternative until anti-TB treatment has been completed

- Rifampicin plus double dose LPV/r or with RTV super boosted (400 mg bid) + LPV
- For other regimens based on 2 NRTIs plus NVP, RPV, DOR, ETV or MVC, consultation with an HIV specialist is recommended



# Post-exposure Prophylaxis (PEP)

### PEP recommended in case of:

Risk	Nature of exposure	Status of source person
Blood	Subcutaneous or intramuscular penetration with iv or im needle, or intravascular device	HIV-positive or recent serostatus unknown, but presence of HIV risk factors
	Percutaneous injury with sharp instrument (lancet), im or sc needle, suture needle Contact > 15 min of mucous membrane or non- intact skin	HIV-positive
Genital secretions	Anal or vaginal sex and not on PrEP or low PrEP adherence	Viraemic HIV-positive or serostatus unknown but presence of HIV risk factors. If source person is on ART, PEP should be started, HIV-VL should be repeated, and, if undetectable, PEP can be stopped
	Receptive oral sex with ejaculation and not on PrEP or low PrEP adherence	Viraemic HIV-positive
Intravenous drug use	Exchange of syringe, needle, preparation material or any other material	HIV-positive

- Rapid testing of the source person for HBV, HCV and HIV (if HIV-status unknown) recommended
- If source person HIV-positive on ART, order resistance testing if HIV-VL detectable
- Individualise PEP according to the source's treatment history and previous resistance tests
- For sexual exposure, if HIV-positive source has documented undetectable HIV-VL, PEP is no longer recommended
- PEP to be started ideally < 4 hours after the exposure, and no later than 48/72 hours
- · Duration of PEP: 4 weeks (unless discontinued due to lack of indication)
- PEP regimens: TDF/FTC or TAF/FTC + RAL bid or qd, or + DRV/b qd.
   TDF/FTC or TAF/FTC+ DTG qd or TAF/FTC/BIC may be also considered as alternatives
- · Full sexual health screen in case of sexual exposure
- Emergency contraception counselling for sexual exposure
- · Follow-up:
  - HIV serology + HBV and HCV, pregnancy test (women) within 48 hours of exposure and test for STIs if appropriate
  - Re-evaluation of PEP indication by HIV expert within 48-72 hours
  - Assess tolerability of PEP regimen
  - Transaminases, HCV-PCR and HCV serology at month 1 if source person HCV-positive (observed or suspected)
  - Follow-up HIV serology: mandatory at the end of PEP and repeat 6-8 weeks later
  - Discuss opportunity to start PrEP

# **Pre-exposure Prophylaxis (PrEP)**

- PrEP should be used in adults at high-risk of acquiring HIV infection when condoms are not used consistently. Before PrEP is initiated, HBV serology status should be documented
- Recommended in HIV-negative men who have sex with men (MSM)
  and transgender individuals when condoms are not used consistently
  with casual partners or with partners with HIV who are not virally suppressed on treatment. A recent STI, use of post-exposure prophylaxis
  or chem-sex may be markers of increased risk for HIV
- May be considered in HIV-negative heterosexual women and men who are inconsistent in their use of condoms and have multiple sexual partners where some may have untreated or inadequately suppressed HIV infection
- PrEP is a medical intervention that provides a high level of protection against HIV acquisition but does not protect against other STIs or pregnancy and should be used in combination with other preventive interventions. PrEP should be supervised by a doctor, experienced with sexual health and use of HIV medicines, possibly as part of a shared care arrangement

### The following procedures are recommended:

- Documented negative fourth generation HIV test a week prior to starting PrEP. In case of suspicion of acute HIV-infection, an RNA test on plasma should also be performed, page 15. During PrEP, a fourth generation HIV test should be repeated at one month and then every 3 months. In stable long-term users who are on 6 monthly prescriptions an interim third generation test that can be performed without a visit to clinic is acceptable
- PrEP should be changed to triple-drug ART without interruption in case of early clinical signs of HIV seroconversion or a positive HIV diagnostic test which may necessitate referral for evaluation to an HIV unit, see ART initiation page 12
- PrEP may continue during pregnancy and breastfeeding if the risk of acquiring HIV persists
- Before PrEP is initiated, HBV serology status should be documented. If HBsAg positive, see Clinical Management and Treatment of HBV and HCV Co-infection
- Counsel that PrEP does not prevent other types of STIs; screen for STI (syphilis, chlamydia, gonorrhoeae, HAV, HCV) when starting PrEP and regularly during use of PrEP, pages 7-9
- Counsel that TDF-based PrEP may impact renal and bone health, see pages 71 and 73-75. Check renal function within the first 3 months of starting PrEP and check renal function and bone health during PrEP according to guidelines on TDF use
- Counsel that PrEP, like other prevention methods, only works when it is taken. An adherence check one month after starting is recommended, and counselling may be required in follow-up
- Counsel that PrEP can be prescribed long-term but that each consecutive PrEP prescription should cover the period to the next visit which will be every 3 months for the majority but could be a maximum of 6 months in stable long-term users (over one year of daily PrEP)

- 3. PrEP regimen
- TDF/FTC 300\*/200 mg 1 tablet qd. In both men and women PrEP should be taken for 7 days before the first exposure and stopped 7 days after the last exposure
- A trial with daily TAF/FTC in MSM and transgender women has shown non inferiority to daily TDF/FTC. No data are available in other high risk groups
- For men only, PrEP may be dosed 'on demand' (double dose of TDF/FTC 2-24 hours before each sexual intercourse, followed by two single doses of TDF/FTC, 24 and 48 hours after the first drug intake; no data for TAF/FTC so far). There are no efficacy data with on demand PrEP with TDF/FTC in women
- Use of generic formulations of TDF/FTC, if and where available, may help to improve the cost-effectiveness of PrEP, which is essential for its use as public health approach
- Data on renal outcomes with use of TDF vs. TAF in those on PrEP with renal impairment is limited, recommendations to follow guidelines on TDF use in persons with HIV, see pages 74-76. Similarly, no data on use of "on demand" vs daily PrEP for renal outcomes
- In certain countries, TDF is labelled as 245 mg rather than 300 mg to reflect the amount of the prodrug (tenofovir disoproxil) rather than the fumarate salt (tenofovir disoproxil fumarate)

# **Adverse Effects of ARVs and Drug Classes**

	Skin	Digestive	Liver	cv	Musculo- skeletal	Genito- urinary	Nervous	Body fat	Metabolic	Other
NRTIs										
ABC	Rash	Nausea* Diarrhoea*		IHD						*Systemic hypersensitivity syndrome (HLA B*57:01 dependent)
ZDV <sup>(ii)</sup>	Nail pigmen- tation	Nausea	Steatosis		Myopathy, Rhabdomy- olysis			Lipoatrophy	Dyslipi- daemia, Hyperlacta- taemia	Anaemia
3TC										
FTC										
TDF(iii)			Hepatitis		↓ BMD, Osteoma- lacia	↓ eGFR, Fanconi syndrome				
TAF(iii)									Weight gain	
NNRTIs										
EFV	Rash		Hepatitis				Neuropsychiatric events including: depression, sleep disturbance, headache		Dyslipi- daemia, Gynaeco- mastia	↓ plasma 25(OH) vitamin D
ETV	Rash									
NVP	Rash*		Hepatitis*							*Systemic hypersensitivity (CD4 count and gender dependent)
RPV	Rash		Hepatitis			↓ eGFR <sup>(iv)</sup>	Depression, Sleep disturbance, Headache			
DOR							Sleep disturbance, Headache			
Pls										
ATV <sup>(v)</sup>		Nausea	Hyperbiliru- binaemia, Jaundice, Cholelithiasis			↓ eGFR, Nephrolithiasis			Dyslipi- daemia	
DRV <sup>(v)</sup>	Rash	and Diarrhoea <sup>(vii)</sup>		IHD		Nephrolithiasis			Dyslipi- daemia	
LPV <sup>(vi)</sup>				IHD		↓ eGFR			Dyslipi- daemia	
Boosting										
RTV		Nausea and diarrhoea				↓ eGFR <sup>(iv)</sup>			Dyslipidae- mia	
COBI		Nausea and diarrhoea				↓ eGFR <sup>(iv)</sup>			Dyslipidae- mia	

INSTI									
RAL		Nausea			Myopathy, Rhabdomy- olysis		Sleep disturbance, Headache	Weight gain	Systemic hypersensitivity syndrome <sup>(viii)</sup>
DTG	Rash	Nausea				↓ eGFR <sup>(iv)</sup>	Sleep disturbance, Headache	Weight gain	Systemic hypersensitivity syndrome (< 1%) Minimal non-significant increase in neural tube defects <sup>(ix)</sup>
EVG/c		Nausea, Diarrhoea				↓ eGFR <sup>(iv)</sup>	Sleep disturbance, Headache	Weight gain	
BIC						↓ eGFR <sup>(iv)</sup>	Sleep disturbance, Headache	Weight gain	
CAB	Injection site reac- tions(x)						Sleep disturbance, Headache		Pyrexia <sup>(xi)</sup>
Entry inhib	itors								
Ibalizumab	Rash	Nausea Diarrhoea					Dizziness Headache		
FTR	Rash	Nausea, Vomiting, Abdominal pain, Diarrhoea					Headache		
MVC			Hepatitis	Postural hypo- tension					
ENF	Injection site nodules								Hypersensitivity

### "Frequent effects" (events expected in at least 10% of treated individuals), in bold

"Severe effects" (events that can put a person's life at risk and represent a medical emergency), in red

- "Neither frequent nor severe effects", in non-bold black
- Still available, but generally not recommended due to toxicity
- iii TDF and TAF are prodrugs of tenofovir. TDF, but not TAF, may have kidney and bone toxicity particularly when co-administered with RTV or COBI boosting. TDF, but not TAF, decreases plasma lipids. TAF, but not TDF, may promote weight gain particularly when co-administered with DTG or BIC, see pages 71, 74, 75, 86
- iv Due to inhibition of renal tubular creatinine secretion without affecting glomerular filtration itself
- V ATV can be used unboosted or boosted with low-dose RTV or COBI ATV-related adverse effects are more common with boosting. DRV can be used boosted with low-dose RTV or COBI. Both low-dose RTV and COBI as boosters may cause minor digestive problems and lipid increases (low-dose RTV more than COBI). IHD reported with ritona-vir-boosted DRV only (no data with COBI-boosted DRV, although lipid effects lower)
- vi Still available but seldom used. Requires RTV-boosting
- vii Frequency and severity differs between individual PIs
- viii DRESS syndrome reported in a few cases, potentially associated to HLA-B\*53
- ix See Treatment of Pregnant Women or Women considering Pregnancy
- x CAB is available in oral or injectable formulations; injection site reactions are an adverse effect of injectable CAB
- xi Pyrexia includes feeling hot or body temperature increased
- Refers to effects seen in relation to hypersensitivity reactions

### Notes:

- 1. The adverse effects listed in the table above are not exhaustive, but represent the most important effects with a likely causal relation. Nausea, diarrhoea and rash are frequently observed in persons on ART, and these symptoms are indicated in the table for drugs where clinical experience suggests a possible causal link
- D4T, ddI, FPV, IDV, SQV and TPR removed. Please refer to EACS v9.1 for details, http://www.eacsociety.org/files/2018\_guidelines-9.1-english.pdf

# Part III Drug-drug Interactions and Other Prescribing Issues

ARVs are recognised to be amongst the therapeutic agents with the highest potential for drug-drug interactions (DDIs) as these drugs can be both a victim (affected by other drugs) and/or a perpetrator (affect other drugs) of DDIs. Given the life-long ART, DDIs are practically unavoidable in persons with HIV and comorbid conditions. Thus, the potential for DDIs should be considered systematically when selecting an ART regimen or when any new medicine is coadministered to existing ART with particular attention to adjust dosage and perform clinical monitoring when needed.

The im administration of the ARVs CAB and RPV presents the advantage of eliminating DDIs occurring at the gastrointestinal level. However, DDIs can still occur at the hepatic level as illustrated below. Bypassing the gastrointestinal tract does not mitigate the magnitude of DDIs with drugs inducing metabolism.

# Drug-Drug Interactions after Oral and Intramuscular Administration of CAB and RPV

Intramuscular administration: - DDIs at the gastrointestinal level are avoided

→ DDIs at the hepatic level can still occur (magnitude of DDIs with inducers is not mitigated)

CYP3A4 UGT1A1/9 drug transporters

Mechanisms of DDIs after ORAL administration

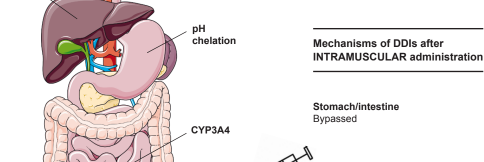
### Stomach/intestine

- Change gastric pH
- Chelation divalent cations
- Inhibition/induction of CYP3A4, drug transporters

### Liver

 Inhibition/induction of CYP3A4, UGT1A1/9, drug transporters

Adapted from Hodge D et al. Clin Pharmacokinet 2021



### Liver

 Inhibition/induction of CYP3A4, UGT1A1/9, drug transporters

### Examples of medications interacting with the oral but not the intramuscular administration of RPV

Antacids; famotidine; lansoprazole; liraglutide; omeprazole; orlistat; pantoprazole; rabeprazole; ranitidine

### Examples of medications interacting with the oral but not the intramuscular administration of CAB

Antacids; calcium; iron; magnesium; multivitamins containing divalent cations; orlistat; strontium ranelate

The DDIs profiles between ARVs and coadministered medicines within a therapeutic class are also presented in the corresponding Co-morbidities section and Viral Hepatitis Co-infection section

Detailed information on DDIs can be found on the University of Liverpool DDIs websites: http://www.hiv-druginteractions.org and http://www.hep-druginteractions.org

Age-related physiological changes and co-morbidities predispose older persons with HIV to inappropriate drug use or dosing in addition to DDIs

Besides highlighting the most common DDIs, this section also provides guidance on how to adjust drug dosing in case of liver or renal impairment, considerations for those with swallowing difficulties and what to consider when prescribing drugs in older persons with HIV including the top ten drug classes to avoid

# **Drug-drug Interactions between ARVs and non-ARVs**

No	n-ARV drugs	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB	CAB/	DTG	EVG/c	RAL	TAF	TDF
NO	II-ARV urugs		AIV/I		DRV/I					NVF	↑4%	FIK	IVIVC	ыс	oral	RPV	סוט	EVG/C	KAL	IAF	IDF
	atorvastatin	↑822%	1	↑290%	1	↑490%	↓2%	↓43%	↓37%	ţ	D10%	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
v	fluvastatin	1	1	1	1	$\leftrightarrow$	$\leftrightarrow$	1	1	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
drugs	pravastatin	1	1	1	↑81%	↑33%	$\leftrightarrow$	↓44%	↓	$\leftrightarrow$	1	↓4%	$\leftrightarrow$	$\leftrightarrow$							
lar	rosuvastatin	↑242%	↑213%	↑93%	↑48%	↑108%	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑69%	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑38%	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Cardiovascular	simvastatin	1	1	1	1	1	$\leftrightarrow$	↓68%	↓	ļ	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
iova	amlodipine	†a	†a	1	1	↑a	$\leftrightarrow$	Ţ	Ţ	ļ	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
ard	diltiazem	†a	† <b>a</b>	1	1	↑a	Е	↓69%	ţΕ	↓	Е	Е	Е	Е	$\leftrightarrow$	Е	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
0	metoprolol	†a	†a	1	1	↑a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
	verapamil	†a	↑a	1	1	† <b>a</b>	Е	Ţ	ţΕ	ļ	E	E	Е	E	$\leftrightarrow$	Е	$\leftrightarrow$	1	$\leftrightarrow$	Е	Е
	warfarin	1	↑ or ↓	1	↓	<b>↓</b>	$\leftrightarrow$	↑ or ↓	1	↑ or ↓	$\leftrightarrow$	↓	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	bupropion	$\leftrightarrow$	1	$\leftrightarrow$	↓	↓57%	$\leftrightarrow$	↓55%	$\leftrightarrow$	↓	$\leftrightarrow$	↑?	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	carbamaze- pine	↑D	↑D	↑D	1	↑D c	D	↓27% Ď36%	D	↓D	D	D	D	D	D	D	D49%	↑ D	Dc	D	$\leftrightarrow$
	citalopram	↑a,b	↑a,b	1	1	↑a,b	$\leftrightarrow$	↓	<b>↓</b>	<b>↓</b>	↔b	↔b	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔b	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	diazepam	1	1	1	1	1	$\leftrightarrow$	↓	↓	↓	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
S	lamotrigine	$\leftrightarrow$	↓32%	$\leftrightarrow$	↓	↓50%	$\leftrightarrow$	Ţ	$\leftrightarrow$	↓1%	$\leftrightarrow$	$\leftrightarrow$									
drugs	midazolam (oral)	1	1	1	1	1	↓18%	↓	↓	ļ	$\leftrightarrow$	$\leftrightarrow$	↑18%	↑15%	↑10%	$\leftrightarrow$	$\leftrightarrow$	1	↓8%	$\leftrightarrow$	$\leftrightarrow$
CNS	mirtazapine	↑b	↑b	1	1	↑ <b>b</b>	$\leftrightarrow$	↓	↓	Ţ	↔b	↔b	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔b	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	paroxetine	<b>↑</b> ↓?	<b>↑</b> ↓?	<b>↑</b> ↓?	↓39%	<b>↑</b> ↓?	$\leftrightarrow$	$\leftrightarrow$	↑3%	$\leftrightarrow$	<b>↑</b> ↓?	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$							
	phenytoin	D	↓D	D	ţD	↓D c	D	↓D	D	D	D	D	D	D	D	D	D d	D	Dc	D	$\leftrightarrow$
	pimozide	1	1	1	1	1	$\leftrightarrow$	1	↓	↓	↔b	↔b	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔b	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	sertraline	1	<b>↓</b>	1	↓49%	↓b	$\leftrightarrow$	↓39%	↓	↓ ↓	$\leftrightarrow$	↓7%	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	triazolam	1	1	1	1	1	$\leftrightarrow$	↓	Ţ	Ţ	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	clarithromycin	↑E a,b	↑E a,b	↑E	1	↑ a,b	1	↓39%	↓39% Ě42%	↓31% Ě26%	Εb	E a,b	Е	E	$\leftrightarrow$	Еb	$\leftrightarrow$	↑E	$\leftrightarrow$	Е	Е
S	fluconazole	↑? a,b	↔ a,b	↑?	$\leftrightarrow$		1	$\leftrightarrow$	E86%	E100%	Εb	E a,b	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	Εb	$\leftrightarrow$	↑?	$\leftrightarrow$	E?	Е
tive	itraconazole	↑E b	↑E b	↑E	↑E	↑E b	1	↓39%	ţΕ	↓61%	Εb	Εb	Е	Е	$\leftrightarrow$	Εb	$\leftrightarrow$	↑E	$\leftrightarrow$	Е	Е
Anti-infectives	rifabutin	↑D e	↑ f	↑D e	↑ f	↑ f	D50% g	↓38% h	↓17% Ď37%	↑17%	D42% i	D30%	j	D38%	$\leftrightarrow$	D	$\leftrightarrow$	↑D e	E19%	Dk	$\leftrightarrow$
Ant	rifampicin	D	D72%	D	D57%	D75% I	D82%	D26%m	D	D58%	D80%	D82%	D	D75%	D59%	D	D54%n	D	D40%0	Dk	D12%
	voriconazole	↑↓ Eb	↑↓ Db	↑E	<b>↓</b>	↑↓ Eb	1	ţΕ	↑14% E36%	ţΕ	E	Е	Е	E61%	$\leftrightarrow$	Е	$\leftrightarrow$	↑E	$\leftrightarrow$	$\leftrightarrow$	Е
	antacids	D	D	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔	$\leftrightarrow$	D	$\leftrightarrow$	$\leftrightarrow$	D	D	$\leftrightarrow$	D	D	Dρ	$\leftrightarrow$	$\leftrightarrow$
	PPIs	D	D	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	D	$\leftrightarrow$	Е	$\leftrightarrow$	$\leftrightarrow$						
	H2 blockers	D	D	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	D	$\leftrightarrow$	Е	$\leftrightarrow$	$\leftrightarrow$						
	alfuzosin	↑ b	↑ b	1	1	↑ b	$\leftrightarrow$	↓	↓	↓	↔ <b>b</b>	↔ b	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔ b	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	beclo- metasone (inhaled)	↑ q	↑ q	↑? <b>q</b>	↓11% <b>r</b>	↑ q	$\leftrightarrow$	↑ q	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
	budesonide (inhaled)	↑s	↑s	† s	↑ s	† s	$\leftrightarrow$	↓	ļ	ļ	$\leftrightarrow$	↑s	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
sno	buprenor- phine	1	↑67%t	1	↓11%t	↑~2%	$\leftrightarrow$	↓50%	↓25%	↓9%	$\leftrightarrow$	↑30%	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑35%	$\leftrightarrow$	$\leftrightarrow$	↑~5%
Miscellaneous	ergot deriva- tives	1	1	1	1	1	Е	1	1	↓	Е	$\leftrightarrow$	E	$\leftrightarrow$	$\leftrightarrow$	Е	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Misc	ethinylestra- diol	↑1%u	↓19%v	↓30%	↓44% <mark>u</mark>	↓42%u	↓2%	w	↑22%	↓20%	↑14%	↑40%x	↓<1%	↑4%	↑2%	$\leftrightarrow$	↑3%	↓25%y	↓2%	↑11%	$\leftrightarrow$
	fluticasone (inhaled)	↑s	↑ s	† s	<b>†</b> s	† s	$\leftrightarrow$	ļ	Ţ	ţ	$\leftrightarrow$	†s	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	methadone	↑?ab	↔ ab	↑?	↓16%	↓53%ab	↓5%	↓52%	↑6%	↓~50%	↓16%ab	↑14%ab	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔ab	↓2%	↑7%	$\leftrightarrow$	$\leftrightarrow$	↑~5%
							↔	↓	<b>↓</b>	, 	$\leftrightarrow$	· ·	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	· ↔	1	$\leftrightarrow$	$\leftrightarrow$	· ·
	salmeterol (inhaled)	1	1	1	1	1	``	*													
		<b>†</b>	↑ ↑	↑ ↑	↑ ↑	1	↔	<b>1</b>	↓37%	<b>↓</b>	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	(inhaled) sildenafil									↓ Dz	↔ D z	↔ D z	↔ D z	↔ D z	$\leftrightarrow$	↔ D z	↔ D e	↑ D z		↔ D z	$\leftrightarrow$

### Colour legend

No clinically significant interaction expected These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

### Legend

↑ Potential elevated exposure of the non-ARV drug
 ↓ Potential decreased exposure of the non-ARV drug

→ No significant effect

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug
ATV/c ATV co-formulated with COBI (300/150 mg qd)
DRV/c DRV co-formulated with COBI (800/150 mg qd)
CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

\* table summarises the drug-drug interactions between HIV therapy and some commonly prescribed co-medicines as well as the drug-drug interactions of particular clinical relevance. This table is not exhaustive.

### **Further Information**

For additional drug-drug interactions and for more detailed pharmacokinetic interaction data and dosage adjustments, please refer to: http://www.hiv-druginteractions.org (University of Liverpool)

### Interactions with ABC, FTC, 3TC, ZDV

ABC: decreased ABC exposure with phenytoin, rifampicin

ABC: decreased methadone exposure
ABC: increased carbamazepine exposure
FTC, 3TC: no clinically relevant interactions expected.

ZDV: decreased ZDV exposure with clarithromycin, rifampicin ZDV: increased ZDV exposure with fluconazole, methadone

ZDV: increased carbamazepine exposure ZDV: decreased phenytoin exposure

### Interactions with ibalizumab

None

### Comments

- a ECG monitoring is recommended.
- b Caution as both drugs can induce QT interval prolongation.
- c Co-administration with LPV/r 800/100 qd or RAL 1200 mg qd is not recommended. If use is unavoidable, give LPV/r 400/100 mg bid or RAL 400 mg bid, with monitoring of response.
- d The European SmPC recommends DTG 50 mg bid in persons without INSTI resistance. The US Prescribing Information recommends that co-administration should be avoided as there are insufficient data to make dosing recommendations.
- e Reduce rifabutin to 150 mg 3 times per week.
- f Reduce rifabutin to 150 mg qd. Monitoring for rifabutin-related toxicities (i.e. uveitis or neutropenia) is advised with daily administration of rifabutin.
- The product label for DOR recommends to increase DOR dosage to 100 mg bid when co-administered with rifabutin. DOR should be kept at 100 mg bid for at least another 2 weeks following cessation of rifabutin due to the persisting inducing effect upon discontinuation of a moderate/ strong inducer.
- h Increase rifabutin to 450 mg daily.
- The RPV dose should be increased to 50 mg qd during co-administration (and decreased to 25 mg qd when rifabutin is stopped). Note, it is recommended to maintain RPV 50 mg qd for at least another 2 weeks following cessation of rifabutin due to the persisting inducing effect upon discontinuation of a moderate/strong inducer.
- j Increase MVC to 600 mg bid in absence of PI. With PI (except TPV/r, FPV/r), give MVC 150 mg bid.
- k Rifamycins decrease the exposure of TAF when given 25 mg qd therefore the label recommends to use TAF 25 mg bid. However, the intracellular tenofovir diphosphate (active entity) concentrations are likely to be higher than those observed with TDF even without rifampicin suggesting that usage of TAF 25 mg qd with rifampicin, rifapentine or rifabutin may be acceptable.
- If no other option use RTV 400 mg bid or double dose LPV/r.
- m EFV should be used at 600 mg qd in presence of rifampicin (in absence of rifampicin, EFV can be used at 400 mg qd or 600 mg qd).
- Administer DTG 50 mg bid in treatment-naïve or INSTI-naïve persons. This dose adjustment should be maintained for 2 weeks after stopping rifampicin as the inducing effect persists after discontinuation of a strong inducer. Alternative to rifampicin should be used where possible for INSTI-experienced persons with certain INSTI-associated resistance substitutions or clinically suspected INSTI resistance.
- RAL 400 or 800 mg bid.
- Al, Mg containing antacids not recommended with RAL 400 mg bid or 1200 mg qd. If co-administration with an antacid is unavoidable, calcium carbonate antacids can be used but only with RAL 400 mg bid.
- q Increase in concentration of active metabolite observed with RTV 100 mg bid alone but without significant effect on adrenal function. Caution is still warranted, use the lowest possible corticosteroid dose and monitor for corticosteroid side effects.
- r DRV/r decreased the exposure of active metabolite (beclometasone-17-monopropionate), no significant effect on adrenal function was
- S Risk of having elevated corticosteroid levels, Cushing's syndrome and adrenal suppression. This risk is present for oral and injected corticosteroid but also for topical, inhaled or eye drops administration.
- t Concentrations of norbuprenorphine increased.
- Alternative or additional contraceptive measures are recommended or, if used for hormone replacement therapy, monitor for signs of oestrogen deficiency.
- v Increase in ethinylestradiol with unboosted ATV.
- w No effect on ethinylestradiol as a combined oral contraceptive, but ethinylestradiol decreased when administered as a vaginal ring. Progestin decreased with both methods. Use with EFV is not recommended.
- x The daily dose of ethinylestradiol should not exceed 30 µg. Caution is advised, particularly in persons with additional risk factors for thromboembolic events.
- y European SmPC states a hormonal contraceptive should contain at least 30 µg ethinylestradiol.
- Z A study suggests a low risk of a clinically relevant pharmacokinetic interaction with low-hyperforin formulations (< 1 mg/day) of St John's Wort (hyperforin is the constituent responsible for induction of CYPs and P-gp). Coadministration may be considered with St John's Wort formulations that clearly state the hyperforin content and which have a total daily hyperforin dose of 1 mg or less.



# **Drug-drug Interactions between Analgesics and ARVs**

Ana	lgesics	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
	aspirin	$\leftrightarrow$	↔b																		
s	celecoxib	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	†a	†a	$\leftrightarrow$	↔b										
esic	diclofenac	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	†a	†a	$\leftrightarrow$	Εb										
analgesics	ibuprofen	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	†a	†a	$\leftrightarrow$	↔b										
ida	mefenamic acid	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	†a	†a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑11%	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔b
oido	naproxen	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	†a	†a	$\leftrightarrow$	↔b										
Non-opioid	nimesulide	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	†a	†a	$\leftrightarrow$	↔b										
Z	paracetamol	$\leftrightarrow$	↓3%	$\leftrightarrow$																	
	piroxicam	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑a	†a	$\leftrightarrow$	↔b										
	alfentanil	1	1	1	1	1	$\leftrightarrow$	1	<b>1</b>	↓	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	buprenorphine	1	↑67%c	1	↓11%c	↑~2%	$\leftrightarrow$	↓50%	↓25%	↓9%	$\leftrightarrow$	↑30%	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑35%	$\leftrightarrow$	$\leftrightarrow$	↑~5%
	codeine	↑d	↑d	↑d	↑d	↑d	$\leftrightarrow$	1	<b>1</b>	↓	$\leftrightarrow$	↑d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	diamorphine	↔e	↓e,f	↔e	↓e,f	↓e,f	$\leftrightarrow$	1	↔e	$\leftrightarrow$	↔e	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$							
	dihydrocodeine	1	<b>↓</b> ↑	1	<b>↓</b> ↑	<b>↓</b> ↑	$\leftrightarrow$	↓↑	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
ics	fentanyl	1	1	1	1	1	$\leftrightarrow$	<b>1</b>	<b>\</b>	↓	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
ges	hydrocodone	↓↑g	↓↑g	↓↑g	↓↑g	↓↑g	$\leftrightarrow$	↓↑h	↓↑ <mark>h</mark>	↓↑ <mark>h</mark>	$\leftrightarrow$	↓↑g	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
Opioid analgesics	hydromorphone	$\leftrightarrow$	Ţ	$\leftrightarrow$	↓	<b>↓</b>	$\leftrightarrow$	1	$\leftrightarrow$												
oid	methadone	↑?i	↔i	↑?	↓16%	↓53%i	↓5%	↓52%	↑6%	↓~50%	↓16%i	↑14%i	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔i	↓2%	↑7%	$\leftrightarrow$	$\leftrightarrow$	↑~5%
O	morphine	↔e	↓e,f	↔e	↓e,f	↓e,f	$\leftrightarrow$	1	↔e	$\leftrightarrow$	↔e	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$							
	oxycodone	1	1	1	1	↑160%	$\leftrightarrow$	1	1	Ţ	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	pethidine	1	<b>↓</b>	1	<b>1</b>	<b>↓</b>	$\leftrightarrow$	↓j	↓j	↓j	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	sufentanil	1	1	1	1	1	$\leftrightarrow$	1	1	Ţ	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	tapentadol	$\leftrightarrow$																			
	tramadol	↑d	↑d	↑d	↑d	↑d	$\leftrightarrow$	↓k	$\leftrightarrow$	↓k	$\leftrightarrow$	↑d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						

### Colour legend

No clinically significant interaction expected
These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

### Legend

Potential elevated exposure of the analgesic
 Potential decreased exposure of the analgesic

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd) DRV/c DRV co-formulated with COBI (800/150 mg qd) CAB/RPV CAB and RPV im long acting injections

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

## Interactions with ABC, FTC, 3TC, ZDV

ABC: decreased methadone exposure

FTC, 3TC: no clinically relevant interactions expected.

ZDV: potential additive haematological toxicity with ibuprofen, naproxen. ZDV: Moderately increased ZDV exposure with methadone; monitor

for toxicity.

### Interactions with ibalizumab

None

### Comments

- a Clinical significance unknown. Use the lowest recommended dose particularly in individuals with risk factors for CVD, those individuals at risk of developing gastrointestinal complications, persons with hepatic or renal impairment, and in elderly persons.
- b Potential risk of nephrotoxicity which is increased if NSAID is used for a long duration, if the person has a pre-existing renal dysfunction, a low body weight or receives other drugs that may increase TDF exposure. Concurrent use of NSAIDs with TDF warrants monitoring of renal function.
- c Concentrations of norbuprenorphine increased.
- d Potential decrease of the analgesic effect due to the reduced conversion to the active metabolite.
- e Inhibition of P-gp by RTV, COBI or ETV could potentiate the effect of opiate in the CNS.
- Concentrations of parent drug decreased but concentrations of active metabolite increased.
- g Concentrations of hydrocodone increased, but concentrations of active metabolites (norhydrocodone and hydromorphone) decreased. The clinical significance of this is unclear.
- h Concentrations of hydrocodone decreased, but concentrations norhydrocodone increased. The clinical significance of this is unclear.
- Both drugs can potentially prolong the QT interval, ECG monitoring recommended.
- j Concentrations of parent drug decreased and concentrations of neurotoxic metabolite increased.
- k Concentrations of parent drug decreased but no change in concentrations of more active metabolite.

### **Further Information**

For additional drug-drug interactions and for more detailed pharmacokinetic interaction data and dosage adjustments, please refer to: http://www.hiv-druginteractions.org (University of Liverpool)



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# **Drug-drug Interactions between Anticoagulants/Antiplatelet Agents and ARVs**

	coagulants ntiplatelets	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
	acenocoumarol	$\leftrightarrow$	ļ	$\leftrightarrow$	Ţ	1	$\leftrightarrow$	↑or↓	1	↓	$\leftrightarrow$	↓	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	apixaban	↑a	†a	†a	†a	†a	$\leftrightarrow$	ļ	↓	↓	$\leftrightarrow$	↑?	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	†a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	argatroban	$\leftrightarrow$																			
	betrixaban	↑b,c	↑b,c	↑c	↑c	↑b,c	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	↔b	↔b	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔b	$\leftrightarrow$	↑c	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
w	dabigatran	↑d	↑e	↑d	↑e	↑?	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	†?	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
lant	dalteparin	$\leftrightarrow$																			
Anticoagulants	edoxaban	↑f	↑f	↑f	↑f	↑f	$\leftrightarrow$	↑f	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
ntice	enoxaparin	$\leftrightarrow$																			
<	fondaparinux	$\leftrightarrow$																			
	heparin	$\leftrightarrow$																			
	phenprocoumon	1	↑or↓g	1	↑or↓	↑or↓	$\leftrightarrow$	↓	↑or↓	Ţ	$\leftrightarrow$	↑or↓	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	rivaroxaban	1	1	1	1	1	$\leftrightarrow$	↓	↓	Ţ	$\leftrightarrow$	↑?	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	warfarin	1	↑or↓g	1	Ţ	↓	$\leftrightarrow$	↑or↓	1	↑or↓	$\leftrightarrow$	↓	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
nts	aspirin	$\leftrightarrow$																			
agents	clopidogrel	↓h	↓h	↓h	↓h	↓h	$\leftrightarrow$	↓h E	↓h	↑i E	$\leftrightarrow$	↓h	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
telet	dipyridamole	1	↓j	$\leftrightarrow$	Ţ	1	$\leftrightarrow$	Ţ	<b>↓</b>	$\leftrightarrow$											
Antiplatelet	prasugrel	↓k	↓k	↓k	↓k	↓k	$\leftrightarrow$	↓k	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
Ant	ticagrelor	1	1	1	1	1	$\leftrightarrow$	↓	1	↓ ·	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						

### Colour legend

No clinically significant interaction expected These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

### Legend

Potential elevated exposure of the anticoagulant/

antiplatelet agent

Potential decreased exposure of the anticoagulant/

antiplatelet agent No significant effect

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd) DRV/c DRV co-formulated with COBI (800/150 mg qd) CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

### Interactions with ABC, FTC, 3TC, ZDV

ABC: may potentially reduce the pharmacodynamic effect of clopidogrel. FTC, 3TC, ZDV: no clinically relevant interactions expected.

### Interactions with ibalizumab

None

### Comments

- US label suggests to use apixaban at a reduced dose (2.5 mg bid) if needed.
- b Both drugs can potentially prolong the QT interval, ECG monitoring recommended.
- C US label recommends to use a reduced initial betrixaban dose of 80 mg followed by 40 mg qd.
- d Dabigatran should be reduced to 100 mg bid in persons with normal renal function and to 75 mg bid in case of moderate renal impairment. Coadministration should be avoided in case of severe renal impairment.
- No significant increase in DRV/r exposure when administered simultaneously with dabigatran in persons with no renal impairment.
- f European label advises to consider a dose reduction of edoxaban from 60 mg to 30 mg, however, US label recommends no dose modification.
- g Unboosted ATV predicted to increase the anticoagulant, monitor INR and adjust the anticoagulant dosage accordingly.
- h Decreased conversion to active metabolite leading to non-responsiveness to clopidogrel. An alternative to clopidogrel should be considered.
- Increase in amount of active metabolite via induction of CYP3A4 and CYP2B6.
- j Unboosted ATV predicted to increase dipyridamole exposure due to UGT1A1 inhibition.
- k Reduced active metabolite, but without a significant reduction in prasugrel activity.

### **Further Information**



# **Drug-drug Interactions between Antidepressants and ARVs**

An	idepressants	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
NaSSA	mirtazapine	↑a	†a	1	1	↑a	$\leftrightarrow$	ļ	ļ	1	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	citalopram	↑ a,b	↑ a,b	1	1	↑ a,b	$\leftrightarrow$	<b>↓</b>	<b>1</b>	<b>1</b>	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	escitalopram	↑ a,b	↑ a,b	1	1	↑ a,b	$\leftrightarrow$	↓	<b>\</b>	1	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	fluoxetine	1	1	1	1	†a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
SSRI	fluvoxamine	1	1	1	1	†a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	Е	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
0,	paroxetine	<b>↑↓?</b>	<b>↑↓?</b>	<b>↑↓?</b>	↓39%	<b>↑↓?</b>	$\leftrightarrow$	$\leftrightarrow$	↑3%	$\leftrightarrow$	↑↓?	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$							
	sertraline	1	↓	1	↓49%	↓a	$\leftrightarrow$	↓39%	<b>\</b>	↓	$\leftrightarrow$	↓7%	$\leftrightarrow$	↑9%	$\leftrightarrow$						
	vortioxetine	↑c	↑c	↑ <b>c</b>	↑c	↑c	$\leftrightarrow$	↑c	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
	desvenlafaxine	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
SNRI	duloxetine	1	↑↓	1	↑↓	↑↓	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
S	milnacipran	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	venlafaxine	†a	↑a	1	1	†a	$\leftrightarrow$	<b>↓</b>	1	1	↔a	↔a	D	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	amitriptyline	1	1	1	1	↑ a,b	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
	clomipramine	↑ a,b	↑ a,b	↑ <b>b</b>	↑ <b>b</b>	↑ a,b	$\leftrightarrow$	1	1	$\downarrow$	$\leftrightarrow$	↑b	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	desipramine	†a	†a	1	1	↑5% <mark>a</mark>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
TCA	doxepin	1	1	1	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
	imipramine	↑ a,b	↑ a,b	↑ <b>b</b>	↑ <b>b</b>	↑ a,b	$\leftrightarrow$	↓	$\downarrow$	$\downarrow$	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	↑b	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	nortriptyline	†a	†a	1	1	†a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	trimipramine	†a	†a	1	1	†a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
TeCA	maprotiline	†a	↑a	1	1	†a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Te	mianserin	† <b>a</b>	↑a	1	1	†a	$\leftrightarrow$	↓	$\downarrow$	$\downarrow$	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	agomelatine	$\leftrightarrow$	↓	$\leftrightarrow$	<b>↓</b>	<b>↓</b>	$\leftrightarrow$														
	bupropion	$\leftrightarrow$	<b>↓</b>	$\leftrightarrow$	<b>↓</b>	↓57%	$\leftrightarrow$	↓55%	$\leftrightarrow$	1	$\leftrightarrow$	↑?	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	nefazodone	1	1	1	1	1	Е	ţΕ	↓E	↓E	Е	Е	Е	Е	$\leftrightarrow$	Е	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
SIS	phenelzine	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Others	reboxetine	1	1	1	1	1	$\leftrightarrow$	↓	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	St John's wort	Dd	Dd	Dd	Dd	Dd	Dd	Dd	Dd	Dd	Dd	Dd	Dd	Dd	$\leftrightarrow$	Dd	De	Dd	D	Dd	$\leftrightarrow$
	tranylcypromine	1	1	1	1	1	$\leftrightarrow$	↓	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	trazodone	↑ a,b	↑ a,b	1	1	↑ a,b	$\leftrightarrow$	↓	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						

### Colour legend

No clinically significant interaction expected
These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

Legend

Potential elevated exposure of the antidepressant
 Potential decreased exposure of the antidepressant

→ No significant effect

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd) DRV/c DRV co-formulated with COBI (800/150 mg qd) CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

NaSSA noradrenergic specific serotonergic antidepressant

**SSRI** selective serotonin reuptake inhibitors

SNRI serotonin and norepinephrine reuptake inhibitors

TCA tricyclic antidepressants
TeCA tetracyclic antidepressants

### Interactions with ABC, FTC, 3TC, ZDV

ABC, FTC, 3TC, ZDV: no clinically relevant interactions expected.

### Interactions with ibalizumab

None

### Comments

- a Caution as both drugs can induce QT interval prolongation.
- b ECG monitoring is recommended.
- c Based on the patient clinical response, a lower dose of vortioxetine may be needed in poor CYP2D6 metabolizers in the presence of a strong CYP3A4 inhibitor.
- d A study suggests a low risk of a clinically relevant pharmacokinetic interaction with low-hyperforin formulations (< 1 mg/day) of St John's Wort (hyperforin is the constituent responsible for induction of CYPs and P-gp). Coadministration may be considered with St John's Wort formulations that clearly state the hyperforin content and which have a total daily hyperforin dose of 1 mg or less.
- e The European SmPC recommends DTG 50 mg bid in persons without INSTI resistance. The US Prescribing Information recommends that co-administration should be avoided as there are insufficient data to make dosing recommendations.

### **Further Information**



# **Drug-drug Interactions between Antihypertensives and ARVs**

Ant	ihypertensives	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
	captopril	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$															
	cilazapril	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$															
	enalapril	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$															
tors	fosinopril	$\leftrightarrow$	1	$\leftrightarrow$	1	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
hibi	lisinopril	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$															
ACE inhibitors	perindopril	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$															
AC	quinapril	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$															
	ramipril	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$															
	trandolapril	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$															
Ŋ	candesartan	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$															
nist	eprosartan	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$															
tago	irbesartan	$\leftrightarrow$	Ţ	$\leftrightarrow$	J.	<b>↓</b>	$\leftrightarrow$	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$							
Angiotensin antagonists	losartan	$\leftrightarrow$	↓a	$\leftrightarrow$	↓a	↓a	$\leftrightarrow$	↑b	↑b	$\leftrightarrow$	↓a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$							
nsir	olmesartan	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$									
jote	telmisartan	$\leftrightarrow$	· ↑	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$									
Ang	valsartan	1	1	<b>↑</b>	<b>1</b>	1	↔	↔	↔	↔	↔	·	$\leftrightarrow$	↔	$\leftrightarrow$	$\leftrightarrow$	↔	↔	↔	↔	$\leftrightarrow$
	atenolol	↑c	↔C	1	↔	↔C	↔	↔	← →	↔		↔		<b>↑</b>	↔	$\leftrightarrow$	<b>1</b>	1		←→	← →
	bisoprolol	↑c	↑C	<u> </u>	<b>1</b>	↑C	↔	<b>1</b>	<b>1</b>	<b>1</b>	←→	←→	←→	↔	↔	$\leftrightarrow$	↔	<u> </u>			$\leftrightarrow$
	carvedilol	↑c	↑↓c	<u> </u>	↑↓	↑↓c	↔	↑↓	↑↓	<b>↓</b>				↔	↔	← →	←→	<u> </u>		←→	
ទ	labetalol	↑C	↓c	→ ·	1 *	↓c	$\leftrightarrow$	1	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
ß blockers	metoprolol	↑c	↑C	1	<b>↑</b>	↑C	↔	<b>↓</b>	<b>↓</b>	↔	↔	↔	←→	↔	↔	↔	↔	1	↔	↔	←→
old	nebivolol	↑C	↑C	<u> </u>	<u> </u>	↑C	↔	↔	←	↔				↔	↔	$\leftrightarrow$	←→	<u> </u>	↔	←→	
8	oxprenolol	↑c	†c	↔	l l	†c	↔			$\leftrightarrow$	$\leftrightarrow$		$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	pindolol				<b>→</b>			<b>↓</b>	<b>↓</b>			<b>↔</b>								$\leftrightarrow$	
		↑c	↑C	<b>↑</b>	<u> </u>	↑c	↔	↔	↔	↔	↔	<b>↔</b>	<b>↔</b>	↔	↔	<b>↔</b>	↔	↑ •	<b>↔</b>		↔
	propranolol amlodipine	↑c	↑C	<b>1</b>	1	↑C	↔	<b>↔</b>	↔	↔	↔	↔	<b>↔</b>	↔	↔	<b>↔</b>	↔	↑ •	↔	↔	↔
S	diltiazem	↑d ↑d	↑d ↑d	<b>↑</b>	<u> </u>	↑e	↔ E	↓69%	↓E	<b>1</b>	↔ E	↔ E	↔ E	↔ E	$\leftrightarrow$	↔ E	↔	↑ •	↔	$\leftrightarrow$	$\leftrightarrow$
Icium channel blockers	felodipine	↑d	↑d	↑ ↑	<u> </u>	↑e	↔	10970		<b>+</b>	↔	↔	↔	↔	$\leftrightarrow$	↔	$\leftrightarrow$ $\leftrightarrow$	↑ ↑	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
old	lacidipine	↑d	↑d	↑ ↑	<u> </u>	†e †e	$\leftrightarrow$	<b>1</b>	<b>↓</b>	<b>↓</b>	↔f	↔f	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔f	$\leftrightarrow$	↑ ↑	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Jue	lercanidipine	↑	↑	<u> </u>	1	1	$\leftrightarrow$	<b>1</b>	<b>1</b>	<b>1</b>	$\leftrightarrow$	←	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	←	$\leftrightarrow$	<u> </u>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
chai	nicardipine	↑d	↑d	<u> </u>	<u> </u>	↑e	E	1	↓E	<b>1</b>	Ef	Ef	E	$\leftrightarrow$	$\leftrightarrow$	Ef	$\leftrightarrow$	<u> </u>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
E	nifedipine	↑d	↑d	1	<u> </u>	↑e	↔	1	↓ <b>∟</b>	1	↔	↔	↔	$\leftrightarrow$	$\leftrightarrow$	↔	$\leftrightarrow$	<u> </u>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Calci	nisoldipine	↑d	↑d	<u> </u>	<u> </u>			1	1	1		$\leftrightarrow$						<u> </u>		$\leftrightarrow$	
0	verapamil	↑d	↑d	<u> </u>	<u> </u>	†e †e	↔ E	<b>1</b>	↓E	<b>↓</b>	↔ E	E	↔ E	↔ E	$\leftrightarrow$	↔ E	$\leftrightarrow$ $\leftrightarrow$	<u> </u>	$\leftrightarrow$	E	↔ E
	amiloride	γ <b>α</b> ↔	γ <b>ω</b>	↔	→ ·	←	↔	<b>→</b>	↔	↔	↔	↔	↔	↔	$\leftrightarrow$	↔		→ ·	$\leftrightarrow$	↔	↔
	bendroflu-																1				
	methiazide chlortalidone	$\leftrightarrow$	<b>↔</b>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$															
		←→	←→	↔	←→	←→	$\leftrightarrow$	<b>↔</b>	↔	←→	$\leftrightarrow$	←→	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
tics	eplerenone	1	1	1	1	1	$\leftrightarrow$	<u></u>	↓ ↓	<u></u>	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	↔						
Diuretics	furosemide hydrochloro-	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	E															
	thiazide	<b>↔</b>	<b>↔</b>	<b>↔</b>	<b>↔</b>	<b>↔</b>	$\leftrightarrow$	<b>↔</b>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	<b>↔</b>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	indapamide	1	1	1	1	1	$\leftrightarrow$	<b>+</b>	<b>1</b>	<u></u>	$\leftrightarrow$	↔	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	torasemide	$\leftrightarrow$	<b>1</b>	$\leftrightarrow$	<u></u>	<b>↓</b>	$\leftrightarrow$	1	1	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	<u></u>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	xipamide	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$															
	clonidine	←→	←→	←→	←→	←→	$\leftrightarrow$	↔	↔	↔	$\leftrightarrow$	←→	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	doxazosin	1	1	1	1	1	$\leftrightarrow$	<u></u>	Ţ	<u></u>	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
(0	hydralazine	$\leftrightarrow$	↔g	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔h										
Others	methyldopa	$\leftrightarrow$	↔g	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
ō	moxonidine	↔	↔	↔	↔	↔	$\leftrightarrow$	↔	↔	↔	$\leftrightarrow$	↔	$\leftrightarrow$	$\leftrightarrow$	↑?						
	prazosin	↑?	↑?	↑?	†?	↑?	$\leftrightarrow$	↓?	↓?	↓?	$\leftrightarrow$	1	←→	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑?	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	sacubitril	1	1	1	1	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	Е	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1
	spironolactone	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$															

### Colour legend

No clinically significant interaction expected
These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

### Legend

Potential elevated exposure of the antihypertensive
 Potential decreased exposure of the antihypertensive

→ No significant effect

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd) DRV/c DRV co-formulated with COBI (800/150 mg qd) CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

**Note:** although some drug interactions are predicted to potentially require a dosage adjustment based on the drug's metabolic pathway, clinical experience with a particular antihypertensive and ARV drug may indicate that dosage adjustments are not an a priori requirement

### Interactions with ABC, FTC, 3TC, ZDV

ABC, FTC, ZDV: no clinically relevant interactions expected. 3TC: increased 3TC exposure with atenolol and amiloride. 3TC: increased exposure of atenolol and amiloride.

### Interactions with ibalizumab

None

### Comments

- a Parent drug concentrations decreased but active metabolite increased.
- b Parent drug concentrations increased but active metabolite decreased.
- c Risk of PR interval prolongation.
- d ECG monitoring recommended.
- Use with caution as both LPV and calcium channel blockers prolong the PR interval. Clinical monitoring is recommended.
- f Caution as both drugs can induce QT interval prolongation.
- Use with caution in persons with a history of postural hypotension or on concomitant medicinal products known to lower blood pressure, and those at increased risk of cardiovascular events.
- h Hydralazine has some nephrotoxic potential. If co-administration is unavoidable, monitor renal function closely.

### **Further Information**



# **Drug-drug Interactions between Anti-malarial Drugs and ARVs**

Ant	imalarial drugs	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
	amodiaquine	1	1	$\leftrightarrow$	1	1	$\leftrightarrow$	↑ a	<b>↓?</b>	↓29% <mark>a</mark>	$\leftrightarrow$										
	artemisinin	1	1	1	1	1	D	ļ	ţD	ţD	D	D	D	D	$\leftrightarrow$	D	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	atovaquone	$\leftrightarrow$	↓10%	$\leftrightarrow$	↓ b	↓74%b	$\leftrightarrow$	↓75% <mark>b</mark>	↓E55% <mark>b</mark>	↓ b	$\leftrightarrow$										
	chloroquine	↔ c,d	↔ c,d	↔ d	↔ d	↔ c,d	$\leftrightarrow$	↔ <b>e</b>	↔f	↔ f	↔ c,g	c,g	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔ c,g	$\leftrightarrow$	↔ d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	clindamycin	1	1	1	1	1	$\leftrightarrow$	↓	<b>\</b>	<b>↓</b>	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
.ngs	doxycycline	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↓?	↓?	↓?	$\leftrightarrow$										
line dı	halofantrine	↑ <b>g</b>	↑ g	1	1	↑ g	$\leftrightarrow$	↓	1	<b>↓</b>	↔ g	↔ c,g	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔ g	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
second line drugs	hydroxy- chloroquine	↑ c,g	↑ c,g	1	1	↑ c,g	$\leftrightarrow$	↔ <b>e</b>	<b>1</b>	1	↔g	↔ c,g	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	⇔g	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
and s	lumefantrine	↑ c,g	↑ c,g	1	↑175%	↑382% c,g	$\leftrightarrow$	↓~40%	1	↓D46%	↔g	↔ g	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔g	↑10%	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
First line and	mefloquine	↑ c,g	↑ c,g	1	1	↓28%c,g	$\leftrightarrow$	Ţ	1	↓	↔ g	↔ g	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔g	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Ē	piperaquine	↑ c,g	↑ c,g	↑ c	↑ c	↑ c,g	Е	<b>\</b>	<b>\</b>	1	Εg	↔ g	Е	Е	$\leftrightarrow$	↔ g	$\leftrightarrow$	↑ c	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	primaquine	↔ g	↔g	$\leftrightarrow$	$\leftrightarrow$	⇔g	$\leftrightarrow$	↔ h	↔h	↔ h	↔ g	↔ g	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔g	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	proguanil	$\leftrightarrow$	↓41%b	$\leftrightarrow$	↑p	↓38%b	$\leftrightarrow$	↓44%b	↓E55%b	↓ b	$\leftrightarrow$										
	pyrimethamine	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	quinine	↑ c,g	↑ c,g	1	1	↓56% <b>c</b> ,g	$\leftrightarrow$	<b>\</b>	1	1	⇔g	↔ c,g	Е	$\leftrightarrow$	$\leftrightarrow$	↔ g	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	sulfadoxine	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						

### Colour legend

No clinically significant interaction expected
These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration

Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

# Legend

Potential elevated exposure of the antimalarial drug
 Potential decreased exposure of the antimalarial drug

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd)
DRV/c DRV co-formulated with COBI (800/150 mg qd)
CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

### Interactions with ABC, FTC, 3TC, ZDV

ABC: no clinically relevant interactions expected.

FTC: increased FTC exposure with pyrimethamine, sulfadoxine.
3TC: increased 3TC exposure with pyrimethamine, sulfadoxine.
ZDV: potential additive haematological toxicity with amodiaquine, atovaquone, primaquine, pyrimethamine, sulfadoxine.

### Interactions with ibalizumab

None

### Comments

- a Liver toxicity.
- b Take with high fat meal, consider dose increase.
- c ECG monitoring is recommended.
- d Chloroquine concentrations may increase, but to a moderate extent. No dose adjustment is required but monitor toxicity.
- Chloroquine/hydroxychloroquine concentrations may increase or decrease. No dose adjustment is required but monitor toxicity and efficacy.
- f Chloroquine concentrations may decrease, but to a moderate extent. No dose adjustment is required but monitor efficacy.
- g Caution as both drugs can induce QT interval prolongation.
- h Increase of haemotoxic metabolites.

### **Further Information**



# **Drug-drug Interactions between Anti-tuberculosis Drugs and ARVs**

	Anti-tuberculosis drugs		ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
First line and second line drugs	amikacin	$\leftrightarrow$	↔ a																		
	bedaquiline	↑ b	↑ b	1	1	↑62% b	$\leftrightarrow$	↓18%	1	↑3%	↔ b	↔ b	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔ b	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	capreomycin	$\leftrightarrow$	↑ c	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑Ea														
	clofazimine	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	Е	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	Е	Е	Е	Е	$\leftrightarrow$	Е	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	cycloserine	$\leftrightarrow$																			
	delamanid	d	d	d	d	d	$\leftrightarrow$	↔ e	$\leftrightarrow$	$\leftrightarrow$	↔f	↔f	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔f	$\leftrightarrow$	d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	ethambutol	$\leftrightarrow$																			
	ethionamide	$\leftrightarrow$																			
	isoniazid	$\leftrightarrow$																			
	kanamycin	$\leftrightarrow$	↔ a																		
	linezolid	$\leftrightarrow$																			
	moxifloxacin	↑ b	↓ b	$\leftrightarrow$	↓	↓b	$\leftrightarrow$	1	1	$\leftrightarrow$	↔ b	↔ b	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔ b	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	para-aminosali- cylic acid	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑E														
	pretomanid	↓b	↓ b	1	↓	↓17% b	$\leftrightarrow$	↓35%	1	↓	↔ b	↔ b	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔ b	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	pyrazinamide	$\leftrightarrow$																			
	rifabutin	↑ D g	†h	↑ D g	↑h	↑ <b>h</b>	D50%i	↓38%j	D37%	↑17%	D42%k	D30%	- 1	D38%	$\leftrightarrow$	D	$\leftrightarrow$	↑ D g	E19%	Dm	$\leftrightarrow$
	rifampicin	D	D72%	D	D57%	D75%n	D82%	D26%s	D	D58%	D80%	D82%	Do	D75%	D59%	D	D54%p	D	D40%q	Dm	D12%
	rifapentine	D	D	D	D	D	D	D	D	D	D	D	D o	D	D	D	Dr	D	D	Dm	$\leftrightarrow$
	streptomycin	$\leftrightarrow$	↔ a																		

### Colour legend

No clinically significant interaction expected
These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

### Legend

Potential elevated exposure of the anti-tuberculosis drug
 Potential decreased exposure of the anti-tuberculosis drug

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd)
DRV/c DRV co-formulated with COBI (800/150 mg qd)

CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

### Interactions with ABC, FTC, 3TC, ZDV

ABC: potentially moderately increased ABC exposure with rifampicin but no a priori dose adjustment required.

FTC: Exposure of FTC and/or capreomycin may increase when co-administered. Monitor renal function as appropriate.

FTC: Exposure of FTC and/or para-aminosalicylic acid may increase when co-administered.

3TC: Exposure of 3TC and/or capreomycin may increase when co-administered. Monitor renal function as appropriate.

3TC: Exposure of 3TC and/or para-aminosalicylic acid may increase when co-administered.

ZDV: Rifampicin decreased ZDV AUC by 47%. Co-administration is not recommended in ZDV's European label, but the US label says routine dose modification is not warranted.

### Interactions with ibalizumab

None

### Comments

- Co-administration should be avoided due to the risk of additive tubular toxicity, but if such use is unavoidable, closely monitor renal function.
- b Both drugs can potentially prolong the QT interval, ECG monitoring recommended.
- Aminoglycosides are nephrotoxic (risk is dose and treatment duration related). Renal function should be monitored as clinically appropriate and the dosage of the ARV adjusted accordingly.

- d Co-administration is expected to increase concentrations of DM-6705, a delamanid metabolite which is associated with QT prolongation. Frequent ECG monitoring is recommended.
- e A higher rate of neuropsychiatric adverse effects (e.g., euphoric mood and abnormal dreams) was observed with delamanid plus EFV com- pared to either drug alone.
- f RPV, FTR and DM-6705 (a delamanid metabolite) can potentially prolong the QT interval, ECG monitoring recommended.
- g Reduce rifabutin to 150 mg 3 times per week.
- Reduce rifabutin to 150 mg qd. Monitoring for rifabutin-related toxicities (i.e. uveitis or neutropenia) is advised with daily administration of rifabutin.
- The product label for DOR recommends to increase DOR dosage to 100 mg bid when co-administered with rifabutin. DOR should be kept at 100 mg bid for at least another 2 weeks following cessation of rifabutin due to the persisting inducing effect upon discontinuation of a moderate/ strong inducer.
- Increase rifabutin to 450 mg qd.
- k The RPV dose should be increased to 50 mg qd during co-administration (and decreased to 25 mg qd when rifabutin is stopped). Note, it is recommended to maintain RPV 50 mg qd for at least another 2 weeks following cessation of rifabutin due to the persisting inducing effect upon discontinuation of a moderate/strong inducer.
- Increase MVC to 600 mg bid in absence of PI. With PI (except TPV/r, FPV/r), give MVC 150 mg bid.
- m Rifamycins decrease TAF exposure when given 25 mg. However, the intracellular tenofovir diphosphate (active entity) concentrations are likely to be higher than those observed with TDF even without rifampicin [1] suggesting that usage of TAF 25 mg qd may be acceptable.
- n If no other option use RTV 400 mg bid or double dose LPV/r.
- O Give MVC 600 mg bid.
- A dose adjustment of DTG to 50 mg bid is recommended in treatmentnaïve or INSTI-naïve persons. This dose adjustment should be maintained for 2 weeks after stopping rifampicin as the inducing effect persists after discontinuation of a strong inducer. Alternatives to rifampicin should be used where possible for INSTI-experienced persons with certain INSTI-associated resistance substitutions or clinically suspected INSTI resistance.
- RAL 400 or 800 mg bid.
- Based on DTG interactions studies with rifabutin and rifampicin, consider administering DTG at 50 mg bid in the presence of rifapentine. This dose adjustment should be maintained for 2 weeks after stopping rifapentine as the inducing effect persists after discontinuation of a strong inducer.
- S Efavirenz should be used at 600 mg qd in presence of rifampicin (in absence of rifampicin, efavirenz can be used at 400 mg qd or 600 mg qd).

### **Further Information**



# **Drug-drug Interactions between Anxiolytics and ARVs**

Anx	iolytics	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
	alprazolam	1	1	1	1	1	$\leftrightarrow$	<b>\</b>	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	chlor-diazepoxide	1	1	1	1	1	$\leftrightarrow$	<b>↓</b>	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
BZD	clonazepam	1	1	1	1	1	$\leftrightarrow$	<b>\</b>	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	lorazepam	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	oxazepam	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
SSRI	escitalopram	†a	†a	1	1	†a	$\leftrightarrow$	<b>\</b>	1	1	↔b	↔b	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔b	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
SS	paroxetine	↑↓?	↑↓?	<b>↑</b> ↓?	↓39%	↑↓?	$\leftrightarrow$	$\leftrightarrow$	↑3%	$\leftrightarrow$	↑↓?	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$							
SNRI	duloxetine	1	↑↓	1	↑↓	$\uparrow\downarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
SN	venlafaxine	↑ b	↑ <b>b</b>	1	1	↑ <b>b</b>	$\leftrightarrow$	<b>\</b>	1	1	↔b	↔b	D	$\leftrightarrow$	$\leftrightarrow$	↔b	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Others	buspirone	1	1	1	1	1	$\leftrightarrow$	1	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
Oth	hydroxyzine	↑a,b	↑a,b	↑a,b	↑a,b	↑a,b	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										

Colour legend

No clinically significant interaction expected
These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration

Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

Legend

↑ Potential elevated exposure of the anxiolytic therapy
Potential decreased exposure of the anxiolytic therapy

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd) DRV/c DRV co-formulated with COBI (800/150 mg qd) CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

BZD benzodiazepines

**SSRI** selective serotonin reuptake inhibitors

**SNRI** serotonin and norepinephrine reuptake inhibitors

Interactions with ABC, FTC, 3TC, ZDV

ABC, FTC, 3TC, ZDV: no clinically relevant interactions expected.

#### Interactions with ibalizumab

None

#### Comments

- a ECG monitoring is recommended.
- b Caution as both drugs can induce QT interval prolongation.

#### **Further Information**



# **Drug-drug Interactions between Bronchodilators (for COPD) and ARVs**

Bro	nchodilators	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
	aclidinium bromide	$\leftrightarrow$																			
LAMA	glycopyrronium bromide	$\leftrightarrow$																			
ΓĀ	tiotropium bromide	$\leftrightarrow$																			
	umeclidinium bromide	1	1	1	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
SAMA	ipratropium	$\leftrightarrow$																			
	formoterol	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	indacaterol	↑b	↑b	↑b	↑b	↑b	$\leftrightarrow$	1	1	1	$\leftrightarrow$	↑b	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
LABA	olodaterol	1	1	1	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
_	salmeterol	1	1	1	1	1	$\leftrightarrow$	1	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	vilanterol	1	1	1	1	1	$\leftrightarrow$	1	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
SABA	salbutamol (albuterol)	$\leftrightarrow$																			
S)	terbutaline	$\leftrightarrow$																			
MX	aminophylline	$\leftrightarrow$	↓	$\leftrightarrow$	1	1	$\leftrightarrow$														
Σ	theophylline	$\leftrightarrow$	↓	$\leftrightarrow$	1	1	$\leftrightarrow$														
PDE4	roflumilast	1	1	1	1	1	$\leftrightarrow$	↓	↓	ţ	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	beclometasone	↑c	↑ <b>c</b>	↑?c	↓11%d	↑c	$\leftrightarrow$	↑c	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
	budesonide	↑e	↑e	↑e	↑e	↑e	$\leftrightarrow$	↓	↓	↓	$\leftrightarrow$	↑e	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
<u>S</u>	ciclesonide	↑f	↑ <b>f</b>	↑f	↑f	↑f	$\leftrightarrow$	↑f	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
	fluticasone	↑e	↑e	↑e	↑e	↑e	$\leftrightarrow$	↓	<b>\</b>	↓	$\leftrightarrow$	↑e	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	mometasone	↑e	↑e	↑e	↑e	↑e	$\leftrightarrow$	<b>↓</b>	↓	↓	$\leftrightarrow$	↑e	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						

#### Colour legend

No clinically significant interaction expected
These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

#### Legend

Potential elevated exposure of the bronchodilator
 Potential decreased exposure of the bronchodilator

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd) DRV/c DRV co-formulated with COBI (800/150 mg qd) CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

ICS inhaled corticosteroids
LABA long-acting β2 agonists

LAMA long-acting muscarinic antagonists MX methylxanthines

PD4 phosphodiesterase 4 inhibitors SABA short-acting β2 agonists

SAMA short-acting muscarinic antagonists

#### Interactions with ABC, FTC, 3TC, ZDV

ABC, FTC, 3TC, ZDV: no clinically relevant interactions expected.

#### Interactions with ibalizumab

None

#### Comments

- a Caution as both drugs can induce QT interval prolongation.
- b Exposure can be increased up to 2-fold however this increase does not raise any concerns based on indacaterol's safety data.
- c Increase in concentration of active metabolite observed with RTV 100 mg bid alone but without significant effect on adrenal function. Caution is still warranted, use the lowest possible corticosteroid dose and monitor for corticosteroid side effects.
- DRV/r decreased the exposure of active metabolite (beclometasone-17-monopropionate), no significant effect on adrenal function was seen.
- e Risk of having elevated corticosteroid levels, Cushing's syndrome and adrenal suppression. This risk is present for oral and injected corticosteroid but also for topical, inhaled or eye drops administration.
- f No dose adjustment required but monitor closely, especially for signs of Cushing's syndrome when using a high dose or prolonged administration.

#### **Further Information**

For additional drug-drug interactions and for more detailed pharmacokinetic interaction data and dosage adjustments, please refer to: http://www.hiv-druginteractions.org (University of Liverpool)

#### Note

Fixed dose combinations are available for LAMA + LABA + ICS, e.g., mometasone + indacaterol + glycopyrronium fluticasone + umeclidinium + vilanterol formoterol + glycopyrronium + beclometasone

budesonide + formoterol + glycopyrronium



# **Drug-drug Interactions between Contraceptives and ARVs**

Cor	traceptives	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
Es	ethinylestradiol (COC, TS, VR)	↑1%a	↓19%b	↓30%	↓44% <mark>a</mark>	↓42% <mark>a</mark>	↓2%	С	↑22%	↓20%	↑14%	↑40%d	↓<1%	↑4%	↑2%	$\leftrightarrow$	↑3%	↓25%e	↓2%	↑11%	$\leftrightarrow$
	desogestrel (COC)	1	↑f,b	1	† <b>g</b>	<b>†</b> 9	$\leftrightarrow$	↓h	↓	↓	$\leftrightarrow$	↔d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑e,f	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	desogestrel (POP)	1	1	1	1	1	$\leftrightarrow$	↓h	<b>↓</b>	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	drospirenone (COC)	↑130%	↑f,b	↑58%g	<b>↑</b> g	<b>†</b> 9	$\leftrightarrow$	↓h	1	ļ	$\leftrightarrow$	↔d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑e,f	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	etonogestrel (IP)	1	1	1	1	↑52%	$\leftrightarrow$	↓63% h	1	1	↑18%	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑ 19-54%	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	etonogestrel (VR)	1	↑~71% i	↑i	↑i	↑i	$\leftrightarrow$	↓~79% h	1	1	$\leftrightarrow$	↔d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑i	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	gestodene (COC)	1	↑f,b	1	<b>†</b> g	<b>†</b> 9	$\leftrightarrow$	↓h	1	1	$\leftrightarrow$	↔d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑e,f	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	levonorgestrel (COC)	↓8%	↑f,b	1	<b>↑</b> g	<b>†</b> 9	↑21%	↓h	<b>↓</b>	1	$\leftrightarrow$	↔d	↓2%	$\leftrightarrow$	↑12%	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Su	levonorgestrel (IP)	1	1	1	1	1	$\leftrightarrow$	↓57%h	1	↑14%	↑28%	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Progestins	levonorgestrel (IUD)	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Ā	levonorgestrel (POP)	1	1	1	1	1	$\leftrightarrow$	↓h	<b>1</b>	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	medroxy-proges- terone (POI)	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑~70%	$\leftrightarrow$	↔n	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	norelgestromin (TS)	1	↑f,b	1	<b>↑</b> g	↑83% <b>g</b>	$\leftrightarrow$	↓h	↓	ļ	$\leftrightarrow$	↔d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑e,f	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	norethisterone (COC)	1	↑f,j	1	↓14%g	↓17%g	$\leftrightarrow$	↓h	↓5%	↓19%	↓11%	↑8% <mark>d</mark>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑e,f	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	norethisterone (POI)	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	norethisterone (POP)	1	↑50%	1	↑50%	↑50%	$\leftrightarrow$	↓h	1	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	norgestimate (COC)	1	↑85% f,b	1	<b>†</b> g	<b>†</b> 9	$\leftrightarrow$	↓64%h	<b>\</b>	↓	$\leftrightarrow$	↔d	$\leftrightarrow$	↑8%	$\leftrightarrow$	$\leftrightarrow$	↓2%	↑126% e,f	↑14%	$\leftrightarrow$	$\leftrightarrow$
	norgestrel (COC)	1	↑f,b	1	<b>†</b> g	<b>†</b> 9	$\leftrightarrow$	↓h	↓	1	$\leftrightarrow$	↔d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑e,f	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	levonorgestrel (EC)	↑k	↑ <b>k</b>	↑ <b>k</b>	↑ <b>k</b>	↑ <b>k</b>	$\leftrightarrow$	↓58%I	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑ <b>k</b>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Other	mifepristone	↑k	↑ <b>k</b>	↑ <b>k</b>	↑ <b>k</b>	↑ <b>k</b>	Ek	↓	1	↓	Ek	$\leftrightarrow$	Ek	Ek	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑ <b>k</b>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	ulipristal	↑k	↑ <b>k</b>	↑k	↑ <mark>k</mark>	↑ <b>k</b>	$\leftrightarrow$	↓m	↓m	↓m	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑ <b>k</b>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$



#### Colour legend

No clinically significant interaction expected These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

#### Legend

Potential elevated exposure of the hormone
 Potential decreased exposure of the hormone

→ No significant effect

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd) DRV/c DRV co-formulated with COBI (800/150 mg qd) CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

Es estrogens

COC combined oral contraceptive EC emergency contraception

IP implant

IUD intrauterine device
POI progestin only injectable
POP progestin only pill
TS transdermal patch
VR vaginal ring

#### Interactions with ABC, FTC, 3TC, ZDV

ABC, FTC, 3TC, ZDV: no clinically relevant interactions expected.

#### Interactions with ibalizumab

None

#### Comments

- Alternative or additional contraceptive measures are recommended or, if used for hormone replacement therapy, monitor for signs of oestrogen deficiency.
- b Unboosted ATV increased ethinylestradiol AUC by 48%. Use no more than 30 μg of ethinylestradiol if co-administered with unboosted ATV and at least 35 μg of ethinylestradiol if co-administered with ATV/r.
- Depending on the contraceptive method, ethinylestradiol concentrations are either not significantly changed (COC) or significantly decreased (VR). Levels of co-administered progestin are markedly decreased. Use with EFV is not recommended as it may impair contraceptive efficacy.
- d Daily dose of ethinylestradiol should not exceed 30 µg. Caution is advised, particularly in persons with additional risk factors for thromboembolic events.
- e European SmPC states a hormonal contraceptive should contain at least 30 µg ethinylestradiol.
- f When used in a combination pill, the estrogen component is reduced to a small extent.
- g When used in a combination pill, the estrogen component is significantly reduced, caution is recommended and additional contraceptive measures should be used.
- h EFV is expected to decrease the progestin exposure and thereby impair the efficacy of the contraceptive method. A reliable method of barrier contraception must be used in addition to hormonal contraceptives.
- Used in combination with ethinylestradiol (0.015 mg/day) which is predicted to be decreased. Since there is no possibility to adjust ethinylestradiol, caution is recommended and additional contraceptive measures should be used.
- Unboosted ATV increased ethinylestradiol AUC by 48% and norethisterone AUC by 110%. Use no more than 30 μg of ethinylestradiol if co-administered with unboosted ATV and at least 35 μg of ethinylestradiol if co-administered with ATV/r.
- k Unlikely to have clinical consequences as hormone is administered as single dose.
- Use 3 mg as a single dose for emergency contraception. Note, doubling the standard dose may be outside the product license in some regions, but a pharmacokinetic study showing that a 3 mg single dose of levonorgestrel compensated for the reduction in levonorgestrel supports this recommendation.
- M Not recommended; non-hormonal emergency contraception (Cu-IUD) should be considered.
- n A modeling study predicted a higher risk of having subtherapeutic medroxyprogesterone concentrations (i.e. <0.1 ng/mL) at week 12 in women with higher BMI on EFV treatment and even higher risk when EFV was given together with rifampicin. The risk of subtherapeutic concentrations is prevented by dosing medroxyprogesterone every 8-10 weeks in women with a higher body weight on EFV and particularly on efavirenz plus rifampicin.

#### Further Information

### **Drug-drug Interactions between Corticosteroids and ARVs**

Cor	ticosteroids	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	віс	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
	beclometasone (inhalation)	↑a	↑a	↑?a	↑11%b	↑ <mark>a</mark>	$\leftrightarrow$	↑a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
	betamethasone	↑c	↑c	↑c	↑ <b>c</b>	↑ <b>c</b>	D	1	<b>↓</b>	1	D	D	D	D	$\leftrightarrow$	D	$\leftrightarrow$	↑c	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	budesonide (inhalation)	†c	↑c	↑¢	↑c	↑¢	$\leftrightarrow$	ļ	<b>↓</b>	1	$\leftrightarrow$	↑¢	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	ciclesonide (inhalation)	↑d	↑d	↑d	↑d	↑d	$\leftrightarrow$	↑d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
eroids	clobetasol (topical)	↑c,e	↑c,e	↑c,e	↑c,e	↑c,e	$\leftrightarrow$	↑c,e	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
coste	dexamethasone	↑c D	D	1	↓ D	1	D	D	D f	D	$\leftrightarrow$	D	$\leftrightarrow$	↑c D	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$				
d corti	flunisolide (inhalation)	<b>†</b> g	↑g	↑g	<b>†</b> g	↑g	$\leftrightarrow$	1	↓	1	$\leftrightarrow$	↑g	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
and/or injected corticosteroids	fluocinolone (topical)	↑c,e	↑c,e	†c,e	↑c,e	↑c,e	$\leftrightarrow$	↑c,e	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
ınd/or	fluticasone (inhalation)	†c	↑c	↑c	↑c	↑c	$\leftrightarrow$	1	<b>\</b>	1	$\leftrightarrow$	↑c	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
oral, topic a	hydrocortisone (oral)	†c	↑c	↑c	↑ <b>c</b>	↑c	$\leftrightarrow$	1	↓	↓	$\leftrightarrow$	↑c	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	hydrocortisone (topical)	$\leftrightarrow$																			
Inhaled,	methyl-prednis- olone	†c	↑c	↑c	↑c	↑c	$\leftrightarrow$	Ţ	<b>\</b>	1	$\leftrightarrow$	↑c	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
_	mometasone (inhalation)	†c	↑¢	↑¢	↑c	↑¢	$\leftrightarrow$	1	1	1	$\leftrightarrow$	↑¢	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	prednisolone (oral)	†c	↑c	↑¢	↑c	↑¢	$\leftrightarrow$	↓20%	<b>\</b>	1	$\leftrightarrow$	↑¢	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	prednisone	↑c	↑c	↑¢	↑c	↑¢	$\leftrightarrow$	↓20%	1	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	E 11%	↑¢	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	triamcinolone	↑c	↑c	↑c	↑c	↑c	$\leftrightarrow$	1	<b>\</b>	1	$\leftrightarrow$	↑c	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						

#### Colour legend

No clinically significant interaction expected
These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

#### Legend

Potential elevated exposure of the corticosteroid
 Potential decreased exposure of the corticosteroid

→ No significant effect

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd) DRV/c DRV co-formulated with COBI (800/150 mg qd) CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

#### Interactions with ABC, FTC, 3TC, ZDV

ABC, FTC, 3TC, ZDV: no clinically relevant interactions expected.

#### Interactions with ibalizumab

None

#### Comments

- a Co-administration of RTV (100 mg bid) increased the concentrations of the active metabolite (beclometasone-17-monopropionate) but no significant effect on adrenal function was seen. Caution is still warranted, use the lowest possible corticosteroid dose and monitor for corticosteroid side effects.
- b DRV/r decreased the exposure of active metabolite (beclometasone-17monopropionate), no significant effect on adrenal function was seen.
- c Risk of having elevated corticosteroid levels, Cushing's syndrome and adrenal suppression. This risk is present for oral and injected corticosteroid but also for topical, inhaled or eye drops administration.
- d No dose adjustment required but monitor closely, especially for signs of Cushing's syndrome when using a high dose or prolonged administration.
- The extent of percutaneous absorption is determined by many factors such as degree of inflammation and alteration of the skin, duration, frequency and surface of application, use of occlusive dressings.
- f Consider using MVC a dose of 600 mg bid with dexamethasone in the absence of a PI or other potent CYP3A4 inhibitors, particularly if dexamethasone is used at a high dose and in case of long-term treatment. Consider decreasing MVC to 150 mg bid with dexamethasone in presence of a protease inhibitor or strong CYP3A4 inhibitor.
- g Use the lowest possible flunisolide dose with monitoring for corticosteroid side effects.

#### Further Information



# **Drug-drug Interactions between COVID-19 Therapies and ARVs**

co	VID-19 Therapy	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
	bamlanivimab/ etesevimab	$\leftrightarrow$																			
nAbs	casirivimab/ imdevimab	$\leftrightarrow$																			
and r	molnupiravir	$\leftrightarrow$																			
rugs	nirmatrelvir/r	↔a	↔ a	↔ a	↔ a	↔ a	Е	↔ b	$\leftrightarrow$	↔ b	Е	$\leftrightarrow$	Еc	Е	$\leftrightarrow$	Е	$\leftrightarrow$	↔ a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Antiviral Drugs and mAbs	remdesivir	$\leftrightarrow$																			
Anti	sotrovimab	$\leftrightarrow$																			
	tixagevimab/ cilgavimab	$\leftrightarrow$																			
	anakinra	$\leftrightarrow$																			
	baricitinib	$\leftrightarrow$																			
	canakinumab	$\leftrightarrow$																			
	convalescent plasma	$\leftrightarrow$																			
seic	COVID-19 vaccines	$\leftrightarrow$																			
Immune Therapies	dexamethasone (low dose*)	↑ d	↑ d	↑ d	↑ d	↑ d	De	↓ f	↓ f	↓ f	Dg	D	Dh	$\leftrightarrow$	$\leftrightarrow$	D	$\leftrightarrow$	↑ d	$\leftrightarrow$	D	$\leftrightarrow$
nune	hydrocortisone	↑ d	↑ d	↑ d	↑ d	↑ d	$\leftrightarrow$	↓ f	↓ f	↓ f	$\leftrightarrow$	↑ d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
<u>Ē</u>	infliximab	$\leftrightarrow$																			
	methyl-prednis- olone	↑ d	↑ d	↑ d	↑ d	↑ d	$\leftrightarrow$	↓ f	↓ f	↓ f	$\leftrightarrow$	↑ d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	ruxolitinib	↑ i	↑i	↑ i	↑ i	↑ i	$\leftrightarrow$	ļ	↓	1	$\leftrightarrow$	↑ i	$\leftrightarrow$	Е	Е						
	sarilumab	$\leftrightarrow$																			
	tocilizumab	$\leftrightarrow$																			

#### Colour legend

No clinically significant interaction expected
These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

#### Legend

Potential elevated exposure of the COVID therapy
Potential decreased exposure of the COVID therapy

D Potential decreased exposure of ARV drug

E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd)
DRV/c DRV co-formulated with COBI (800/150 mg qd)
CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

 Evaluation of the DDI risk refers to a dexamethasone dose of 6 mg qd and does not apply to higher doses of dexamethasone.

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

mAbs monoclonal antibodies

#### Interactions with ABC, FTC, 3TC, ZDV

ABC, FTC, 3TC: no clinically relevant interactions expected.ZDV: potential additive haematological toxicity with anakinra, baricitinib, canakinumab, ruxolitinib, sarilumab, tocilizumab.

#### Interactions with ibalizumab

None

#### Comments

- a RTV or COBI containing regimens are continued with no dosage modification. Inform about potential occurrence of adverse effects.
- b Ritonavir bid is expected to counteract the inducing effect of EFV, NVP.
- c Consider using MVC at a dose of 150 mg bid.
- d Product labels for dexamethasone, hydrocortisone and methylprednisolone do not recommend co-administration of strong CYP3A4 inhibitors but this is unlikely to be clinically significant given the low dose of corticosteroids used in COVID-19 treatment.
- Consider increasing DOR to 100 mg bid during treatment for COVID-19 and for approximately 2 weeks after the end of treatment.
- f Doubling the dose of dexamethasone, hydrocortisone or methylprednisolone is recommended.
- Dexamethasone is a dose dependent CYP3A4 inducer and may decrease RPV concentrations. Although the level of induction at the dose recommended for COVID (6 mg/day) is likely to be relatively modest, it is advised either using hydrocortisone (IV, 200 mg/day) or, alternatively, giving dexamethasone but doubling the dose of RPV to 50 mg qd. This dose should be maintained for 2 weeks after the end of treatment as any reduction in RPV concentrations may persist for up to 14 days after stopping dexamethasone.
- h Consider using MVC at a dose of 600 mg bid with dexamethasone in the absence of a PI or other potent CYP3A4 inhibitors. Consider decreasing MVC to 150 mg bid with dexamethasone in presence of a PI or strong CYP3A4 inhibitor. These dose adjustments should be considered during treatment for COVID-19 and for approximately 2 weeks after the end of treatment.
- The ruxolitinib European product label advises reducing ruxolitinib dose by half and administering bid. Monitor closely for cytopenia and titrate ruxolitinib based on safety and efficacy.

#### **Further Information**



# **Drug-drug Interactions between Hormone Replacement Therapy (HRT) and ARVs**

	mone repla- nent therapy	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
	estradiol	↑a	↓ b	↑ a	↓ b	↓ b	$\leftrightarrow$	↓b	↓ b	↓b	$\leftrightarrow$	↑a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑ a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
len	drospirenone	↑ a,c	↑ a	↑ a	↑a	↑a	$\leftrightarrow$	↓b	↓b	↓b	$\leftrightarrow$	↔ a,d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
ogestogen	dydrogesterone	↑a	↑ a	↑ a	↑ a	↑ a	$\leftrightarrow$	↓ b	↓ b	↓b	$\leftrightarrow$	↔ a,d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
& Pro	levonorgestrel	↑a	↑ a	↑ a	↑ a	↑ a	$\leftrightarrow$	↓ b	↓ b	↓b	$\leftrightarrow$	↔ a,d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑ a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Estrogen	medroxypro- gesterone (oral)	↑ a	↑ a	↑ a	↑ a	↑ a	$\leftrightarrow$	↓ b	↓ b	↓ b	$\leftrightarrow$	↔ a,d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑ a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Est	norethisterone	↑a	↑ a	↑ a	↑a	↑ a	$\leftrightarrow$	↓ b	↓ b	↓ b	$\leftrightarrow$	↔ a,d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	norgestrel	↑a	↑ a	↑ a	↑ a	↑ a	$\leftrightarrow$	↓ b	↓ b	↓b	$\leftrightarrow$	↔ a,d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑ a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$

#### Colour legend

No clinically significant interaction expected
These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

#### Legend

↑ Potential elevated exposure of the hormone

Potential decreased exposure of the hormone

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd)
DRV/c DRV co-formulated with COBI (800/150 mg qd)
CAB/RPV CAB and RPV im long acting injections
(PK and/or QT interactions shown are with RPV)

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

#### Interactions with ABC, FTC, 3TC, ZDV

ABC, FTC, 3TC, ZDV: no clinically relevant interactions expected.

#### Interactions with ibalizumab

None

#### Comments

- a The clinical significance of increased estradiol exposure in terms of overall risk of deep vein thrombosis, pulmonary embolism, stroke and myocardial infarction in postmenopausal women receiving substitution hormones in unknown. The use of estrogen alone or in combination with a progestogen should be used at the lowest effective dose and for the shortest duration consistent with treatment goals and risks for individual women. Postmenopausal women should be re-evaluated.
- b Monitor for signs of estrogen deficiency.
- c Coadministration is contraindicated in the US product label due to the potential for hyperkalaemia. The European product label recommends clinical monitoring for hyperkalaemia.
- d No effect on progestogen but potential increase in estrogen exposure.

#### **Further Information**

# **Drug-drug Interactions between Immunosuppressants (for SOT) and ARVs**

	nuno- pressants	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
cs	prednisone	1	1	1	1	1	$\leftrightarrow$	↓20%	1	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	E11%	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
5	azathioprine	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$								
AM	mycophenolate	$\leftrightarrow$	ţа	$\leftrightarrow$	↓a	↓a	$\leftrightarrow$	↓a	$\leftrightarrow$	↓ <mark>a</mark> D13%	$\leftrightarrow$	↑ Eb									
CN	cyclosporine	†a	†a	†a	†a	†a	Е	↓a	↓a	↓a	Е	$\leftrightarrow$	Е	Е	$\leftrightarrow$	Е	$\leftrightarrow$	†a	$\leftrightarrow$	Е	Eb
ō	tacrolimus*	↑a,c	†a,c	† <mark>a</mark>	†a	↑a,c	ţа	↓a	↓a	↓a	↔C	↔C	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔C	$\leftrightarrow$	†a	$\leftrightarrow$	$\leftrightarrow$	↔b
mTOR	everolimus	1	1	1	1	1	$\leftrightarrow$	↓a	↓a	↓a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
E	sirolimus	1	1	1	1	1	ţа	↓a	↓a	↓a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	↔b						
	anti-thymocyte globulin	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$								
Other	basiliximab	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$								
	belatacept	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$								

#### Colour legend

No clinically significant interaction expected
These drugs should not be co-administered
Potential clinically significant interaction that is likely to require addi-

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

#### Legend

Potential elevated exposure of the immunosuppressant Potential decreased exposure of the immunosuppressant

→ No significant effect

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd)
DRV/c DRV co-formulated with COBI (800/150 mg qd)
CAB/RPV CAB and RPV im long acting injections
(PK and/or QT interactions shown are with RPV)

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

AM antimetabolite
CNI calcineurin inhibitors
CS corticosteroids
mTOR mTOR inhibitors

#### Interactions with ABC, FTC, 3TC, ZDV

ABC: potential decrease in mycophenolate exposure.
 ZDV: potential risk of additive haematoxicity with azathioprine.
 ZDV: potential alteration in mycophenolate exposure, monitor plasma concentrations.

#### Interactions with ibalizumab

None

#### Comments

- a TDM of immunosuppressant is recommended.
- b Monitor renal function.
- Both drugs can potentially prolong the QT interval, ECG monitoring recommended.

#### **Further Information**



<sup>\*</sup> available as prolonged release formulation

# **Drug-drug Interactions between Pulmonary Antihypertensives and ARVs**

	monary antihy- tensives	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
	ambrisentan	1	1	1	1	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$								
ERA	bosentan	†a	†a	†a	†a	↑a	D	1	↓	Ţр	D	1	D	D	$\leftrightarrow$	D	D	†a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	macitentan	1	1	1	1	1	$\leftrightarrow$	1	1	<b>\</b>	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
E2	sildenafil	1	1	1	1	1	$\leftrightarrow$	↓	↓	<b>↓</b>	↓3%	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
PDE5	tadalafil	1	1	1	1	1	$\leftrightarrow$	↓	↓	<b>\</b>	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
SGC	riociguat	1	1	1	1	1	$\leftrightarrow$	1	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	epoprostenol	$\leftrightarrow$																			
<b>₽</b>	iloprost	$\leftrightarrow$																			
	treprostinil	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$												
F	selexipag	↔C	↔C	↔C	↔C	↑120%d	$\leftrightarrow$	↔C	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										

#### Colour legend

No clinically significant interaction expected These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

#### Legend

Potential elevated exposure of the pulmonary antihypertensive Potential decreased exposure of the pulmonary antihypertensive

(PK and/or QT interactions shown are with RPV)

→ No significant effect

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd) DRV/c DRV co-formulated with COBI (800/150 mg qd) CAB/RPV CAB and RPV im long acting injections

ERA endothelin receptor antagonists

Ipr IP receptor agonists
PA prostacyclin analogues

PDE5 phosphodiesterase type 5 inhibitors sGC soluble guanylate cyclase stimulators

#### Interactions with ABC, FTC, 3TC, ZDV

ABC, FTC, 3TC, ZDV: No clinically relevant interactions expected.

#### Interactions with ibalizumab

None

#### Comments

- Co-administration is not recommended in the European labels, but the US labels suggest the following dose modifications: When starting bosentan in persons already on PI/b or EVG/c use a bosentan dose of 62.5 mg qd or every other day. Discontinue bosentan at least 36 h prior to starting PI/b or EVG/c and restart after at least 10 days at 62.5 mg qd or every other day.
- b Potential additive liver toxicity.
- Exposure of parent drug increased but exposure of active metabolite unchanged.
- d This change is unlikely to be clinically relevant.

#### **Further Information**



# **Drug-drug Interactions between Viral Hepatitis Drugs and ARVs**

	al hepatitis igs	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	віс	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
	elbasvir/ grazoprevir	1	†376% †958%	1	↑66% ↑650%	↑271% ↑1186%	↓4% ↑7%	↓54% ↓83%	1	1	↑7% ↓2%	<b>↔</b> ↑	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↓2% ↓19%	↑118% ↑436%	↓19% ↓11%	$\leftrightarrow$	↓7% ↓14%
	glecaprevir/ pibrentasvir	1	↑553% ↑64%	1	↑397%	↑338% ↑146%	$\leftrightarrow$	ļ	↓	1	E 84%	1	Е	E	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑205% ↑57% E47%	E47%	$\leftrightarrow$	E29%
DAAs	sofosbuvir	$\leftrightarrow$	$\leftrightarrow$	1	↑34%	$\leftrightarrow$	$\leftrightarrow$	↓6%	$\leftrightarrow$	$\leftrightarrow$	↑9%	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↓5% D27%	$\leftrightarrow$	↓6%
HCV D	sofosbuvir/ ledipasvir	↑ a	↑8% ↑113%a	↑ a	↑34% ↑39% <mark>a</mark>	↔a	↑4% ↓8%	↓6% ↓34%a	$\leftrightarrow$	$\leftrightarrow$	↑10% ↑8% <mark>a</mark>	1	Е	↑7% ↓13%	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑36% ↑78%a	↓5% ↓9% D~20%	E32%	Ea
	sofosbuvir/ velpatasvir	↔a	↑22% ↑142%a	↔a	↓28% ↓16%a	↓29% ↑2% <mark>a</mark>	$\leftrightarrow$	↓3% ↓53%	1	1	↑16% ↓1%	1	Е	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↓8% ↓9%	↑ a	↑24% ↓2%	$\leftrightarrow$	Ea
	sofosbuvir/ velpatasvir/ voxilaprevir	1	↑40% ↑93% ↑331%	↑a	↓28% ↓5% ↑143%b	1	$\leftrightarrow$	1	1	1	$\leftrightarrow$	1	Е	↑9% ↓4% ↓9%	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑22% ↑16% ↑171%a	$\leftrightarrow$	E	Ea
HDV	Bulevirtide	1	1	1	1	1	Е	1	1	$\leftrightarrow$	E	$\leftrightarrow$	Е	$\leftrightarrow$	$\leftrightarrow$	Е	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$

#### Colour legend

No clinically significant interaction expected
These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration

Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

#### Legend

↑ Potential elevated exposure of the hepatitis therapy
 ↓ Potential decreased exposure of the hepatitis therapy

→ No significant effect

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd)
DRV/c DRV co-formulated with COBI (800/150 mg qd)
CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

Numbers refer to decreased or increased AUC as observed in drug-drug interaction studies.

First/second numbers refer to AUC changes for EBR/GZR or GLE/PIB or SOF/LDV or SOF/VEL.

First/second/third numbers refer to AUC changes for SOF/VEL/VOX

#### Interactions with ABC, FTC, 3TC, ZDV

ABC, FTC, 3TC, ZDV: no clinically relevant interactions expected.

#### Interactions with ibalizumab

None

#### Comments

- a Monitoring of renal function recommended due to increase of tenofovir concentration if the regimen contains TDF.
- b Study details are with DRV/r qd. DRV bid has not been studied and should be used with caution as voxilaprevir concentrations may increase more than with DRV qd (this would be of further significance in cirrhotic patients). Monitoring of renal function recommended due to increase of tenofovir concentrations if the regimen contains TDF.

#### **Further Information**



# **Administration of ARVs in Persons with Swallowing Difficulties**

Drug	Formulation	Crush tablets	Open capsules	Comment
NRTIs				
ABC	tablet (300 mg) solution (20 mg/mL)	yes		Bitter taste. Crushed tablets can be added to small amount of semi-solid food or liquid, all of which should be consumed immediately
FTC	capsule (200 mg) solution (10 mg/mL)	no	yes	Dissolve in ≥ 30 mL of water, contains Na 460 µmol/mL Bioequivalence: 240 mg solution = 200 mg capsule; adjust dosage accordingly
3TC	tablet (150, 300 mg) solution (10 mg/mL) <sup>(vii)</sup>	yes		Crushed tablets can be added to small amount of semi-solid food or liquid, all of which should be consumed immediately
TDF	tablet (300 <sup>(i)</sup> mg) granules (33 mg/g)	yes		Better: dissolve in ≥ 1 dL of water/orange or grape juice (bitter taste) Mix granules in a container with soft food not requiring chewing (e.g. yoghurt or applesauce). Granules must not be mixed with liquids
ZDV	capsule (100, 250 mg)	no	no	Sticky, bitter taste
	oral solution (10 mg/mL), iv infusion (10 mg/mL)			Better: use oral solution or iv 6 mg/kg per day in glucose 5%
TAF/FTC	tablet (25/200 mg and 10/200 mg) <sup>(v)</sup>	yes		Crushing of tablets is not recommended in the product information. However based on data with the fixed-dose combination tablet (TAF/FTC/DRV/c), crushing of tablets does not impact significantly TAF/FTC pharmacokinetics (of note: TAF bioavailability is reduced by 20% (crushing) but this decrease is unlikely to be clinically significant)(viii)
TDF/FTC	tablet (300 <sup>0</sup> /200 mg)	yes		Better: dissolve in ≥ 1 dL of water/orange or grape juice (bitter taste)
ABC/3TC	tablet (600/300 mg)	no		Use solution of individual compounds
ZDV/3TC	tablet (300/150 mg)	yes		Disperse in ≥ 15 mL water, alternative: use solution of individual compounds
ABC/3TC/ZDV	tablet (300/150/300 mg)	no		Use solution of individual compounds
NNRTIs				
DOR	tablet (100 mg)	no		Tablet must be swallowed whole
TDF/3TC/DOR	tablet (300/300/100 mg)	no		Tablet must be swallowed whole
EFV	tablet (600 mg)	yes		Tablets may be divided for ease of swallowing. Capsules can be opened and the content administered with a small amount of food using the capsule
	capsule (50, 100, 200 mg)	no	yes	sprinkle method of administration
ETV	tablet (200 mg)	no		Disperse in ≥ 5 mL water. The glass should be rinsed with water several times and each rinse completely swallowed to ensure the entire dose is consumed
NVP	tablet (100, 200, 400 mg) <sup>(ii)</sup> suspension (10 mg/mL)	yes <sup>(ii)</sup>		Dissolve in water
RPV	tablet (25 mg)	no		Crushing of tablets and dispersion into a liquid is not recommended. RPV is insoluble in water over a wide pH range
TDF/FTC/EFV	tablet (300 <sup>(i)</sup> /200/600 mg)	no		Tablets must be swallowed whole
TAF/FTC/RPV	tablet (25/200/25 mg)(v)	no		Tablets should be swallowed whole and should not be chewed, crushed or split
TDF/FTC/RPV	tablet (300 <sup>0</sup> /200/25 mg)	no		Crushing of tablets and dispersion into a liquid is not recommended. RPV is insoluble in water over a wide pH range
Pls	<u> </u>			
ATV	capsule (100, 150, 200, 300 mg) oral powder (50 mg)	no	no	Do not open the capsule, swallow whole
ATV/c	tablet (300/150 mg)	no		Tablets should be swallowed whole and should not be chewed, broken, cut or crushed
DRV	tablet (75,150, 400, 600, 800 mg) solution (100 mg/mL)	yes		Take with food. Crushed tablets can be added to small amount of semi-solid food or liquid, all of which should be consumed immediately
DRV/c	tablet (800/150 mg)	yes		Crushing of tablets is not recommended in the product information. However, based on data with the fixed-dose combination tablet (TAF/FTC/DRV/c), crushing of tablets does not impact significantly DRV/c pharmacokinetics(viii)
LPV/r	tablet (200/50 mg) solution (80/20 mg/mL)	no		42% alcohol, do not dilute with water (risk of precipitation), rinse with milk (no water); take with food, bitter taste.  Not recommended for use with polyurethane feeding tubes due to potential incompatibility. Feeding tubes that are compatible with ethanol and propylene glycol, such as silicone and polyvinyl chloride (PVC) feeding tubes, can be used.
RTV	tablet (100 mg) oral suspension (100 mg) solution (80 mg/mL)	no		43% alcohol, do not dilute solution (risk of precipitation), rinse with milk (no water); bitter taste; take with food.  Not recommended for use with polyurethane feeding tubes due to potential incompatibility. Feeding tubes that are compatible with ethanol and propylene glycol, such as silicone and polyvinyl chloride (PVC) feeding tubes, can be used.
TAF/FTC/DRV/c	tablet (10/200/800/150 mg)(v)	yes		Crushing of tablets has no significant effect on the pharmacokinetics of the components of the tablet (of note: TAF bioavailability is reduced by 20% (crushing) but this decrease is unlikely to be clinically significant. TAF bioavailability is not changed when splitting the pill) (viii)



Drug	Formulation	Crush tablets	Open capsules	Comment
Others			Julios	
CAB	tablet (30 mg)	no		Tablets must be swallowed whole
CAB/RPV LA	injectable	NA	NA	
DTG	tablet (10, 25, 50 mg) dispersable tablet (5 mg)	yes		Tablets may be split or crushed and added to a small amount of semi-solid food or liquid, all of which should be consumed immediately
FTR	tablet (600 mg)	no		The prolonged released tablet should be swallowed whole
Ibalizumab	injectable	NA	NA	
MVC	tablet (25, 75, 150, 300 mg) oral solution (20 mg/mL)	yes		While the company does not have any specific kinetic information, crushing the tablet is not expected to negatively affect the bioavailability
RAL <sup>(iii)</sup>	tablet (400, 600 mg) chewable tablets (25, 100 mg) granule oral suspension (100 mg)	yes		The bioavailability of the chewable tablet is higher: 300 mg chewable tablet (= 400 mg film-coated tablet)
RPV/DTG	tablet (25/50 mg)	no		Tablets should be swallowed whole and should not be chewed, crushed or split
TAF/FTC/BIC	tablet (25/200/50 mg)(v)	no		Tablets should be swallowed whole and should not be chewed, crushed or split
TAF/FTC/EVG/c	tablet (10/200/150/150 mg)(*)	yes		Crushing of tablets is not recommended in the product information. However, based on data with the fixed-dose combination tablet TAF/FTC/DRV/c), crushing of tablets does not impact significantly TAF/FTC pharmacokinetics (of note: TAF bioavailability is reduced by 20% (crushing) but this decrease is unlikely to be clinically significant) <sup>(vii)</sup> . Similarly, crushing of /TDF/FTC/EVG/c did not have a significant effect of the pharmacokinetics of EVG/c <sup>(v)</sup> Dissolving TAF/FTC/EVG/c tablet in tap water did not significantly alter the pharmacokinetics of TAF, FTC and EFV/c.
TDF/FTC/EVG/c	tablet (300 <sup>(1)</sup> /200/150/150 mg)	yes		Crushing of tablets does not significantly modify the pharmacokinetic profiles <sup>(N)</sup>
ABC/3TC/DTG(vi)	tablet (600/300/50 mg)	yes		Tablets may be split or crushed and added to a small amount of semi-solid food or liquid, all of which should be consumed immediately
Prophylaxis/treatme	ent of opportunistic infections			
azithromycin	tablet (250, 500 mg) suspension (40 mg/mL)	no		
cotrimoxazole	tablet (400/80 mg, forte 800/160 mg) solution (40/8 mg/mL)	yes; forte difficult		Dilute solution 3-5 times with water (high osmolality)
fluconazole	capsule (50, 200 mg) suspension (40 mg/mL)	no	yes	
pyrimethamine	tablet (25 mg)	yes		Take with food
valganciclovir	tablet (450 mg) solution (50 mg/mL)	no	no	Difficult to dissolve
rifampicin	tablet (450, 600 mg)	yes		Take on empty stomach
	capsule (150, 300 mg)	no	yes	
	suspension (20 mg/mL)			
rifabutin	capsule (150 mg)	no	yes	Mix with apple sauce, syrup (insoluble in water)
isoniazid	tablet (100, 150 mg)	yes		Take on empty stomach
pyrazinamide	tablet (500 mg)	yes		
ethambutol	tablet (100, 400 mg)	yes		Difficult to dissolve Better: use iv solution
rifampicin/isoniazid	tablet (150/100, 150/75 mg)	yes		Take on empty stomach
rifater (rifampicin, isoniazid, pyrazinamide)	tablet (120/50/300 mg)	yes		Take on empty stomach
rimstar (rifampicin, isoniazid, pyrazinamide, ethambutol)	tablet (150/75/400/275 mg)	yes		Take on empty stomach

For recommendations on prophylaxis/treatment of opportunistic infections, see Part VI Opportunistic Infections

- In certain countries TDF is labelled as 245 mg rather than 300 mg to reflect the amount of the prodrug (tenofovir disoproxil) rather than the fumarate salt (tenofovir disoproxil fumarate). The 245 mg dose is equivalent to 7.5 scoops of granules
- i Extended release effect lost. Note: NVP 400 mg qd (immediate release) can lead to sub-therapeutic trough levels in individuals with higher body
- weight (≥ 90 kg) compared to NVP 200 mg bid. Therefore, NVP bid administration should be preferred in individuals with higher body weight

  Crushing tablets is not recommended in the product information, however absorption of RAL was not compromised when the drug was crushed, dissolved in 60 mL warm water and administered by gastrostomy tube. In addition, RAL drug absorption has been shown to be higher in persons taking RAL 400 mg bid by chewing the tablets as compared to swallowing the intact tablets

- iv Crushing tablets is not recommended in the product information however the pharmacokinetic profiles of TDF/FTC/EVG/c were not significantly modified when the fixed-dose combination tablet (Stribild) was crushed and administered with food or with drip feed compared to the administration of the whole tablet
- v TAF is used at 10 mg when co-administered with drugs that inhibit P-gp. TAF is used at 25 mg when co-administered with drugs that do not inhibit P-gp
- vi The pharmacokinetic profiles of ABC/3TC/DTG were not modified to a clinically significant extent when the fixed-dose combination tablet (Triumeq) was crushed and administered suspended in water or in enteral nutrition (of note: crushing leads to a 26% increase in DTG exposure)
- vii The bioavailability of 3TC solution has been shown to be significantly reduced in a dose dependent manner by sorbitol present in other liquid formulations (e.g. ABC, NVP, cotrimoxazole)
- viii Crushing of tablets is not recommended in the product information, however the individual pharmacokinetic profiles of TAF/FTC/ DRV/c were not significantly modified when the fixed-dose combination tablet (Symtuza) was administered crushed or split compared to the whole tablet



# **Dose Adjustment of ARVs for Impaired Hepatic Function**

NRTIs	
ABC	Child-Pugh Class A: 200 mg bid (use oral solution) Child-Pugh Class B or C: contraindicated
FTC	No dosage adjustment
3TC	No dosage adjustment
TAF	No dosage adjustment
TAF/FTC	No dosage adjustment
TDF	No dosage adjustment
TDF/FTC	No dosage adjustment
ZDV	Reduce dose by 50% or double the interval between doses if Child-Pugh Class C
NNRTIs	
EFV	No dosage adjustment; use with caution in persons
TDF/FTC/EFV	with hepatic impairment
ETV	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data
NVP	Child-Pugh Class B or C: contraindicated
RPV	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data
TAF/FTC/RPV	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data
TDF/FTC/RPV	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data
TDF/3TC/DOR	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data
DOR	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data

Pls						
ATV	Child-Pugh Class A: no dose adjustment					
	Child-Pugh Class B: 300 mg qd (unboosted)					
	Child-Pugh Class C: not recommended					
ATV/c	Child-Pugh Class A: no dosage adjustment					
	Child-Pugh Class B or C: not recommended					
COBI	Refer to recommendations for the primary PI					
DRV	Child-Pugh Class A or B: no dosage adjustment					
	Child-Pugh Class C: not recommended					
DRV/c	Child-Pugh Class A or B: no dosage adjustment					
	Child-Pugh Class C: not recommended					
TAF/FTC/DRV/c	Child-Pugh Class A or B: no dosage adjustment					
	Child-Pugh Class C: not recommended					
LPV/r	No dosage recommendation; use with caution in persons with hepatic impairment					
RTV	Refer to recommendations for the primary PI					
Al						
FTR	No dosage adjustment					
FI						
ENF	No dosage adjustment					
EI						
Ibalizumab	No dosage adjustment					
CCR5 Inhibitor						
MVC	No dosage recommendations. Concentrations will likely be increased in persons with hepatic impairment					
INSTI						
RAL	No dosage adjustment					
EVG	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data					
DTG	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data					
BIC	Child-Pugh Class A or B: no dosage adjustment					
	Child-Pugh Class C: no data, not recommended					
TAF/FTC/EVG/c	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data					
TDF/FTC/EVG/c	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data					
ABC/3TC/DTG	Use separate compounds and refer to those adjustments					
TAF/FTC/BIC	Child-Pugh Class A or B: no dosage adjustment					
	Child-Pugh Class C: no data					
CAB	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data					

Note: Hepatic dysfunction is a good indication for TDM as clinical experience with these dose adjustments is very limited



# **Dose Adjustment of ARVs for Impaired Renal Function**

	eGFR <sup>(i)</sup> (	(mL/min)			Haemodialysis(ii)			
	≥ 50	30-49	10-29	< 10	- Hacimodialysis**			
NRTIs								
Individual agents								
ABC(iii)	300 mg q12h or 600 mg q24h		No dose adjustment required					
FTC <sup>(v)</sup>	200 m	ng q24h	200 mg q72h	200 mg q96h	200 mg q24h(iv)			
3TC(v)	300 mg q24h	150 mg q24h	100 mg q24h(vi)	50-25 mg q24h(vi)	50-25 mg q24h(iv, vi)			
TDF <sup>(vii)</sup>	300 <sup>(viii)</sup> mg q24h	300 <sup>(viii)</sup> mg q48h	Not recommended (300(viii) mg q72-96h, if no alternative	Not recommended (300(viii) mg q7d, if no alternative)	300 <sup>(viii)</sup> mg q7d <sup>(iv)</sup>			
TAF(ix,x)		25 <sup>(xi)</sup> mg q24h	ii iio aitorriaaro	No data	25 mg q24h(iv)			
ZDV	300 mg q12h		stment required	100 mg q8h	100 mg q8h <sup>(iv)</sup>			
Combinations	3 4			3 4	3 4			
ABC(iii)/3TC(v)	600/300 mg q24h							
ZDV/3TC	300/150 mg q12h		l Ise indivi	dual drugs				
ABC/3TC/ZDV	300/150/300 mg q12h	_	OGC IIIdivi	addi diago				
TAF(ix)/FTC(v)	• • • • • • • • • • • • • • • • • • • •	) mg q24h	Use individ	ual drugs <sup>(xv)</sup>	25/200 mg q24 <sup>(iv)</sup>			
TDF(vii)/FTC(v)	300 <sup>(viii)</sup> /200 mg q24h	300(viii)/200 mg q48h		Use individual drugs				
NNRTIs								
EFV	600 mg q24h		No dose adjus	tment required				
ETV	200 mg q12h		No dose adjus	tment required				
NVP	200 mg q12h	No	dose adjustment requ	ired	Additional 200 mg <sup>(iv)</sup>			
RPV	25 mg q24h		No dose adjus	tment required				
TAF(ix)/FTC(v)/RPV	25 <sup>(xi)</sup> /200/2	25 mg q24h	Use individ	ual drugs <sup>(xv)</sup>	25/200/25 mg q24h(iv)			
TDF <sup>(vii)</sup> /FTC <sup>(v)</sup> /RPV	300 <sup>(vii)</sup> /200/25 mg q24h		Use indivi	dual drugs				
DOR	100 mg q24h	N	o dose adjustment requ	uired; < 10: no PK data	xix)			
TDF <sup>(vii)</sup> /3TC <sup>(v)</sup> /DOR	300(viii)/300/100 mg q24h		Use indivi	dual drugs				
Pls <sup>(vii)</sup>								
ATV/c	300/150 mg q24h Do not initiate if eGFR < 70 mL/min if used with TDF *	No dose adjustment i	Not recommended					
ATV/r	300/100 mg q24h	No dose adjustment i		Not recommended				
DRV/r	800/100 mg q24h 600/100 mg q12h	No dose adjustment i	required <sup>(xiii)</sup>					
DRV/c	800/150 mg q24h Do not initiate if eGFR < 70 mL/min if used with TDF *	No dose adjustment i	required <sup>(xiii)</sup>		Not evaluated			
TAF(ix)/FTC(v)/DRV/c	10/200/800/150 mg q	ղ24h	Use individual drugs					
LPV/r	400/100 mg q12h	No dose adjustment i	equired <sup>(xiii)</sup>					
Other ART								
RAL	1 x 400 mg tablet q12h or 2 x 600 mg tablets q24h	No dose adjustment i	required <sup>(xiii)</sup>					
DTG	50 mg q24h	No dose adjustment i	required <sup>(xiii)</sup>					
3TC(V)/DTG	300/50 mg q24h	Use individual drugs						
ABC(iii)/3TC(v)/DTG	600/300/50 mg q24h	Use individual drugs	cvi)					
RPV/DTG	25/50 mg q24h	No dose adjustment i	required <sup>(xiii)</sup>					
TAF(ix)/FTC(v)/BIC	25/200/50 mg q24h	No dose adjustment		eGFR > 15 - < 30 mL/	No adjustment if on			
		required <sup>(xviii)</sup>	min or if eGFR < 15 mL/min without o		HD, however, use should generally be avoided and only used if potential benefits outweigh potential risks(xviii)			
TAF <sup>(ix)</sup> /FTC <sup>(v)</sup> /EVG/c	10/200/150/150 mg q	q24h	Not recommended <sup>(xii)</sup>		10/200/150/150 mg q24h <sup>(iv)</sup>			
TDF <sup>(vii)</sup> /FTC <sup>(v)</sup> /EVG/c	300\frac{\piniii}{200/150/150} mg q24h Do not initiate if eGFR < 70 mL/min	Not recommended						



CAB	30 mg q24h	No dose adjustment required <sup>(xvii)</sup>				
CAB LA RPV LA	400/600 mg 1x/4 w 600/900 mg 1x/8 w	No dose adjustment required <sup>(xvii)</sup>				
MVC: co-administered without CYP3A4 inhibitors(xtv)	300 mg q12h	No dose adjustment required <sup>(xiii)</sup>				
MVC: co-administered with CYP3A4 inhibitors(xiv)	If eGFR < 80 mL/min 150 mg q24h <sup>(xiv)</sup>					
Ibalizumab	2000 mg loading dose followed by 800 mg every 2 weeks. No dose adjustment required					
FTR	600 mg q12h	No dose adjustment required				

- eGFR: Use CKD-EPI formula; the abbreviated modification of diet in renal disease (aMDRD) or the Cockcroft-Gault (CG) equation may be used as an alternative; see https://www.chip.dk/Tools-Standards Clinical-risk-scores
- ii For Continuous Ambulatory Peritoneal Dialysis (CAPD) dosing for hemodialysis may be used. However, elimination of drugs in ČAPD varies depending on CAPD conditions. TDM therefore is recommended
- iii Potential cardiovascular risk of ABC may increase cardiovascular risk associated with renal failure
- iv After dialysis
- Large bodily accumulation in impaired renal function. Although affinity for mitochondrial DNA polymerase is low and clinical toxicity in patients with severe renal impairment is rare, long-term mitochondrial toxicity is possible and must be monitored (polyneuropathy, pancreatitis, lactate acidosis, lipodystrophy, metabolic disturbances) 150 mg loading dose
- TDF and (boosted) PIs are associated with nephrotoxicity; consider vii alternative ART if pre-existing CKD, risk factors for CKD and/or decreasing eGFR, see ARV-associated Nephrotoxicity and Kidney Disease: Definition. Diagnosis and Management
- In certain countries TDF is labelled as 245 mg rather than 300 mg to reflect the amount of the prodrug (tenofovir disoproxil) rather than the fumarate salt (tenofovir disoproxil fumarate)
- Limited clinical data documented limited accumulation in hemodialyix sis. However, there is no long-term data on residual kidney function and bone toxicity. No data for eGFR < 10 mL/min but no dialysis Only licenced for HBV
- 10 mg if co-administered with a boosting agent (inhibition of P-glycoprotein, P-gp)

- xii TAF/FTC/EVG/c as a single tablet regimen should generally be avoided in persons with end-stage renal disease on chronic dialysis. However, TAF/FTC/EVG/c may be used with caution if the potential benefits are considered to outweigh potential risks. One clinlical study has demonstrated safety of TAF/FTC/EVG/c for persons with HIV on chronic dialysis
- Limited data available in persons with renal impairment; pharmacokinetic analysis suggests no dose adjustment required
- See summary of product characteristics for specific recommendations; use with caution if eGFR ≤ 30 mL/min. 10 mg if co-administered with a boosting agent (inhibition of P-glycoprotein, P-gp) TAF/FTC and TAF/FTC/RPV single tablet regimens should gener-
- ally be avoided in persons with end-stage renal disease on chronic dialysis. However, these combinations may be used with caution if the potential benefits are considered to outweigh potential risks
- ABC/3TC/DTG as a single tablet regimen should generally be avoided in persons with end-stage renal disease on chronic haemodialysis. A recent case series study found that use of ABC/3TC/DTG appears to be a safe and effective option in persons with HIV on chronic dialysis, however these findings need to be confirmed in a larger trial
- In persons with HIV with eGFR < 30 mL/min, co-administration with a strong CYP3A4 inhibitor (e.g. ketoconazole, posaconazole) should be used only if the benefit outweighs the risk
- xviii
- According to the product label
  Doravirine is modestly removed by haemodialysis so that no dosage xix adjustment is needed
- Due to lack of COBI data in persons with renal impairment

For recommendations on ART use in persons with HIV undergoing renal transplantation, see Solid Organ Transplantation



# Selected Non-ARV Drugs Requiring Dosage Adjustment in Renal Insufficiency

Therapeutic class and drugs	CL <sub>CRT</sub> threshold for adjustment <sup>a,b</sup>	Additional information <sup>c</sup>
ANTIBACTERIALS <sup>d</sup>		
Fluoroquinolones		
Ciprofloxacin	≤ 60 mL/min	
Delafloxacin	< 30 mL/min	iv dosage: 200 mg every 12 hours; oral dosage: 450 mg every 12 hours
Levofloxacin	≤ 50 mL/min	
Ofloxacin	≤ 50 mL/min	
Cephalosporins		
Cefpodoxime	≤ 40 mL/min	
Ceftazidime	≤ 50 mL/min	
Cefepime	≤ 50 mL/min	
Penicillins	= 50 IIIL/IIIIII	
Amoxicillin/clavulanate	≤ 30 mL/min	
Benzylpenicillin (parenteral)	≤ 60 mL/min	
Piperacillin/tazobactam	≤ 40 mL/min	
Aminoglycosides		
Amikacin	≤ 70 mL/min	Dose dependent oto- and nephrotoxicity. Avoid in renal insufficiency if alternatives available otherwise perform TDM
Gentamicin	≤ 70 mL/min	uves available otherwise perform 1 DW
Tobramycin	≤ 70 mL/min	
Miscellaneous		
Nitrofurantoin		Avoid if CL <sub>CRT</sub> < 60 mL/min
Trimethoprim-sulfamethoxazole	≤ 30 mL/min	
Vancomycin	≤ 50 mL/min	Dose dependent nephrotoxicity. TDM recommended
Antimycotics	·	
Fluconazole	≤ 50 mL/min	No adjustment in single dose therapy
Antivirals		
Ribavirin	≤ 50 mL/min	
Valaciclovir	variable	Dose adjustment depends on indication and person characteristics (< 30, < 50 or < 75 mL/min)
Antimycobacterials		
Ethambutol	≤ 30 mL/min	
Antithrombotics		
Apixaban	< 50 mL/min	Dose adjustment depends on indication and person characteristics. It may be required for $CL_{CRT}$ < 50 mL/min. Avoid if $CL_{CRT}$ < 15 mL/min
Dabigatran	≤ 50 mL/min	Contraindicated if CL <sub>CRT</sub> < 30 mL/min
Edoxaban	≤ 50 mL/min	Avoid if CL <sub>CRT</sub> < 15 mL/min
Enoxaparin	< 30 mL/min	Dose adjustment depends on indication and person characteristics.
Rivaroxaban	< 50 mL/min	Dose adjustment depends on indication and person characteristics. It may be required for $CL_{CRT}$ < 50 mL/min. No dose adjustment if recommended dose is 10 mg qd Avoid if $CL_{CRT}$ < 15 mL/min
BETA BLOCKERS		
Atenolol	≤ 35 mL/min	
Sotalol	≤ 60 mL/min	
ACE INHIBITORS		
Enalapril	≤ 80 mL/min	Dose adjustment for starting dose
Lisinopril	≤ 80 mL/min	Dose adjustment for starting dose
Perindopril	< 60 mL/min	
Ramipril	< 60 mL/min	
CARDIOTONIC AGENT	· OO IIIE/IIIIII	
Digoxin	≤ 100 mL/min	Dose adjustment for maintenance and loading dose. Avoid in renal insufficiency
	3 100 IIIL/IIIII	if alternatives
ANTIDIABETICS		
Biguanide		
Metformin	< 60 mL/min	Contraindicated if CL <sub>CRT</sub> < 30 mL/min
GLP1-agonist		
Exenatide	≤ 50 mL/min	Avoid if CL <sub>CRT</sub> < 30 mL/min



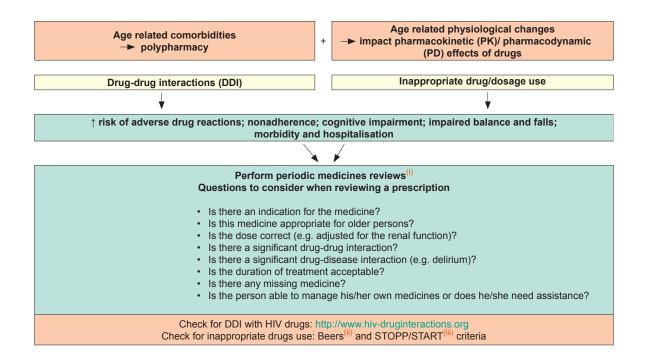
DPP4-inhibitors		
Alogliptin	≤ 50 mL/min	
Saxagliptin	< 45 mL/min	
Sitagliptin	< 45 mL/min	
Vildagliptin	< 50 mL/min	
SGLT2-inhibitors		
Canagliflozin	< 60 mL/min	Should not be initiated if $CL_{CRT}$ < 60 mL/min. Dose adjustment if $CL_{CRT}$ falls below 60 mL/min during treatment, and stop if $CL_{CRT}$ < 45 mL/min (lack of efficacy)
Dapagliflozin	-	Should not be initiated if $CL_{CRT}$ < 60 mL/min. Stop if $CL_{CRT}$ < 45 mL/min (lack of efficacy)
Empagliflozin	< 60 mL/min	Should not be initiated if $CL_{CRT}$ < 60 mL/min. Dose adjustment if $CL_{CRT}$ falls below 60 mL/min during treatment, and stop if $CL_{CRT}$ < 45 mL/min (lack of efficacy)
GOUT MEDICATION		
Allopurinol	≤ 50 mL/min	
Colchicine	≤ 50 mL/min	Dose dependent toxicity. Routine monitoring of colchicine adverse reactions recommended
ANTIPARKINSON DRUG		
Pramipexole	≤ 50 mL/min	Dose adjustment depends on indication
ANALGESICS		
NSAIDs	-	Avoid chronic use in persons with any stage of renal insufficiency
Morphine	-	Risk of respiratory depression in persons with renal insufficiency due to accumulation of 6-morphine-glucuronide (highly active metabolite). Avoid if alternatives; or titration to adequate pain control with close monitoring for signs of overdose
Oxycodone	< 50 mL/min	Initial dosage: reduced dose at initiation and further titration to adequate pain control and close monitoring for signs of overdose
Tramadol	< 30 mL/min	Increase dosing interval to 8-12 hours. Maximum daily dose 200 mg
ANTIEPILEPTICS		
Eslicarbazepine	30-60 mL/min	Start with a dose of 200 mg qd or 400 mg every other day for 2 weeks followed by 400 mg qd  Not recommended in case of severe renal impairment
Gabapentin	< 80 mL/min	
Levetiracetam	< 80 mL/min	
Pregabalin	< 60 mL/min	
PSYCHOLEPTIC		
Lithium	< 90 mL/min	Reduced dose and slow titration. TDM recommended. Avoid if CL <sub>CRT</sub> < 30 mL/min
DISEASE-MODIFYING ANTI-RHEU	JMATIC DRUGS (DMARDs)	
Methotrexate (low dose)	< 60 mL/min	Dose dependent toxicity. Contraindicated if CL <sub>CRT</sub> < 30 mL/min

- Legend

  a Renal function estimated for dosage adjustment mostly based on Cockcroft formula (CL<sub>CRT</sub>: creatinine clearance)

  b For persons with creatinine clearance < 15 mL/min or persons on dialysis, a nephrologist should be consulted
- The drug package insert should be consulted for specific dose adjustments
- No dose adjustment on antibacterial loading dose

# **Prescribing in Older Persons with HIV**



i-iii The Beers and STOPP criteria are tools established by experts in geriatric pharmacotherapy to detect and reduce the burden of inappropriate prescribing in older persons (note: these tools were established for persons > 65 years old given that PK and PD effects may be more apparent after this age cut-off). Inappropriate medicines include, for instance, those which in older persons with certain diseases can lead to drug-disease interactions, are associated with a higher risk of adverse drug reactions in olderpersons, medicines that predictably increase the risk of falls in the older persons or those to be avoided in case of organ dysfunction. The START criteria consist of evidence-based indicators of potential prescribing omission in older persons with specific medical conditions

# **Selected Top 10 Drug Classes To Avoid in Older Persons with HIV**

Drug class	Problems/alternatives
First generation antihistamines e.g., clemastine, diphenhydramine, doxylamine, hydroxyzine	Strong anticholinergic properties, risk of impaired cognition, delirium, falls, peripheral anticholinergic adverse reactions (dry mouth, constipation, blurred vision, urinary retention).  Alternatives: cetirizine, desloratadine, loratadine
Tricyclic antidepressants e.g., amitryptiline, clomipramine, doxepin, imipramine, trimipramine	Strong anticholinergic properties, risk of impaired cognition, delirium, falls, peripheral anticholinergic adverse reactions (dry mouth, constipation, blurred vision, urinary retention).  Alternatives: citalopram, escitalopram, mirtazapine, venlafaxine
Benzodiazepines Long and short acting benzodiazepines e.g., clonazepam, diazepam, midazolam Non-benzodiazepines hypnotics e.g., zolpidem, zopiclone	Elderly are more sensitive to their effect, risk of falls, fractures, delirium, cognitive impairment, drug dependency. Use with caution, at the lowest dose and for a short duration.  Alternatives: non-pharmacological treatment of sleep disturbance/sleep hygiene.
Atypical antipsychotics e.g., clozapine, olanzapine, quetiapine	Anticholinergic adverse reactions, increased risk of stroke and mortality (all antipsychotics).  Alternatives: aripiprazole, ziprasidone
Urological spasmolytic agents e.g., oxybutynin, solifenacin, tolterodine	Strong anticholinergic properties, risk of impaired cognition, delirium, falls, peripheral anticholinergic adverse reactions (dry mouth, constipation, blurred vision, urinary retention).  Alternatives: non-pharmacological treatment (pelvic floor exercises).
Stimulant laxatives e.g., senna, bisacodyl	Long-term use may cause bowel dysfunction. Alternatives: fibres, hydration, osmotic laxatives
NSAIDs e.g., diclofenac, indomethacin, ketorolac, naproxen	Avoid regular, long-term use of NSAIDs due to risk of gastrointestinal bleeding, renal failure, worsening of heart failure.  Alternatives: paracetamol, weak opioids
Digoxin Dosage > 0.125 mg/day	Avoid doses higher than 0.125 mg/day due to risk of toxicity.  Alternatives for atrial fibrillation: beta-blockers
Long acting sulfonylureas e.g., glyburide, chlorpropamide	Can cause severe prolonged hypoglycemia. Alternatives: metformin or other antidiabetic classes
Cold medications Most of these products contain antihistamines (e.g., diphenhydramine) and decongestants (e.g., phenylephrine, pseudoephedrine)	First generation antihistamines can cause central and peripheral anticholinergic adverse reactions as described above. Oral decongestants can increase blood pressure.

**Legend**NSAID nonsteroidal anti-inflammatory drug



# **Dosage Recommendations for Hormone Therapy when Used at High Doses for Gender Transitioning**

		HIV Drugs	Starting Dose	Average Dose	Maximum Dose
Estro-		No predicted effect a	2 mg/day	4 mg/day	8 mg/day
gens	Estradiol oral	Inhibits metabolism b,f	1 mg/day	2 mg/day	4 mg/day
		Induces metabolism c	Increase estradiol dosage as	s needed based on clinical effects	and monitored hormone levels.
	Estradiol gel	No predicted effect a	0.75 mg bid	0.75 mg tid	1.5 mg tid
	(preferred for >40 y and/or	Inhibits metabolism b,f	0.5 mg bid	0.5 mg tid	1 mg tid
	smokers)	Induces metabolism c	Increase estradiol dosage a	s needed based on clinical effects	and monitored hormone levels.
	Estradiol patch	No predicted effect a	25 μg/day	50-100 μg/day	150 μg/day
	(preferred for	Inhibits metabolism b,f	25 μg/day*	37.5-75 μg/day	100 μg/day
	>40 y and/or smokers)	Induces metabolism c	Increase estradiol dosage a	s needed based on clinical effects	and monitored hormone levels.
	Conjugated	No predicted effect a	1.25-2.5 mg/day	5 mg/day	10 mg/day
	estrogen†	Inhibits metabolism b,f	0.625-1.25 mg/day	2.5 mg/day	5 mg/day
		Induces metabolism c		s needed based on clinical effects	and monitored hormone levels.
	Ethinylestra-	No predicted effect a	No interaction expected, but	not recommended due to thrombo	otic risks
	diol	Inhibits metabolism b,f	Not recommended		
		Induces metabolism c	Not recommended		
Andro-	Spironolactone	No predicted effect a	50 mg/day	150 mg/day	400 mg/day
gen		Inhibits metabolism d	No interaction expected. No	dose adjustment required.	
Block- ers ‡		Induces metabolism e	No interaction expected. No	dose adjustment required.	
•	Finasteride	No predicted effect a	2.5 mg/day	2.5 mg/day	5 mg/day
		Inhibits metabolism d	Finasteride has a large safe	ty margin. No dose adjustment req	uired.
		Induces metabolism e	Increase finasteride dosage	as needed based on clinical effect	s and monitored hormone levels.
	Cyproterone acetate	No predicted effect a	50 mg/day	150 mg/day	150 mg/day
		No predicted effect a	25 mg/day	75 mg/day	75 mg/day
		Induces metabolism e	Increase cyproterone dosag	e as needed based on clinical effe	cts and monitored hormone levels.
	Goserelin	No predicted effect a	3.6 mg/month	3.6 mg/month	3.6 mg/month
		Inhibits metabolism d	No interaction expected. No	dose adjustment required.	
		Induces metabolism e	No interaction expected. No	dose adjustment required.	
	Leuprorelin	No predicted effect a	3.75 mg/month	3.75 mg/month	3.75 mg/month
	acetate	Inhibits metabolism d	No interaction expected. No	dose adjustment required.	
		Induces metabolism e	No interaction expected. No	dose adjustment required.	
	Triptorelin	No predicted effect a	3.75 mg/month	3.75 mg/month	3.75 mg/month
		Inhibits metabolism d	No interaction expected. No	dose adjustment required.	
		Induces metabolism e	No interaction expected. No	dose adjustment required.	
Andro-	Testosterone	No predicted effect a	12.5-25 mg in the morning	50 mg in the morning	100 mg in the morning
jens	topical gel 1%	Inhibits metabolism d	12.5-25 mg in the morning	25-50 mg in the morning	50-100 mg in the morning
		Induces metabolism e	Increase testosterone dosag	ge as needed based on clinical effe	ects and monitored hormone levels.
	Testosterone	No predicted effect a	Not applicable	50-100 mg/week	Not applicable
	enanthate or	Inhibits metabolism d	Not applicable	25-50 mg/week	Not applicable
	cypionate	Induces metabolism e	Increase testosterone dosag	ge as needed based on clinical effe	ects and monitored hormone levels.
	Testosterone undecanoate	No predicted effect a	Not applicable	750 mg IM, repeat after 4 weeks and then every 10 weeks	Not applicable
		Inhibits metabolism d	Not applicable	375-500 mg IM, repeat after 4 weeks and then every 10 weeks	Not applicable
		Induces metabolism e	Increase testosterone dosag	ge as needed based on clinical effe	ects and monitored hormone levels.
	Testosterone	No predicted effect a	Not applicable	250 mg/2-3 weeks	Not applicable
	mixed esters	Inhibits metabolism d	Not applicable	125 mg/2-3 weeks	Not applicable
		Induces metabolism e	Increase testosterone dosac	ge as needed based on clinical effe	ects and monitored hormone levels.

#### Comments

- a ARVs with no predicted effect: CAB, DOR, RPV, MVC, BIC, DTG, RAL, ABC, FTC, 3TC, TAF, TDF, ZDV
- b ARVs predicted to inhibit estrogen metabolism: ATV alone, ATV/c, DRV/c, EVG/c
- c ARVs predicted to induce estrogen metabolism: ATV/r, DRV/r, LPV/r, EFV, ETV, NVP
- d ARVs predicted to inhibit androgen blocker and androgen metabolism: ATV alone, ATV/c, DRV/c, EVG/c, ATV/r, DRV/r, LPV/r
- e ARVs predicted to induce androgen blocker and androgen metabolism: EFV, ETV, NVP
- f FTR inhibits only estrogens
- Matrix type transdermal patch can be cut in order to reduce the amount of hormone delivered/day
- † Conjugated estrogen is associated with high thromboembolic risk and therefore should be avoided
- Androgen deprivation treatment may prolong the QT interval. Caution should be taken when using with ARVs that can potentially prolong the QT interval (i.e., ATV alone, ATV/r, ATV/c, FTR, LPV/r, RPV)

#### Recommendations for dose changes

- Dose changes in presence of inhibitors of estrogen metabolism are based on the assumption
  that the magnitude of the DDI is expected to be less pronounced for transdermal or topical
  applications than for oral drug administration as the first-pass metabolism is avoided
- Dose changes in presence of inhibitors of testosterone metabolism are based on the assumption that the magnitude of the DDI is expected to be less pronounced for topical and intramuscular applications than for oral drug administration as the first-pass metabolism is avoided

# Part IV Prevention and Management of Co-morbidities

Successful management of persons with HIV goes beyond provision of effective ART, with increasing focus attributed to the appropriate management of co-morbidities in order to ensure the best outcomes. Recognised co-morbidities that disproportionately affect people with HIV include mental health issues (particularly depression and anxiety disorders), cardiovascular, pulmonary, hepatic, metabolic, neoplastic, renal, bone, central nervous system disorders as well as sexual dysfunction (including age-related changes such as menopause). Collectively, these conditions can significantly impact the physical and mental health of people with HIV as they grow older. Recognising that older persons comprise a significant proportion of many populations living with HIV, the current version of the Guidelines suggests HIV-specific age cut-offs for screening for many of these co-morbidities as well as greater focus on prevalent conditions such as weight gain and obesity and age-related conditions such as frailty.

Potential contributors to co-morbidity pathogenesis include a higher prevalence of recognised risk factors, potential toxicities from ART exposure, and HIV infection (or co-infections with CMV and HCV) contributing to immune dysfunction/dysregulation, chronic immune activation and inflammation. Taking this into consideration, particular focus should be paid to cessation of smoking, which contributes to many of the co-morbidities described.

The COVID-19 pandemic has brought many challenges to the care of persons with HIV, including interruption or significant changes to routine healthcare provision. In this setting, it is of particular importance that healthcare professionals other than HIV specialists, who are involved in the care of people with HIV and who are not familiar with the use of ART, should consult their HIV specialist colleagues before introducing or modifying any treatments for co-morbidities. As intervals between visits to HIV clinics are increasingly extended, or even interrupted, persons with HIV may need more frequent review by their primary care doctor and we would encourage establishment of formal shared-care arrangements to optimise management of co-morbidities and prevent unwanted drug-drug interactions.

Many HIV doctors are not specialists in managing co-morbidities and, although general guidance on management of common co-morbidities is included in these Guidelines, HIV doctors should seek expert advice where appropriate in the prevention and management of such conditions. Situations where consultation is generally recommended are indicated within this document.

In particular, as individuals with treated HIV age, some may experience multiple co-morbidities occurring together, which may contribute to frailty and disability. Such circumstances may require a comprehensive "geriatric-type" multidimensional, multidisciplinary assessment aimed at appropriately capturing the composite of medical, psychosocial and functional capabilities and limitations of older persons with HIV. A suggested approach for the management of older persons with HIV are included in this version of the Guidelines.

Depending on future clinical research findings, and the constantly evolving challenges presented by the COVID-19 pandemic these recommendations will be regularly updated as required, <a href="http://www.eacsociety.org">http://www.eacsociety.org</a> and in the EACS Guidelines App.

The current recommendations highlight co-morbidities that are seen frequently in the routine care of persons with HIV and those for which specific issues should be considered.

# **Opioid Addiction, Pharmacological Treatment**

Opioid substitution therapy (OST), also called opioid agonist therapy (OAT) is used to prevent withdrawal symptoms in persons who discontinue long term use of analgesics that act on opioid receptors or as a treatment for people with opioid use disorder. OST includes conventional treatments such as methadone maintenance therapy and buprenorphine maintenance therapy.

Comorbid mental health disorders can interfere with the adherence to OST, and result in poorer outcomes of addiction treatment.

#### Characteristics of drugs used as OST(i)

Feature	Methadone	Buprenorphine		
Dose required to prevent withdrawal symptoms according to degree of opioid dependency	Linear relationship (from 10-300 mg per day)	Linear relationship for persons with less opioid dependency only – ceiling effect (max daily dose 24 mg)		
Interaction with ARVs	Methadone plasma concentrations are reduced if used together with:  • NVP & EFV: ↓ 50%  • LPV/r: ↓ 50%  • No clinically significant alterations of methadone PK with other commonly used ART agents	Buprenorphine (B) and active metabolite norbuprenorphine (N) plasma concentrations are reduced if combined with NNRTIs and increased if combined with some PIs or INSTIs • EFV: ↓ up to 50% (B) and 70% (N) • ETV: ↓ 25% (B) • ATV/r: ↑ 50-100% (B&N) • DRV/r: ↑ 50% (N) • Caution: B reduces ATV; do not use without RTV or COBI boosting • EVG/c, ↑ 35-42% (B&N) (BIC, CAB, DOR, DTG, FTR, RAL, RPV & LPV/r do not affect B & N metabolism)		
	<b>Caution:</b> withdrawal symptoms if combined with AF drug toxicity if such ARVs are interrupted – reverse	RV that decreases plasma concentration and risk of if ARVs increase plasma concentration		
Risk of overdose	Yes	See <sup>(iii)</sup>		
Causing QT prolongation on ECG	Yes (dose-response relationship)(ii)	No		
Risk of obstipation	High	High		
Type of administration	Tablet or liquid	Tablet applied sublingual		
Risk of further impairment in persons with existing liver impairment	Yes	Yes		

- i See Drug-drug Interactions between Analgesics and ARVs
- ii ECG recommended for daily methadone doses exceeding 50 mg; special caution with concomitant use of other drugs known to cause QT prolongation (e.g. certain ARVs (such as LPV/r, RPV, FTR), amiodarone, astemizole, azithromycin, clarithromycin, chloroquine, citalopram, domperidone, escitalopram, fluconazole and moxifloxacin)
- Buprenorphine is commonly used as a fixed-dose combination with naloxone. Risk of overdose of buprenorphine may be reduced with the use of fixed dose combination with naloxone



# **Cancer: Screening Methods**®

Problem	Persons	Procedure	Evidence of benefit	Screening interval	Additional comments
Anal cancer	MSM and persons with HPV-associated dysplasia <sup>(ii)</sup>	Digital rectal exam ± anal cytology	Unknown; advocated by some experts	1-3 years	If anal cytology abnormal, anoscopy
Breast cancer	Women 50-74 years(iii)	Mammography	↓ Breast cancer mortality	1-3 years	
Cervical cancer	Women > 21 years	PAP smear or liquid based cervical cytology test	↓ Cervical cancer mortality	1-3 years	HPV genotype testing may aid PAP/liquid based cervical screening
Colorectal cancer	Persons 50-75 years or with a life expectancy > 10 years	According to local screening programme practice. Colonoscopy every 10 years if willing/able. If unable, annual faecal immunochemistry test (FIT) for occult blood, or multitarget stool DNA (MT-sDNA) testing every 3 years, or computed tomography colonography (CTC) every 5 years	Colorectal cancer mortality	Depending on screening method used	
HepatoCellular Carcinoma (HCC)	HCC screening should follow current EASL guidelines* see pages 8, 81 and 115(**)	Ultrasound (and alpha- foetoprotein)	Earlier diagnosis allowing for improved ability for surgical eradication	Every 6 months	* Risk factors for HCC in this population include family history of HCC, ethnicity (Asians, Africans), HDV and age > 45 years. EASL guidelines propose using the PAGE-B score in Caucasians to assess the HCC risk, however this score has not been validated in persons with HIV
Prostate cancer	Men > 50 years with a life expectancy >10 years	PSA <sup>(v)</sup>	Use of PSA is controversial	1-2 years	Pros: ↑ early diagnosis and modest ↓ prostate cancer specific mortality. Cons: overtreatment, adverse effects of treatment on quality of life
Lung Cancer	Age 50-80 years old who are at high risk of lung cancer (at least a 20 pack-year smoking history, and are either current smokers or former smokers having quit within the past 15 years)	Low-dose helical CT (where local screening programs are available)	↓ Lung cancer related mortality	Every year	Evidence confirmed in large RCT, but persons with HIV not included and there may be a higher false positive rate among people with HIV

- Screening recommendations derived from the general population.
  - These screenings should preferably be done as part of national general population screening programmes.

    Careful examination of skin should be performed regularly to detect cancers such as Kaposi's sarcoma, basal cell carcinoma and malignant melanoma
- ii Includes Anal Intraepithelial Neoplasia (AIN), Penile Intraepithelial Neoplasia (PIN), Cervical Intraepithelial Neoplasia (CIN), Vaginal Intraepithelial Neo-plasia (VAIN) and Vulval Intraepithelial Neoplasia (VIN).
- iii US and Australian national Guidelines recommend an upper age limit of 74 years, whilst some other national Guidelines suggest 70 years.
- IV HCC screening is indicated in all cirrhotic HBV or HCV co-infected persons (even if HCV infection has been cured and HBV replication is medically suppressed) in a setting where treatment for HCC is available. Although the cost-effectiveness of HCC screening in persons with F3 fibrosis is uncertain, surveillance may be considered based on an individual risk assessment (https://easl.eu/publication/easl-clinical-practice-guidelines-management-of-hepato-cellular-carcinoma/). In HBV-positive non-cirrhotics, HCC screening should follow current EASL guidelines. Risk factors for HCC in this population include family history of HCC, ethnicity (Asians, Africans), HDV and age > 45 years. EASL guidelines propose using the PAGE-B score in Caucasians to assess the HCC risk, however this score has not been validated in persons with HIV, see pages 81 and 115
- V Whilst prostate cancer screening with PSA can reduce prostate cancer specific mortality, the absolute risk reduction is very small. Given limitations in the design and reporting of the randomized trials, there remain important concerns that the benefits of screening are outweighed by the potential harms to quality of life, including the substantial risks for over-diagnosis and treatment complications.



# **Cancer: Treatment Monitoring**

- Careful attention must be paid to potential drug-drug interactions between systemic anti-cancer therapy and ART.
   Advice is available at www.hiv-druginteractions.org and www.cancer-druginteractions.org
- Chemotherapy and radiotherapy are associated with an unpredictable decline in CD4 counts even in persons stable on ART, OI prophylaxis should therefore be considered at any CD4 count threshold in persons undergoing cancer treatment with chemotherapy and radiotherapy
- Persons affected by KS treated with either liposomal doxorubicin or paclitaxel are not at increased risk of CD4 count decline and standard OI prophylaxis Guidelines should be followed, see pages 123-139
- One month after the end of the chemo- or radiotherapy treatment we recommend repeating CD4 counts and following standard OI recommendations, see pages 123-139
- · Persons undergoing autologous or allogenic stem cell transplantation should follow standard national/local guidance for anti-infective prophylaxis

#### Specific OI prophylaxis recommended in persons undergoing cancer treatment

- PCP prophylaxis, see page 126
- · Fungal prophylaxis, fluconazole 50 mg qd
  - Although the evidence for azole antifungal prophylaxis originates from haematological malignancy in HIV seronegative populations, we recommend use of antifungal prophylaxis in persons with HIV on chemotherapy or radiotherapy especially those affected by haematological malignancies. Fluconazole is the agent of choice because of the favorable interaction profile despite lack of activity against invasive Aspergillosis, see Drug-drug interactions between ARVs and Non-ARVs, page 27
- HSV/VZV prophylaxis, see pages 93 and 130
- NTM prophylaxis only in those with a detectable plasma HIV-VL, see page 125



### **Lifestyle Interventions**

Adults who adhere to Guidelines which promote a healthy diet and physical activity have lower rates of cardiovascular morbidity and mortality than those who do not. In adults without overt cardiovascular risk factors counselling interventions result in improvements in health-promoting behaviors and a positive but small benefit in preventing CVD. In adults with cardiovascular risk factors, counselling interventions have a moderate benefit in preventing CVD. Most important among lifestyle interventions is the recommendation of smoking cessation. All adults should be advised to stop smoking; the benefit of smoking cessation is substantial.

This table may be used as an example, but referring to individual national Guidelines would be just as appropriate.

# Dietary counselling

- Dietary intervention should not interfere with the dietary requirements necessary for appropriate absorption of ART drugs (e.g. maintaining sufficient calorie intake for RPV).
- · Keep caloric intake balanced with energy expenditure
- Limit intake of saturated fat, cholesterol and refined carbohydrates
- Reduce total fat intake to < 30% and dietary cholesterol to < 300 mg/day</li>
- Emphasise intake of vegetables, fruit and grain products with fibre
- · Cut back on beverages and foods with added sugar
- Choose and prepare foods with little or no salt. Adequate intakes of salt in adults have been estimated mostly around 3 g/day
- Emphasise consumption of fish, poultry (without skin) and lean meat
- Consider referral to dietician, one-week food and drink diary to discover 'hidden' calories
- · Avoid binge eating ('yo-yo dieting')
- In persons with HIV-related wasting and dyslipidaemia, address wasting first and consider referral to dietician
- Persons who are obviously overweight should be motivated to lose weight. Starvation diets are not recommended (immune defence mechanisms potentially decreased).
   Malnutrition has to be addressed where observed.
   Normal BMI range: 18.5-24.9; Overweight: 25.0-29.9, Obesity: > 30.0 kg/m²

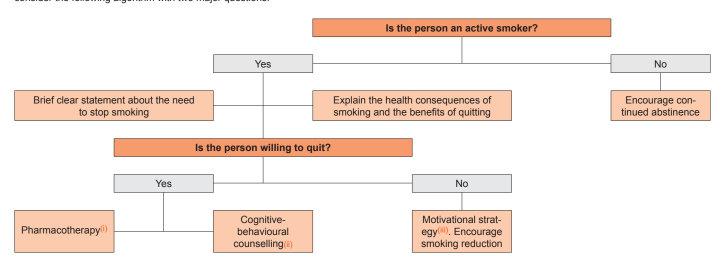
- The following questions are helpful to determine average alcohol intake
  - How often do you drink alcohol: never, ≤ 1/month, 2-4x/month, 2-3x/week, > 4x/week
  - 2. If you drink alcohol, how much typically at a time: 1-2, 3-4, 5-6, 7-9, > 10 drinks
  - How many times do you have 6 or more alcoholic drinks at one occasion: never, < 1/month, 1x/month, 1x/week, more or less daily
- Intake of alcohol should be restricted to no more than one drink per day for women and two drinks per day for men (< 20-40 g/day)</li>
- In particular, persons with hepatic disease, see NAFLD, adherence problems, inadequate CD4 count increase, tumours, past tuberculosis, diarrhoea and other conditions associated with high alcohol intake should be motivated to decrease or stop alcohol intake

# Exercise promotion

- Promote active lifestyle to prevent and treat obesity, hypertension and diabetes
- Encourage self-directed moderate level physical activity (take the stairs, walk to work, cycling, swimming, hiking, etc.)
- Emphasise regular moderate-intensity exercise rather than vigorous exercise
- Achieve cardiovascular fitness (e.g. 30 minutes brisk walking > 5 days a week)
- · Maintain muscular strength and joint flexibility

# **Smoking cessation**

Persons with HIV who smoke tobacco should be made aware of the substantial health benefits of smoking cessation which include reducing the risk of tobacco-related diseases, slowing the progression of existing tobacco related disease, and improving life expectancy by an average of 10 years. Regularly consider the following algorithm with two major questions:



Adapted from the European Smoking Cessation Guidelines and Calvo-Sanchez M., et al, 2015

- i Pharmacotherapy: Nicotine replacement therapy: nicotine substitution (patch, chewing gum, spray), varenicline and bupropion are approved by the EMA. Bupropion is contraindicated with epilepsy and varenicline may induce depression. Bupropion may interact with PIs and NNRTIs, see Drug-drug Interactions between ARVs and Non-ARVs
- ii Cognitive-behavioral intervention: Use specific available resources
- iii Motivational strategy: Identify potential health risks of the smoker and to stratify both acute (e.g. exacerbations of COPD) and long-term (e.g. infertility, cancer) risks. Explain the personal benefits of stopping smoking. Identify the barriers or obstacles that might impede the success of a quit attempt. Smoking cessation interventions should be delivered repeatedly, as long as the person is not willing/ready enough to quit smoking

At this moment, neither EMA nor FDA approve e-cigarettes as a smoking cessation agent. In persons with HIV there is no data on long-term outcomes and it is not possible to add any more specific recommendations. EACS follows the statement issued by the CDC in 2018

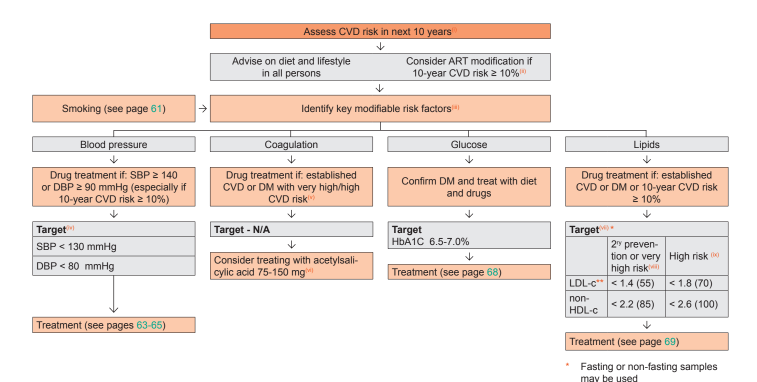
There is inadequate evidence to determine the effect of e-cigarettes on achievement of smoking cessation as well as the harms of e-cigarettes when used as a smoking cessation tool



# **Prevention of Cardiovascular Disease (CVD)**

#### Principles:

The intensity of efforts to prevent CVD depends on the underlying risk of CVD, which can be estimated. The preventive efforts are diverse in nature and require involvement of a relevant specialist, in particular if the risk of CVD is high and always in persons with a history of CVD.



- Use the Framingham equation or whatever system local National Guidance recommends; a risk equation developed from HIV populations without history of CVD is available: see https://www.chip.dk/Tools-Standards/Clinical-risk-scores. This assessment and the associated considerations outlined in this figure should be repeated annually in all persons under care, see page 8, to ensure that the various interventions are initiated in a timely way
- ii Options for ART modification include:
  - (1) Replace with NNRTI or INSTI known to cause less metabolic disturbances and/or lower CVD risks, see page 16
  - (2) Consider replacing ZDV or ABC with TDF or use an NRTIsparing regimen
- iii Of the modifiable risk factors outlined, drug treatment is reserved for certain subgroups where benefits are considered to outweigh potential harm. Of note, there is a combined benefit of various interventions in target groups identified. Per 10 mmHg reduction in systolic blood pressure, per 1 mmol/L (39 mg/dL) reduction in TC and with use of acetylsalicylic acid, each reduces risk of IHD by 20-25%; the effect is additive. Observational studies suggest that smoking cessation results in about 50% less risk of IHD and this is additive to other interventions
- iv Age 65+: Target 130-139 SBP 70-79 DBP Age 18-65: 120-129 SBP 70-79 DBP
- Ambulatory blood pressure monitoring is recommended using home BP Persons with DM in the absence of clear contraindications and established CVD or other target organ damage (any proteinuria, UA/C > 3, eGFR < 30 mL/min, left ventricular hypertrophy, or retinopathy) or ≥ 3 major risk factors (age, hypertension, dyslipidemia, smoking, obesity) or early T1DM (> 20 years) or DM ≥ 10 years plus any other risk factor

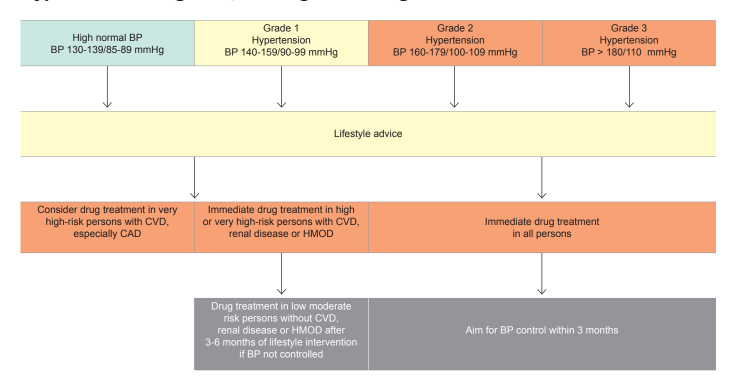
In acute settings (Post-MI, ischemic, stroke or stent insertion) dual antiplatelet therapy is recommended for up to 1 year

and ≥ 50% reduction from

baseline

- vii Target levels are to be used as guidance and are not definitive expressed as mmol/L with mg/dL in parenthesis. In case LDL-c cannot be measured or calculated because of high triglyceride levels, the non-HDL-c (TC minus HDL-c) target should be used. Target levels for TG are usually < 1.7 mmol/L (150 mg/dL) but the independent contribution from TG to CVD risk is uncertain
- viii Very high-risk persons: Documented atherosclerotic CVD (ASCVD), either clinical [ACS (MI or unstable angina), stable angina, coronary revascularization (PCI, CABG, and other arterial revascularization procedures), stroke and TIA, and peripheral arterial disease] or unequivocal on imaging [significant plaque on coronary angiography or CT scan (multivessel coronary disease with two major epicardial arteries having > 50% stenosis), or on carotid ultrasound]. DM with target organ damage, or at least three major risk factors, or early onset of T1DM of long duration (> 20 years). Severe CKD (eGFR < 30 mL/min). A calculated SCORE ≥ 10% for 10-year risk of fatal CVD. Familial hypercholesterolemia with ASCVD or with another major risk factor</p>
- High-risk persons: Markedly elevated single risk factors, in particular TC > 8 mmol/L (> 310 mg/dL), LDL-c > 4.9 mmol/L (> 190 mg/dL), or BP ≥ 180/110 mmHg. Familial hypercholesterolemia without other major risk factors. Persons with DM without target organ damage, a with DM duration ≥ 10 years or another additional risk factor. Moderate CKD (eGFR > 30 < 60 mL/min). A calculated SCORE ≥ 5% and < 10% for 10-year risk of fatal CVD

# **Hypertension: Diagnosis, Grading and Management**

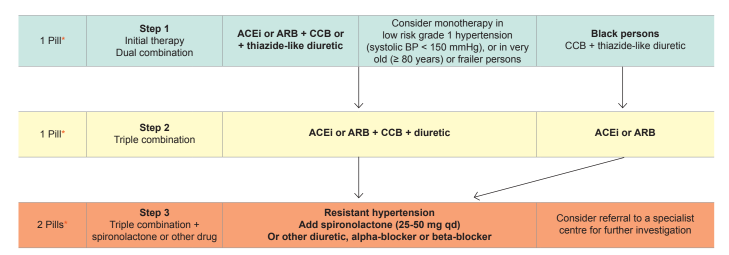


Adapted from: The Task Force for the management of arterial hypertension of the European Society of Cardiology (ESC) and the European Society of Hypertension (ESH) European Heart Journal (2018) 39, 3021–3104.

Initiation of blood pressure-lowering treatment (lifestyle changes and medication) at different initial office blood pressure levels.

BP = blood pressure; CAD = coronary artery disease; CVD = cardiovascular disease; HMOD = hypertension-mediated organ damage.

# **Hypertension: Drug Sequencing Management**



#### **Beta-blockers**

Consider beta-blockers at any treatment step, when there is a specific indication for their use, e.g. heart failure, angina, post-MI, atrial fibrillation, or younger women with, or planning, pregnancy

Adapted from: The Task Force for the management of arterial hypertension of the European Society of Cardiology (ESC) and the European Society of Hypertension (ESH) European Heart Journal (2018) 39, 3021–3104

ACEi: angiotensin converting enzyme inhibitor, ARB: angiotensin receptor blocker, CCB: calcium channel blocker

\* Where combination pill is not available single tablets should be used

# **Drug-drug Interactions between Antihypertensives and ARVs**

															CAB	CAB/					
Ant	ihypertensives	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	oral	RPV	DTG	EVG/c	RAL	TAF	TDF
	captopril	$\leftrightarrow$																			
	cilazapril	$\leftrightarrow$																			
w	enalapril	$\leftrightarrow$																			
itor	fosinopril	$\leftrightarrow$	1	$\leftrightarrow$	1	1	$\leftrightarrow$														
ACE inhibitors	lisinopril	$\leftrightarrow$																			
i i	perindopril	$\leftrightarrow$																			
¥	quinapril	$\leftrightarrow$																			
	ramipril	$\leftrightarrow$																			
	trandolapril	$\leftrightarrow$																			
ts	candesartan	$\leftrightarrow$																			
onis	eprosartan	$\leftrightarrow$																			
ıtagı	irbesartan	$\leftrightarrow$	↓	$\leftrightarrow$	<b>↓</b>	<b>↓</b>	$\leftrightarrow$	1	1	$\leftrightarrow$	↓	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$							
nar	losartan	$\leftrightarrow$	↓a	$\leftrightarrow$	↓a	↓a	$\leftrightarrow$	↑ <mark>b</mark>	↑ <b>b</b>	$\leftrightarrow$	↓a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$							
ensi	olmesartan	$\leftrightarrow$	1	$\leftrightarrow$																	
Angiotensin antagonists	telmisartan	$\leftrightarrow$	1	$\leftrightarrow$																	
An	valsartan	1	1	1	1	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$								
	atenolol	↑c	↔C	1	$\leftrightarrow$	↔C	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	1	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	bisoprolol	↑c	↑c	1	1	↑c	$\leftrightarrow$	<b>1</b>	<b>1</b>	<b>1</b>	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	carvedilol	↑c	↑↓c	1	↑↓	↑↓c	$\leftrightarrow$	↑↓	↑↓	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$							
ers	labetalol	↑c	†c	$\leftrightarrow$	<b>↓</b>	†c	$\leftrightarrow$	<b>\</b>	1	$\leftrightarrow$											
β blockers	metoprolol	↑c	↑c	1	1	↑c	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
βbl	nebivolol	↑c	↑c	1	1	↑c	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
	oxprenolol	↑c	†c	$\leftrightarrow$	<b>↓</b>	†c	$\leftrightarrow$	<b>\</b>	↓	$\leftrightarrow$											
	pindolol	↑c	↑c	1	1	↑c	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
	propranolol	↑c	↑c	1	1	↑c	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
	amlodipine	↑d	↑d	1	1	↑e	$\leftrightarrow$	<b>↓</b>	1	<b>↓</b>	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
ers	diltiazem	↑d	↑d	1	1	↑e	Е	↓69%	ţΕ	1	Е	Е	Е	Е	$\leftrightarrow$	Е	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
cium channel blockers	felodipine	↑d	↑d	1	1	↑e	$\leftrightarrow$	1	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
el b	lacidipine	↑d	↑ <mark>d</mark>	1	1	<b>↑e</b>	$\leftrightarrow$	<b>↓</b>	1	<b>↓</b>	↔f	↔f	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔f	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
ann	lercanidipine	1	1	1	1	1	$\leftrightarrow$	<b>↓</b>	1	<b>↓</b>	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
n ch	nicardipine	↑d	↑ <mark>d</mark>	1	1	<b>↑e</b>	Е	<b>↓</b>	↓E	<b>↓</b>	Ef	Ef	Е	$\leftrightarrow$	$\leftrightarrow$	Ef	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	nifedipine	↑d	↑d	1	1	↑e	$\leftrightarrow$	<b>↓</b>	1	Ţ	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
Ca	nisoldipine	↑d	↑d	1	1	↑e	$\leftrightarrow$	1	↓	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	verapamil	↑d	↑d	1	1	†e	Е	1	ţΕ	↓	E	Е	Е	Е	$\leftrightarrow$	Е	$\leftrightarrow$	1	$\leftrightarrow$	Е	Е
	amiloride	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$														
	bendroflu- methiazide	$\leftrightarrow$																			
	chlortalidone	$\leftrightarrow$																			
cs	eplerenone	1	1	1	1	1	$\leftrightarrow$	<b>1</b>	↓	<b>↓</b>	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
Diuretics	furosemide	$\leftrightarrow$	Е																		
Dif	hydrochloro- thiazide	$\leftrightarrow$																			
	indapamide	1	1	1	1	1	$\leftrightarrow$	<b></b>	<b>1</b>	<b>↓</b>	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	torasemide	$\leftrightarrow$	<u> </u>	$\leftrightarrow$	<b>1</b>	<u> </u>	$\leftrightarrow$	<b>↑</b>	1	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	<b>1</b>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	xipamide	$\leftrightarrow$																			
	clonidine	$\leftrightarrow$																			
	doxazosin	1	1	1	1	1	$\leftrightarrow$	<b>1</b>	1	↓	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	hydralazine	$\leftrightarrow$	↔g	$\leftrightarrow$	↔h																
S	methyldopa	$\leftrightarrow$	↔g	$\leftrightarrow$																	
Others	moxonidine	$\leftrightarrow$	↑?																		
	prazosin	↑?	↑?	↑?	↑?	†?	$\leftrightarrow$	↓?	↓?	↓?	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑?	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	sacubitril	1	1	1	1	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	Е	$\leftrightarrow$	1						
	spironolactone	$\leftrightarrow$																			



#### Colour legend

No clinically significant interaction expected These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/ monitoring or dosage adjustment is unlikely to be required

#### Legend

Potential elevated exposure of the antihypertensive Potential decreased exposure of the antihypertensive

No significant effect

D Potential decreased exposure of ARV drug Ε Potential elevated exposure of ARV drug

ATV co-formulated with COBI (300/150 mg qd) ATV/c DRV co-formulated with COBI (800/150 mg qd) DRV/c CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

Note: although some drug interactions are predicted to potentially require a dosage adjustment based on the drug's metabolic pathway, clinical experience with a particular antihypertensive and ARV drug may indicate that dosage adjustments are not an a priori requirement

#### Interactions with ABC, FTC, 3TC, ZDV

ABC, FTC, ZDV: no clinically relevant interactions expected. 3TC: increased 3TC exposure with atenolol and amiloride.

3TC: increased exposure of atenolol and amiloride.

#### Interactions with ibalizumab

#### Comments

- Parent drug concentrations decreased but active metabolite increased.
- Parent drug concentrations increased but active metabolite decreased.
- Risk of PR interval prolongation.
- ECG monitoring recommended.
- Use with caution as both LPV and calcium channel blockers prolong the PR interval. Clinical monitoring is recommended.
- Caution as both drugs can induce QT interval prolongation.
- Use with caution in persons with a history of postural hypotension or on concomitant medicinal products known to lower blood pressure, and those at increased risk of cardiovascular events.
- Hydralazine has some nephrotoxic potential. If co-administration is unavoidable, monitor renal function closely.

#### **Further Information**



# **Type 2 Diabetes: Diagnosis**

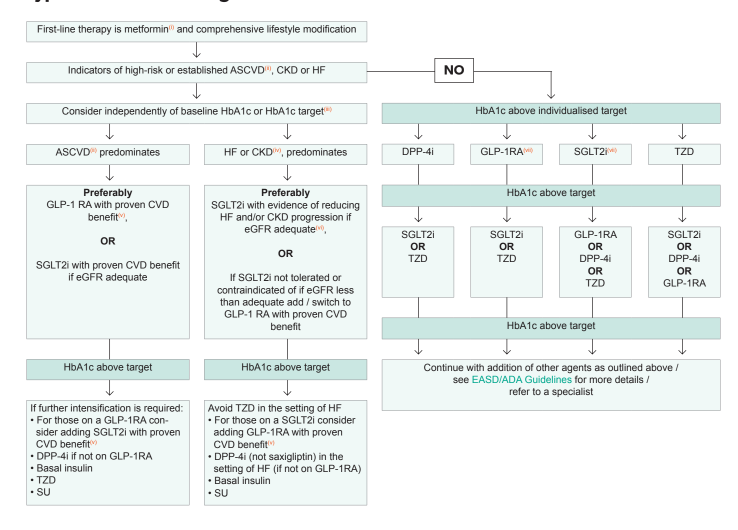
#### Diagnostic criteria<sup>(i)</sup>

	Fasting plasma glucose mmol/L (mg/dL) <sup>(ii)</sup>	Oral glucose tolerance test (OGTT) 2-h value mmol/L (mg/dL) <sup>(iii)</sup>	HbA1c <sup>(iv)</sup> (mmol/mol)		
Diabetes	≥ 7.0 (126) OR→	≥ 11.1 (200)	≥ 6.5% (≥ 48)		
Impaired glucose tolerance (IGT)	< 7.0 (126) AND→	7.8 – 11.0 (140-199)	Prediabetes		
Impaired fasting glucose (IFG)	5.7– 6.9 AND (100-125)	< 7.8 (140)	5.7-6.4% (39-47)		

- As defined by WHO
- An abnormal finding should be repeated before confirming the diagnosis

  Recommended in persons with HIV with fasting blood glucose of 5.7 6.9 mmol/L (100-125 mg/dL) as it may identify persons with overt diabetes
- Do not use HbA1c in presence of haemoglobinopathies, increased erythrocyte turnover and severe liver or kidney dysfunction. Falsely high values are measured under supplementation with iron, vitamin C and E as well as older age (age > 70: HbA1c + 0.4%). HbA1c values in treated persons with HIV, particularly when on ABC, tend to underestimate type 2 diabetes. Both IGT and IFG increase CVD morbidity and mortality and increase the risk of developing diabetes by 4-6-fold. These persons should be targeted for lifestyle intervention, and their CVD risk factors must be evaluated and treated

## **Type 2 Diabetes: Management**



- i Metformin may worsen lipoatrophy. Consider lower dose in persons with HIV with mild to moderate CKD or in individuals receiving DTG
- ii Established atherosclerotic cardiovascular disease (ASCVD) or indicators of high ASCVD (age ≥ 55 years + left ventriciular hypertrophy or coroanary, carotid, lower extermity artery stenosis > 50%)
- iii No data for any oral anti-diabetic agents in terms of CVD prevention in persons with HIV. Choice of drugs dependent on a variety of individual- & disease-specific factors; no clinically significant drug-drug-interaction or adverse effects on CD4 counts expected. Always consider individualised HbA1c targets, which depend on e.g. disease duration, life expectancy, risk for hypoglycemia, individual preference
- iv Heart failure (HF) defined as reduced ejection fraction < 45%, chronic kidney disease (CKD): eGFR > 30 < 60 mL/min or UA/C > 30 mg/mmoL, particularly UA/C > 300 mg/mmoL
- v Proven CVD benefit means it has label indication of reducing CVD events
- vi Empagliflozin, canagliflozin and dapagliflozin have shown reduction in HF and to reduce CKD progression
- vii Compelling need to minimise weight gain or promote weight loss use GLP-1RA or SGLT2i. GLP-1RA with good efficacy for weight loss: semaglutide > liraglitide > dulaglutide > exenatide > lixisenatide

# **Dyslipidaemia**

**Principles:** Higher LDL-c levels increase risk of CVD and reduction diminishes this risk (see table below for drugs used on this indication). For triglycerides (TG), there is no goal, but < 1.7 mmol/L (< 150 mg/dL) indicates lower risk and higher levels indicate a need to look for other risk factors. Statin treatment is recommended as the first drug of choice to reduce CVD risk in high-risk individuals with hypertriglyceridemia [TG > 2.3 mmol/L (> 200 mg/dL)]. Confirmation of hypertriglyceridemia needs to be verified with fasting lipid testing. Very high TG (> 10 mmol/L or > 900 mg/dL) increase risk of pancreatitis, fibrates should be used.

Less calories, more exercise, reducing bodyweight, and stopping smoking tend to improve (increase) HDL. Eating fish, reducing calories, saturated fat

and alcohol intake reduce triglyceride levels. Reducing dietary saturated fat intake improves LDL-levels; if not effective, consider change of ART, then consider lipid-lowering medicine, see page 62.

Statins should be used by all those with established vascular disease and in persons who are not at LDL-c goals considering their level of CVD risk, irrespective of lipid levels, see Treatment goal for LDL-c for very high and high CVD risk persons. In high risk persons with statin intolerance, drugdrug interactions between high intensity statins and ART, or those unable to reach LDL-c goals on statins and/or ezetimibe, a PCSK9 inhibitor should be considered.

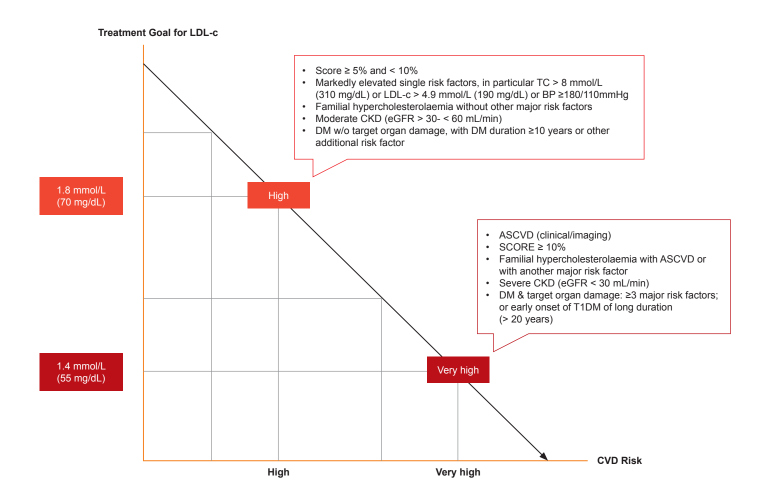
#### Drugs used to lower LDL-c

Drug class	Drug	Dose	Adverse effects	Advice on use of lipid lowering therapy together with ART			
				use with PI/r	use with NNRTIs		
Statin <sup>(i,viii)</sup>	Atorvastatin <sup>(ii)</sup>	10-80 mg qd	Gastrointestinal symptoms, headache, insomnia, rhabdomyolysis (rare)	Start with low dose(V) (max daily dose: 10 mg (ATV/r); 20 mg (LPV/r); 40 mg (DRV/r)	Consider higher dose <sup>(vi)</sup>		
	Fluvastatin(iii)	20-80 mg qd	and toxic hepatitis	Consider higher dose(vi)	Consider higher dose(vi)		
	Pravastatin <sup>(iii)</sup>	20-80 mg qd		Consider higher dose(vi,vii)	Consider higher dose(vi)		
	Rosuvastatin <sup>(ii)</sup>	5-40 mg qd		Start with low dose(V) (max daily dose: 10 mg (ATV/r, LPV/r) 20 mg (DRV/r)	Start with low dose(v)		
	Simvastatin <sup>(ii)</sup>	10-40 mg qd		Contraindicated			
	Pitavastatin(viii)	1-4 mg qd		No interaction expected			
Intestinal cholesterol absorption inhibitor \( \big( \text{i,ix} \)	Ezetimibe(iv)	10 mg qd	Gastrointestinal symptoms	No interaction expected			
PCSK9-inhibitors <sup>(X)</sup>	Evolocumab	140 mg 2 weekly or 420 mg monthly	Nil	No interaction expected			
	Alirocumab	75 mg or 150 mg 2 weekly					

- A statin is preferred first-line therapy; different statins have variable intrinsic LDL-c lowering ability
- ii, iii, iv Target levels for LDL-c, see page 70. In persons where LDL-c targets are difficult to achieve, consult/refer to specialist Expected range of reductions of LDL-c: ii 1.5-2.5 mmol/L (60-100 mg/dL), iii 0.8-1.5 mmol/L (35-60 mg/dL), iv 0.2-0.5 mmol/L (10-20 mg/dL)
- v, vi The ARV may v inhibit (statin toxicity, ↓ dose) or vi induce (= less effect of statin, ↑ dose gradually to achieve expected benefit ii, iii) the excretion of the statin
- vii Exception: If used with DRV/r, start with lower dose of pravastatin viii Pitavastatin has as yet no morbidity/mortality trial data to support its use but may have advantages of reducing immune activation and arterial inflammation, fewer drug-drug interactions, more HDL increase and less adverse glucose effect than other statins
- This agent can be used for persons intolerant of statins or added to a statin when LDL-c reduction is inadequate despite maximally tolerated statin
- x Data in persons with HIV available for evolocumab



# **Treatment Goals for LDL-c for Very High and High CVD Risk Persons**



Adapted from: 2019 ESC/EAS Guidelines for the management of dyslipidaemia: lipid modification to reduce cardiovascular risk. Eur Heart J 2020 Jan 1;41(1):111-188.

Treatment algorithm for pharmacological low-density lipoprotein cholesterol lowering. Treatment goals for low-density lipoprotein cholesterol, very-high and high CVD risk.

ASCVD = atherosclerotic cardiovascular disease; BP = blood pressure; CKD = chronic kidney disease; CVD = cardiovascular disease; DM = diabetes mellitus; eGFR = estimated glomerular filtration rate; LDL-C = low-density lipoprotein cholesterol; PCSK9 = proprotein convertase subtilisin/kexin type 9; SCORE = Systematic Coronary Risk Estimation; T1DM = type 1 DM; T2DM = type 2 DM; TC = total cholesterol

#### Moderate CVD risk:

Young persons (T1DM < 35 years; T2DM < 50 years) with DM duration < 10 years, without other risk factors. Calculated SCORE > 1 % and < 5% for 10-year risk of fatal CVD/. LDL-c goal 2.6 mmol/L (100 mg/dL)

#### Low CVD risk

Calculated SCORE < 1% for 10-year risk of fatal CVD. LDL-c goal 3.0 mmol/L (116 mg/dL)



# **Bone Disease: Screening and Diagnosis**

Condition	Characteristics	Risk factors	Diagnostic tests
Osteoporosis  • Postmenopausal women and men age ≥ 50 years with BMD T-score ≤ -2.5 at hip or lumbar spine  • Premenopausal women and men age < 50 years with BMD Z-score ≤ -2 at hip or lumbar spine and fragility fracture	Reduced bone mass and altered bone quality Increased risk of fractures in persons with HIV Asymptomatic until fractures occur Aetiology multifactorial Loss of BMD observed with ART initiation (mainly during 1st year) Greater loss of BMD with initiation of certain ARVs()	Consider classic risk factors <sup>(ii)</sup> and estimate fracture risk using FRAX in people ≥ 40 years  Consider DXA in any person with ≥ 1 risk of: <sup>(iii)</sup> 1. Postmenopausal women 2. Men ≥ 50 years 3. High risk for falls <sup>(iv)</sup> 4. Those with high fracture risk (> 20% 10-year major osteoporotic fracture risk based on FRAX assessment without DXA) 5. History of low impact fracture 6. Clinical hypogonadism (symptomatic, see Sexual Dysfunction) 7. Oral glucocorticoid use (minimum 5 mg/d prednisone equivalent for > 3 months)	DXA scan In those with classic risk factors who require DXA, where feasible consider DXA scan prior to ART initiation or soon after initiation.  Add DXA result to FRAX® to refine fracture risk prediction (http://www. shef.ac.uk/FRAX)  • May underestimate risk in persons with HIV  • Consider using HIV as a cause of secondary osteoporosis™  • Trabecular Bone Score (TBS: derived from DXA scan result) may also be added to FRAX® risk prediction.  Rule out causes of secondary osteoporosis if BMD low(vi)  Lateral spine X-rays (lumbar and thoracic) if osteoporosis on DXA, or significant height loss (≥ 4 cm) or kyphosis develops. (DXA-based vertebral fracture assessment can be used as an alternative to lateral spine X-ray)
Osteomalacia	Defective bone mineralisation     Associated with vitamin D deficiency     Increased risk of fractures and bone pain     Vitamin D deficiency may cause proximal muscle weakness	Dark skin     Dietary deficiency     Avoidance of sun exposure     Malabsorption     Obesity     Renal phosphate wasting(vii)	Measure serum 25(OH) vitamin D, see page 72. If deficient or insufficient, check PTH levels and consider vitamin D replacement if clinically indicated, see page 72    ng/mL   nmol/L     Deficiency   < 10   < 25     Insufficiency   < 20   < 50     X-rays and bone biopsy can also help in the diagnosis
Osteonecrosis	Infarct of epiphyseal plate of long bones resulting in acute bone pain     Rare but increased prevalence in persons with HIV	Risk factors:  • Low CD4 count  • Glucocorticoid exposure  • IVDU  • Alcohol  • Blood coagulation disorders	MRI

Greater loss of BMD observed with initiation of regimens containing TDF and some Pls.\* Additional loss and gains in BMD observed with switch to and away from TDF-containing ARV regimens, respectively. Clinical relevance to fracture risk not determined. TAF is associated with less bone loss than TDF

Consider replacing TDF if:

- Osteoporosis / progressive bone loss
- History of fragility fracture
- \* There are limited data on use of PIs and changes after their replacement.
- ii Classic risk factors: older age, female gender, hypogonadism, family history of hip fracture, low BMI (≤ 19 kg/m²), smoking, physical inactivity, history of low trauma fracture, alcohol excess (> 3 units/day), glucocorticoid exposure (minimum prednisone 5 mg/qd or equivalent for > 3 months)
- iii If BMD T-score normal (≥ -1), repeat after 3-5 years in risk groups 1, 2 and, 3; no need for re-screening with DXA in risk groups 4, 5 and 6 unless risk factors change and only rescreen group 7 if glucocorticoid use ongoing

- iv Falls Risk Assessment Tool (FRAT), see https://www2.health.vic.gov.au/ about/publications/policiesandguidelines/falls-risk-assessment-tool
- v If including BMD within FRAX, entering yes in the secondary cause box will not be considered in the FRAX algorithms, as it is assumed that secondary osteoporosis affects fracture risk solely through BMD. However, if the contribution of HIV infection to fracture risk is partially independent of BMD, fracture probability may be underestimated by FRAX
- vi Causes of secondary osteoporosis include hyperparathyroidism, vitamin D deficiency, hyperthyroidism, malabsorption, hypogonadism or amenorrhoea, diabetes mellitus, and chronic liver disease
- vii Use of tenofovir disoproxil fumarate (TDF) is associated with cases of renal phosphate wasting. For diagnosis and management of renal phosphate wasting, see Indications and Tests for Proximal Renal Tubulopathy (PRT)

# **Vitamin D Deficiency: Diagnosis and Management**

Vitamin D	Test	Therapy <sup>(i)</sup>
Deficiency: < 10 ng/mL (< 25 nmol/L) <sup>(ii)</sup> Insufficiency: < 20 ng/mL (< 50 nmol/L)	Serum 25-hydroxy vitamin D (25(OH) vitamin D) If deficient, consider checking parathyroid hormone (PTH), calcium, phosphate(iii), alkaline phosphatase	If vitamin D deficient, replacement recommended. Various regimens suggested <sup>(iv)</sup> Supplementation with vitamin D may reduce bone loss with initiation of ART, see page 71 Consider re-checking 25(OH) vitamin D levels 3 months after replacement. After replacement, maintenance with 800-2,000 IU vitamin D daily
Vitamin D insufficiency is prevalent (>80%) in some cohorts of populations with and without HIV – may not be directly associated with HIV  Factors associated with lower vitamin D:	Check vitamin D status in persons with history of:  • low bone mineral density and/or fracture  • high risk for fracture  Consider assessment of vitamin D status in persons with other factors	Replacement and/or supplementation of vitamin D is recommended for persons with both vitamin D insufficiency <sup>(vi)</sup> and one of the following:  • osteoporosis  • osteomalacia  • increased PTH (once the cause has been identified)  Consider re-testing after 6 months of vitamin D intake
<ul> <li>Dark skin</li> <li>Dietary deficiency</li> <li>Avoidance of sun exposure</li> <li>Malabsorption</li> <li>Obesity</li> <li>Chronic kidney disease</li> <li>Some ARVs<sup>(v)</sup></li> </ul>	associated with lower vitamin D levels (see left column)	

- i Can be provided according to national recommendations/availability of preparations (oral and parenteral formulations). Combine with calcium where there is insufficient dietary calcium intake. Consider that in some countries food is artificially fortified with vitamin D
- Vitamin D insufficiency has a prevalence of up to 80% in HIV cohorts and was associated with increased risk for osteoporosis, type 2 diabetes, mortality and AIDS events. However, causal association not proven for all outcomes. Consider seasonal differences (in winter approximately 20% lower than in summer)
- iii Consider that hypophosphataemia can be associated with TDF therapy. This phosphate loss through proximal renal tubulopathy may be independent of low vitamin D, see page 76. A combination of low calcium + low phosphate +/- high alkaline phosphatase may indicate osteomalacia and vitamin D deficiency
- iv Expect that 100 IU vitamin D daily leads to an increase in serum 25(OH) vitamin D of approximately 1 ng/mL. Some experts prefer a loading dose of e.g. 10,000 IU vitamin D daily for 8-10 weeks in persons with vitamin D deficiency. The principal goal is to achieve a serum level > 20 ng/mL (50 nmol/L) and to maintain normal serum PTH levels. Combine with calcium where potential for insufficient dietary calcium intake. The therapeutic aim is to maintain skeletal health; vitamin D supplementation has not been proven to prevent other co-morbidities in persons with HIV
- V The role of HIV-therapy or specific drugs remains unclear. Some studies suggest an association of EFV with reductions in 25(OH)D but not 1,25(OH)D. PIs may also affect vitamin D status by inhibiting conversion of 25(OH)D to 1,25(OH)D
- vi The implications of vitamin D levels that are below the physiological reference range but not markedly reduced and the value of supplementation in that situation are not completely understood



# **Approach to Fracture Reduction**

### Reducing risk of fractures

# Persons at high risk of fractures:

- · Frail or sarcopenic persons
- Previous fracture, particularly if recent
- Low BMD

- Aim to decrease falls by addressing frailty and fall risks
- Consider bisphosphonate<sup>(ii)</sup>
  - Treatment based on fracture history and FRAX score (see section on Bone Disease Screening and Diagnosis).
  - Ensure adequate calcium and vitamin D intake
- Consider choice of ARV in those at high risk of fractures<sup>(iv)</sup>
  - No significant interactions between bisphosphonates and ARVs
- Optimal management of frailty includes optimising nutrition, exercise (aerobic and resistance training), see section on frailty, page 110
- In complicated cases (e.g. young men, premenopausal women, recurrent fracture despite bone protective therapy), refer to osteoporosis specialist
- If on bisphosphonate treatment, repeat DXA after 2 years. Persons without response to treatment refer to
  osteoporosis specialist for second line treatment. Re-assess need for continued treatment after 3-5 years
- i Falls Risk Assessment Tool (FRAT), See page 72 for diagnosis and management of vitamin D deficiency https://www2.health.vic.gov.au/about/publications/policiesandguidelines/ falls-risk-assessment-tool
- ii Bisphosphonate treatment with either of alendronate 70 mg once weekly po; risedronate 35 mg once weekly po; ibandronate 150 mg po once a month or 3 mg iv every 3 months; zoledronate 5 mg by iv infusion once yearly
- iii See page 72 for diagnosis and management of vitamin D deficiency
- iv See page 71; some ARVs can affect BMD but relationship to increased fractures are not well defined. Consider relative risk/benefit of using these agents in persons with high fracture risk



# **Kidney Disease: Definition, Diagnosis and Management**

#### Diagnosis of kidney disease eGFR(i) > 60 mL/min > 60 mL/min, but > 30 - ≤ 60 mL/ ≤ 30 mL/min accelerated decline of eGFR<sup>3</sup> UA/C(iii) < 3 Regular follow-up · Check risk factors for CKD and nephrotoxic medicines including ART UA/C(iii) 3-30 · Check risk factors for CKD, use of nephrotoxic medicines Proteinuria (mg/mmol) · Discontinue or adjust drug dosages where appropriate including ART and potential artificial decline in eGFR<sup>(i)</sup> · Perform renal ultrasound Discontinue or adjust drug dosages where appropriate(v) · Urgent referral to nephrologist · Perform renal ultrasound • In persons with HIV with ESRD consider transplantation · If haematuria present with any level of proteinuria refer to evaluation, see page 113 nephrologist · Refer to nephrologist if new CKD or progressive decline in $UA/C^{(ii)} > 30$

<sup>\*</sup> Defined as decrease in eGFR of 5 mL/min per year for ≥3 consecutive years or confirmed 25% eGFR decline from baseline

Prevention of progressive	Comment
renal disease	
1. ART	Start ART immediately where HIV-associated nephropathy (HIVAN)  (vii) or HIV immune complex disease strongly suspected. Immunosup- pressive therapy may have a role in immune complex diseases. Renal biopsy to confirm histological diag- nosis recommended Consider discontinuing or replacing TDF** by non-tenofovir drug or by TAF*** if:  • UP/C 15-50 mg/mmol (see tubu- lopathy section) • eGFR > 60 mL/min, but decrease in eGFR by 5 mL/min per year for at least 3 consecutive years or confirmed 25% eGFR decline from baseline • co-morbidities with a high risk of CKD (i.e. diabetes and hypertension) • body weight < 60 kg • use of a PI/b as a third agent Discontinue or Replace TDF** by non-tenofovir drug or by TAF*** if: • eGFR ≤ 60 mL/min • UP/C > 50 mg/mmol • nephrotoxic comedication • previous TDF toxicity (proximal renal tubulopathy)  ** Expert opinion pending clinical data  ***There are limited data on use of TAF with low eGFR, particularly eGFR ≤ 10 mL/min
Start ACE inhibitors or angiotensin-II receptor antagonists if:     A. Hypertension and/or b. Proteinuria	
3. General measures: a. Avoid nephrotoxic drugs includi	CKD and proteinuria are independent ent risk factors for CVD
NSAID b. Lifestyle measures (smoking, weight, diet) c. Treat dyslipidaemia(viii) and diabetes(ix)	
<ul> <li>d. Adjust drug dosages where necessary<sup>(v)</sup></li> </ul>	

- i For eGFR: Use CKD-EPI formula based on serum creatinine, gender, age and ethnicity because eGFR quantification is validated > 60 mL/min. The abbreviated modification of diet in renal disease (aMDRD) or the Cockcroft-Gault (CG) equation may be used as an alternative https://www.chip.dk/Tools-Standards/Clinical-risk-scores.

  Definition CKD: eGFR < 60 mL/min for > 3 months (see https://kdigo.
  - Definition CKD: eGFR ≤ 60 mL/min for ≥ 3 months (see https://kdigo.org/wp-content/uploads/2017/02/KDIGO\_2012\_CKD\_GL.pdf. If not previously known to have CKD, confirm pathological eGFR within 2 weeks. Several medications and/or dietary elements or supplements may artificially increase serum creatinine and thus reduce eGFR without affecting UP/C, including the use of creatinine and protein supplements. Renal function should be reassessed after ceasing dietary supplements and/or, where available, using cystatin C-based eGFR measurements (in stable persons on ART). Use of DTG, BIC, RPV, RAL, COBI and RTV boosted PIs is also independently associated with increases in serum creatinine and reduction of eGFR (10-15 mL/ min or up to 25%) due to inhibition of proximal tubular creatinine transporters and/or intestinal transporters without impairing actual glomerular filtration. Consider a new set point after 1-2 months. Use of NSAID and recreational drugs may also affect renal perfusion and thereby cause transient creatinine increase.
- ii Urinalysis: use urine dipstick to screen for haematuria. To screen for proteinuria, use urine dipstick and if ≥ 1+ check urine albumin/creatinine (UA/C) to screen for glomerular disease or protein/creatinine (UP/C) to screen for both glomerular and tubular disease, see iii and ARV-nephrotoxicity. Proteinuria is defined as persistent if confirmed on ≥ 2 occasions > 2-3 weeks apart
- iii UA/C largely detects glomerular disease and can be used for screening for HIV-associated renal disease and in those with DM but is not appropriate for screening for tubular proteinuria secondary to drug nephrotoxicity (e.g. TDF), where UP/C should be used, see Indications and Tests for Proximal Renal Tubulopathy and ARV-nephrotoxicity. KDIGO screening values for UA/C are: < 3, 3-30 and > 30 mg/mmoL and for UP/C: < 15, 15-50, > 50 mg/mmol. UA/C and UP/C ratio are calculated as urine protein albumin (or protein) (mg/L) / urine creatinine (mmol/L); may also be expressed as mg/mg. Conversion factor for mg to mmol creatinine is x 0.000884
- iv Repeat eGFR and urinalysis as per screening table, see page 8
- See Dose Adjustment of ARVs for Impaired Renal Function
- vi Joint management with a nephrologist
- vii HIVAN suspected if black ethnicity & UAP/C > 30 mg/mmol & no haematuria
- viii See page 69-70
- ix See page 68
- x Different models have been developed for calculating a 5-years CKD risk score while using different nephrotoxic ARVs integrating HIV-independent and HIV-related risk factors

# **ARV-associated Nephrotoxicity**

Renal abnormality*	ARV	Management	
Proximal tubulopathy with any combination of:  1. Proteinuria: urine dipstick ≥ 1, or confirmed increase in UP/C > 15 mg/mmol <sup>(i)</sup> 2. Progressive decline in eGFR and eGFR ≤ 90 mL/min <sup>(ii)</sup> 3. Phosphaturia <sup>(iii)</sup> : confirmed hypophosphataemia secondary to increased urine phosphate leak  4. Glucosuria in non-diabetics	TDF**	Assessment:  • Tests for proximal renal tubulopathy/renal Fanconi syndrome(iii) (less frequent in Black persons with HIV)  • Consider renal bone disease if hypophosphataemia of renal origin: measure 25(OH) vitamin D, PTH, DXA  Replace TDF by non-tenofovir drug or TAF*** if:  • Documented tubular proteinuria and/or glucosuria  • Progressive decline in eGFR and no other cause  • Confirmed hypophosphataemia of renal origin and no other cause  • Osteopenia/osteoporosis in the presence of increased urine phosphate leak	
Nephrolithiasis:  1. Crystalluria 2. Haematuria <sup>(iv)</sup> 3. Leukocyturia 4. Loin pain 5. Acute renal insufficiency	IDV ATV (DRV)	Assessment:  • Urinalysis for crystalluria/stone analysis  • Exclude other cause for nephrolithiasis  • Renal tract imaging including CT scan  Consider stopping IDV/ATV if:  • Confirmed renal stones  • Recurrent loin pain +/- haematuria	
Interstitial nephritis:  1. Progressive decline in eGFR(ii)  2. Tubular proteinuria(iii)/ haematuria  3. Eosinophiluria (if acute)  4. Leukocyte casts	IDV ATV	Assessment: • Renal ultrasound • Refer to nephrologist  Consider stopping IDV/ATV if: • Progressive decline in eGFR and no other cause	
Progressive decline in eGFR, but none of the above <sup>(ii)</sup>	TDF** PI/r(vi)	Complete assessment: Risk factors for CKD <sup>(v)</sup> (see Kidney Disease: Definition, Diagnosis and Management) PRT, UA/C, UP/C (see Kidney Disease: Definition, Diagnosis and Mangement and Indications and Tests for Proximal Renal Tubulopathy (PRT) Renal tract ultrasound, see page 74  Consider stopping ARVs with potential nephrotoxicity if: Progressive decline in eGFR and no other cause <sup>(v)</sup>	

- \* Use of DTG, BIC, RPV, COBI and PI/b is associated with an increase in serum creatinine/reduction of eGFR (10-15 mL/min or up to 25%) due to inhibition of proximal tubular creatinine transporters without impairing actual glomerular filtration: consider new set point after 1-2 months
- \*\* TAF has shown lower tenofovir-related renal adverse effects due to lower systemic tenofovir exposure. Switch-studies from TDF to TAF and certain PIs suggest potential reversion of renal toxicity, however, longterm experience with TAF is lacking
- \*\*\* There are limited data on use of TAF with low eGFR, particularly eGFR ≤ 10 mL/min
- i UP/C in spot urine detects total urinary protein including protein of glomerular or tubular origin. The urine dipstick analysis primarily detects albuminuria as a marker of glomerular disease and is inadequate to detect tubular disease
- ii For eGFR: use CKD-EPI formula based on serum creatinine, gender, age and ethnicity because eGFR quantification is validated > 60 mL/min. The abbreviated modification of diet in renal disease (aMDRD) or the Cockcroft-Gault (CG) equation may be used as an alternative; see https://www.chip.dk/Tools-Standards/Clinical-risk-scores
- iii See Indications and Tests for Proximal Renal Tubulopathy (PRT)
- iv Microscopic haematuria is usually present
- Different models have been developed for calculating 5-year CKD risk score while using different nephrotoxic ARVs integrating HIV-independent and HIV-related risk factors
- vi RTV used as a boosting agent may induce nephrosclerosis



# **Indications and Tests for Proximal Renal Tubulopathy (PRT)**

Indications for proximal renal tubulopathy tests	Proximal renal tubulopathy tests <sup>(iv)</sup> , including	Replace TDF by non-tenofovir drug or TAF* alternative drug if:
<ul> <li>Progressive decline in eGFR<sup>(i)</sup> &amp; eGFR ≤ 90 mL/min &amp; no other cause and/or</li> <li>Confirmed hypophosphataemia<sup>(ii)</sup> and/or</li> <li>Confirmed increase in UP/C<sup>(iii)</sup></li> <li>Renal insufficiency even if stable (eGFR ≤ 60 mL/min)</li> <li>Tubular proteinuria<sup>(v)</sup></li> </ul>	Blood phosphate and urinary phosphate excretion(vi) Blood glucose and glucosuria Serum bicarbonate and urinary pH(vii) Blood uric acid level and urinary uric acid excretion(viii) Serum potassium and urinary potassium excretion	Confirmed proximal renal tubulo- pathy with no other cause

- i For eGFR: use CKD-EPI formula. The abbreviated MDRD (Modification of Diet in Renal Disease) or the Cockcroft-Gault (CG) equation may be used as an alternative, see https://www.chip.dk/Tools-Standards/Clinical-risk-scores
- ii Serum phosphate < 0.8 mmol/L or according to local thresholds; consider renal bone disease, particularly if alkaline phosphatase increased from baseline: measure 25(OH) vitamin D, PTH</p>
- UP/C in spot urine, detects total urinary protein, including protein of glomerular or tubular origin. The urine dipstick analysis primarily detects albuminuria as a marker of glomerular disease and is inadequate to detect tubular disease
- It is uncertain which tests discriminate best for TDF renal toxicity. Proximal tubulopathy is characterised by: proteinuria, hypophosphataemia, hypokalaemia, hypouricaemia, renal acidosis, glucosuria with normal blood glucose level. Renal insufficiency and polyuria may be associated. Most often, only some of these abnormalities are observed
- Tests for tubular proteinuria include retinol binding protein, α1- or β2-microglobulinuria, urine cystatin C, aminoaciduria
- vi Quantified as fractional excretion of phosphate (FEPhos): (PO<sub>4</sub>(urine) / PO<sub>4</sub>(serum) / (Creatinine(urine) / Creatinine(serum) in a spot urine sample collected in the morning in fasting state. Abnormal > 0.2 (> 0.1 with serum phosphate < 0.8 mmol/L)
- vii S-bicarbonate < 21 mmol/L and urinary pH > 5.5 suggests renal tubular acidosis
- viii Fractional excretion of uric acid (FEUricAcid): (UricAcid(urine) / UricAcid(serum) / (Creatinine(urine) / Creatinine(serum) in a spot urine sample collected in the morning in fasting state; abnormal > 0.1
- \* There are limited data on use of TAF with eGFR ≤ 10 mL/min



# **Dose Adjustment of ARVs for Impaired Renal Function**

	eGFR <sup>(i)</sup> (	(mL/min)			Haemodialysis(ii)
	≥ 50	30-49	10-29	< 10	- Hacimounarysis**
NRTIs					
Individual agents					
ABC <sup>(iii)</sup>	300 mg q12h or 600 mg q24h		No dose adjus	tment required	
FTC <sup>(v)</sup>	200 m	ig q24h	200 mg q72h	200 mg q96h	200 mg q24h <sup>(iv)</sup>
3TC(v)	300 mg q24h	150 mg q24h	100 mg q24h <sup>(vi)</sup>	50-25 mg q24h <sup>(vi)</sup>	50-25 mg q24h <sup>(iv, vi)</sup>
TDF(vii)	<u> </u>		Not recommended	Not recommended	<u> </u>
	300 <sup>(viii)</sup> mg q24h	300 <sup>(viii)</sup> mg q48h	(300 <sup>(viii)</sup> mg q72-96h, if no alternative	(300 <sup>(viii)</sup> mg q7d, if no alternative)	300 <sup>(viii)</sup> mg q7d <sup>(iv)</sup>
TAF(ix,x)		25 <sup>(xi)</sup> mg q24h		No data	25 mg q24h <sup>(iv)</sup>
ZDV	300 mg q12h	No dose adjus	tment required	100 mg q8h	100 mg q8h <sup>(iv)</sup>
Combinations					
ABC(iii)/3TC(v)	600/300 mg q24h				
ZDV/3TC	300/150 mg q12h		Use indivi	dual drugs	
ABC/3TC/ZDV	300/150/300 mg q12h			-	
TAF(ix)/FTC(v)	25 <sup>(xi)</sup> /200	mg q24h	Use individ	ual drugs(xv)	25/200 mg q24 <sup>(iv)</sup>
TDF(vii)/FTC(v)	300(viii)/200 mg g24h	300 <sup>(viii)</sup> /200 mg g48h		Use individual drugs	
NNRTIS	000 7200 mg q24m	7200 mg q+0m		OSC III aivia dai ai ags	
EFV	600 mg g24h		No dogo odina	tmont required	
	600 mg q24h			tment required	
ETV	200 mg q12h	No		tment required	Additional 200 mon(iv
NVP	200 mg q12h	INO	dose adjustment requi		Additional 200 mg <sup>(iv)</sup>
RPV	25 mg q24h	25 041		tment required	05/000/05 04/ (50
TAF(ix)/FTC(v)/RPV		25 mg q24h		ual drugs <sup>(xv)</sup>	25/200/25 mg q24h <sup>(iv)</sup>
TDF(vii)/FTC(v)/RPV	300 <sup>(vii)</sup> /200/25 mg q24h		Use indivi		6.4.4
DOR	100 mg q24h	No dose adjustment required; < 10: no PK da		· · · · · · · · · · · · · · · · · · ·	XIX)
TDF(vii)/3TC(v)/DOR	300 <sup>(m)</sup> /300/100 mg q24h		Use indivi	dual drugs	
Pls <sup>(vii)</sup>					
ATV/c	300/150 mg q24h Do not initiate if eGFR < 70 mL/min if used with TDF *	No dose adjustment required <sup>(xiii)</sup>			Not recommended
ATV/r	300/100 mg q24h	No dose adjustment r	No dose adjustment required <sup>(xiii)</sup>		Not recommended
DRV/r	800/100 mg q24h 600/100 mg q12h	No dose adjustment r	•		
DRV/c	800/150 mg q24h Do not initiate if eGFR < 70 mL/min if used with TDF *	No dose adjustment required <sup>(xiii)</sup>			Not evaluated
TAF(ix)/FTC(v)/DRV/c	10/200/800/150 mg q	24h	Use individual drugs		
LPV/r	400/100 mg q12h	No dose adjustment r	equired <sup>(xiii)</sup>		
Other ART		,	<u> </u>		
RAL	1 x 400 mg tablet q12h or 2 x 600 mg tablets q24h	No dose adjustment r	required <sup>(xiii)</sup>		
DTG	50 mg q24h	No dose adjustment r	required <sup>(xiii)</sup>		
3TC(V)/DTG	300/50 mg q24h	Use individual drugs	•		
ABC(iii)/3TC(v)/DTG	600/300/50 mg q24h	Use individual drugs	vi)		
RPV/DTG	25/50 mg q24h	No dose adjustment r	equired <sup>(xiii)</sup>		
TAF <sup>(ix)</sup> /FTC <sup>(v)</sup> /BIC	25/200/50 mg q24h	No dose adjustment required <sup>(xviii)</sup>	Not recommended if	nL/min without chronic	No adjustment if on HD, however, use should generally be avoided and only used if potential benefits outweigh
TAF <sup>(x)</sup> /FTC <sup>(v)</sup> /EVG/c	10/200/150/150 mg q	  24h	Not recommended <sup>(xii)</sup>		potential risks <sup>(xviii)</sup> 10/200/150/150 mg q24h <sup>(iv)</sup>
TDF <sup>(vii)</sup> /FTC <sup>(v)</sup> /EVG/c	300 <sup>\text{\text{\text{out}}}</sup> /200/150/150 mg q24h Do not initiate if eGFR < 70 mL/min	Not recommended			YETII .



CAB	30 mg q24h	No dose adjustment required <sup>(xvii)</sup>
CAB LA RPV LA	400/600 mg 1x/4 w 600/900 mg 1x/8 w	No dose adjustment required <sup>(xxii)</sup>
MVC: co-administered without CYP3A4 inhibitors(xtv)	300 mg q12h	No dose adjustment required <sup>(xiii)</sup>
MVC: co-administered with CYP3A4 inhibitors(xiv)	If eGFR < 80 mL/min 150 mg q24h <sup>(xiv)</sup>	
Ibalizumab	2000 mg loading dose followed by 800 mg every 2 weeks. No dose adjustment required	
FTR	600 mg q12h No dose adjustment required	

- eGFR: Use CKD-EPI formula; the abbreviated modification of diet in renal disease (aMDRD) or the Cockcroft-Gault (CG) equation may be used as an alternative; see https://www.chip.dk/Tools-Standards Clinical-risk-scores
- ii For Continuous Ambulatory Peritoneal Dialysis (CAPD) dosing for hemodialysis may be used. However, elimination of drugs in ČAPD varies depending on CAPD conditions. TDM therefore is recommended
- iii Potential cardiovascular risk of ABC may increase cardiovascular risk associated with renal failure
- iv After dialysis
- Large bodily accumulation in impaired renal function. Although affinity for mitochondrial DNA polymerase is low and clinical toxicity in patients with severe renal impairment is rare, long-term mitochondrial toxicity is possible and must be monitored (polyneuropathy, pancreatitis, lactate acidosis, lipodystrophy, metabolic disturbances) 150 mg loading dose
- TDF and (boosted) PIs are associated with nephrotoxicity; consider vii alternative ART if pre-existing CKD, risk factors for CKD and/or decreasing eGFR, see ARV-associated Nephrotoxicity and Kidney Disease: Definition. Diagnosis and Management
- In certain countries TDF is labelled as 245 mg rather than 300 mg to reflect the amount of the prodrug (tenofovir disoproxil) rather than the fumarate salt (tenofovir disoproxil fumarate)
- Limited clinical data documented limited accumulation in hemodialyix sis. However, there is no long-term data on residual kidney function and bone toxicity. No data for eGFR < 10 mL/min but no dialysis Only licenced for HBV
- 10 mg if co-administered with a boosting agent (inhibition of P-glycoprotein, P-gp)

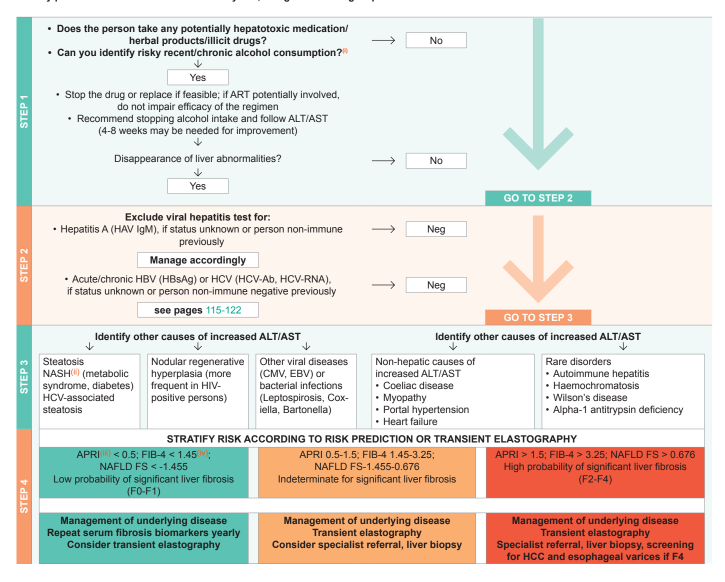
- χij TAF/FTC/EVG/c as a single tablet regimen should generally be avoided in persons with end-stage renal disease on chronic dialysis.

  However, TAF/FTC/EVG/c may be used with caution if the potential benefits are considered to outweigh potential risks. One clinlical study has demonstrated safety of TAF/FTC/EVG/c for persons on chronic
- Limited data available in persons with renal impairment; pharmacokinetic analysis suggests no dose adjustment required
- See summary of product characteristics for specific recommendations; use with caution if eGFR ≤ 30 mL/min. 10 mg if co-administered with a boosting agent (inhibition of P-glycoprotein, P-gp) TAF/FTC and TAF/FTC/RPV single tablet regimens should gener-
- ally be avoided in persons with end-stage renal disease on chronic dialysis. However, these combinations may be used with caution if the potential benefits are considered to outweigh potential risks
- ABC/3TC/DTG as a single tablet regimen should generally be avoided in persons with end-stage renal disease on chronic haemodialysis. A recent case series study found that use of ABC/3TC/DTG appears to be a safe and effective option in persons on chronic dialysis, however these findings need to be confirmed in a larger trial
- In persons with eGFR < 30 mL/min, co-administration with a strong CYP3A4 inhibitor (e.g. ketoconazole, posaconazole) should be used only if the benefit outweighs the risk
- xviii According to the product label
- xix Doravirine is modestly removed by heamodialysis so that no dosage adjustment is needed
- Due to lack of COBI data in persons with HIV with renal impairment

For recommendations on ART use in persons with HIV undergoing renal transplantation, see Solid Organ Transplantation, page 113

# Work-up and Management of persons with Increased ALT/AST

Identify potential cause of increased liver enzymes, using the following steps:



See pages 80-81 and 83-84

- i > 20 g in women, > 30 g in men
- ii Non-Alcoholic Steatohepatitis, see NAFLD
- iii APRI, AST to Platelet Ratio Index = (AST in IU/L) / (AST Upper Limit of Normal in IU/L)/ (Platelets in 109/L)
- iv FIB-4 = Age [years] x AST [U/L])/([platelet [10°/L]) x ALT<sup>1/2</sup> [U/L]). For NAFLD aetiology FIB-4 cut offs are as follows: < 1.30 (low risk), > 2.67 high risk. FIB-4 cut off < 2.0 (instead of < 1.30) should be considered in persons aged > 65 years

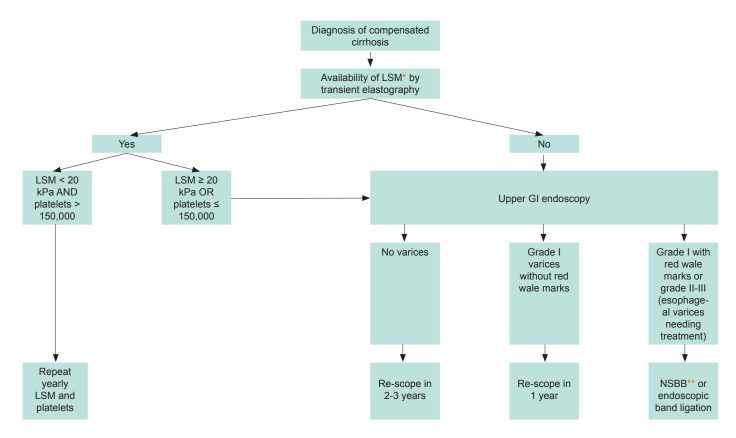
# **Liver Cirrhosis: Classification and Surveillance**

# Child-Pugh classification of the severity of cirrhosis

	Points <sup>(i)</sup>		
	1	2	3
Total bilirubin, mg/dL (µmol/L)	< 2 (< 34)	2-3 (34-50)	> 3 (> 50)
Serum albumin, g/L (µmol/L)	> 35 (> 507)	28-35 (406-507)	< 28 (< 406)
INR	< 1.7	1.7-2.20	> 2.20
Ascites	None	Mild/Moderate (diuretic responsive)	Severe (diuretic refrac- tory)
Hepatic encephalopathy	None	Grade I-II (or suppressed with medicine)	Grade III-IV (or refractory)

i 5-6 points: Class A 7-9 points: Class B 10-15 points: Class C

# Algorithm for surveillance for varices and primary prophylaxis



Based on Baveno VII consensus (EASL)

- \* LSM, liver stiffness measurement;
- \*\* NSBB, non-selective beta-blocker: prefer carvedilol 6.25-25 mg/day

Persons with compensated cirrhosis without varices on screening endoscopy should have endoscopy repeated every 2 years with ongoing liver injury, obesity or alcohol use or every 3 years if liver injury is quiescent, e.g., after viral clearance, alcohol abstinence

Hepatic Venous Pressure Gradient (HVPG) when available, allows a direct measure of portal hypertension and prognostic stratification of persons with compensated cirrhosis

HVPG < 6 mmHg: no portal hypertension

HVPG 6-9 mmHg: portal hypertension non clinically significant

HVPG  $\geq$  10 mmHg: clinically significant portal hypertension

In primary and secondary prophylaxis for variceal bleeding HVPG measurement allows to monitor efficacy of beta-blockers



# **Liver Cirrhosis: Management**

Management of persons with cirrhosis should be done in collaboration with experts in liver diseases. More general management guidance is described below. For dosage adjustment of antiretrovirals, see Dose Adjustment of ARVs for Impaired Hepatic Function

In end-stage liver disease (ESLD), use of EFV may increase risk of CNS symptoms.

ART, if otherwise indicated, also provides net benefit to cirrhotic persons. See Diagnosis and Management of Hepatorenal Syndrome (HRS)

### 200 Diagnosis and management of riopaters and cynt

#### Management of hypervolaemic hyponatreaemia (Na⁺ concentracion≤130 mmol/L)

- 1. Fluid restriction: 1000-1500 mL/
- 2. Hold diuretics
- 3. Consider albumin infusion
- At present, the use of vaptans should be limited to controlled clinical studies

# Management strategy of hepatic encephalopathy (HE)

# General management

- Identify and treat precipitating factor (GI haemorrhage, infection, pre-renal azotaemia, constipation, sedatives)
- In patients with severe hyperacute disease with HE and highly elevated arterial ammonia who are at risk of cerebral oedema, nutritional protein support can be deferred for 24-48 h until hyperammonemia is controlled
- Recommend enteral or parenteral nutritional support in critically ill patients

#### Specific therapy

Lactulose 30 cm³ po every 1-2 hours until bowel evacuation, then adjust to a dosage resulting in 2-3 formed bowel movements per day (usually 15-30 cm³ po bid)
Lactulose enemas (300 cm³ in 1L of water) in persons who are unable to take it po. Lactulose can be discontinued once the precipitating factor has resolved
Rifaximin 550 mg po bid is an effective add-on therapy to lactulose for prevention of overt hepatic encephalopathy recurrence

# Management strategy in uncomplicated ascites

# General management

- Treat ascites once other complications have been treated
- Avoid NSAIDs

Prophylaxis (Norfloxacin 400 mg po qd) should be given to persons at high risk of spontaneous bacterial peritonitis (SBP)

- 1) Persons with cirrhosis and gastrointestinal bleeding
- Perosns who have had one or more episodes of SBP. (Recurrence rates of SBP within one year have been reported to be close to 70%)
- Persons in which ascitic fluid protein is < 1.5 g/dL along with
  - Impaired renal function: creatinine ≥1.2 mg/dL (106 μmol/L), blood urea nitrogen ≥ 25 mg/dL (8.9 mmoL/L), or serum sodium ≤ 130 mEq/L (130 mmoL/L)
  - Liver failure: Child-Pugh score ≥ 9 with bilirubin
     ≥ 3 mg/dL

# Specific management

- Salt restriction: 1-2 g/day. Liberalise if restriction results in poor food intake
- Large volume paracentesis as initial therapy only in persons with tense ascites
- Administer iv albumin (= 6-8 g/L ascites removed)

# Follow-up and goals

- Adjust diuretic dosage every 4-7 days
- Weigh the person at least weekly and BUN, uric acid (UA) as surrogate for volume status s-creatinine, and electrolytes measured every 1-2 weeks while adjusting dosage
- Double dosage of diuretics if: weight loss < 2 kg a week and BUN, UA, creatinine and electrolytes are stable
- Halve the dosage of diuretics or discontinue if: weight loss ≥ 0.5 kg/day or if there are abnormalities in BUN, UA, creatinine or electrolytes
- Maximum diuretic dosage: spironolactone (400 mg qd) and furosemide (160 mg qd)

### **Nutrition of cirrhotic persons**

### Caloric requirements

 30-35 Kcal/kg/day and a protein intake of 1.2-1.5 g/kg/day of normal body weight

#### Protein requirements

- Protein restriction is not recommended
- Type: rich in branched chain (nonaromatic) amino acids
- Some studies support that parenteral proteins carry less risk of encephalopathy since not converted by colonic bacteria into NH<sub>3</sub>

#### Micronutrients

Mg and Zn

#### Analgesia in persons with hepatic failure

- Acetaminophen can be used; caution on daily dose (max 2 g/day)
- NSAIDs generally avoided; predispose persons with cirrhosis to develop GI bleeding. Persons with decompensated cirrhosis are at risk for NSAID-induced renal insufficiency
- Opiate analgesics are not contraindicated but must be used with caution in persons with pre-existing hepatic encephalopathy

## Screening for HepatoCellular Carcinoma (HCC)

- HCC screening is indicated in all cirrhotic HBV or HCV co-infected persons (even if HCV infection has been cured and HBV replication is medically suppressed) in a setting where treatment for HCC is available. Although the cost-effectiveness of HCC screening in persons with F3 fibro- sis is uncertain, surveillance may be considered based on an individual risk assessment https://easl.eu/publications/clinical-practice-guide-lines/
- In HBV-positive non-cirrhotics, HCC screening should follow current EASL guidelines. Risk factors for HCC in this population include family history of HCC, ethnicity (Asians, Africans), HDV and age > 45 years. EASL guidelines propose using the PAGE-B score in Caucasians to assess the HCC risk, however this score has not been validated in persons with HIV; see pages 8, 59 and 115. Table on fibrosis cut-offs, page 121
- Ultrasound (US), with or without alpha-foetoprotein (AFP). every 6 months.
   AFP should not be used alone. AFP is a suboptimal surveillance tool because of low sensitivity and specificity

### When to refer for liver transplantation Best to refer early as disease progresses rapidly

= MELD<sup>(1)</sup> score 12 (listing at 15)

Decompensated cirrhosis (at least one of the following complications)

- Ascites
- · Hepatic encephalopathy
- Variceal bleeding
- Spontaneous bacterial peritonitis
- Hepatorenal syndrome
- Hepatopulmonary syndrome
- NASH cirrhosis<sup>(ii)</sup>
- HCC

See Solid Organ Transplantation (SOT)

- Unit for both s-creatinine and s-bilirubin is mg/dL. MELD score = 10 {0,957 Ln (serum creatinine (mg/dL)) + 0.378 Ln (total bilirubin (mg/dL)) + 1.12 Ln (INR) + 0.643}, see http://www.mdcalc.com/meld-score-model-for-end-stage-liver-disease-12-and-older/
- Particularly with metabolic decompensations

# Non-Alcoholic Fatty Liver Disease (NAFLD)

Non-alcoholic fatty liver disease (NAFLD) is characterized by fatty infiltration of the liver (hepatic steatosis involving > 5% of hepatocytes) either on liver histology or imaging.

To be diagnosed with NAFLD, a person must not have a history of heavy alcohol use or another condition that may be causing the liver condition (such as HCV).

It is often associated with components of the metabolic syndrome: obesity, type 2 diabetes, dyslipidemia and hypertension.

Experts proposed redefining NAFLD as metabolic-associated fatty liver disease (MAFLD) to provide a positive rather than exclusionary diagnosis. However, the role of contemporary ART on MAFLD (in particular regarding an association with weight gain) remains unknown.

There are two types of NAFLD:

- 1. Non-alcoholic fatty liver (NAFL), fatty infiltration but no inflammation
- 2. Non-alcoholic steatohepatitis (NASH), with fatty infiltration along with liver inflammation (hepatocyte injury with or without fibrosis)

The prevalence of NAFLD is higher in persons with HIV (30 - 40%) than in the general population. Nearly half of the persons with HIV who undergo evaluation for unexplained liver test abnormalities are found to have NAFLD.

#### Non-Alcoholic SteatoHepatitis (NASH)

- · Early NASH: no or mild (F0-F1) fibrosis
- Fibrotic NASH: significant (≥ F2) or advanced (≥ F3, bridging) fibrosis
- · NASH-cirrhosis (F4)
- · HCC (can occur in the absence of cirrhosis and histological evidence of NASH)

#### Diagnosis

- · Ultrasound is the preferred first-line diagnostic procedure for imaging of
- Whenever imaging tools are not available or feasible, serum biomarkers and scores are an acceptable alternative for the diagnosis
- Where available and in experienced centres, transient elastography with controlled attenuation parameter could be used to diagnose HIV-associated

- NAFLD, although no optimal cut-off has been established yet. Few studies have validated CAP cut-off in HIV-associated NAFLD using different values (248 dB/m or 285 dB/m)
- A quantitative estimation of liver fat can only be obtained by MR spectroscopy as well as MRI-PDFF. This technique is of value in clinical trials and experimental studies but is expensive and not recommended in the clinical setting
- NASH has to be diagnosed by a liver biopsy showing steatosis, hepatocyte ballooning and lobular inflammation

### Consideration of ARV drugs

· Consider use of metabolic neutral ART regimens in individuals at risk of or with NAFLD (e.g. risk of weight gain induced by INSTI or TAF)

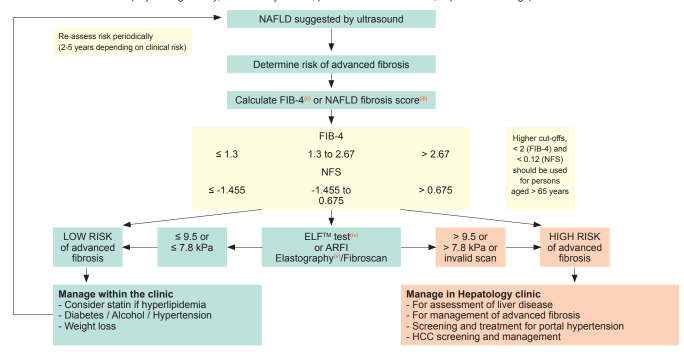
### **Treatment of NAFLD**

- · Lifestyle modification and weight reduction is the cornerstone of treatment
- Dietary restriction PLUS progressive increase in aerobic exercise/resistance training: Caloric restriction (500-1,000 /day) targeting 7-10% weight loss in persons with central obesity and/or overweight; 150-200 min/ week of moderate intensity aerobic physical activities in 3-5 sessions
- A Mediterranean diet should be advised to improve steatosis and insulin
- Pharmacotherapy should be reserved for individuals with NASH, particularly for those with significant fibrosis ≥ F2 and individuals with less severe disease, but at high risk of faster disease progression (i.e. with diabetes, metabolic syndrome, persistently increased ALT, high necroinflammation)
- Management of NASH should be discussed with hepatologists. Options with proven efficacy include pioglitazone, vitamin E and bariatric surgery. In the specific setting of HIV-associated NAFLD, tesamorelin and vitamin E have demonstrated efficacy, however larger studies are needed. Researchers advocate for inclusion of persons with HIV in ongoing global trials of new antifibrotic molecules for NASH
- Statins may be safely used but have demonstrated no impact on NAFLD thus far. The same is true for n-3 polyunsaturated fatty acids

# Diagnostic Flow-chart to Assess and Monitor Disease Severity in Case of Suspected **NAFLD and Metabolic Risk Factors**

# Persons with HIV at risk of NAFLD®

(any among obesity, metabolic syndrome, persistent elevation of ALT, exposure to d-drugs)



These recommendations are largely inspired by the EASL-EASD-EASO Clinical Practice Guidelines for the Management of Non-Alcoholic Fatty Liver Disease: European Association for the Study of the Liver (EASL), European Association for the Study of Diabetes (EASD) and European Association for the Study of Obesity

NAFLD, Non-alcoholic fatty liver disease
FIB-4 = Age ([years] x AST [U/L]) / ([platelet [109/L]) x ALT [U/L])
NFS, Non-alcoholic fatty liver disease Fibrosis Score = -1.675 + 0.037 x age (years) + 0.094 x BMI (kg/m²) + 1.13 x impaired fasting glucose/diabetes mellitus<sup>(v)</sup> (yes = 1/ no = 0) + 0.99 x AST/ALT ratio-0.013 x platelet (x10°)-0.66 x albumin(g/dL)

ELF™ test, Enhanced Liver Fibrosis Test is a blood test that provides an estimate of liver fibrosis severity by measuring Hyaluronic Acid (HA), Amino-terminal propeptide of type III

procollagen (PIIINP), Tissue inhibitor of metalloproteinase 1 (TIMP-1)

ARFI elastography, Acoustic Radiation Force Impulse



# Diagnosis and Management of Hepatorenal Syndrome / Acute Kidney Injury (HRS-AKI)

Diagnosis	<ul> <li>Cirrhosis; acute liver failure; acute-on-chronic liver failure</li> <li>Increase in serum creatinine ≥ 0.3 mg/dl (≥ 26.5 μmoL/L) within 48 h or ≥ 50% from baseline value according to ICA consensus document and/or         Urinary output ≤ 0.5 mL/kg bodyweight ≥ 6h</li> <li>No full or partial response, after at least 2 days of diuretic withdrawal and volume expansion with albumin (recommended dose of albumin is 1g/kg of body weight per day to a maximum of 100 g/day)</li> <li>Absence of shock</li> <li>No current or recent treatment with nephrotoxic drugs</li> <li>Absence of parenchymal disease as indicated by proteinuria &gt; 500 mg/day, microhematuria (&gt; 50 red blood cells per high power field, urinary injury biomarkers (if available) and/or abnormal renal ultrasonography         Suggestion of renal vasoconstriction with FENa of &lt; 0.2% (with levels &lt; 0.1% being highly predictive)</li> </ul>			
Recommended therapy	Liver transplant (priority dependent on MELD score, see page 81). If person is on transplant list, MELD score should be updated daily and communicated to transplant centre, see Solid Organ Transplantation (SOT)			
Alternative (bridging therapy)	Vasoconstrictors	terlipressin	0.5-2.0 mg iv every 4-6 hours	
	or octreotide  100-200 μg sc tid  → Goal to increase mean arterial pressure by 15 mmHg			
	+ midodrine 5-15 mg po tid			
	and iv albumin (both for at least 7 days)		50-100 g iv qd	



# **Dose Adjustment of ARVs for Impaired Hepatic Function**

NRTIs	
ABC	Child-Pugh Class A: 200 mg bid (use oral solution) Child-Pugh Class B or C: contraindicated
FTC	No dosage adjustment
3TC	No dosage adjustment
TAF	No dosage adjustment
TAF/FTC	No dosage adjustment
TDF	No dosage adjustment
TDF/FTC	No dosage adjustment
ZDV	Reduce dose by 50% or double the interval between doses if Child-Pugh Class C
NNRTIs	
EFV	No dosage adjustment; use with caution in persons
TDF/FTC/EFV	with hepatic impairment
ETV	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data
NVP	Child-Pugh Class B or C: contraindicated
RPV	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data
TAF/FTC/RPV	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data
TDF/FTC/RPV	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data
TDF/3TC/DOR	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data
DOR	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data

Pls	
ATV	Child-Pugh Class A: no dose adjustment
	Child-Pugh Class B: 300 mg qd (unboosted)
	Child-Pugh Class C: not recommended
ATV/c	Child-Pugh Class A: no dosage adjustment
	Child-Pugh Class B or C: not recommended
COBI	Refer to recommendations for the primary PI
DRV	Child-Pugh Class A or B: no dosage adjustment
	Child-Pugh Class C: not recommended
DRV/c	Child-Pugh Class A or B: no dosage adjustment
	Child-Pugh Class C: not recommended
TAF/FTC/DRV/c	Child-Pugh Class A or B: no dosage adjustment
	Child-Pugh Class C: not recommended
LPV/r	No dosage recommendation; use with caution in persons with hepatic impairment
RTV	Refer to recommendations for the primary PI
Al	
FTR	No dosage adjustment
FI	
ENF	No dosage adjustment
EI	
Ibalizumab	No dosage adjustment
CCR5 Inhibitor	
MVC	No dosage recommendations. Concentrations will likely be increased in persons with hepatic impairment
INSTI	
RAL	No dosage adjustment
EVG	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data
DTG	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data
BIC	Child-Pugh Class A or B: no dosage adjustment
	Child-Pugh Class C: no data, not recommended
TAF/FTC/EVG/c	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data
TDF/FTC/EVG/c	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data
ABC/3TC/DTG	Use separate compounds and refer to those adjustments
TAF/FTC/BIC	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data
CAB	Child-Pugh Class A or B: no dosage adjustment Child-Pugh Class C: no data

Note: Hepatic dysfunction is a good indication for TDM as clinical experience with these dose adjustments is very limited  $\frac{1}{2} \frac{1}{2} \frac{$ 

# **Lipodystrophy and Obesity: Prevention and Management**

Lipodystrophy	Lipohypertrophy <sup>(i)</sup>
<ul> <li>Prevention</li> <li>Avoid d4T and ZDV or pre-emptively switch. No evidence of benefit by switching other antiretrovirals</li> <li>Avoid excessive weight loss due to diet and exercise</li> <li>In ART-naïve persons, limb fat usually increases with initiation of ART not containing d4T or ZDV, reflecting "return-to-health" type of response</li> </ul>	Prevention No proven strategy No contemporary ART has been specifically associated with increased visceral adiposity An excess of visceral fat has been reported in HIV vs. non-HIV non-obese persons for the same BMI Weight reduction or avoidance of weight gain may decrease visceral fat Avoid corticosteroids with RTV or COBI-boosted drugs as it may cause Cushing syndrome or adrenal insufficiency, see Drug-Drug Interactions between Corticosteroids and ARVs
<ul> <li>Management</li> <li>Modification of ART: Switch away from d4T or ZDV         <ul> <li>Increase in total limb fat ~400-500 g/year (in the first two years)</li> <li>Risk of toxicity from new drug, see Adverse Effects of ARVs &amp; Drug Classes</li> </ul> </li> <li>Surgical intervention         <ul> <li>Offered for cosmetic relief of (facial) lipoatrophy</li> </ul> </li> </ul>	Management Diet and exercise may reduce visceral adiposity; Limited data, but not consistently associated with improvement in insulin sensitivity and blood lipids No prospective trials in persons with HIV to indicate degree of diet and/or exercise needed to maintain reduction in visceral fat Pharmacological interventions to treat lipohypertrophy have not been proven to provide long-term effects and may introduce new complications; Growth hormone (not approved for this indication in Europe) Decreases visceral adipose tissue May worsen insulin resistance Tesamorelin (not approved in Europe; approved for this indication by FDA Metformin (not approved for this indication in Europe) Decreases visceral adipose tissue in insulin resistant persons May worsen subcutaneous lipoatrophy Surgical therapy can be considered for localised lipomas/buffalo humps Duration of effect variable

i Lipohypertrophy may occur as localised lipomas in the subcutaneous region or as increased visceral adiposity, both intra-abdominally and/or in the epicardium. Lipohypertrophy may occur without obesity.

Increased visceral adiposity is defined by waist circumference:

- for men: ≥ 94 cm (≥ 90 cm for Asian men) is high, and > 102 cm is very high
- for women: ≥ 80 cm is high and > 88 cm is very high
- ii Tesamorelin (growth hormone releasing factor) was shown to reduce visceral adipose tissue volume but this effect was lost on discontinuation.



# Weight gain and Obesity

	Weight Gain	Obesity	Comments	
Definition	It is a physiological phenomenon associated with aging. Body weight of an average European adult is estimated to increase by 0.3 - 0.5 kg per year Definition is lacking. An increase > 5% of weight is often used, as opposed to the magnitude of weight loss recommended in lifestyle interventions as initial treatment of cardiometabolic conditions	BMI-based definitions (WHO): Overweight: BMI 25 to < 30 kg/m² Class I obesity: BMI 30 to < 35 kg/m² Class II obesity: BMI 35 to < 40 kg/m² Class III obesity: BMI ≥ 40 kg/m² For Asian populations, overweight is defined as BMI 23 to 27.5 kg/m² and obesity ≥ 27.5 kg/m²	Weight gain and obesity represent a continuum associated with negative health outcomes	
Consequences	Increased risk of DM, hypertension, dyslipidemia, and CVD	Body image disturbance Increased risk of DM, hypertension, CVD, some cancers, obstructive sleep apnea, cholecystitis, erectile dysfunction, non-alcoholic fatty liver disease, osteoarthritis, depression, and neurocognitive impairment		
Contributing factors	Excess alcohol consumption	Sedentary lifestyle Altered sleep pattern Intake of excess or poor-quality calories (e.g., saturated fats, processed sugars) Excess alcohol consumption Some medications (e.g., psychotropic drugs, steroids, anti-diabetic drugs)		
Impact of ART	Initiation of ART increases weight as part of a retu INSTI and TAF may induce greater weight gain th	See Adverse effects of ARVs and drug classes		
Aim of intervention	Emphasise the importance of behaviour goals rati An objective of 5 - 10% weight loss may have ber ↑ 5% HDL cholesterol ↓ 5 mmHg systolic and diastolic BP in hyperten ↓ 0.5% (decrease 2.55 mmol/mol) HbA1c in DN • Improving sleep apnoea			
Management	Motivation to change: Discuss support systems (e.g. family, friends), motivating factors, and barriers to change Discuss benefits of making changes Set realistic and achievable lifestyle changes			
Lifestyle recommendations	Consider behavioral intervention (motivational interviewing, stimulus control or cognitive restructuring) along with self-monitoring; intensify behavioral intervention if several unsuccessful weight loss attempts		See Lifestyle Interventions	
General principles	Treat underlying or associated conditions There are several drugs specifically recommended for those with a BMI ≥ 30 kg/m² or ≥ 25 kg/m² and weight-related complications (DM, hypertension) (e.g. orlistat, phentermine/topiramate, lorcaserin, naltrexone/bupropion, liraglutide). These drugs should be prescribed by an endocrinologist or obesity expert. All of them may have adverse effects and drug-drug interactions with ART		Consider TDM (therapeutic drug monitoring) in obese persons.  † risk of virological failure with long acting CAB/RPV in obese persons	
Bariatric surgery		Medical devices or endoscopic procedures (e.g intragastric balloon, aspiration therapy, endoscopic sleeve gstrroplasty) or bariatric surgery should be considered in persons with a BMI ≥ 40 kg/m² or ≥ 35 kg/m² with obesity-related co-morbidities refractory to serious attempts at lifestyle changes and should be coordinated through an established, specialistled obesity programme.	Consider therapeutic drug monitoring and drug dose adjustment post-bariatric surgery	



# Hyperlactataemia and Lactic Acidosis: Diagnosis, Prevention and Management

Risk factors	Prevention/Diagnosis	Symptoms
HCV/HBV co-infection     Use of ribavirin     Liver disease	Routine monitoring of serum lactate levels not recommended - does not predict risk of lactic acidosis	Hyperlactataemia: unexplained nausea, abdominal pain, hepatomegaly, elevated ALT and/or AST, weight loss
• Low CD4 count	Measurement of serum lactate, bicarbonate &	Acidaemia: asthenia, dyspnoea, arrhythmias
• Pregnancy	arterial blood gases + pH indicated in case of	Guillain-Barré-like syndrome
Female sex	symptoms suggestive of hyperlactataemia	
Obesity	<ul> <li>Close monitoring for symptoms if &gt; 1 risk factor</li> </ul>	

# Management

Serum lactate (mmoL/L)	Symptoms	Action
> 5 <sup>(i)</sup>	Yes / No	<ul> <li>Repeat test under standardised conditions to confirm &amp; obtain arterial pH and bicarbonate(i)</li> <li>If confirmed, exclude other causes         <ul> <li>Arterial pH ↓ and/or bicarbonate ↓(i): Stop NRTIs</li> <li>Arterial pH and/or bicarbonate normal: Consider switch from high to low-risk NRTI &amp; monitor carefully OR stop NRTIs</li> </ul> </li> </ul>
2-5	Yes	Exclude other causes; if none found: watchfully follow up OR consider switch from high to low-risk NRTI, OR stop NRTI
2-5	No	Repeat test If confirmed, watchfully follow up
< 2		None

i Lactic acidosis is a rare but life-threatening situation usually associated with symptoms; high risk if serum lactate > 5 and especially > 10 mmoL/L

Management of lactic acidosis (irrespective of serum-lactate level)

Admit the person. Stop NRTIs. Provide iv fluids. Vitamin supplementation can be used (vitamin B complex forte 4 mL bid, riboflavin 20 mg bid, thiamine 100 mg bid; L-carnitine 1000 mg bid), although benefit is not proven



# **Travel**

General precautions	Delay travel until clinically stable and treatment established     Provide drug prescription and referral letter for emergencies     Provide medical certificate for import of personal medicines/syringes     Carry ARVs split between suitcase and hand luggage     Beware of fake drugs
ART	Maintain hours of medicines (e.g. 23.00 local time) when switching time zones, shortening the interval to the next dose when flying east
Acknowledge increased susceptibility <sup>(1)</sup> of persons with HIV	<ul> <li>1. Observe food hygiene</li> <li>Particularly important for travellers visiting friends and relatives (VFR)</li> <li>Bacterial enterocolitis e.g. diarrhoeagenic <i>E. coli, Salmonella, Shigella, Campylobacter</i></li> <li>Opportunistic intestinal parasitosis Cryptosporidium, Cyclospora, Cystoisospora, Microsporidia</li> <li>2. Prevent insect bites</li> <li>Repellents (DEET ≥ 30%), spray clothing with insecticide (permethrin)</li> <li>Sleep under insecticide-treated bednet</li> <li>Malaria chemoprophylaxis/emergency standby treatment<sup>(ii)</sup> (to be taken with meals)</li> <li>Yellow fever, see page 90</li> <li>Leishmaniasis beware of sand flies (dogs)</li> </ul>

Advice on travel restrictions, see http://www.hivtravel.org

- Higher intestinal susceptibility due to HIV-associated GALT destruction, low CD4 count. More severe malaria with CD4 count < 350 cells/µL</li>
   According to malaria risk at travel destination and national guidelines
- ii According to malaria risk at travel destination and national guidelines adherence counselling is particularly important in persons visiting friends and relatives. See Drug-drug Interactions between Anti-malarial Drugs and ARVs

# **Drug-drug Interactions between Anti-malarial Drugs and ARVs**

Ant	imalarial drugs	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
	amodiaquine	1	1	$\leftrightarrow$	1	1	$\leftrightarrow$	↑ a	↓?	↓29% <mark>a</mark>	$\leftrightarrow$										
	artemisinin	1	1	1	1	1	D	ļ	ţD	ţD	D	D	D	D	$\leftrightarrow$	D	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	atovaquone	$\leftrightarrow$	↓10%	$\leftrightarrow$	↓ b	↓74%b	$\leftrightarrow$	↓75% <mark>b</mark>	↓E55% <mark>b</mark>	↓ b	$\leftrightarrow$										
	chloroquine	↔ c,d	↔ c,d	↔ d	↔ d	↔ c,d	$\leftrightarrow$	↔ <b>e</b>	↔f	↔ f	↔ c,g	c,g	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔ c,g	$\leftrightarrow$	↔ d	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	clindamycin	1	1	1	1	1	$\leftrightarrow$	↓	<b>\</b>	<b>↓</b>	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
.ngs	doxycycline	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↓?	↓?	↓?	$\leftrightarrow$										
line dı	halofantrine	↑ <b>g</b>	↑ g	1	1	↑ g	$\leftrightarrow$	↓	1	<b>↓</b>	↔ g	↔ c,g	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔ g	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
second line drugs	hydroxy- chloroquine	↑ c,g	↑ c,g	1	1	↑ c,g	$\leftrightarrow$	↔ <b>e</b>	<b>1</b>	1	↔g	↔ c,g	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	⇔g	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
and s	lumefantrine	↑ c,g	↑ c,g	1	↑175%	↑382% c,g	$\leftrightarrow$	↓~40%	1	↓D46%	↔g	↔ g	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔g	↑10%	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
First line and	mefloquine	↑ c,g	↑ c,g	1	1	↓28%c,g	$\leftrightarrow$	<b>\</b>	1	↓	↔ g	↔ g	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔g	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Ē	piperaquine	↑ c,g	↑ c,g	↑ c	↑ c	↑ c,g	Е	<b>\</b>	<b>\</b>	1	Εg	↔ g	Е	Е	$\leftrightarrow$	↔ g	$\leftrightarrow$	↑ c	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	primaquine	↔ g	↔g	$\leftrightarrow$	$\leftrightarrow$	⇔g	$\leftrightarrow$	↔ h	↔h	↔ h	↔ g	↔ g	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔g	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	proguanil	$\leftrightarrow$	↓41%b	$\leftrightarrow$	↑p	↓38%b	$\leftrightarrow$	↓44%b	↓E55%b	↓ b	$\leftrightarrow$										
	pyrimethamine	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	quinine	↑ c,g	↑ c,g	1	1	↓56% <b>c</b> ,g	$\leftrightarrow$	<b>\</b>	1	1	⇔g	↔ c,g	Е	$\leftrightarrow$	$\leftrightarrow$	↔ g	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	sulfadoxine	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						

# Colour legend

No clinically significant interaction expected
These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration

Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

# Legend

Potential elevated exposure of the antimalarial drug
 Potential decreased exposure of the antimalarial drug

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd)
DRV/c DRV co-formulated with COBI (800/150 mg qd)
CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

### Interactions with ABC, FTC, 3TC, ZDV

ABC: no clinically relevant interactions expected.

FTC: increased FTC exposure with pyrimethamine, sulfadoxine.
3TC: increased 3TC exposure with pyrimethamine, sulfadoxine.
ZDV: potential additive haematological toxicity with amodiaquine, atovaquone, primaquine, pyrimethamine, sulfadoxine.

### Interactions with ibalizumab

None

# Comments

- a Liver toxicity.
- b Take with high fat meal, consider dose increase.
- c ECG monitoring is recommended.
- d Chloroquine concentrations may increase, but to a moderate extent. No dose adjustment is required but monitor toxicity.
- Chloroquine/hydroxychloroquine concentrations may increase or decrease. No dose adjustment is required but monitor toxicity and efficacy.
- f Chloroquine concentrations may decrease, but to a moderate extent. No dose adjustment is required but monitor efficacy.
- g Caution as both drugs can induce QT interval prolongation.
- h Increase of haemotoxic metabolites.

# Further Information

For additional drug-drug interactions and for more detailed pharmacokinetic interaction data and dosage adjustments, please refer to: http://www.hiv-druginteractions.org (University of Liverpool)



# **Vaccination**

- Vaccinate according to national guidelines for healthy population, preferably after having achieved suppressed viraemia and immune reconstitution (CD4 count > 200 cells/µL)
- Consider repeating vaccinations performed at CD4 count < 200 cells/µL (< 14%) or unsuppressed viraemia once adequate immune reconstitution is achieved (HIV-VL undetectable and CD4 count > 200 cells/µL)
- As vaccine responses may be significantly lower in persons with HIV
   (i.e. lower seroconversion rates, faster titer decline), do not use
   rapid schedules (e.g. rabies, tick-borne encephalitis, HAV/HBV) and
   consider antibody titres to assess their effectiveness if vaccinated
   at CD4 count < 200 cells/µL or unsuppressed viremia (e.g. rabies,
   tick-borne encephalitis, HAV, meningococci). Be attentive to observe
   boosters and all post-exposure measures (particularly after potential
   rabies exposure)</li>
- Avoid polysaccharide vaccination
- For background data, see http://www.bhiva.org/vaccination-guidelines. aspx

- For attenuated live vaccines(i)
  - (in addition to restrictions for general population):
    - \*Varicella, measles, mumps, rubella, yellow fever
       Contraindicated if CD4 count < 200 cells/µL (14%) and/or AIDS.</li>
       Impaired protection after vaccination with unsuppressed viraemia.
       Administer immunoglobulins if exposed and not yet vaccinated
  - Oral live typhoid

Preferred if CD4 count > 200 cells/µL (> 14%). Contraindicated if CD4 count < 200 cells/µL (14%): then give inactivated parenteral polysaccharide vaccine

Infection	Vaccination rationale	Comment
Influenza Virus	Higher rate of pneumonia. Explicitly recommended in all persons with HIV	Yearly, use 4-valent vaccine if available
Human Papilloma Virus (HPV)	Shared risk with HIV of contracting infection. Higher rate of cervical and anal cancer	Vaccinate with 3 doses between ages 9 and 45 (health insurance coverage differs by country according to age, sex, sexual orientation).  Use 9-valent vaccine if available.  Persons treated for high grade dysplasia could benefit from a full course vaccination for secondary prevention
Hepatitis B Virus (HBV)	Shared risk with HIV of contracting infection. Untreated HIV accelerates progression of liver disease	Vaccinate if seronegative. Repeat doses until anti-HBs antibodies $\geq 10 \ IU/L$ / $\geq 100 \ IU/L$ according to national Guidelines. In order to reach $\geq 100 \ IU/L$ in non-responders repeat 3 doses if anti-HBs < 10 IU/L, 1 dose if anti-HBs < 100 IU; 0 consider double dose (40 $\mu g$ ) or use more immunogenic vaccines in particular with low CD4 count and high HIV-VL. No benefit for intradermal application. See page 115
Hepatitis A Virus (HAV)	According to risk profile (travel, close contact with children, MSM, IVDU, active hepatitis B or C infection, chronic liver disease)	Vaccinate if seronegative. Consider checking antibody titres in persons with high risk. Weaker immune response expected with HAV/HBV co-vaccine. See page 115
Neisseria meningitidis	According to risk profile (travel, close contact with children, MSM)	Use conjugated 4-valent vaccine (for serotypes A, C, W-135, Y; 2 doses 1-2 months apart) if available. Booster every five years if exposure continues. Polysaccharide vaccine no longer recommended. Vaccination against Meningococcus serotype B according to national Guidelines
Streptococcus pneumoniae	Higher rate and severity of invasive disease. Vaccine explicitly recommended for all persons with HIV	One dose of a conjugated vaccine: PCV-13, PCV-15 or PCV-20a for all persons according to availability and national guidelines, also if pre-vaccinated with PPV-23 polysaccharide vaccine. For patients vaccinated with PCV-13 or PCV-15 one dose of PPV-23 at least 2 months after the conjugate vaccine may be considered in some national guidelines for all persons with HIV
Varicella Zoster Virus (VZV)	Higher rate and severity of both chicken- pox and zoster	Perform serology if exposure history negative. Vaccinate if seronegative. For contraindications, see*. To prevent shingles, preferably use adjuvant recombinant sub-unit vaccine over live-attenuated vaccine according to national guidelines
Yellow Fever Virus	Mandatory for travel to selected countries (provide exemption letter if no true risk of exposure)	Contraindicated if past or current haematological neoplasia or thymus affection (thymoma, resection/radiation) For other contraindications, see*. Booster q 10 years
Rabies		For persons with CD4 count < 200 cells/µL or unsuppressed viremia consider pre-exposure vaccination with 3 doses (0, 7, 28 days) and titre control 14 days later. In case of exposure: full post-exposure prophylaxis including rabies immunoglobulins (RIG). If pre-exposure rabies vaccination administered when CD4 > 200 cells/µL: Post-exposure prophylaxis as for immunocompetent (one dose day 0 and day 3, without RIG)
Severe Acute Respiratory Syndrome 2 (SARS-CoV-2)	Low CD4 count and non-suppressed HIV-VL may increase the risk of acquiring SARS-CoV-2 infection and/or progressing to severe COVID-19	In a pandemic situation, all persons with HIV should be vaccinated according to the national guidelines irrespective of CD4 count and HIV-VL. Advanced HIV infection (CD4 count < 200 cells/µL) and persons with detectable HIV viremia have poorer humoral immune responses and are candidates for COVID-19 booster doses

- i Administer live vaccines simultaneously or with an interval of 4 weeks
- ii In case of non-response, ART should contain TDF or TAF
- iii Conjugated vaccines are more immunogenic, induce memory cells, respond to boosting and reduce mucosal colonisation



# **Sexual and Reproductive Health**

Screening questions about sexual and reproductive health and sexual function should be routinely asked at HIV consultation.

Effective Measures to Reduce Sexual transmission of HIV										
Measure	Comment									
ART for HIV-positive partner	Undetectable equals untransmissible (U=U) from 6 months of fully suppressive ART if no active STIs     Consider in e.g. sero-different couples <sup>(i)</sup>									
Pre-exposure prophylaxis (PrEP)	Effective in HIV-negative persons with high risk sexual situations, see Pre-exposure prophylaxis (PrEP)									
Post-exposure prophylaxis (PEP)	Consider after situations of unprotected anal or vaginal intercourse, if one partner has detectable HIV-VL and the other partner is seronegative     Start as soon as possible and within 48/72 hours post sexual exposure     See Post-exposure prophylaxis (PEP)									
Male condom or female condom use	Effective in treated and untreated persons									

U=U should be discussed with all persons with HIV, at diagnosis and when starting/switching ART. The evidence is now clear that a person with an undetectable VL do not transmit HIV sexually. Large studies of sexual HIV transmission among thousands of sero-different couples, one partner of which was living with HIV and the other was not, were undertaken in recent years. In those studies, there was not a single case of linked sexual transmission of HIV from a virally suppressed person with HIV to their HIV-negative partner. However, a person can only know whether he or she is virally suppressed by taking a VL test.

i see page 12

### Reproductive health

All persons should be asked about their reproductive goals at HIV diagnosis and in follow-up and receive appropriate and ongoing reproductive counselling. Providing contraception and reproductive counselling to women living with HIV is essential if pregnancy is not currently desired.

#### Conception:

Reproductive health issues should be preferentially discussed with all partners, particularly in sero-different couples. See Drug-drug Interactions between Contraceptives and ARVs

# Approaches for sero-different couples who want to have children:

Ensuring the partner living with HIV is on fully suppressive ART should be a primary goal for people who wish to conceive. Screening for STIs (and treatment, if required) of both partners is strongly recommended if conception is planned.

For ART in women living with HIV wishing to conceive, see pages 18-19

The following list represents selected measures with increasing safety for sero-different couples without active STIs:

- Intercourse without condoms during times of maximum fertility (determined by ovulation monitoring), if the partner living with HIV has undetectable HIV-VI
- PrEP in the absence of HIV viral suppression e.g. during the first 6 months of ART or if uncertainty about HIV-positive partner's adherence see Pre-exposure Prophylaxis (PrEP)
- Vaginal syringe injection of seminal fluid during times of maximum fertility if the male partner is HIV-negative
  - Sperm washing, with or without intra-cytoplasmic sperm injection, is no longer recommended because of effectiveness of ART in avoiding HIV transmission at conception in male persons with HIV with undetectable HIV-VL

# Contraception

Women living with HIV of childbearing age should be offered contraception counselling. If hormonal contraceptives are preferred options, EFV should be avoided as it can impair the efficacy of the contraceptive method. Boosted regimens can be used with some contraceptive methods, see <a href="Drug-Drug Interactions">Drug-Drug Interactions</a> between Contraceptives and ARVs. Otherwise intra-uterine device should be offered as the preferred option due to its high effectiveness, well established safety and no DDIs. STI and HIV transmission risk should be carefully discussed along with contraception counseling

#### Menopause

#### Education

Healthcare providers should present accessible information on menopause to women and encourage the use of self-assessment tools (eg. Menopause Rating Scale (MRS), Greene Climacteric Scale (GCS), see also Mental Health, Depression: Screening and Diagnosis, Anxiety Dosorders: Screening and Diagnosis

### Screening

We recommend yearly, pro-active assessment of menopausal symptoms in women living with HIV aged > 40 years using a validated menopause symptom questionnaire, such as the MRS or GCS

### General health risk assessment for women age > 40 years

- Cancer, see Cancer screening methods
- ii Assessment of bone mineral density (BMD), see Bone Disease: Screening and Diagnosis
  - Assess risk factors for low BMD. If BMD is normal at initial assessment, consider fracture risk using FRAX® every 3-5 years
  - Consider DXA in women with 10-year major osteoporotic fracture risk
     20% based on FRAX® regardless of menopausal status
  - Consider DXA in women with prior history of low impact fracture regardless of menopausal status
  - Reassess DXA in those with osteoporosis after 2 years if on treatment to ensure response and reassess need for continued treatment after 3-5 years
- iii CVD risk assessment yearly, especially in women with vasomotor symptoms, see Prevention of Cardiovascular Disease
- Wental health screen for anxiety and depression, consider screening tools such as GAD-2, see also Mental Health, Depression: Screening and Diagnosis

# Treatment for menopausal women

- Topical (vaginal) hormone replacement therapy (HRT) should be considered in all woman given the positive effects on sexual health and urogenital symptoms
- Vi Systemic HRT should be considered in women experiencing vasomotor, mood or urogenital symptoms.
- vii Transdermal estrogen (with progesterone if a woman has a uterus) is the preferred HRT option due to the lower thromboembolic risk. See Drugdrug interactions between HRT and ARVs
- iii Women with premature ovarian insufficiency should be offered HRT until at least the expected age of menopause (eg. aged 50-52 years) to reduce longer term morbidity and mortality risk



# Special considerations regarding transgender people

HIV and general medical care, including sexual health services, are often not designed to cover the specific needs of transgender people. Transgender people are often not included in gender-specific health care surveillance programmes.

Using a two-stage question helps both individual care and the development of appropriate services.

- (i) What is your current sex?
- (ii) Is this the same sex you were given at birth?

### Sex, gender and sexuality

Although sex is sometimes wrongly decided at birth, it is also independent of sexuality. Specific care for people who are transgender includes medical issues linked to biology (for example cervical screen for some trans men) and social factors (linked to the design of services in a clinic setting, appropriate naming, gender-neutral facilities).

Sexuality cannot be assumed by either sex or gender

#### In general:

- · ART is equally effective for trans and cis gender people
- · Access to and management of gender affirming hormones
- See dosage recommendation for hormone therapy when used at high doses for gender transitioning
- Support for good sexual health and access to reproductive services are equally important for trans people
- · There are minimal data about STIs

# Sexual dysfunction

Guidelines for treatment of sexual dysfunction in the general population are available. Refer to specialist where appropriate, see Sexual Dysfunction and Treatment of Sexual Dysfunction



# STI screening and treatment

STI screening should be offered to all sexually active persons at the time of HIV diagnosis, annually thereafter or at any time STI symptoms are reported and during pregnancy. More frequent screening at three-month intervals is warranted for persons at particularly high risk of STIs, including those with multiple or anonymous partners. Frequent HIV screening is also essential for those on PrEP, see Pre-exposure Prophylaxis (PrEP)

Diagnosis procedures should follow local or national guidelines. More comprehensive advice can be found at https://iusti.org/treatment-guidelines/

The following STIs should be universally considered in persons with HIV and their sexual partner(s):

	Therapy	Comment
Chlamydia infection	Consider doxycycline (100 mg po bid 7-10 days, contraindicated in pregnancy) for urethritis and cervicitis (1)  Preferred if rectal infection  Or alternatively: azithromycin 1 g po as a single dose  If rectal infection a test of cure (TOC) should be performed  For Lymphogranuloma venereum (LGV) doxycycline (100 mg po bid for 21 days)  Alternatives:  erythromycin (500 mg po qid(1)) or levofloxacin (500 mg po qd) for 7 days (or erythromycin 500 mg po qid(1)) for 21 days in case of LGV)	May cause therapy-resistant proctitis in HIV-positive MSM     Screening recommended at genital, rectal and pharyngeal sites according to exposure     Rectal and pharyngeal infections are usually asymptomatic     Consider co-infections with Neisseria gonorrhoeae     Avoid sexual activity for 7 days post treatment initiation     Individuals should only resume having sex after symptoms have resolved and sex partners have been treated     The same treatment for LGV is recommended for asymptomatic individuals and contacts of individuals with LGV
Gonorrhoea	Ceftriaxone (1 g im as a single dose) <sup>(i)</sup>	Can cause proctitis, prostatitis and epididymitis Screening recommended at genital, rectal and pharyngeal sites according to exposure Rectal and pharyngeal infections are usually asymptomatic Often asymptomatic in women Avoid sexual activity for 7 days post treatment initiation Individuals should only resume having sex after symptoms have resolved and sex partners have been treated Fluoroquinolone resistance is highly prevalent in all regions Note ceftriaxone 1 g im as a single dose is based on recent BHIVA recommendations, https://www.bhiva.org/guidelines. IUSTI Guidelines recommend 500 mg im with azithromycin 2 g as a single dose, however these recommendations have not been updated in several years, https://iusti.org/regions/guidelines/
HBV infection HCV infection	See detailed information on HIV/HCV or HIV/HBV co-infections, pages 116-117	Interruption of TDF, 3TC or FTC can lead to HBV reactivation     Clusters of acute HAV and HCV infection in HIV-positive MSM across Europe     See Vaccination
HPV infection	There are several treatment modalities for the management of genital warts with no evidence to suggest one approach is better than another approach. Consider operative removal by laser surgery, infrared coagulation, cryotherapy, etc.  Management of both pre-invasive cervical lesions as well as peri- and intra-anal lesions should follow local or national guidelines	Infection is mostly asymptomatic; relapse of genital warts is frequent     Cervical PAP smear test recommended in all HIV-positive women     Anal HPV screening and cytology should be considered in all persons with HIV practicing anal sex     Consider high resolution anoscopy in case of suspicious cytological findings (rectal palpation or external inspection is not sufficient)     See Vaccination
HSV infection	Primary infection: aciclovir (400-800 mg po tid), famciclovir (250-500 mg po tid) or valaciclovir (1000 mg po bid) for 7-10 days  Recurrent episodes: aciclovir (400 mg po tid) or valaciclovir (500 mg po bid) for 5-10 days  Suppressive management: Chronic suppressive therapy is usually offered to persons who experience six or more clinical episodes per year or who experience significant anxiety or distress related to their clinical recurrences. Chronic suppression: aciclovir (400-800 mg bid or tid) or famciclovir 500 mg bid or valaciclovir 500 mg po bid	Treatment of HSV2 alone does not prevent HIV-transmission and only modestly prevents HIV disease progression
Syphilis	Penicillin is the gold standard for the treatment of syphilis in both pregnant and non-pregnant individuals.  Primary/secondary syphilis: benzathine penicillin G (2.4 million IU im as single dose). In early syphilis adjunctive treatment with prednisolone (20-60 mg po daily for 3 days) prevents optic neuritis, uveitis and Jarisch-Herxheimer reaction  Alternative regimen include doxyycline (100 mg po bid for 14 days)  Late latent syphilis and syphilis of unknown duration: benzathine penicillin (2.4 million IU im weekly on days 1, 8 and 15); the alternative doxycycline (100 mg po bid for 4 weeks) is considered less effective  Neurosyphilis: penicillin G (6 x 3 - 4 million IU iv for at least 2 weeks)  There is no evidence to give a general recommendation on prednisolone use in this condition  Alternative regimen: ceftriaxone (2 g iv daily for 10 to 14 days) if the person can be safely treated with other beta-lactam drugs. Doxycycline (200 mg po bid) for 21 days is also an alternative approach, but should be reserved for exceptional circumstances. This regimen has very limited supporting data <sup>(0)</sup>	Expect atypical serology and clinical courses     Consider cerebrospinal fluid (CSF) testing in persons with neurological symptoms (evidence for intrathecally-produced specific antibodies, pleocytosis, etc.) or late latent syphilis     Successful therapy clears clinical symptoms and decreases VDRL test four-fold within 6-12 months

- i Refer to local Guidelines
- ii Rarely used



# **Sexual Dysfunction**

When sexual complaints exist:	What is the exact nature of the problem? In which phase(s) of the sexual response cycle does the problem occur?	1. Desire (lack of sexual desire or libido; desire discrepancy with partner; aversion to sexual activity) 2. Arousal (difficulties with physical and/or subjective sexual arousal; difficulties or inability to achieve or sustain an erection of sufficient rigidity for sexual intercourse (men); i.e. erectile dysfunction; lack or impaired nocturnal erections (men); difficulties lubricating (women); difficulties sustaining arousal) 3. Orgasm (difficulties experiencing orgasm) 4. Pain (pain with sexual activity; difficulties with vaginal/anal penetration—anxiety, muscle tension; lack of sexual satisfaction and pleasure)									
	Self-assessment of sexual function (questionnaires):	Men International Index of Erectile Function, see Rosen RC, Riley A, Wagner G et al Women Female Sexual Function Index (FSFI), see https://www.fertstert.org/article/S0015-0282%2809%2902741-1/fulltext									
Check for endo- crine causes:	Signs of hypogonadism	Men  - Look for signs of testosterone insufficiency (main: decreased or absent nocturnal erections, decrease in testes size, decreased volume of ejaculate, hot flushes, sweats, reduction of body hair and beard; others: reduced sexual arousal and libido, decreased frequency of sexual thoughts and fantasies, decreased genital sensitivity, erectile dysfunction, loss of vitality; fatigue; loss of muscle mass and muscle strength)  - If signs or symptoms of hypogonadism are present ask for hormonal assessment: lutropin hormone (LH), follicle stimulating hormone (FSH), total testosterone; sex hormone-binding globulin evaluation to calculate free testosterone, see http://www.issam.ch/freetesto.htm	If hypogonadism is present (total testosterone < 300 ng/dL or calculated free testosterone below normal): refer to endocrinologist or andrologist  If hypogonadism is not present: check for other causes								
		Women - Look for signs of estradiol insufficiency/menopause (amenor-rhoea or missed menstrual periods, vaginal dryness, hot flashes, night sweats, sleep disturbances, emotional lability, fatigue, recurrent urogenital infections) - If symptoms of menopause are present ask for hormonal assessment: LH, FSH, estradiol	If symptoms of menopause are present: refer to endocrinologist or gynaecologist  If hypogonadism is not present: check for other causes								
Check for other causes:	Psychological or sociological problems	Stigma, body image alteration, depression, fear of infecting an HIV-negative partner, anxiety, awareness of a chronic disease, condom use	Refer to clinical psychologist								
	Infections	Men - Urogenital infections (note: if complete sexual response possible, e.g. with another partner, with masturbation or nocturnal erections, then no major somatic factors are involved)	Refer to urologist, andrologist, cardiologist								
		Women - Urogenital infections	Refer to gynaecologist								
	Relevant medicines, recreational drugs, alcohol, smoking and other lifestyle factors	Drugs associated with sexual dysfunction: 1) Psychotropics – Men and Women (antidepressants, antiepileptics, antipsychotics, benzodiazepines), 2) Lipid-lowering drugs - Men (statins, fibrates), 3) Antihypertensives - Men (ACE-inhibitors, betablockers, alfablockers), 4) Others - Men and Women (omeprazole, spironolactone, metoclopramide, finasteride, cimetidine); 5) Men and Women - contribution from ART is controversial and benefit from switching studies is not proven	Consider therapy changes								



# **Treatment of Sexual Dysfunction**

Men	Women
Treatment of erectile dysfunction	Sexual pain
Primarily oral PDE5-inhibitors (sildenafil, tadalafil, vardenafil).  • All at least 30 minutes before initiation of sexual activity  • Use lower dose if on Pl/b  - sildenafil (25 mg every 48 hours)  - tadalafil 5 mg initial dose with maximum dose 10 mg in 72 hours  - vardenafil 2.5 mg maximum dose in 72 hours  Cave: Poppers have a synergistic effect with PD5-blockers which can lead to profound hypotension thus concurrent use is not recommended  • Tadalafil also licensed for use as an everyday ongoing therapy	Counselling Local hormone therapy Pelvic physiotherapy Vaginal/rectal suppositories Topical lidocaine Capsaicin Vestibulectomy
Treatment of premature ejaculation	Low desire
<ul> <li>Consider behavioural interventions and/or psychosexual counselling, SSRIs, tricyclic antidepressants, clomipramine and topical anaesthetics</li> <li>Use lower dose of clomipramine and other tricyclic antidepressants if on PI/r, see Drug-drug interactions between antidepressants and ARV</li> <li>Dapoxetine, a short-acting SSRI, is the only drug approved for ondemand treatment of premature ejaculation in Europe. Dapoxetine is contraindicated with boosted ARVs</li> <li>Treatment must be maintained as recurrence is highly likely following withdrawal of medicine</li> </ul>	Counselling Hormonal therapy Bupropion Flibanserin (contraindicated with boosted ARVs due to risk of hypotension)
	Low arousal
	Counselling Hormonal therapy PDE5 inhibitors (e.g., sildenafil)
	Orgasmic dysfunction
	Mindfulness, sex therapy Hormonal therapy Bupriopion PDE inhibitors (e.g., sildenafil) Yohimbine hydrocholoride (concomitant use of boosted ARVs may increase BP)



# **Mental Health: Depression and Anxiety Disorders**

# **Depression: Screening and Diagnosis**

# Significance

- · A higher prevalence of depression is reported in persons with HIV described in 20-40% versus 7% in general population
- · Significant disability and poorer HIV treatment outcomes are associated with depression
- · Depressive disorders are often associated with a significant anxiety and poor overall wellbeing

#### Screening and diagnosis of depression How to diagnose? Screening of all persons recom-· Screen every 1-2 years Symptoms - evaluate regularly mended in view of the high preva-· Two questions A. At least 2 weeks of depressed mood lence of depression 1. Have you often felt depressed, OR sad or without hope in the last B. Loss of interest Populations at particularly high OR few months? 2. Have you lost interest in activities C. Diminished sense of pleasure risk · Positive history of depression in that you usually enjoy? **PLUS** Rule out other medical conditions 4 out of 7 of the following: Depressive episode in personal (such as hypothyroidism, 1. Weight change of ≥ 5% in one month or a persistent change of appetite hypogonadism, Cushing's 2. Insomnia or hypersomnia on most days history syndrome, vitamin B12 deficiency) Older age 3. Changes in speed of thought and movement Adolescence Rule out depressive symptoms 4. Fatigue Persons with history of drug adsecondary to ART and non-ART 5. Feelings of guilt and worthlessness diction, psychiatric, neurologic or medication (such as EFV) 6. Diminished concentration and decisiveness severe somatic co-morbidity · Assessment of the risk of suicide 7. Suicidal ideation or a suicide attempt( Use of EFV should be done with the following questions Use of neurotropic and recreational · Are these just ideas? As part of investigation of neuro-Are they intrusive and how many? · How much control do you have cognitive impairment, see page 104 Socially isolated (particular over these ideas? relevance during COVID-19 · Have you made a plan? pandemic) · Are you about to take action?

i EFV has been associated with a higher risk of suicidal ideation



# **Depression: Management**

Degree of depression	Number of symptoms (see page 96: A, B or C + 4/7)	Treatment	Consultation with expert
No	< 4	No	
Mild	4	Problem-focused consultation     Consider antidepressant treatment <sup>(i)</sup> Recommend physical activity	<ul> <li>Always if treating doctor is unfamiliar with use of antidepressants</li> <li>If depression not responding to treatment</li> <li>If person has suicidal ideation</li> <li>In case of complex situations such as drug addiction, anxiety disorders,</li> </ul>
Intermediate	5-6	Start antidepressant treatment (i,ii,iii)	personality disorders, dementia, acute severe life events • Clinical improvement with antidepressants may take up to 4 weeks;
Severe	> 6	Refer to expert (essential)(iv)	there is no need to change antidepressants before this time.  Dose increment of antidepressant may be considered

- i See Drug-drug Interactions between Antidepressants and ARVs
- iii There is an increased risk of suicide and serious traffic accident in the first 15 days of antidepressant treatment; frequent monitoring in groups 5 and 6 is required during this period
- iii In groups 4, 5 and 6, psychotherapeutic follow-up (e.g. cognitive behavioral therapy CBT) may be indicated (consult with expert advice)
- iv Mental health professionals should always be consulted if there is a risk of suicide

If a person is diagnosed with depression switching off EFV to another third ARV drug according to switch rules is recommended



# Classification, Doses, Safety and Adverse Effects of Antidepressants

Mechanisms & classification	Start dose	Standard dose	Lethality in overdose	Insomnia and agitation <sup>(ii)</sup>	Sedation	Nausea or GI effects	Sexual dysfunction	Weight gain	
	mg	/day							
Selective seroto	nin-reuptake inhib	itors (SSRIs) <sup>(i)</sup>							
paroxetine	10-20	20-40	Low	+	-/+	+	++	++	
sertraline	25-50	50-150	Low	+	-/+	+	+	+(iii)	
citalopram	10-20	20-40	Low	+	-/+	+	+	+(iii)	
escitalopram	5-10	10-20	Low	+	-/+	+	+	+(iii)	
Mixed or dual-ac	tion reuptake inhi	bitors							
venlafaxine	37.5-75	75-225	Moderate	++	-/+	+	+	-/+	
Mixed-action nev	wer agents								
mirtazapine	30	30-60	Low	-/+	++	-/+	-/+	++	

- none
- + moderate
- ++ severe
- For many persons, SSRI induction may be associated with adverse effects (GI tract, dizziness, anxiety, panic attacks). Commencing at lower doses (i.e. 10, 25 & 10 mg for paroxetine, sertraline and citalopram, respectively) and increasing to the above starting doses after 4 to 7 days if tolerated may reduce such effects
- ii Insomnia is associated with DTG and other INSTI containing ART regimens and with the use of some antidepressants. Clinicians should be aware when prescribing DTG and INSTI and antidepressants together
- iii Weight gain may be significant but gradual and insidious



# **Drug-drug Interactions between Antidepressants and ARVs**

Ant	idepressants	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
NaSSA	mirtazapine	†a	†a	1	1	↑a	$\leftrightarrow$	ļ	ļ	1	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	citalopram	↑ a,b	↑ a,b	1	1	↑ a,b	$\leftrightarrow$	↓	<b>\</b>	↓	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	escitalopram	↑ a,b	↑ a,b	1	1	↑ a,b	$\leftrightarrow$	↓	1	<b>↓</b>	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
_	fluoxetine	1	1	1	1	†a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
SSRI	fluvoxamine	1	1	1	1	†a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	Е	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	paroxetine	<b>↑↓?</b>	<b>↑↓?</b>	<b>↑↓?</b>	↓39%	<b>↑↓?</b>	$\leftrightarrow$	$\leftrightarrow$	↑3%	$\leftrightarrow$	<b>↑↓?</b>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$							
	sertraline	1	1	1	↓49%	↓a	$\leftrightarrow$	↓39%	$\downarrow$	$\downarrow$	$\leftrightarrow$	↓7%	$\leftrightarrow$	↑9%	$\leftrightarrow$						
	vortioxetine	↑c	↑c	↑ <b>c</b>	↑c	↑c	$\leftrightarrow$	↑c	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
	desvenlafaxine	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
SNRI	duloxetine	1	↑↓	1	↑↓	↑↓	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
S	milnacipran	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	venlafaxine	↑a	†a	1	1	↑ <mark>a</mark>	$\leftrightarrow$	↓	1	↓ ↓	↔a	↔a	D	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	amitriptyline	1	1	1	1	↑ a,b	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
	clomipramine	↑ a,b	↑ a,b	↑ <b>b</b>	↑b	↑ a,b	$\leftrightarrow$	↓	1	↓	$\leftrightarrow$	↑b	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	desipramine	↑a	↑a	1	1	↑5% <mark>a</mark>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
TCA	doxepin	1	1	1	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
	imipramine	↑ a,b	↑ a,b	↑ <b>b</b>	↑b	↑ a,b	$\leftrightarrow$	↓	1	ļ	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	↑b	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	nortriptyline	↑a	↑a	1	1	↑ <mark>a</mark>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	trimipramine	↑a	†a	1	1	↑ <mark>a</mark>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
TeCA	maprotiline	↑a	†a	1	1	↑ <mark>a</mark>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
<u>e</u>	mianserin	↑a	↑a	1	1	†a	$\leftrightarrow$	↓	1	↓ ↓	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	agomelatine	$\leftrightarrow$	ļ	$\leftrightarrow$	<b>↓</b>	<b>↓</b>	$\leftrightarrow$														
	bupropion	$\leftrightarrow$	1	$\leftrightarrow$	$\downarrow$	↓57%	$\leftrightarrow$	↓55%	$\leftrightarrow$	$\downarrow$	$\leftrightarrow$	↑?	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	nefazodone	1	1	1	1	1	Е	ţΕ	↓E	↓E	Е	Е	Е	Е	$\leftrightarrow$	Е	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
ers	phenelzine	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Others	reboxetine	1	1	1	1	1	$\leftrightarrow$	1	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	St John's wort	Dd	Dd	Dd	Dd	Dd	Dd	Dd	Dd	Dd	Dd	Dd	Dd	Dd	$\leftrightarrow$	Dd	De	Dd	D	Dd	$\leftrightarrow$
	tranylcypromine	1	1	1	1	1	$\leftrightarrow$	↓	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	trazodone	↑ a,b	↑ <b>a</b> ,b	1	1	↑ a,b	$\leftrightarrow$	↓ ·	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						

# Colour legend

No clinically significant interaction expected
These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

# Legend

Potential elevated exposure of the antidepressant
 Potential decreased exposure of the antidepressant

→ No significant effect

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd) DRV/c DRV co-formulated with COBI (800/150 mg qd) CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

NaSSA noradrenergic specific serotonergic antidepressant

**SSRI** selective serotonin reuptake inhibitors

SNRI serotonin and norepinephrine reuptake inhibitors

TCA tricyclic antidepressants
TeCA tetracyclic antidepressants

# Interactions with ABC, FTC, 3TC, ZDV

ABC, FTC, 3TC, ZDV: no clinically relevant interactions expected.

# Interactions with ibalizumab

None

### Comments

- a Caution as both drugs can induce QT interval prolongation.
- b ECG monitoring is recommended.
- c Based on the patient clinical response, a lower dose of vortioxetine may be needed in poor CYP2D6 metabolizers in the presence of a strong CYP3A4 inhibitor.
- d A study suggests a low risk of a clinically relevant pharmacokinetic interaction with low-hyperforin formulations (< 1 mg/day) of St John's Wort (hyperforin is the constituent responsible for induction of CYPs and P-gp). Coadministration may be considered with St John's Wort formulations that clearly state the hyperforin content and which have a total daily hyperforin dose of 1 mg or less.
- e The European SmPC recommends DTG 50 mg bid in persons without INSTI resistance. The US Prescribing Information recommends that co-administration should be avoided as there are insufficient data to make dosing recommendations.

# **Further Information**

For additional drug-drug interactions and for more detailed pharmacokinetic interaction data and dosage adjustments, please refer to: http://www.hiv-druginteractions.org (University of Liverpool)

# **Anxiety Disorders: Screening and Diagnosis**

# Significance

- Studies which included a diagnostic interview report a high prevalence of anxiety disorders in persons with HIV<sup>®</sup>
- Specific anxiety disorders include the following:
  - panic disorder (10% in persons with HIV)
  - generalized anxiety disorder (5.6% persons with HIV)
  - · social anxiety disorder (9% persons with HIV)
- · post-traumatic stress disorder (PTSD)
- · Significant disability and poorer HIV treatment outcomes are associated with anxiety
- · Anxiety disorders are often associated with substance use behavior

#### Screening and diagnosis of anxiety How to diagnose? Consider screening all persons Generalised Anxiety Disorder-2 If GAD-2 cut-off score of ≥ 3, ask the following questions to diagnose with HIV recommended at each (GAD-2) Screening tool<sup>(1)</sup>: General Anxiety Disorder: clinic visit (in view of the high 'Over the last 2 weeks, how often prevalence of anxiety) have you been bothered by the excessive anxiety for more days than not over 6 months following problems?' difficulty controlling worry Populations at particularly high Feeling nervous, anxious or on associated with at least three of these symptoms (restlessness, fatigue, risk difficulty concentrating, irritability, muscle tension, sleep disturbances) Positive history of anxiety · Not being able to stop or control significant life impairment disorders in family not attributable to another substance or medical condition worrying Anxious personality not being better explained by another medical disorder Alcohol excess Score each question and calculate As part of investigation of cognitive Seek expert advice to diagnose panic disorders, social phobia and PTSD 0. Not at all impairment, see page 104 Rule out hyperthyroidism, hypoglycemia and hyperadrenocorticism. Multiple stressful life events 1. Several days (particular relevance during 2. More than half the days Exclude caffeine excess and use of stimulants (such as cocaine, crystal COVID-19 pandemic) 3. Nearly every day meth, amphetamines)

i GAD-2 score is a validated screening tool in persons with HIV, https://www.hiv.uw.edu/page/mental-health-screening/gad-2



# **Anxiety Disorders: Management**

Degree of anxiety disorders	GAD-2 Score	Treatment	Consultation with expert
Minimal	< 3	Relaxation techniques	
Significant	≥ 3	Recommend relaxation techniques Consider benzodiazepines, mainly clonazepam or lorazepam for a short period of time (less than 4 weeks) Consider antidepressant treatment with SSRI(I) Consider psychotherapeutic intervention: Cognitive Behavioral Therapy Cognitive Behavioral Stress Management Mindfulness-based Cognitive Therapy Peer Support Counseling	<ul> <li>Always if treating doctor is unfamiliar with use of antidepressants</li> <li>If anxiety not responding to treatment</li> <li>If person has suicidal ideation</li> <li>In case of complex situations such as drug addiction, anxiety disorders, personality disorders, dementia, acute severe life events</li> <li>Clinical improvement with antidepressants may take up to 4 weeks; there is no need to change antidepressants before this time Dose increment of antidepressant may be considered</li> </ul>
Generalized anxiety disorder		Start antidepressant treatment with SSRI and benzodiazepine if needed (to reduce anxiety faster) <sup>(i, ii)</sup> Refer to mental health expert to start psychotherapeutic intervention	

i See Drug-drug Interactions between Anxiolytics and ARVs
 ii Mental health professionals should always be consulted if there is a risk of suicide

# **Classification, Doses and Adverse Effects of Anxiolytics**

Mechanisms & classification	Starting dose	Usual therapeutic daily dose	Lethality in overdose	Insomnia and/or agitation	Sedation	Nausea or GI effects	Sexual dysfunction	Weight gain
Benzodiazepines	3							
alprazolam			no (unless if combined with other CNS drugs)	++	+++	++	++	++
chlordiazepoxide	5 mg qd	10-100 mg	no (unless if combined with other CNS drugs)	frequency unknown	++	rare	rare	frequency unknown
clonazepam	0.25 mg bid	1-2 mg	no (unless if combined with other CNS drugs)	+	++	rare	+	+
oxazepam	zepam 10 mg tid 30-60 mg n		no (unless if combined with other CNS drugs)	frequency unknown	++	rare	rare	no
Selective serotor	nin reuptake inhib	itors						
escitalopram	10 mg qd	10-20 mg	no (unless if combined with other CNS drugs)	++	++	+++	++	+
paroxetine	20 mg qd	20-60 mg	no (unless if combined with other CNS drugs)	++	++	+++	+++	++
Serotonin and no	orepinephrine reu	ptake inhibitors			I		_	
duloxetine	30 mg qd	30-60 mg	yes (at > 1000 mg)	++	+++	+++	++	+
venlafaxine	75 mg qd	75-225 mg	yes	+++	+++	+++	++	++
Others								,
buspirone	5 mg bid or tid	15-60 mg (60 mg)	no	++	+++	++	no	frequency unknown
hydroxyzine	12.5 - 12.5 - 25 mg	25-100 mg (100 mg)	no	frequency unknown	+++	frequency unknown	no	no

Frequencies of adverse effects as reported in clinical studies, frequencies are not placebo-corrected. Rare (> 1/10,000 to < 1/1000): rare Uncommon (> 1/1000 to < 1/1000): +

Common (> 1/100 to < 1/10): ++\*
Very common (> 1/10): +++

The information on the starting dose and side effects is mostly issued from the European product label of the individual drug



# **Drug-drug Interactions between Anxiolytics and ARVs**

Anx	ciolytics	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
	alprazolam	1	1	1	1	1	$\leftrightarrow$	<b>↓</b>	<b>↓</b>	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	chlor-diazepoxide	1	1	1	1	1	$\leftrightarrow$	<b>1</b>	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
BZD	clonazepam	1	1	1	1	1	$\leftrightarrow$	<b>↓</b>	<b>1</b>	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	lorazepam	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	oxazepam	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
SSRI	escitalopram	†a	↑a	1	1	†a	$\leftrightarrow$	<b>\</b>	<b>1</b>	1	↔b	↔b	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔b	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
SS	paroxetine	↑↓?	↑↓?	↑↓?	↓39%	↑↓?	$\leftrightarrow$	$\leftrightarrow$	↑3%	$\leftrightarrow$	↑↓?	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$							
SNRI	duloxetine	1	↑↓	1	↑↓	$\uparrow\downarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
S	venlafaxine	↑ b	↑ b	1	1	↑ <b>b</b>	$\leftrightarrow$	1	1	1	↔b	↔b	D	$\leftrightarrow$	$\leftrightarrow$	↔b	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Others	buspirone	1	1	1	1	1	$\leftrightarrow$	1	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
Oth	hydroxyzine	↑a,b	↑a,b	↑a,b	↑a,b	↑a,b	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										



No clinically significant interaction expected These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/ monitoring or dosage adjustment is unlikely to be required

### Legend

Potential elevated exposure of the anxiolytic therapy Potential decreased exposure of the anxiolytic therapy

No significant effect

D Potential decreased exposure of ARV drug Ε Potential elevated exposure of ARV drug

ATV co-formulated with COBI (300/150 mg qd) ATV/c DRV co-formulated with COBI (800/150 mg qd) DRV/c CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

BZD benzodiazepines

SSRI selective serotonin reuptake inhibitors

SNRI serotonin and norepinephrine reuptake inhibitors

# Interactions with ABC, FTC, 3TC, ZDV

ABC, FTC, 3TC, ZDV: no clinically relevant interactions expected.

#### Interactions with ibalizumab

#### Comments

- ECG monitoring is recommended.
- Caution as both drugs can induce QT interval prolongation.

### **Further Information**

For additional drug-drug interactions and for more detailed pharmacokinetic interaction data and dosage adjustments, please refer to: http://www.hiv-druginteractions.org (University of Liverpool)



# Algorithm for Diagnosis and Management of Cognitive Impairment without Obvious Confounding Conditions

#### **Abbreviations** CSF cerebrospinal fluid GDR genotypic drug resistance test HAD HIV-associated dementia LOQ Limit of quantification MND mild neurocognitive disorder MRI brain magnetic resonance imaging NP neuropsychological opportunistic infections Ols **RCT** randomised controlled trial The person or their Initial assessments(ii) relatives complaining of, or care giver Problems suspected noting cognitive problems - without Evaluation for obvious confounding depression and conditions( anxiety and possible treatment(iii Problems persisting but depression or anxiety excluded or optimally managed NP examination(iv) Cognitive impairment(v) Neurological examination Brain MRI CSF examination(vi) Additional causes of cognitive impairment other than HIV excluded and/or managed Diagnosis: HIV-associated cognitive impairment Off ART On ART CSF viral escape Other situations(viii) Start ART(ix), (refer to Optimise ART(ix) Systemic Likely general guidelines) by CSF failure with ART and plasma GDR plasma toxicity testing HIV-RNA>CSF HIV-RNA V Repeat CSF exam-Refer to Switch ination and other general from evaluations as by guidelines EFV or consider clinical judgement other ART toxicities

## Obvious confounding conditions:

- 1. Severe psychiatric conditions
- Use of anticholinergic drugs with high burden score for cognitive impairment (e.g. amitriptyline, chlorpromazine)
- 3. Abuse of psychotropic drugs
- 4. Alcohol abuse
- Sequelae from previous CNS-OIs, pre-treatment cognitive disease or other neurological diseases
- 6. Current CNS-Ols or other neurological diseases

#### The following questions may be used to guide initial assessments (other screening assessments are acceptable)

- Do you experience frequent memory loss (e.g. do you forget the occurrence of special events even the more recent ones, appointments, etc.)?
- Do you feel that you are slower when reasoning, planning activities, or solving problems?
- Do you have major difficulties paying attention (e.g. to a conversation, book or film)?

Answering "yes" to one or more of these questions may suggest the presence of cognitive disorders, although not necessarily linked to HIV.

- iii See Depression: Screening and Diagnosis and Anxiety Disorders: Screening and Diagnosis
- iv NP examination should include tests exploring the following cognitive domains: fluency, executive functions, speed of information processing, attention/working memory, verbal and visual learning, verbal and visual memory, motor skills plus assessment of daily functioning
- Cognitive impairment is defined by impairment in cognitive function on the above neuro-psychological test where performance is compared to age and education-matched appropriate controls and is considered clinically significant
- vi Neurological examination, brain MRI and CSF examination are required to exclude other pathologies (consultation with neurologist specialist may be required) and to further characterise possible HIVassociated cognitve impairment by including assessment of CSF HIV-RNA level and, where appropriate, evidence for genotypic drug resistance (GDR) in a paired CSF and plasma sample

# vii CSF escape definition:

Either CSF HIV-RNA above LOQ and plasma HIV-RNA below LOQ; or HIV-RNA above LOQ in both CSF and plasma, with CSF HIV-RNA greater than plasma HIV-RNA.

In CSF escape:

- Avoid dual ART therapies
- Include dual nucleoside backbones in ART regimens where possible
- Avoid ATV (boosted or unboosted) due to association with CSF escape in retrospective cohorts
- Avoid RAL 1200 mg qd due to lack of evidence in CSF escape
- Consider DTG 50 mg bid in cases with documented or suspected INSTI resistance
- viii Including situations that do not fulfill the CSF escape definition, but can benefit from ART optimisation
- Avoid EFV because of its possible effects on cognitive function and potentially confounding CNS effects due to neuropsychiatric effects

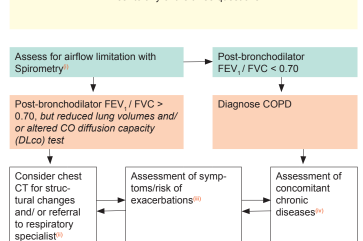
# **Chronic Lung Disease**

# Screen for chronic lung disease:

Do you have ANY of the following on a regular basis:

- a) shortness of breath when walking up a slight hill or hurrying on flat ground;
- b) cough and/or sputum;
- c) recurrent wheezing

"Yes" to any of the three questions

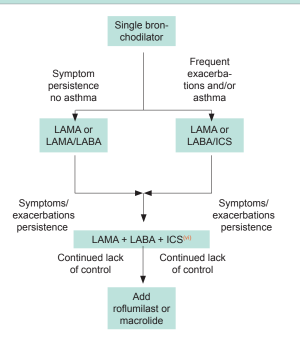


"Yes" to "shortness of breath on light exercise or at rest"

"No" Repeat questions annually

Make comprehensive assessment particularly for the risk of concomitant CVD including pulmonary hypertension

## Treatment of COPD®



**LABA:** long-acting β2-agonist

LAMA: long-acting muscarinic antagonist

ICS: inhaled corticosteroid

Reassess and adjust regularly according to the response to treatment in terms of dyspnea and/or acute exacerbations

# There are 3 lifesaving interventions in COPD:

- 1. Smoking cessation
- 2. Chronic oxygen when stable (non-exacerbated) resting  ${\rm SpO_2} \le 88\%$  (or  ${\rm PaO_2} \le 55~{\rm mmHg}$ )
- 3. Non-invasive ventilation (NIV) in individuals with persistent hypercapnic respiratory failure after an acute exacerbation

- i Risk assessment for spirometry should be undertaken in the setting of COVID-19
- ii Based on expert opinion, also consider interstitial lung disease, CT scan may help to identify people with interstitial lung disease and lung cancer
- iii Assessment of either dyspnoea using mMRC, see https://www.verywell-health.com/guidelines-for-the-mmrc-dyspnea-scale-914740 or symptoms using CAT™, see http://www.catestonline.org/ and history of exacerbations (including prior hospitalisations)
- iv COPD itself has significant extra-pulmonary (systemic) effects including weight loss, nutritional abnormalities and skeletal muscle dysfunction
- v Each pharmacological treatment should be individualised and guided by the severity of symptoms, risk of exacerbations, adverse effects, co-morbidities, drug availability and cost, and the individual's response, preference and ability to use various drug delivery devices. Inhaler technique needs to be assessed regularly. Long-term use of high dose ICS and/or use of oral glucocorticoids has no evidence of benefits in COPD and increase the risk of pneumonia. The addition of medium dose ICS to LABA or LAMA or LABA/LAMA is recommended in individuals with history of frequent exacerbations and/or asthma and/or eosinophilia (> 3%), or anyway in individuals not adequately controlled by LAMA/LABA combination. ICS should be avoided in subjects with eosinopenia (< 1%) Antibiotics should be used to treat acute exacerbation or in case of high CRP and purulent sputum (PCT is a more questionable biomarker). Azithromycin may also be considered in non-smokers, not well controlled with maximal inhaled drug dosage.</p>
- vi LAMA/LABA/ICS are now available in a fixed dose combination. This drug combination improves clinical control of COPD and increases life expectancy

With the exception of low dose beclometasone, do not use inhaled glucocorticoids with boosted ART regimens, see Drug-drug Interactions between Corticosteroids and ARVs.

Influenza, SARS-CoV-2 and pneumococcal vaccination decrease rates of lower respiratory tract infections, see Vaccination. Pertussis vaccination is also suggested in people with COPD

# **Drug-drug Interactions between Bronchodilators (for COPD) and ARVs**

Bro	nchodilators	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
	aclidinium bromide	$\leftrightarrow$																			
LAMA	glycopyrronium bromide	$\leftrightarrow$																			
LA	tiotropium bromide	$\leftrightarrow$																			
	umeclidinium bromide	1	1	1	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
SAMA	ipratropium	$\leftrightarrow$																			
	formoterol	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
4	indacaterol	↑ <mark>b</mark>	↑b	↑b	↑b	↑b	$\leftrightarrow$	1	↓	1	$\leftrightarrow$	↑b	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
LABA	olodaterol	1	1	1	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
_	salmeterol	1	1	1	1	1	$\leftrightarrow$	1	↓	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	vilanterol	1	1	1	1	1	$\leftrightarrow$	1	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
SABA	salbutamol (albuterol)	$\leftrightarrow$																			
Ŝ	terbutaline	$\leftrightarrow$																			
MX	aminophylline	$\leftrightarrow$	↓	$\leftrightarrow$	1	1	$\leftrightarrow$														
Σ	theophylline	$\leftrightarrow$	↓	$\leftrightarrow$	1	1	$\leftrightarrow$														
PDE4	roflumilast	1	1	1	1	1	$\leftrightarrow$	1	↓ ·	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	beclometasone	↑ <b>c</b>	↑ <b>c</b>	↑?c	↓11%d	↑c	$\leftrightarrow$	↑ <b>c</b>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
	budesonide	↑e	↑e	↑e	↑e	↑e	$\leftrightarrow$	1	1	1	$\leftrightarrow$	↑e	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
SOI	ciclesonide	↑ <b>f</b>	↑ <b>f</b>	↑f	↑f	↑f	$\leftrightarrow$	↑ <b>f</b>	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										
	fluticasone	↑e	↑e	↑e	†e	↑e	$\leftrightarrow$	1	Ţ	1	$\leftrightarrow$	↑e	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
	mometasone	↑e	↑e	↑e	↑e	↑e	$\leftrightarrow$	1	↓	1	$\leftrightarrow$	↑e	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						

Colour legend

No clinically significant interaction expected These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/ monitoring or dosage adjustment is unlikely to be required

Legend

Potential elevated exposure of the bronchodilator Potential decreased exposure of the bronchodilator

No significant effect

D Potential decreased exposure of ARV drug Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd) DRV/c DRV co-formulated with COBI (800/150 mg qd) CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

ICS inhaled corticosteroids LABA long-acting β2 agonists

LAMA long-acting muscarinic antagonists MX methylxanthines PD4 phosphodiesterase 4 inhibitors SABA short-acting β2 agonists SAMA short-acting muscarinic antagonists Interactions with ABC, FTC, 3TC, ZDV

ABC, FTC, 3TC, ZDV: no clinically relevant interactions expected.

# Interactions with ibalizumab

None

### Comments

- Caution as both drugs can induce QT interval prolongation.
- Exposure can be increased up to 2-fold however this increase does not raise any concerns based on indacaterol's safety data.
- Increase in concentration of active metabolite observed with RTV 100 mg bid alone but without significant effect on adrenal function. Caution is still warranted, use the lowest possible corticosteroid dose and monitor for corticosteroid side effects.
- DRV/r decreased the exposure of active metabolite (beclometasone-17-monopropionate), no significant effect on adrenal function was seen.
- Risk of having elevated corticosteroid levels, Cushing's syndrome and adrenal suppression. This risk is present for oral and injected corticosteroid but also for topical, inhaled or eye drops administration.
- No dose adjustment required but monitor closely, especially for signs of Cushing's syndrome when using a high dose or prolonged administration.

# **Further Information**

For additional drug-drug interactions and for more detailed pharmacokinetic interaction data and dosage adjustments, please refer to: http://www.hiv-druginteractions.org (University of Liverpool)

Fixed dose combinations are available for LAMA + LABA + ICS, e.g., mometasone + indacaterol + glycopyrronium fluticasone + umeclidinium + vilanterol

formoterol + glycopyrronium + beclometasone budesonide + formoterol + glycopyrronium



# **Drug-drug Interactions between Pulmonary Antihypertensives and ARVs**

	monary ihypertensives	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
	ambrisentan	1	1	1	1	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$								
ERA	bosentan	†a	†a	†a	†a	↑a	D	ļ	1	↑p	D	1	D	D	$\leftrightarrow$	D	D	†a	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	macitentan	1	1	1	1	1	$\leftrightarrow$	↓	1	1	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
E5	sildenafil	1	1	1	1	1	$\leftrightarrow$	ļ	1	↓	↓3%	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
PDE5	tadalafil	1	1	1	1	1	$\leftrightarrow$	ļ	<b>1</b>	↓	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
SGC	riociguat	1	1	1	1	1	$\leftrightarrow$	↓	1	<b>↓</b>	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
	epoprostenol	$\leftrightarrow$																			
<b>ĕ</b>	iloprost	$\leftrightarrow$																			
	treprostinil	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	1	$\leftrightarrow$												
<u>P</u>	selexipag	↔C	↔C	↔C	↔C	↑120%d	$\leftrightarrow$	↔C	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$										

### Colour legend

No clinically significant interaction expected
These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

#### Legend

↑ Potential elevated exposure of the pulmonary antihypertensive Potential decreased exposure of the pulmonary antihypertensive

→ No significant effect

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd) DRV/c DRV co-formulated with COBI (800/150 mg qd) CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

ERA endothelin receptor antagonists

Ipr IP receptor agonists
PA prostacyclin analogues

PDE5 phosphodiesterase type 5 inhibitors sGC soluble guanylate cyclase stimulators

### Interactions with ABC, FTC, 3TC, ZDV

ABC, FTC, 3TC, ZDV: No clinically relevant interactions expected.

### Interactions with ibalizumab

None

#### Comments

- Co-administration is not recommended in the European labels, but the US labels suggest the following dose modifications: When starting bosentan in persons already on PI/b or EVG/c use a bosentan dose of 62.5 mg qd or every other day. Discontinue bosentan at least 36 h prior to starting PI/b or EVG/c and restart after at least 10 days at 62.5 mg qd or every other day.
- b Potential additive liver toxicity.
- Exposure of parent drug increased but exposure of active metabolite unchanged.
- d This change is unlikely to be clinically relevant.

# **Further Information**

For additional drug-drug interactions and for more detailed pharmacokinetic interaction data and dosage adjustments, please refer to: http://www.hiv-druginteractions.org (University of Liverpool)



### **Managing Older Persons with HIV**

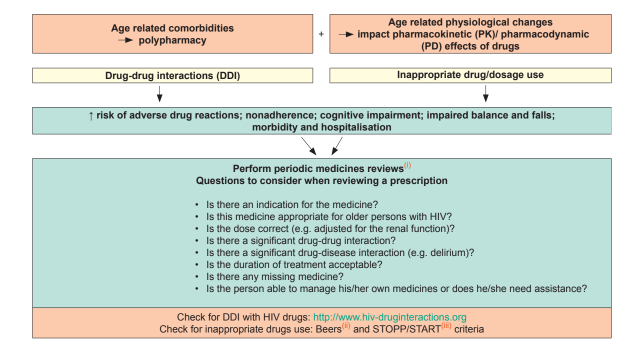
Physical function, frailty and geriatric syndromes have shown to better predict survival and quality of life among older people in the general population than co-morbidity alone. Managing older people with HIV has to move from the management of each condition separately to a multidimensional assessment focused on preserving physical function, aimed at promoting health aging and quality of life. This section will focus on important geriatric issues: polypharmacy, frailty and falls.

### **Polypharmacy**

Polypharmacy is defined as the concurrent use of > 5 drugs, a cut-off that has been associated with an increased risk of adverse health outcomes. In HIV medicine, the term polypharmacy most often refers to non-HIV medications given in addition to ARVs.

The complexity of medication burden should be considered owing to its clinical consequences: the higher risk of drug-drug interactions and adverse events, the risk of non-adherence to HIV and non-HIV medications, and the risk of hospitalisation, falls, other geriatric syndromes and death. Polypharmacy is often unavoidable when treating a patient with multiple co-morbid conditions making the use of polypharmacy appropriate in this context whereas "unnecessary or inappropriate polypharmacy" is deleterious and should be avoided. Interventions to prevent unnecessary/inappropriate polypharmacy include medication reconciliation and medication review. The concept of 'deprescribing' or the planned and supervised process of dose reduction or stopping of medication that may cause harm, or no longer provide benefit has gained increasing attention as a means to reduce unnecessary/inappropriate polypharmacy in older persons with HIV. A freely-accessible resource to help deprescribe can be found at medstopper.com.

### **Prescribing in Older Persons with HIV**



i-iii The Beers and STOPP criteria are tools established by experts in geriatric pharmacotherapy to detect and reduce the burden of inappropriate prescribing in older persons (note: these tools were established for persons > 65 years old given that PK and PD effects may be more apparent after this age cut-off). Inappropriate medicines include, for instance, those which in older persons with certain diseases can lead to drug-disease interactions, are associated with a higher risk of adverse drug reactions in olderpersons, medicines that predictably increase the risk of falls in the older persons or those to be avoided in case of organ dysfunction. The START criteria consist of evidence-based indicators of potential prescribing omission in older persons with specific medical conditions



# **Selected Top 10 Drug Classes To Avoid in Older Persons with HIV**

Drug class	Problems/alternatives
First generation antihistamines e.g., clemastine, diphenhydramine, doxylamine, hydroxyzine	Strong anticholinergic properties, risk of impaired cognition, delirium, falls, peripheral anticholinergic adverse reactions (dry mouth, constipation, blurred vision, urinary retention).  Alternatives: cetirizine, desloratadine, loratadine
<b>Tricyclic antidepressants</b> e.g., amitryptiline, clomipramine, doxepin, imipramine, trimipramine	Strong anticholinergic properties, risk of impaired cognition, delirium, falls, peripheral anticholinergic adverse reactions (dry mouth, constipation, blurred vision, urinary retention).  Alternatives: citalopram, escitalopram, mirtazapine, venlafaxine
Benzodiazepines Long and short acting benzodiazepines e.g., clonazepam, diazepam, midazolam Non-benzodiazepines hypnotics e.g., zolpidem, zopiclone	Elderly are more sensitive to their effect, risk of falls, fractures, delirium, cognitive impairment, drug dependency. Use with caution, at the lowest dose and for a short duration.  Alternatives: non-pharmacological treatment of sleep disturbance/sleep hygiene.
Atypical antipsychotics e.g., clozapine, olanzapine, quetiapine	Anticholinergic adverse reactions, increased risk of stroke and mortality (all antipsychotics).  Alternatives: aripiprazole, ziprasidone
Urological spasmolytic agents e.g., oxybutynin, solifenacin, tolterodine	Strong anticholinergic properties, risk of impaired cognition, delirium, falls, peripheral anticholinergic adverse reactions (dry mouth, constipation, blurred vision, urinary retention).  Alternatives: non-pharmacological treatment (pelvic floor exercises).
Stimulant laxatives e.g., senna, bisacodyl	Long-term use may cause bowel dysfunction. Alternatives: fibres, hydration, osmotic laxatives
NSAIDs e.g., diclofenac, indomethacin, ketorolac, naproxen	Avoid regular, long-term use of NSAIDs due to risk of gastrointestinal bleeding, renal failure, worsening of heart failure.  Alternatives: paracetamol, weak opioids
<b>Digoxin</b> Dosage > 0.125 mg/day	Avoid doses higher than 0.125 mg/day due to risk of toxicity.  Alternatives for atrial fibrillation: beta-blockers
Long acting sulfonylureas e.g., glyburide, chlorpropamide	Can cause severe prolonged hypoglycemia. Alternatives: metformin or other antidiabetic classes
Cold medications Most of these products contain antihistamines (e.g., diphenhydramine) and decongestants (e.g., phenylephrine, pseudoephedrine)	First generation antihistamines can cause central and peripheral anticholinergic adverse reactions as described above. Oral decongestants can increase blood pressure

**Legend**NSAID nonsteroidal anti-inflammatory drug



### **Frailty**

Frailty is defined as a clinical syndrome associated with decreased reserve, high vulnerability to stressors and associated with risk of negative health-related outcomes including mortality. Frailty should be regarded as a distinct entity to the disease or condition that may be contributing to it. This geriatric syndrome is more prevalent than expected in persons with HIV compared to HIV-negative matched controls and may occur at an earlier age. Early identification and management of frailty is a priority since it is potentially reversible. Older persons with HIV aged 50 years and over should be offered screening for frailty using a validated rapid frailty instrument. An algorithm to identify those persons who may benefit from a frailty assessment is detailed below.

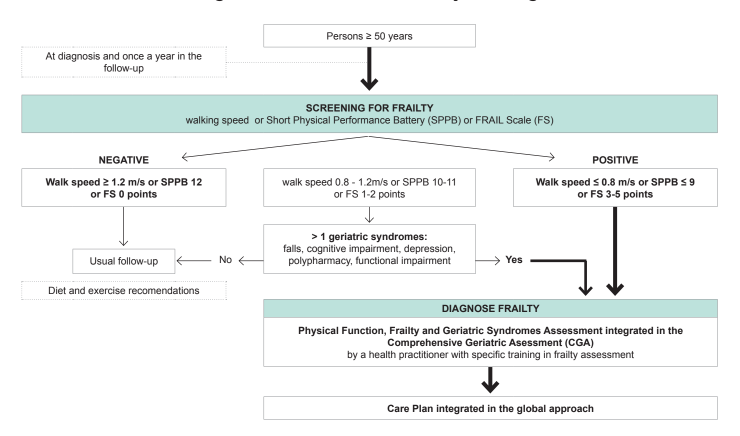
#### **Screening for Frailty**

Screening for frailty in persons with HIV above 50 years of age should be considered. The age cut-off was chosen as the incidence of frailty in persons with HIV has been shown to increase above this age. Evidence of benefit is still unknown. It is advocated by some experts.

Screening has to be performed using validated tools for this purpose and can be provided by any trained health staff (nurses, general practitioners, etc.). In the absence of a gold standard, the instrument to screen frailty we suggest is the FRAIL Scale (FS) because it is easy, cheap and quick to develop but other validated tools, such as walking speed measurement or Short Physical Performance Battery (SPPB) can also be used.

FRAIL SCALE								
How much time during the previous 4 weeks did you feel tired?	All the time, most of the time = 1 point							
Do you have any difficulty walking up 10 steps alone without resting and without aids?	Yes = 1 point							
Do you have any difficulty walking several hundred meters alone with/without aids?	Yes = 1 point							
How many illnesses do you have from this list?: hypertension, DM, cancer, chronic lung disease, heart attack, congestive heart failure, angina, asthma, arthritis, stroke and kidney disease.	> 5 = 1 point							
Have you had weight loss of 5% or more?	Yes = 1 point							

### **Algorithm Recommended for Frailty Screening**



Adapted from Brañas F, et al. European Geriatric Medicine. 2019;10(2):259-265



#### **Formal Frailty Assessment and Management**

	How to diagnose frailty	
	Frailty Phenotype	Frailty Index
Clinical definition	Clinical syndrome based on presence of specific signs and symptoms	Based on accumulation of deficits
How to assess	Assessed by five specific features:  1. self-reported weight loss (a)  2. self-reported exhaustion (b)  3. low levels of physical activity as measured by Minnesota Leisure physical activity questionnaire (c)  4. measured 4 m walk speed time (d)  5. measured grip strength (e)	A frailty index is calculated based on the number of health deficits out of > 30 assessed health deficits  Health variables, including signs and symptoms of disease, laboratory measures, and self-reported data  Data routinely collected in medical records can be included if they characterise age-related, acquired health deficits which cover a range of physiologic systems
How to interpret	Categorical variables Total score of 5 items: 0 deficits = fit 1-2 deficits = pre-frail 3 + deficits = frail	Continuous variables Index ranges from 0 to 1: ≤ 0.25 = fit 0.25 - 0.4 = frail > 0.4 = most frail
	How to address frailty	

How to address frailty

Promote Comprehensive Geriatric Assessment (CGA), aimed at personalising interventions according to benefits/priorities for a given person through a multidisciplinary diagnostic and treatment process, that identifies medical, psychosocial, and functional limitations aimed at maximising overall health with ageing and the improvement of quality of life

#### Recommendations

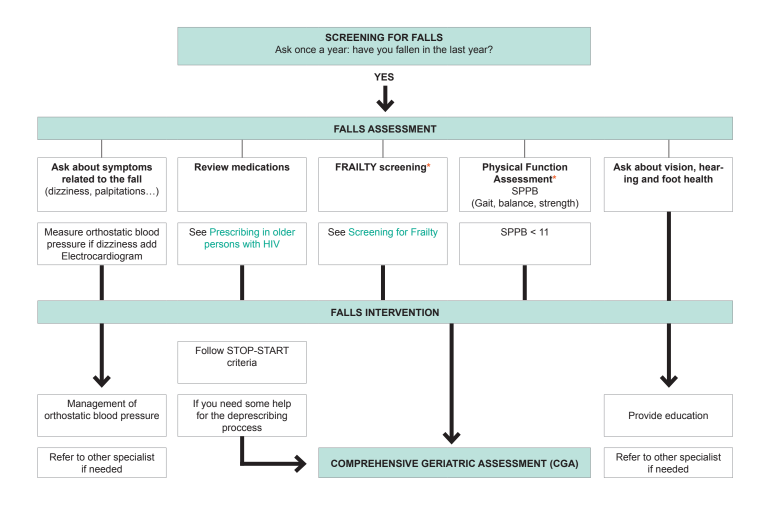
In persons with HIV who are frail:

- 1. Sustain and recover physical function impairment and sarcopenia prescribing physical activity with a resistance training component
- 2. Address polypharmacy by reducing or deprescribing any inappropriate/superfluous medications, see Prescribing in older persons with HIV
- 3. Screen for, and address modifiable causes of fatigue
- 4. For persons exhibiting unintentional weight loss, screen for reversible causes and consider food fortification and protein/caloric supplementation
- 5. Prescribe vitamin D for individuals deficient in vitamin D, see page 72
- (a) Self-reported unintentional weight loss was considered present if exceeding 4.5 kg or ≥5% of body weight in the last year
- (b) **Exhaustion** is present if the participant answers "occasionally" or "most of the time" to both of the following statements (questions from the Center for Epidemiologic Studies Depression Scale): During the last week, how often have you felt that 1. everything you did was an effort, or 2. you could not 'get going'
- (c) Low physical activity as considered present if the participant's physical activity is lower than 383 kcal/week in men and 270 kcal/week in women which is equivalent to < 2.5 hours/week in men and < 2 hours/week in women using the Minnesota Leisure Time Activity Questionnaire
- (d) Walk speed time is measured by a 4-meter walking test in usual pace (one trial). A deficit is assigned according to the following gender-specific criteria
  - Men: height ≤ 173 cm and speed ≤ 0.6531 m/s; height > 173 cm and speed ≤ 0.762 m/s
  - Women: height ≤ 159 cm and speed ≤ 0.6531 m/s; height > 159 cm and speed ≤ 0.762 m/s
- (e) **Maximum grip strength** can be assessed using a handheld dynamometer the mean value of three consecutive measurements of the dominant hand (adjusted by sex and BMI quartile based on CHS population):
  - Men: BMI ≤ 24 kg and strength < 29 kg; BMI 24.1–26 and strength < 30 kg; BMI 26.1–28 and strength < 30 kg; BMI > 28 and strength < 32 kg</p>
  - Women: BMI ≤ 23 and strength < 17 kg; BMI 23.1–26 and strength < 17.3 kg; BMI 26.1–29 and strength < 18 kg; BMI > 29 and strength < 21 kg</p>



#### **Falls**

A fall is defined as an event which results in a person coming to rest inadvertently on the ground or floor or other lower level. Falls are a common geriatric syndrome in persons with HIV, as the prevalence is estimated to be between 25% and 30%, affect independent movement and mobility in older people and therefore their quality of life



\* If FRAIL Scale is > 1 point, as the person has 1 geriatric syndrome (falls), then is not necessary to perform the SPPB in the HIV clinic as the CGA is already recommended

# **Solid Organ Transplantation (SOT)**

#### **General features**

- · HIV infection is not a contraindication for transplantation consideration.
- Experts in HIV medicine should preferably be members of the multidisciplinary team, responsible for the pre-transplant evaluation, and take primary responsibility for the management of the HIV infection and the prevention and treatment of OIs

#### Organ criteria for SOT

 Persons with HIV should be considered for organ transplantation using the same indications as used in HIV-negative persons. Persons with HIV with HCC can be evaluated for liver transplantation if they fulfill the Milan criteria<sup>(i)</sup>

#### Organ donation

- Persons with HIV can receive organs from living (renal) and deceased (all types of SOT) HIV-negative donors
- In some European countries the use of organs from HIV-positive donors is allowed but the efficacy and safety of this approach is currently being evaluated in the context of research studies

#### **HIV-infection criteria for SOT**

According to most international guidelines, persons with HIV should fulfill the following criteria to be considered for SOT

- 1. Clinical criteria. No active Ols or HIV-related cancers. Individuals with PML, chronic crypto/microsporidiosis, multi-drug resistant fungal or mycobacterial infections, NHL and visceral KS to be excluded. For non-HIV-related cancers same criteria apply as in the general HIV-negative population
- 2. Immunological criteria. CD4 > 200 cells/µL for all SOT except for liver transplantation where CD4 > 100 cells/µL. Persons with previous opportunistic infections should have a CD4 > 200 cells/µL
- Virological criteria. Full control of HIV replication prior to and after transplantation should be confirmed/predicted in all cases
- **4. Drug abuse**. Abstinence period: alcohol = 6 months; heroin/cocaine = 2 years. Former IVDUs can be in methadone programme

#### Preparing for transplantation

#### Antiretroviral therapy

- Choice of ART components should avoid drugs known to cause organ dysfunction or drugs with a high potential for drug-drug interactions if at all possible, see Drug-drug Interactions between Immunosuppressants (for SOT) and ARVs
- Using a pharmacological booster (RTV or COBI) and some of the NNRTIs are best avoided, see Drug-drug Interactions between Immunosuppressants (for SOT) and ARVs
- For individuals nearing indication for transplantation, ART should be modified to ensure this if at all possible
- · Unboosted INSTIs plus 2 NRTIs are the preferred regimens
- If the individual has not yet started ART and transplantation is considered, ART should be commenced as soon as possible and preferably before the transplantation is started

#### Viral hepatitis co-infection

In liver transplant candidates, every effort should be made to treat the underlying viral hepatitis independently of MELD score, see pages 115-120. Use of DAAs in persons with HCV co-infection may improve their liver function, and possibly lead to them being removed from the transplant waiting list

#### Prevention of infections

While screening and treatment for latent TB is recommended in all persons with HIV, see page 137, it is particularly important in persons pre-and post-transplantation due to the additional use of immunosuppressants.
 Immunisation regimens and pre-transplant diagnostic protocols are the same as in HIV-negative SOT recipients

#### Follow-up after transplantation

#### Antiretroviral therapy

- · Same recommendations in individuals under preparation for transplantation
- Additionally, ARVs may exacerbate immunosuppressive agents' adverse drug effects (kidney impairment, bone marrow suppression, drug-induced liver injury, etc.). Therefore, careful consideration of which drugs to use is essential see Adverse Effects of ARVs & Drug Classes
- TAF is preferred to TDF, when available, to reduce additive nephrotoxicity to immunosuppressant agents

#### Primary and secondary disease-specific prevention

- Transplant recipients living with HIV should receive the same surveillance, immunisation prophylaxis and pre-emptive regimens as HIV-negative SOT recipients
- Screening and treatment for latent TB is a priority, see page 137

#### Viral hepatitis co-infection

- The efficacy and safety of DAAs in liver transplant recipients living with HIV with HCV recurrence is the same as in HIV-negative recipients
- Anti HBV treatment should follow the same schedules of HIV-negative persons

#### Screening for co-morbidities and frailty

Persons with HIV undergoing SOT have higher risk for some comorbidities including, CVD, DM, bone disease (osteoporosis and aseptic necrosis of the femur) and frailty, see Prevention of Cardiovascular Disease (CVD), Type 2 Diabetes Mellitus: Diagnosis, Type 2 Diabetes Mellitus: Management, Bone Disease: Screening and Diagnosis and Managing Frailty in Older People Living with HIV

#### Immunosuppressive regimens

- Same as in HIV-negative transplant recipients. The risk of acute rejection is however double of that of HIV-negative SOT recipients and, therefore, requires close monitoring
- Special attention to interaction with ART, see Drug-drug Interactions between Immunosuppressants (for SOT) and ARVs
- Using a pharmacological booster (RTV or COBI) and some of the NNRTIs should be used with caution and requiring close monitoring of immunosuppressive drugs, see Drug-drug Interactions between Immunosuppressants (for SOT) and ARVs
- i Milan criteria: solitary tumor smaller than 5 cm or 2 3 tumors of < 3 cm in the absence of macrovascular tumor invasion and extrahepatic metastases



# **Drug-drug Interactions between Immunosuppressants (for SOT) and ARVs**

	nuno- pressants	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	BIC	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
S	prednisone	1	1	1	1	1	$\leftrightarrow$	↓20%	<b>1</b>	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	E11%	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
5	azathioprine	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$								
AM	mycophenolate	$\leftrightarrow$	ţа	$\leftrightarrow$	↓a	↓a	$\leftrightarrow$	↓a	$\leftrightarrow$	↓ <mark>a</mark> D13%	$\leftrightarrow$	↑ Eb									
CN	cyclosporine	†a	†a	†a	†a	†a	Е	↓a	↓a	↓a	Е	$\leftrightarrow$	Е	Е	$\leftrightarrow$	E	$\leftrightarrow$	†a	$\leftrightarrow$	E	Eb
5	tacrolimus*	†a,c	↑a,c	†a	†a	↑a,c	↓a	↓a	↓a	↓a	↔C	↔C	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↔C	$\leftrightarrow$	†a	$\leftrightarrow$	$\leftrightarrow$	↔b
mTOR	everolimus	1	1	1	1	1	$\leftrightarrow$	↓a	↓a	↓a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$						
m	sirolimus	1	1	1	1	1	ţа	↓a	↓a	↓a	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	↔b						
	anti-thymocyte globulin	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$								
Other	basiliximab	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$								
	belatacept	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$								

#### Colour legend

No clinically significant interaction expected These drugs should not be co-administered Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/ monitoring or dosage adjustment is unlikely to be required

#### Legend

Potential elevated exposure of the immunosuppressant Potential decreased exposure of the immunosuppressant

No significant effect

D Potential decreased exposure of ARV drug Ε Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd) DRV/c DRV co-formulated with COBI (800/150 mg qd) CAB/RPV CAB and RPV im long acting injections (PK and/or QT interactions shown are with RPV)

Numbers refer to increased or decreased AUC as observed in drug-drug interaction studies

AM antimetabolite CNI calcineurin inhibitors CS corticosteroids mTOR mTOR inhibitors

#### Interactions with ABC, FTC, 3TC, ZDV

ABC: potential decrease in mycophenolate exposure.

ZDV: potential risk of additive haematoxicity with azathioprine.

potential alteration in mycophenolate exposure, monitor plasma concentrations.

#### Interactions with ibalizumab

#### Comments

- TDM of immunosuppressant is recommended.
- Monitor renal function.
- Both drugs can potentially prolong the QT interval, ECG monitoring recommended

#### **Further Information**

For additional drug-drug interactions and for more detailed pharmacokinetic interaction data and dosage adjustments, please refer to: http://www.hiv-druginteractions.org (University of Liverpool)



<sup>\*</sup> available as prolonged release formulation

# Part V Clinical Management and Treatment of Viral Hepatitis Co-infections

Every person with HCV/HIV co-infection should receive DAA therapy to eradicate HCV, regardless of liver fibrosis stage. Cure of HCV infection substantially reduces the risk for hepatic and extrahepatic complications and eliminates onward HCV transmission. DAAs achieve similar cure rates and tolerability in HCV/HIV co-infected compared to HCV mono-infected persons. Therefore, treatment indication and regimens are the same as in HCV mono-infected persons. All persons with HBV/HIV co-infection should receive ART including TDF or TAF, unless history of tenofovir intolerance. All HBsAg-positive persons should be screened for Hepatitis Delta (HDV)

# General Recommendations for Persons with Viral Hepatitis/HIV Co-infection

#### Screening at baseline

- 1. HCV should be screened for HCV at time of HIV diagnosis and annually thereafter. Screening should use an anti-HCV antibody test. A positive result should be followed by HCV-RNA. and genotype determination which is not mandatory if pangenotypic drugs are to be used. Alternatively, HCV core-antigen testing can be performed to establish chronic HCV infection. Persons engaging in activities associated with increased risk of HCV transmission. should be tested for HCV infection every 3 to 6 months. Persons suspected of recently acquired primary HCV infection with a negative anti-HCV antibody test should be tested for HCV-RNA. HCV-RNA or HCV core-antigen testing is also recommended in persons with ongoing risk behavior for HCV re-infection after successful treatment or spontaneous clearance at 3 to 6-monthly intervals.
- HAV and HBV should be screened for HAV and HBV. Persons who are anti-HBc positive and HBsAg negative, in particular those with elevated liver transaminases, should be screened for HBV-DNA in addition to HBsAg to rule out occult HBV infection
- HDV antibodies should be screened for in all HBsAg positive persons.
- 4. Persons with viral hepatitis co-infection should be assessed for concurrent causes of liver disease such as alcohol consumption, cardiac disease, renal impairment, autoimmunity, genetic or metabolic liver diseases (e.g. genetic haemochromatosis, diabetes mellitus or obesity) and drug-induced hepatotoxicity
- 5. Status of liver damage should be assessed in all persons with viral hepatitis co-infection with a complete blood count, ALT, AST, GGT, ALP, hepatic synthetic function (e.g. coagulation, albumin, cholinesterase) and staging of fibrosis (e.g. FibroScan, liver biopsy, serum fibrosis markers<sup>™</sup>, see Table on cut-off values of non-invasive tests for the detection of advanced fibrosis and cirrhosis)

#### Screening for complications

- 6. HCC screening is indicated in all cirrhotic HBV or HCV co-infected persons (even if HCV infection has been cured and HBV replication is medically suppressed) in a setting where treatment for HCC is available. Although the cost-effectiveness of HCC screening in persons with F3 fibrosis is uncertain, surveillance may be considered based on an individual risk assessment, see page 59. In HBV-positive non-cirrhotics, HCC screening should follow current HCC EASL guidelines (https://easl.eu/publication/easl-clinical-practice-guidelines-management-of-hepatocellular-carcinoma/). Risk factors for HCC in this population include family history of HCC, ethnicity (Asians, Africans), HDV co-infection and age > 45 years. EASL guidelines propose using the PAGE-B score in Caucasians to assess the HCC risk, however this score has not been validated in people with HIV, see pages 9, 59 and 81
- Screening for oesophageal varices upon diagnosis of cirrhosis in co-infected persons is also indicated (every 2-3 years thereafter according to presence of ongoing liver disease if negative for oesophageal varices at initial screening), see page 80

#### **End Stage Liver Disease (ESLD)**

- Persons with HIV and liver cirrhosis require the same measures for the treatment of oesophageal varices, hepatorenal syndrome, hepatic encephalopathy or ascites as HIV-negative persons, see page 80-81 and Diagnosis and Management of Hepatorenal Syndrome / Acute Kidney Injury (HRS-AKI)
- Persons with viral hepatitis/HIV co-infection suffering from ESLD warrant particular attention in the management of liver insufficiency, see Dose Adjustment of ARVs for Impaired Hepatic Function. ART in cirrhotic persons improves overall survival
- Persons with HCC or a MELD-score > 12<sup>(w)</sup>, CD4 count > 100 cells/µL and options for efficacious and durable ART should be evaluated for liver transplantation (OLTX), see Solid Organ Transplantation (SOT)
- Renal complications are frequent, see page 74 and Diagnosis and Management of Hepatorenal Syndrome / Acute Kidney Injury (HRS-AKI)

#### Vaccination, see page 90

- 12. Persons lacking anti-HAV IgG antibodies or anti-HBs antibodies should be offered vaccination for the respective virus to prevent infection regardless of their CD4 count. The response to the HBV vaccine is influenced by the CD4 count and level of HIV-VL. In persons with low CD4 count (< 200 cells/µL) and ongoing HIV replication, ART should be initiated first, prior to respective vaccination. Because of the lack of data on the impact of immunisation in isolated anti-HBc IgG positive persons (HBsAg negative, anti-HBc positive and anti-HBs negative profile), vaccination is not recommended in this population. However, if anti-HBc results are not available, HBV vaccination is recommended in all HBs-Ag negative persons</p>
- 13. In persons vaccinated for HBV with insufficient response (anti-HBs < 10 IU/L), re-vaccination should be considered. Double-dose (40 µg) at 3-4 time points (months 0, 1, 2 and 6) may help to improve response rates to the HBV vaccine. Persons who fail to seroconvert after HBV vaccination and remain at risk for HBV should have annual serological tests for evidence of HBV infection. TDF based cART has been associated with prevention of HBV infection in these persons and ART including TDF or TAF is recommended</p>

#### Prevention/Support

- Psychiatric, psychological, social and medical support should be made available to persons with alcohol intake to stop drinking
- 15. Substitution therapy (opioid replacement therapy) in persons with active drug use as a step towards cessation of active drug use should be encouraged. Help provided (e.g. through needle and syringe exchange programs) reduces the risk of re-infection including parenteral viral transmission (harm reduction strategy), see Opioid Addiction
- 16. Since HBV and HIV, and occasionally HCV, are transmitted sexually, adequate counselling including the use of condoms is advisable. Information on the risk of HCV transmission due to mucosal traumatic sexual practices associated with a high likelihood of blood contact or ongoing IDU, "chemsex" (sex under the influence of recreational drugs taken predominantly intravenously immediately before and/or during sexual contacts)<sup>(N)</sup>, should be provided and risk reduction should be discussed
- 17. İn women of childbearing age, HCV treatment should be initiated prior to conception because of limited safety data in pregnancy, and to reduce the risk of MTCT of HCV. HBV therapy should be continued throughout pregnancy.
- Screening intervals to detect recently acquired HCV infection should be adapted to individual risk assessments and local epidemiology as described in the Recommendations on recently acquired and early chronic hepatitis C in MSM from the European treatment network for HIV, hepatitis and global infectious diseases consensus panel
- ii Anti HCV-Antibodies: turn positive 1-6 months after infection; late seroconversions have been described; may rarely be lost due to immunosuppression
- iii There is no standard conversion formula for converting the amount of HCV-RNA reported in copies/mL to the amount reported in IU/mL. The conversion factor ranges from about one to five HCV-RNA copies per IU/ mL
- iv Risk for percutaneous HCV transmission by sharing equipment for injection drug use; risk for mucosal HCV transmission including fisting, receptive condomless anal intercourse, sharing equipment during nasally administered drug use, sharing sex toys, sharing anal douching equipment, and engaging in sexual intercourse causing rectal trauma with bleeding; the presence of ulcerative sexually transmitted infections (STIs) increases the risk of HCV transmission
- V Serum fibrosis markers include APRI, FIB-4, Hyaluronic acid, Fibrometer, Fibrotest, Forns, Hepascore and other indices. The combination of blood biomarkers, of liver stiffness measurement and blood tests or repeated assessments may improve accuracy EASL recommendations on treatment of Hepatitis C 2020 EASL-The Home of Hepatology (free registration needed to get access) and page 121
- vi MELD calculation, see page 81

# Treatment and Monitoring of Persons with HBV/HIV Co-infection

#### Treatment indication

- All persons with HBV/HIV co-infection should receive ART that includes TDF or TAF unless history of tenofovir intolerance
- Stopping anti-HBV active ART should be avoided in persons with HIV/ HBV co-infection because of the high risk of severe hepatitis flares and decompensation following HBV reactivation hepatitis

#### **Treatment selection**

- If TDF or TAF is strictly contraindicated, entecavir may be prescribed in
- persons with no prior 3TC exposure and together with fully active ART Persons with liver cirrhosis and low CD4 count require careful surveillance in the first months after starting ART in order not to overlook immune reconstitution syndrome and subsequent liver decompensation due to flares of liver enzymes (for management of cirrhotic persons, see pages 80-84). Please note that diagnosis of cirrhosis may be difficult in persons already on HBV treatment
- Caution is warranted to switch from a TDF/TAF-based regimen to drugs with a lower genetic barrier, e.g. FTC or 3TC, in particular in 3TC-pretreated cirrhotic persons as viral breakthrough due to archived YMDD mutations is likely to happen. This has also been described in individuals with previous 3TC HBV-resistance who have been switched from TDF to
- Prior to ART simplification with a regimen without TDF/TAF, HBV status should be re-checked
- For HBV/HIV co-infected persons with BMD changes or CKD, see recommendations for Dose Adjustment of ARVs for Impaired Renal Function and pages 71-76

#### **Treatment goal**

The optimal treatment duration for nucleos(t)ide analogues with anti-HBV activity has not yet been determined and experts recommend life-long therapy. In those on ART where the nucleoside backbone needs changing, anti-HBV therapy may be stopped cautiously after confirmed HBsAg-seroconversion. In persons with liver cirrhosis, stopping of effective anti-HBV treatment is not recommended, in order to avoid liver decompensation due to flares of liver enzymes

#### **Treatment monitoring**

- Liver blood tests should be performed every 3 months during the first year and every 6-12 months thereafter
- HBV-DNA should be determined every 3-6 months during the first year and every 12 months thereafter HBsAg should be checked at 12 months intervals at least until loss of HBsAq(

#### **HBV** reactivation

- 11. In HBs-Ag negative, anti-HBc positive persons undergoing immunosuppression:
  - Those treated with severe immunosuppressive therapy (chemotherapy for lymphoma/leukaemia or stem-cell or solid-organ transplantation) should receive TDF/TAF therapy to prevent HBV reactivation. For persons with other markers of possible HBV exposure including isolated anti-HBs positivity (without a history of vaccination) careful monitoring for HBV reactivation is required
  - In persons treated with B-cell-depleting agents (rituximab, ofatumumab, natalizumab, alemtuzumab, ibritumomab) TDF/TAF should be part of the ART. If TDF/TAF is contraindicated, second line options include ETV, 3TC and FTC. However, cases of reactivation due to 3TC resistance have been described
  - In those not treated with HBV-active ART who receive other immunosuppressive therapy (e.g. TNF alpha inhibitor), careful monitoring with HBV-DNA and HBsAg is required for HBV reactivation. If this is not possible, addition of TDF/TAF is recommended
- Quantitative HBsAg < 1000 IU/mL predicts HBsAg loss

# Treatment and Monitoring of Persons with HCV/HIV Co-infection

#### **Treatment indication**

- Every person with HCV/HIV co-infection must be considered for DAAbased anti-HCV treatment regardless of liver fibrosis stage
- Due to similar HCV cure rates and tolerability in HCV/HIV co-infected persons as in HCV mono-infected persons under DAA therapy, treatment indication and regimens are to be the same as in HCV monoinfection

#### **Treatment selection**

- 3 DAA combinations are now standard of care for chronic HCV infection, see Tables HCV Treatment Options in HCV/ HIV Co-infected Persons. IFN-based therapies and first generation PIs (boceprevir and telaprevir) are not recommended because of insufficient efficacy and increased toxicities.
- Selection of DAA combinations is based upon stage of liver fibrosis, HCV GT<sup>0</sup>, pre-treatment history and resistance-associated substitutions (RAS) if tested
- Due to drug-drug interactions in particular with HIV and HCV PIs, careful checking for interactions is urgently recommended prior to starting HCV therapy, see Drug-drug Interactions between Viral Hepatitis Drugs and ARVs or http://www.hep-druginteractions.org
- 6. Resistance testing, if available, should be considered before re-treatment of persons who failed after a PI-and/or NS5A inhibitor-containing agent. The triple combination of SOF/VEL/VOX for 12 weeks is the treatment of choice for re-treatment, especially if resistance testing is not available. In persons with complex mutations patterns SOF+GLE/PIB + RBV for 12-16 weeks can also be considered. In case of unavailability of SOF/VEL/VOX or SOF + GLE/PIB other regimens with at least two active DAAs could be combined with the preferential use of one drug with high genetic barrier to resistance and with extended treatment durations and potentially addition of RBV. In patients with decompensated cirrhosis SOF/VEL + RBV for 24 weeks is the only available option for re-treatment in case of contraindication to liver transplantation

#### Treatment goal

- 7. The primary aim of HCV treatment is SVR<sub>12</sub> defined as undetectable HCV-RNA 12 weeks after the end of therapy (evaluated using sensitive molecular tests) or HCV core antigen levels where HCV- RNA assays are not available or not affordable. SVR<sub>12</sub> corresponds to a definitive cure of HCV infection in the vast majority of cases
- i If pangenotypic regimens are foreseen, HCV GT determination is not mandatory before starting treatment. HCV GT determination should be considered in persons at risk of reinfection in order to differentiate between relapse and re-infection in case of reemergence of HCV RNA after therapy

See online video lectures HCV/HIV Co-infection from the EACS online course Management of HIV and Co-infections

#### Treatment monitoring

- 8. In persons with advanced fibrosis (≥ F3) differential blood count, creatinine, liver enzymes, bilirubin, albumin and INR measurement after 2-4 weeks of therapy is recommended. In HBsAg negative persons with positive anti-HBc, monitoring of ALT and HBV-DNA in case of ALT elevation is recommended
- In persons with impaired renal function undergoing SOF based treatment creatinine should also be monitored
- 10. HCV-RNA measurement during therapy should only be performed in order to assess compliance and/or break-through in persons experienced to oral DAAs; HCV-RNA should be measured at end-of-treatment and at week 12 or 24 after treatment cessation (to assess SVR). In persons receiving all oral DAA therapy, no association between viral load at any given time-point during therapy and SVR has yet been found. If HCV-RNA determination is not available SVR can be identified by a negative HCV core antigen 24 weeks after treatment end
- 11. HIV-VL every 12 weeks

#### Post-Treatment monitoring

- Surveillance for HCC and for oesophageal varices should be continued if the respective indications were present pre-treatment, despite achieving SVR, see pages 9, 59, 80 and 81
- All persons with concurrent causes of liver disease should undergo periodical clinical assessments
- 14. Increase in body weight and changes in lipid and glucose metabolism have been described after SVR. Thus, surveillance, counseling and treatment for obesity and metabolic alterations should be enforced after SVR, see page 85

#### Treatment of recently acquired HCV infection

- 15. IFN-containing HCV regimens are no longer recommended
- 16. HCV treatment immediately after diagnosis is recommended in persons with ongoing risk behavior to reduce onward transmission. IFN-free treatment with DAAs is recommended as in treatment naïve persons without cirrhosis (except for those with pre-existing cirrhosis), see page 118
- 17. If treatment is not indicated immediately, HCV-RNA should be re-measured 4 weeks later. Treatment is recommended in persons without a decrease of 2\*log<sub>10</sub> of HCV-RNA at 4 weeks compared with initial HCV-RNA, due to the very low probability of spontaneous clearance, and in persons with persistent serum HCV-RNA 12 weeks after diagnosis of recently acquired HCV, see Management of Recently acquired HCV in Persons with HIV Co-infection
- 18. For more detailed information on the management of recently acquired HCV infection we refer to the Recommendations on Recently acquired and early chronic hepatitis C in MSM from the European treatment network for HIV, hepatitis and global infectious diseases consensus panel

# **HCV Treatment Options in HCV/HIV Co-infected Persons**

HCV GT	Treatment regimen	Treatment durat	ion & RBV usage			
		Non-cirrhotic	Compensated cirrhotic	Decompensated cirrhotics CTP class B/C		
1 & 4	EBR/GZR	12 week	12 weeks <sup>(i)</sup>			
	GLE/PIB	8 weeks	8-12 weeks <sup>(ii)</sup>	Not recommended		
	SOF/VEL	12 week	12 weeks			
	SOF/LDV +/- RBV	8-12 weeks without RBV(iii)	12 weeks with RBV <sup>(iv)</sup>	12 weeks with RBV(ix)		
2	GLE/PIB	8 weeks	8-12 weeks <sup>(ii)</sup>	Not recommended		
	SOF/VEL	12 wee	ks	12 weeks with RBV(ix)		
3	GLE/PIB	8 weeks <sup>(v)</sup>	8-12 weeks <sup>(ii,v)</sup>	Not recommended		
	SOF/VEL +/- RBV	12 weeks <sup>(vi)</sup>	12 weeks with RBV(vii)	12 weeks with RBV(ix)		
	SOF/VEL/VOX	-	12 weeks	Not recommended		
5 & 6	GLE/PIB	8 weeks	8-12 weeks <sup>(ii)</sup>	Not recommended		
	SOF/LDV +/- RBV	12 weeks +/- RBV <sup>(viii)</sup>	12 weeks with RBV(iv)	12 weeks with RBV(ix)		
	SOF/VEL	12 we	12 weeks			

For HCV treatment options to be used if preferred options are not available, please see version 10.1 of the EACS Guidelines

EBR =elbasvir

GLE = glecaprevir

GZR =grazoprevir

**LDV** = ledipasvir

PIB = pibrentasvir

RBV =ribavirin

SOF = sofosbuvir

VEL = velpatasvir

VOX =voxilaprevir

RAS =resistance associated substitutions

- In persons with GT1a with baseline HCV-RNA < 800,000 IU/mL and/or absence of NS5A RASs, as well as in treatment-naïve persons with GT4 with HCV-RNA < 800,000 IU/mL In GT 1b treatment-naïve persons with F0-F2 fibrosis 8 weeks can be considered
- ii 8 weeks treatment can be considered in treatment naïve persons
- iii 8 weeks treatment without RBV only in treatment-naïve persons with F < 3 and baseline HCV-RNA < 6 million IU/mL
- iv RBV can be omitted in treatment-naïve or -experienced persons with compensated cirrhosis without baseline NS5A RAS. In persons intolerant to RBV, treatment may be prolonged to 24 weeks
- v Treatment duration in HCV GT3 who failed previous treatment with IFN and RBV +/- SOF or SOF and RBV should be 16 weeks
- vi In treatment experienced persons RBV should be added unless NS5A RASs are excluded; if these persons are intolerant to RBV, treatment may be prolonged to 24 weeks without RBV
- vii If RAS testing is available and demonstrates absence of NS5A RAS Y93H, RBV can be omitted in treatment naive people with compensated cirrhosis
- viii In treatment experienced (exposure to IFN/RBV/SOF) persons, add RBV treatment for 12 weeks or prolong treatment to 24 weeks without RBV
- ix In persons intolerant to RBV, treatment may be prolonged to 24 weeks



# **Drug-drug Interactions between Viral Hepatitis Drugs and ARVs**

	al hepatitis igs	ATV/c	ATV/r	DRV/c	DRV/r	LPV/r	DOR	EFV	ETV	NVP	RPV	FTR	MVC	віс	CAB oral	CAB/ RPV	DTG	EVG/c	RAL	TAF	TDF
	elbasvir/ grazoprevir	1	↑376% ↑958%	1	↑66% ↑650%	↑271% ↑1186%	↓4% ↑7%	↓54% ↓83%	1	1	↑7% ↓2%	<b>↔</b> ↑	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↓2% ↓19%	↑118% ↑436%	↓19% ↓11%	$\leftrightarrow$	↓7% ↓14%
	glecaprevir/ pibrentasvir	1	↑553% ↑64%	1	↑397%	↑338% ↑146%	$\leftrightarrow$	1	1	1	E 84%	1	Е	E	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑205% ↑57% E47%	E47%	$\leftrightarrow$	E29%
DAAs	sofosbuvir	$\leftrightarrow$	$\leftrightarrow$	1	↑34%	$\leftrightarrow$	$\leftrightarrow$	↓6%	$\leftrightarrow$	$\leftrightarrow$	↑9%	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↓5% D27%	$\leftrightarrow$	↓6%
HCV D	sofosbuvir/ ledipasvir	↑ a	↑8% ↑113%a	↑ a	↑34% ↑39% <mark>a</mark>	↔a	↑4% ↓8%	↓6% ↓34%a	$\leftrightarrow$	$\leftrightarrow$	↑10% ↑8% <mark>a</mark>	1	Е	↑7% ↓13%	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑36% ↑78% <mark>a</mark>	↓5% ↓9% D~20%	E32%	Ea
	sofosbuvir/ velpatasvir	↔a	↑22% ↑142%a	↔a	↓28% ↓16%a	↓29% ↑2%a	$\leftrightarrow$	↓3% ↓53%	ļ	1	↑16% ↓1%	1	Е	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↓8% ↓9%	↑ a	↑24% ↓2%	$\leftrightarrow$	Еa
	sofosbuvir/ velpatasvir/ voxilaprevir	1	↑40% ↑93% ↑331%	↑ a	↓28% ↓5% ↑143%b	1	$\leftrightarrow$	1	1	1	$\leftrightarrow$	1	Е	↑9% ↓4% ↓9%	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	↑22% ↑16% ↑171%a	$\leftrightarrow$	Е	Ea
HDV	Bulevirtide	1	1	1	1	1	E	1	1	$\leftrightarrow$	E	$\leftrightarrow$	Е	$\leftrightarrow$	$\leftrightarrow$	E	$\leftrightarrow$	1	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$

#### Colour legend

No clinically significant interaction expected
These drugs should not be co-administered

Potential clinically significant interaction that is likely to require additional monitoring, alteration of drug dosage or timing of administration Potential interaction likely to be of weak intensity. Additional action/monitoring or dosage adjustment is unlikely to be required

#### Legend

↑ Potential elevated exposure of the hepatitis therapy
↓ Potential decreased exposure of the hepatitis therapy

→ No significant effect

D Potential decreased exposure of ARV drug
E Potential elevated exposure of ARV drug

ATV/c ATV co-formulated with COBI (300/150 mg qd) DRV/c DRV co-formulated with COBI (800/150 mg qd) CAB/RPV CAB and RPV im long acting injections

(PK and/or QT interactions shown are with RPV)

Numbers refer to decreased or increased AUC as observed in drug-drug interaction studies.

First/second numbers refer to AUC changes for EBR/GZR or GLE/PIB or SOF/LDV or SOF/VEL.

First/second/third numbers refer to AUC changes for SOF/VEL/VOX

#### Interactions with ABC, FTC, 3TC, ZDV

ABC, FTC, 3TC, ZDV: no clinically relevant interactions expected.

#### Interactions with ibalizumab

None

#### Comments

- a Monitoring of renal function recommended due to increase of tenofovir concentration if the regimen contains TDF.
- b Study details are with DRV/r qd. DRV bid has not been studied and should be used with caution as voxilaprevir concentrations may increase more than with DRV qd (this would be of further significance in cirrhotic patients). Monitoring of renal function recommended due to increase of tenofovir concentrations if the regimen contains TDF.

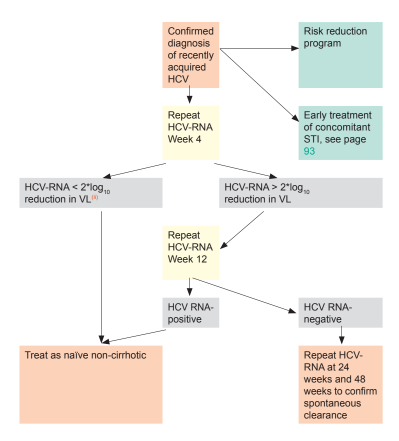
#### **Further Information**

For additional drug-drug interactions and for more detailed pharmacokinetic interaction data and dosage adjustments, please refer to: http://www.hiv-druginteractions.org (University of Liverpool)



# **Management of Recently Acquired HCV Infection**

- 1. DAA based HCV treatment immediately after diagnosis is recommended in persons with ongoing risk behavior
- 2. If immediate treatment is not indicated, the algorithm below should be used



- i HCV-RNA < 2\*log<sub>10</sub> reduction at week 4 is considered as early chronic HCV infection (eg: 2\*log<sub>10</sub> reduction = reduction from 100,000 to 1000 IU/mL)
- ii See also Recommendations on Recently acquired and early chronic hepatitis C in MSM from the European treatment network for HIV, hepatitis and global infectious diseases consensus panel

# **Cut-off Values of Non-invasive Tests for the Detection of Advanced Fibrosis** and Cirrhosis

HIV/Hepatitis C co-infection (according to EASL recommendations on Treatment of Hepatitis C 2020)

Test	Stage of fibrosis	Cut off	Sensitivity (%)	Specificity (%)	Positive predictive value (%)	Negative predictive value (%)
Fibroscan	F3*	10 kPa	72	80	62	89
	F4*	13 kPa	72-77	85-90	42-56	95-98
APRI	F4	2	48	94	n.a.	n.a.
		1	77	75	n.a.	n.a.
Fib-4	F4	3.25	55	92	n.a.	n.a.
		1.45	90	58	n.a.	n.a.

These cut-offs were derived from different studies and the optimal values might vary between populations and must be interpreted together with the individual clinical assessment

#### HIV/Hepatitis B co-infection

Test	Stage of fibrosis	Cut off	Sensitivity (%)	Specificity (%)	Positive predictive value (%)	Negative predictive value (%)
Fibroscan	F3	7.6 kPa	85	87	77	92
	F4	9.4 kPa	92	94	79	98
APRI	F4	2	35	89	26	92
		1	65	75	22	95



<sup>\*</sup>The distinction between F3 and F4 is often imprecise and must be interpreted in the individual clinical context

### **Hepatitis D and E infection**

#### **Hepatitis Delta Virus (HDV)**

- 1. HDV antibodies should be screened for in all HBsAg positive persons
- 2. In persons with positive HDV antibodies, HDV-RNA should be measured in order to assess activity of the disease
- In persons with chronic HDV co-infection and significant liver fibrosis (≥ F2), long-term (at least 12 months) treatment with PEG-IFN might be considered
  in association with TDF-based ART
- 4. Non-invasive fibrosis markers (transient elastography and serum markers) should be used with caution in HIV/HBV co-infected persons with chronic HDV infection as there are no well-established thresholds
- 5. Because of its anti-HBV activity, TDF/TAF should be added, as part of ART, to PEG-IFN in order to reduce HBV-DNA load
- 6. Bulevirtide (2mg/d; s.c.) in combination with TDF/TAF is recommended in HDV-RNA positive persons with compensated liver disease and should be used where available. The optimal duration of treatment remains unclear. Treatment should be performed in centers with sufficient experience
- 7. People with HIV and HDV infection should be referred to university centers for treatment and if possible enrolled in trials on new drugs active against HDV
- 8. Treatment efficacy should be monitored with HBV-DNA and HDV-RNA measurements, when available, and with follow-up of biochemical and liver fibrosis estimates
- 9. Persistent off-treatment HDV-RNA negativity and anti-HBs seroconversion are the ideal goals of antiviral treatment for HDV even if they can only be obtained in a minority of persons. Histological remission of liver disease is a less ambitious but more likely achievable goal
- 10. In persons with HDV and ESLD or HCC, liver transplantation from HBsAg negative donors should be strongly considered. Transplant with anti-HBV prophylaxis post-OLTX cures HBV and HDV infection

#### Hepatitis E Virus (HEV)

- 11. Screening for HEV infection is warranted in persons with symptoms consistent with acute hepatitis, unexplained flares of aminotransferases (even if suspected drug induced liver injury), unexplained elevated liver function tests, neuralgic amyotrophy, Guillain-Barrè, encephalitis or proteinuria
- 12. Screening should include anti-HEV IgG and IgM and HEV-RNA in blood and, if possible, in stool
- 13. Treatment with RBV (600 mg daily) may be considered in cases of severe acute HEV, acute-on-chronic liver failure, extrahepatic HEV related disease or in persons with persisting HEV replication three months after first detection of HEV-RNA. RBV should be given for a duration of 12 weeks followed by HEV-RNA measurements in serum and stool. If HEV-RNA is undetectable in both, RBV can be stopped. In persons in whom HEV-RNA is still detectable in serum and/or stool, RBV may be continued for an additional three months. In the setting of chronic HEV infection in immunosuppressed persons, reduction in immunosuppression should be considered



# Part VI Opportunistic Infections and COVID-19

This section provides:

- Recommendations for timing on ART initiation in persons with OIs without prior ART exposure
- · Overview of IRIS and recommendations on its management
- · Overview of the most important aspects in management of the most frequent OIs occurring in persons with HIV in Europe
- Overview of management of COVID-19 in persons with HIV

See online videos Tuberculosis and HIV Co-infection-Part 1 and Tuberculosis and HIV Co-infection-Part 2 from the EACS online course Management of HIV and Co-infections

# When to start ART in persons with Opportunistic Infections (OIs)

	Initiation of ART	Comments
General recommendation	As soon as possible within 2 weeks after starting treatment for the opportunistic infection	
Tuberculosis	As soon as possible within two weeks of starting TB treatment, regardless of CD4 count	For details, see ART in TB/HIV Co-infection section, page 20
- TB meningitis	ART should be delayed for 4 weeks, but can be initiated within the first 2 weeks in persons with TB meningitis and CD4 < 50 (100) cells/µL	Corticosteroids are recommended as adjuvant treatment for TB meningitis
Cryptococcal meningitis	Defer initiation of ART for at least 4 weeks (WHO recommends a delay of 4-6 weeks and some specialists recommend a delay of 6-10 weeks in severe cryptococcal meningitis)	Corticosteroids are not recommended as adjuvant treatment



# **Immune Reconstitution Inflammatory Syndrome (IRIS)**

Definition	
Paradoxical IRIS	<b>Paradoxical worsening</b> of symptoms during the ART-induced immune-reconstitution period in association with inflammatory signs (by physical exam, imaging or tissue biopsy), after exclusion of the expected course of a treated/untreated OI or drug toxicities
Unmasking IRIS	<b>New onset</b> of symptoms during the ART-induced immune-reconstitution period in association with inflammatory signs (by physical exam, imaging or tissue biopsy), after exclusion of the expected course of a treated/untreated OI or drug toxicities
Prevention	
Cryptococcal meningitis:	
paradoxical IRIS	Start therapy with amphotericin B plus flucytosine and defer start of ART for at least 4 weeks.
unmasking IRIS	Determine serum cryptococcal antigen in persons newly HIV-diagnosed with CD4 counts < 100 cells/µL. If cryptococcal antigen is detected, exclude active cryptococcal disease, and, in particular, examine CSF to rule out cryptococcal meningitis. If meningitis is ruled out, start pre-emptive therapy. For details, see below the specific section on cryptococcal disease
Tuberculosis	
paradoxical IRIS	Simultaneous initiation of ART and prophylactic prednisone in persons with CD4 cell count < 100 cells/μL, who started anti-TB treatment within 30 days prior to ART, may reduce risk of TB-IRIS by 30%. <b>Prednisone</b> dose: 40 mg qd po for 2 weeks, followed by 20 mg qd po for 2 weeks
Treatment	
treatment In cases where anti-inflammatory treatment	veeks with continuation of specific treatment for the OI, without discontinuing ART and without anti-inflammatory ent is contemplated by the physician, corticosteroids or non-steroidal anti-inflammatory agents can be used. e or specific administration schedules in the specific conditions
TB-IRIS	Start of systemic corticosteroids is recommended (e.g., <b>prednisone</b> 1.5 mg/kg/day po for 2 weeks, then 0.75 mg/kg/day for 2 weeks)
Life-threatening CNS-IRIS:	
TB-meningitis	Prednisone (1.5 mg/kg/day po for 2 weeks, then tapering)
PML	<b>Methylprednisolone</b> (1 g/day iv for 3-5 days or dexamethasone 0.3 mg/kg/day iv for 3-5 days), then oral tapering



# **Primary Prophylaxis of Ols According to Stage of Immunodeficiency**

#### CD4 count threshold / indication

CD4 count < 200 cells/µL, CD4 percentage < 14%, recurrent oral thrush, or relevant concomitant immunosuppression\*

Prophylaxis against Pneumocystis jirovecii Pneumonia (PcP) & Toxoplasma gondii infection

Stop: if CD4 count > 100 cells/µL and HIV-VL undetectable over 3 months

\* e.g. use of corticosteroids > 20 mg prednisone equivalent per day for > 2 weeks, cancer chemotherapy, biological agents such as rituximab and others. Decisions on installation and discontinuation in these situations have to be taken individually

	Drug	Dose	Comments
Positive or negative serology for Toxoplasmosis	trimethoprim- sulfamethoxazole (TMP-SMX)	80/400 mg qd po or 160/800 mg qd po or 160/800 mg x 3/week po	
Negative serology for toxoplasmosis	pentamidine	300 mg in 6 mL sterile water x 1 inhalation/month	Does not prevent the rare extrapulmonary manifestations of <i>P. jirovecii</i>
Negative serology for toxoplasmosis	dapsone	100 mg qd po	Check for G6PD-deficiency
Negative serology for toxoplasmosis	atovaquone suspension	1500 mg qd (with food)	
Positive serology for toxoplasmosis	dapsone	200 mg/week po	Check for G6PD-deficiency
	+ pyrimethamine	75 mg/week po	
	+ folinic acid	25-30 mg/week po	
Positive serology for toxoplasmosis	atovaquone suspension +/- pyrimethamine + folinic acid	1500 mg qd po (with food) 75 mg/week po 25-30 mg/week po	
Positive cryptococcal serum antigen and CD4 count < 100 cells/μL	fluconazole	800 mg qd po for 2 weeks followed by 400 mg qd po for 8 weeks	Asymptomatic individual and cryptococ- cal meningitis, pulmonary or other site infection ruled out

#### CD4 count < 50 cells/µL

Prophylaxis against Non-Tuberculous Mycobacteria (NTM) (M. avium complex, M. genavense, M. kansasii)

Prophylaxis is not recommended if ART is started

Prophylaxis may be considered for persons with CD4 counts < 50 cells/µL who remain viremic on ART (drug resistant HIV with no option to achieve virologic control); exclude disseminated MAC disease before starting

Regimens listed are alternatives	azithromycin	1200-1250 mg/week po	Check for interactions with ARVs, see	
	or clarithromycin	500 mg bid po	Drug-drug Interactions between ARVs and Non-ARVs	
	or rifabutin	300 mg qd po	Check for interactions with ARVs, see Drug-drug Interactions between ARVs and Non-ARVs  Active TB should be ruled out before starting rifabutin	



# Primary Prophylaxis, Treatment and Secondary Prophylaxis/Maintenance Treatment of Individual Ols

#### Pneumocystis jirovecii Pneumonia (PcP)

#### Primary prophylaxis

Start: if CD4 count < 200 cells/ $\mu$ L, CD4 percentage < 14%, oral thrush or relevant concomitant immunosuppression, see Primary Prophylaxis of Ols Stop: if CD4 count > 100 cells/ $\mu$ L and HIV-VL undetectable over 3 months

	Drug	Dose	Comments
Negative or positive serology for toxoplasmosis	TMP-SMX	80/400 mg qd po or 160/800 mg qd po or 160/800 mg x 3/week po	
Negative serology for toxoplasmosis	pentamidine	300 mg in 6 mL sterile water x 1 inhalation/month	Does not prevent the rare extrapulmonary manifestations of <i>P. jirovecii</i>
Negative serology for toxoplasmosis	dapsone	100 mg qd po	Check for G6PD-deficiency
Negative serology for toxoplasmosis	atovaquone suspension	1500 mg qd po (with food)	
Positive serology for toxoplasmosis	dapsone	200 mg/week po	Check for G6PD-deficiency
	+ pyrimethamine	75 mg/week po	-
	+ folinic acid	25-30 mg/week po	
Positive serology for toxoplasmosis	atovaquone suspension +/- pyrimethamine + folinic acid	1500 mg qd po (with food) 75 mg/week po 25-30 mg/week po	

#### Treatment

Treat at least 21 days, then secondary prophylaxis until CD4 count > 100 cells/ $\mu$ L and HIV-VL undetectable over 3 months Diagnosis:

**Definitive diagnosis:** Cough and dyspnoea on exertion AND microorganism identification by cytology / histopathology of induced sputum (sensitivity up to 80%), broncho-alveolar lavage (sensitivity > 95%) or bronchoscopic tissue biopsy (sensitivity > 95%)

Presumptive diagnosis: CD4 count < 200 cells/µL AND dyspnoea / desaturation on exertion and cough AND radiology compatible with PcP AND no evidence for bacterial pneumonia AND response to PcP treatment. SARS-CoV-2 pneumonia can resemble PcP and should therefore be included in the differential diagnoses

	Drug	Dose	Comments
Preferred therapy	TMP-SMX	5 mg/kg tid TMP iv/po + 25 mg/kg tid SMX iv/po	Monitor myelotoxicity (mainly neutro- penia), kidney function and electrolytes (mainly high potassium)
	+ prednisone if PaO <sub>2</sub> < 10 kPa or < 70 mmHg, or alveolar/arterial O <sub>2</sub> gradient > 35 mmHg. Start prednisone preferentially 15-30 min before treatment	40 mg bid po 5 days 40 mg qd po 5 days 20 mg qd po 10 days	Benefit of corticosteroids if started within 72 hours after start of treatment
Alternative therapy for moderate to severe	primaquine	30 mg (base) qd po	Check for G6PD deficiency
PcP	+ clindamycin	600-900 mg tid iv/po	
	or pentamidine	4 mg/kg qd iv (infused over 60 min.)	
	For each regimen: + prednisone if PaO <sub>2</sub> < 10 kPa or < 70 mmHg, or alveolar/ arterial O <sub>2</sub> gradient > 35 mmHg. Start prednisone preferentially 15-30 min before TMP/SMX	40 mg bid po 5 days 40 mg qd po 5 days 20 mg qd po 10 days	Benefit of corticosteroids if started within 72 hours after start of treatment  Some studies support the addition of caspofungin or other echinocandins to
			standard treatment in persons with moderate-severe PcP (can be considered, but not mandatory)
Alternative therapy for mild to moderate	primaquine	30 mg (base) qd po	Check for G6PD deficiency
PcP	+ clindamycin	600-900 mg tid po	
	or		
	atovaquone suspension	750 mg bid po (with food)	
	or		
	dapsone	100 mg qd po	Check for G6PD deficiency
	+ trimethoprim	5 mg/kg tid po	In case of rash: reduce dose of TMP (50%), use antihistamines



#### Secondary prophylaxis / Maintenance treatment

Ston: if CD4 count > 100 cells/ul, and HIV-VI, undetectable over 3 months

	Drug	Dose	Comments
Negative or positive serology for toxoplasmosis	TMP-SMX	80/400 mg qd po or 160/800 mg x 3/week po	
Negative serology for toxoplasmosis	pentamidine	300 mg in 6 mL sterile water x 1 inhalation/month	Not to use in the rare case of extrapul- monary manifestations of <i>P. jirovecii</i>
Negative serology for toxoplasmosis	dapsone	100 mg qd po	Check for G6PD-deficiency
Negative serology for toxoplasmosis	atovaquone suspension	1500 mg qd po (with food)	
Positive serology for toxoplasmosis	dapsone	200 mg/week po	Check for G6PD-deficiency
	+ pyrimethamine	75 mg/week po	
	+ folinic acid	25-30 mg/week po	
Positive serology for toxoplasmosis	atovaquone suspension +/- pyrimethamine + folinic acid	1500 mg qd po (with food) 75 mg/week po 25-30 mg/week po	

#### Toxoplasma gondii Encephalitis

#### Primary prophylaxis

Start: if CD4 count < 200 cells/µL, or CD4 percentage < 14%, oral thrush, or relevant concomitant immunosuppression (see above)

Stop: if CD4 count > 100 cells/µL and HIV-VL undetectable over 3 months

	Drug	Dose	Comments
Preferred prophylaxis	TMP-SMX	80/400 mg qd po or 160/800 mg qd po or 160/800 mg x 3/week po	All regimens are also effective against PcP
Alternative prophylaxis	atovaquone suspension	1500 mg qd po (with food)	
	dapsone + pyrimethamine + folinic acid	200 mg/week po 75 mg/week po 25-30 mg/week po	Check for G6PD-deficiency
	atovaquone suspension + pyrimethamine + folinic acid	1500 mg qd po (with food) 75 mg/week po 25-30 mg/week po	
Treatment			

Treat 6 weeks, then secondary prophylaxis until CD4 count > 200 cells/µL and HIV-VL undetectable over 6 months Diagnosis:

Definitive diagnosis: clinical symptoms, typical neuroradiology AND cytological / histological detection of organism in tissue Presumptive diagnosis: clinical symptoms, typical neuroradiology AND response to empirical treatment. This is the standard in most clinical settings

	Drug	Dose	Comments
Preferred therapy	pyrimethamine	Day 1: 200 mg qd po, <b>then</b> • If ≥ 60 kg; 75 mg qd po • If < 60 kg: 50 mg qd po	Monitor for myelotoxicity of pyrimethamine, mostly neutropenia
	+ sulfadiazine	<ul> <li>If ≥ 60 kg: 3000 mg bid po/iv</li> <li>If &lt; 60 kg: 2000 mg bid po/iv</li> </ul>	Sulfadiazine is associated with crystalluria and may lead to renal failure and urolithiasis. Good hydration is essential. Check renal function and urine sediment for microhematuria and crystalluria
	+ folinic acid	10-15 mg qd po	
Alternative therapy	pyrimethamine + clindamycin + folinic acid	Day 1: 200 mg qd po, then • If ≥ 60 kg: 75 mg qd po • If < 60 kg: 50 mg qd po 600-900 mg qid po/iv 10-15 mg qd po	Monitor for myelotoxicity of <b>pyrimethamine</b> , mostly neutropenia Additional PcP prophylaxis is necessary
	or TMP-SMX	5 mg TMP/kg bid iv/po 25 mg SMX/kg bid iv/po	Preferred intravenous regimen if oral route not possible Monitor myelotoxicity (mainly neutropenia), kidney function and electrolytes (mainly high potassium)
	or pyrimethamine + atovaquone + folinic acid	Day 1: 200 mg qd po, <b>then</b> If ≥ 60 kg; 75 mg qd po If < 60 kg: 50 mg qd po 1500 mg bid po (with food) 10-15 mg qd po	Monitor for myelotoxicity of pyrimethamine, mostly neutropenia
	or sulfadiazine + atovaquone	If ≥ 60 kg: 3000 mg bid po/iv     If < 60 kg: 2000 mg bid po/iv     1500 mg bid po (with food)	Sulfadiazine is associated with crystalluria and may lead to renal failure and urolithiasis. Good hydration is essential. Check renal function and urine sediment for microhematuria and crystalluria
	or pyrimethamine + azithromycin + folinic acid	Day 1: 200 mg qd po, then  • If ≥ 60 kg; 75 mg qd po  • If < 60 kg: 50 mg qd po  900-1200 mg qd po  10-15 mg qd po	Monitor for myelotoxicity of pyrimethamine, mostly neutropenia



#### Secondary prophylaxis / Maintenance therapy

Stop: if CD4 count > 200 cells/µL and HIV-VL undetectable over 6 months

	Drug	Dosage	Comments
Regimens listed are alternatives	sulfadiazine + pyrimethamine + folinic acid	2000-3000 mg bid - qid po 25-50 mg qd po 10-15 mg qd po	
	or clindamycin + pyrimethamine + folinic acid	600 mg tid po 25-50 mg qd po 10-15 mg qd po	Additional PcP prophylaxis is necessary
	or atovaquone suspension + pyrimethamine + folinic acid	750-1500 mg bid po (with food) 25-50 mg qd po 10-15 mg qd po	
	or atovaquone suspension	750-1500 mg bid po (with food)	
	or TMP-SMX	160/800 mg bid po	

#### Cryptococcosis - disease caused by Cryptococcus neoformans

Cryptococcal meningitis is the most frequent manifestation of cryptococcosis. Cryptococcal infection can also cause a pneumonitis which may be difficult to distinguish from Pneumocystis pneumonia. Infection may also involve other organs or may be disseminated

Primary prophylaxis: One large RCT in Africa showed that an enhanced infection prophylaxis in severely immunosuppressed persons (CD4 < 50 cells/ µL) including TMP-SMX 160/800 mg for 12 weeks, isoniazid 300 mg/day for 12 weeks, fluconazole 100 mg/day for 12 weeks, azithromycin 500 mg/day for 5 days and albendazole 400 mg single dose may decrease overall opportunistic infections (including cryptococcal meningitis) and mortality. Due to the different epidemiology of opportunistic infections in Africa and in Europe these results may not be extrapolated to European countries

 $\label{eq:decomposition} \textbf{Diagnosis:} \ \ \text{Positive microscopy, OR detection of antigen in serum or CSF OR culture from CSF, blood or urine.} \ \ \text{Serum cryptococcal antigen should be performed in all newly HIV-diagnosed persons with CD4 counts < 100 cells/<math>\mu$ L. See Pre-emptive therapy below

Treatment (Cryptococcal meningitis and disseminated cryptococcosis): 14 days induction therapy, then 8 weeks consolidation therapy, then second-

	Drug	Dose	Comments
Pre-emptive therapy	fluconazole	800 mg qd po for 2 weeks followed by 400 mg qd po for 8 weeks	In case of: - positive cryptococcal serum antigen - asymptomatic individual with CD4 < 100 cells/µL - cryptococcal meningitis, pulmonary or other site infection ruled out
Induction therapy	liposomal amphotericin B + flucytosine	3 mg/kg qd iv 25 mg/kg qid po	- Perform repeated lumbar puncture (LP) until opening pressure is < 20 cm H <sub>2</sub> 0:  150 period of the control of
	or amphotericin B deoxycholate + flucytosine	0.7 mg/kg qd iv 25 mg/kg qid po	- If CSF culture is sterile, switch to oral regimen  - Repeated LPs or CSF shunting are essential to effectively manage increased intracranial pressure which is associated with better survival  - Corticosteroids have no effect in reducing increased intracranial pressure, could be detrimental and are contraindicated  - Flucytosine dosage must be adapted to renal function  - Defer start of ART for at least 4 weeks, since early initiation of ART is associated with decreased survival  - Due to substantial nephrotoxicity amphotericin B deoxycholate should only be used, if liposomal amphotericin B is not available  - Flucytosine may not be available in all European countries. Consider replacing it by fluconazole 800 mg qd during the induction phase  - In resource-limited settings, a large RCT suggested that i) one week of amphotericin B + flucytosine followed by one week of fluconazole 1200 mg qd or ii) two weeks of fluconazole 1200 mg qd or ii) two weeks of fluconazole 1200 mg qd plus flucytosine may be acceptable induction regimens. Another more recent trial showed that a single-dose liposomal amphotericin B (10 mg/kg/d) combined with flucytosine and fluconazole 1200 mg qd for a total of 14 days was noninferior to one week of amphotericin B + flucytosine followed by one week of fluconazole 1200 mg qd.

Consolidation therapy	fluconazole	400 mg qd po (single loading dose of 8 on 1st day)	8 weeks See Drug-drug Interactions between ARVs and Non-ARVs			
Secondary prophylaxis / Maintena	nce therapy					
At least 12 months  Consider to stop: if CD4 count >100 cells/µL and HIV-VL undetectable over 3 months						
Drug Dose Comments						
	fluconazole	200 mg qd po	See Drug-drug Interactions between			

#### Candidiacie

Candidiasis						
Oropharyngeal Candidiasis						
Diagnosis: typical clinical appearance. See	Diagnosis: typical clinical appearance. See Drug-drug Interactions Between ARVs and Non-ARVs, for all azole therapies					
	Drug	Dose	Comments			
Preferred alternatives	fluconazole	150-200 mg qd po	Once or until improvement (5-7 days)			
	nystatin	3-6 lozenges at 400000 units (aprox. 4-6 mL of oral suspension)/day	7-14 days			
	or amphotericin B	oral suspension 1-2 g bid - qid				
Oesophagitis						
<b>Definitive diagnosis:</b> macroscopic inspecti <b>Presumptive diagnosis:</b> if recent onset of			om the mucosal surface			
	Drug	Dose	Comments			
Preferred alternatives	fluconazole	400 mg qd po or 400 mg loading dose, then 200 mg qd po	3 days 10-14 days			
	consider posaconazole or voriconazole or caspofungin and other echinocandins	400 mg bid po 200 mg bid po 70 mg iv qd day 1, then 50 mg qd	In cases of refractory disease, treat according to resistance testing. Adapt <b>posaconazole</b> and <b>voriconazole</b> dose according to MIC's of candida and drug trough levels			

#### Histoplasmosis (Histoplasma capsulatum)

Diagnosis: antigen detection in blood, urine or broncho-alveolar fluid, OR positive microscopy, OR mycological culture of blood, urine, broncho-alveolar fluid, CSF or tissue biopsy, OR PCR in blood or other clinical samples. Aspergillus galactomannan assays may be helpful to diagnose disseminated infections as cross reactivity occurs.

Note: CSF, which shows typically a lymphatic pleocytosis, is usually microscopy and culture negative. Detection of histoplasma antigen or antibody is more sensitive. A clinical diagnosis is possible, if disseminated histoplasmosis is present and CNS infection is not explained by another cause. Fluconazole should not be used for treatment of histoplasmosis. Little clinical evidence is available for the use of voriconazole or posaconazole. Be aware of interactions of azoles with ARVs, see Drug-drug Interactions Between ARVs and Non-ARVs. Measurement of plasma concentration of itraconazole is advised to guide optimal treatment, and itraconazole oral suspension should be preferred due to better bioavailability. Serum itraconazole

trough concentration should be at least 1 mcg/mL if measured by high-performance liquid chromatography (HPLC) Dose Severe disseminated histoplasmosis Induction therapy: liposomal amphotericin B 3 mg/kg qd iv For 2 weeks or until clinical improvement Consolidation therapy: 200 mg tid po for 3 days, For at least 12 months itraconazole then 200 mg bid po Moderate disseminated histoplasmosis itraconazole 200 mg tid po for 3 days, For at least 12 months then 200 mg bid po

5 mg/kg qd iv

200 mg bid - tid po

#### Secondary prophylaxis / Maintenance therapy

Stop: if CD4 count > 150 cells/µL and HIV-VL undetectable over 6 months, negative fungal blood cultures, histoplasma serum antigen < 2 µg/L or negative PCR, if available, and > 1 year treatment

Consider long-term suppressive therapy in severe cases of meningitis and in cases of relapse despite adequate treatment

Induction therapy:

itraconazole

liposomal amphotericin B

Consolidation therapy:

200 mg qd po itraconazole



Histoplasma meningitis

For 4-6 weeks

For at least 12 months and until resolu-

tion of abnormal CSF findings

#### Talaromycosis (Talaromyces (former Penicillium marneffei))

Consider diagnosis in persons with HIV who live/lived in Asia

Diagnosis: antigen detection in blood, urine or broncho-alveolar fluid, OR positive microscopy, OR mycological culture of blood, urine, broncho-alveolar fluid, CSF or tissue biopsy or PCR in blood OR other clinical samples.

Aspergillus galactomanan assays may be helpful to diagnose disseminated infections as cross reactivity occurs

	Drug	Dose	Comments
Severe disseminated talaromycosis	Induction therapy: liposomal amphotericin B	3 mg/kg qd iv	For 2 weeks or until clinical improvement
	Consolidation therapy: itraconazole	200 mg tid po for 3 days, then 200 mg bid po	For at least 10 weeks (followed by secondary prophylaxis)
Moderate talaromycosis	itraconazole	200 mg tid po for 3 days, then 200 mg bid po	For 8 weeks (followed by secondary prophylaxis)

#### Secondary prophylaxis / Maintenance therapy

Secondary prophylaxis: itraconazole 200 mg qd po

Stop: if CD4 count > 100 cells/µL and HIV-VL undetectable over 6 months, negative fungal blood cultures or negative PCR/ negative antigen

#### Herpes simplex virus (HSV) infections

Diagnosis: PCR of skin lesions/CSF/biopsy is the preferred diagnostic method. Tissue antigen detection may be used. Clinical appearance of skin/mucosal lesions is not reliable

During treatment: monitor renal function, adjust drug dose in renal impairment

	Drug	Dose	Comments
Initial and recurrent genital / mucocutaneous HSV			See Sexual and Reproductive Health section, page 91
Severe mucocutaneous lesions	aciclovir	5 mg/kg tid iv	After lesions begin to regress, switch to oral treatment for 21-28 days or longer, until lesions have healed
Encephalitis and retinitis	aciclovir	10 mg/kg tid iv	14-21 days
Aciclovir resistant mucocutaneous HSV infection	foscarnet	90 mg/kg bid iv	Until clinical response If <b>foscarnet</b> is not available, cidofovir 5 mg/kg once weekly can be used. Topical <b>cidofovir</b> and <b>foscarnet</b> can be used for external lesions

#### Varicella zoster virus (VZV) infections

Diagnosis: typical clinical appearance with/without serological testing. PCR of skin lesions/CSF/biopsy is the preferred diagnostic method. Tissue antigen detection may be used

During treatment: monitor renal function, adjust drug dose in renal impairment

	Drug	Dose	Comments
Primary Varicella infection (Chickenpox)	valaciclovir	1000 mg tid po	Chickenpox: 5-7 days,
and Herpes Zoster (Shingles): Not disseminated	or famciclovir or aciclovir	500 mg tid po 800 mg x 5/day po	Shingles: 7-10 days
Herpes Zoster: Disseminated	aciclovir	10 mg/kg tid iv	10-14 days (or until clinical improvement)
Encephalitis (including vasculitis), retinitis	aciclovir	10-15 mg/kg tid iv	14-21 days If retinitis, consult ophthalmologist



#### Cytomegalovirus (CMV) infections

#### **Treatment**

Diagnosis of retinitis: clinical appearance of typical retinal lesions AND response to therapy. PCR of aqueous and vitreous humor optional Diagnosis of esophagitis/colitis: endoscopic presence of ulcerations AND typical histopathological picture (cellular / nuclear inclusion bodies) Diagnosis of encephalitis/myelitis: clinical appearance AND positive PCR in CSF AND other pathology excluded. Antibody testing and PCR in blood not useful for diagnosis of end-organ disease

During treatment: monitor renal function, adjust drug dose in renal impairment

	Drug	Dose	Comments	
Retinitis, immediate sight-threatening	ganciclovir	5 mg/kg bid iv	3 weeks, then secondary prophylaxis	
lesions	or foscarnet	90 mg/kg bid iv	Foscarnet used as alternative therapy if toxicity or resistance to ganciclovir. Most experts would add intravitreal injections of ganciclovir (2 mg) or foscarnet (2.4 mg) for 1-4 doses over 7-10 days in combination with systemic CMV treatment	
Retinitis, small peripheral retinal lesions	valganciclovir	900 mg bid po (with food)	2-3 weeks, then secondary prophylaxis	
	or foscarnet	90 mg/kg bid iv		
Oesophagitis/Colitis	ganciclovir	5 mg/kg bid iv	3-6 weeks, until symptoms resolved, then secondary prophylaxis (switch to oral <b>valganciclovir</b> once tolerated)	
	or foscarnet	90 mg/kg bid iv		
	or valganciclovir	900 mg bid po (with food)	In milder disease if oral treatment tolerated	
Encephalitis/Myelitis	ganciclovir foscarnet	5 mg/kg bid iv 90 mg/kg bid iv	Treat until symptoms resolved and CMV replication in CSF has cleared (negative PCR DNA-CMV in CSF) Treatment is individualised according to clinical symptoms and response to treatment. Some guidelines recommend ganciclovir combined with foscarnet especially in relapse	
Secondary Prophylaxis / Maintenance th	erapy: Cytomegalovirus (CMV) F	Retinitis		
Stop: Inactive lesions treated for at least 3 months and CD4 count > 100 cells/µL and HIV-VL undetectable over 3 months				
Regimens	valganciclovir (preferred regimen)	900 mg qd po (with food)		
	or ganciclovir	6 mg/kg qd (x 5 days/ week) iv		
	or foscarnet	90-120 mg/kg qd (x 5 days/ week) iv		

#### Progressive Multifocal Leukoencephalopathy (PML)

Treatment

Definitive diagnosis (laboratory): evidence of JCV-DNA in CSF AND presence of compatible clinical-radiological picture

Definitive diagnosis (histology): typical histological findings within situ evidence of JCV-DNA antigen or JCV-DNA AND presence of compatible

Presumptive diagnosis: compatible clinical-radiological picture if JCV-DNA in CSF negative or not performed. JCV-DNA in plasma may complement PML

diagnosis, particularly if CSF not available. Ivialy also be a market of disease progression		
Person off-ART	Initiate ART immediately (following general guidelines for treatment, see Initial Combination Regimen for ART-naïve Adults, INSTI may reasonably be preferred, given the importance of rapid immune reconstitution in PML. Attention should be made to development of IRIS, see IRIS section	
Person on-ART, HIV-VL failure	Optimise ART (following general guidelines for treatment, see Virological Failure), INSTI may reasonably be preferred, given the importance of rapid immune reconstitution in PML. Attention should be made to development of IRIS, see IRIS section	
Person on-ART, treated for weeks- months or on effective ART	Continue current ART	
	Note: There is no specific treatment for JCV infection that proved to be effective in PML outside of anecdotal case reports. Therefore, there is no recommendation to use the following drugs which previously or occasionally were used in PML: Alpha-IFN, cidofovir, corticosteroids (except for treatment of IRIS-PML, see IRIS section, cytarabine, iv immunoglobulins, mefloquine, mirtazapine. Newer immune-based approaches have shown some efficacy, including Interleukin-7, infusion of polyomavirus-specific HLA-matched T-cells, anti-PD1 inhibitors (pembrolizumab, nivolumab), but no conclusive data, e.g., from clinical trials or broader clinical experience, are currently supporting their recommendation for clinical use. If used, participation in treatment protocols is strongly encouraged	



#### Bacillary Angiomatosis (Bartonella henselae, Bartonella quintana)

Treatment			
Diagnosis: typical histology			
	Drug	Dose	Comments
	doxycycline	100 mg bid po	Until improvement (until 2 months)
	or clarithromycin	500 mg bid po	Possible interactions with ARVs, see Drug-drug Interactions between ARVs and Non-ARVs



#### Infections with Non-Tuberculous Mycobacteria (NTM) (M. avium complex, M. genavense, M. kansasii)

#### Primary prophylaxis

#### Primary prophylaxis

Prophylaxis is not recommended if ART is started

Prophylaxis may be considered for persons with CD4 counts < 50 cells/µL who remain viremic on ART (drug resistant HIV with no option to achieve virologic control); exclude disseminated MAC disease before starting

	Drug	Dose	Comments
Regimens listed are alternatives	azithromycin	1200-1250 mg/week po	Check for interactions with ARVs, see
	or clarithromycin	500 mg bid po	Drug-drug Interactions between ARVs and Non-ARVs
	or rifabutin	300 mg qd po	Check for interactions with ARVs, see Drug-drug Interactions between ARVs and Non-ARVs. Active TB should be ruled out before starting <b>rifabutin</b>

Diagnosis: clinical appearance and cultures of blood, lymph nodes, bone marrow or other usually sterile specimen. For any treatment regimen, check interactions with ARVs, see Drug-drug Interactions between ARVs and Non-ARVs

Active TB should be ruled out before starting anti-TB drugs (rifampicin, rifabutin, ethambutol, isoniazid)

Mycobacterium avium-intracellulare complex (MAC)			
Preferred therapy	clarithromycin + ethambutol +/- rifabutin	500 mg bid po 15-20 mg/kg qd po 300 mg qd po (or 150 mg qd if PI/b)	rifabutin can be considered in case of severe disease, if resistance to macrolides or ethambutol is suspected, or in case of high bacterial load (> 2*log of CFU/mL of blood). rifabutin is indicated if ART is not given
	rifabutin can be replaced by: + levofloxacin/ moxifloxacin or + amikacin	500 mg qd po/400 mg qd po 10-15 mg/kg qd iv	levofloxacin/ moxifloxacin or amikacin can be considered as a 4th drug for disseminated or severe/refractory disease (no data on additional clinical benefit)
	azithromycin + ethambutol	500 mg qd po 15-20 mg/kg qd po	Consider additional drugs as above
Mycobacterium kansasii			
	rifampicin + isoniazid + ethambutol	600 mg qd po (or <b>rifabutin</b> 300 mg qd po) 300 mg qd po 15-20 mg/kg qd po	12 months after negative culture
	or rifampicin + clarithromycin + ethambutol	600 mg qd po (or <b>rifabutin</b> 300 mg qd po) 500 mg bid po 15-20 mg qd po	12 months after negative culture
Secondary prophylaxis / Maintenance the	rapy for MAC		
Stop: if CD4 count > 100 cells/µL and HIV-V	L undetectable over 6 months and	MAC treatment for at least 12 mor	nths
Mycobacterium avium (MAC) infection	clarithromycin	500 mg bid po	

#### Cryptosporidiosis (C. parvum, C. hominis)

Regimens listed are alternatives

Diagnosis of cryptosporidiosis is made in persons with chronic diarrhea, mostly in cases with CD4 count < 100 cells/µL, by immunofluorescence, acid fast stain, cryptosporidium antigen or PCR of stools or tissue. If the diarrhea lasts > 4 weeks, the diagnosis of cryptosporidiosis is an AIDS defining illness

15-20 mg/kg qd po

15-20 mg/kg qd po

500 mg qd po

Mainstay of therapy is the induction of ART to restore immune competence with CD4 count > 100 cells/µL

+ ethambutol

azithromycin

+ ethambutol

Additional measures are symptomatic treatment, rehydration and electrolyte management

The following antiprotozoal therapies can be used additively to ART in severe cases, but are not sufficient to achieve protozoal eradication without immune restoration

Drug	Dose	Comments
nitazoxanide	500-1000 mg bid po	14 days
or		
paromomycin	500 mg qid po	14-21 days



#### Cystoisosporiasis (Cystoisospora belli, formerly Isospora belli)

#### Treatment

**Diagnosis** of cystoisosporiasis is made in persons with chronic, mostly watery diarrhoea by UV fluorescence or microscopy of stools, duodenal aspirates or intestinal tissue biopsy. If the diarrhea lasts > 4 weeks, the diagnosis of cystoisosporiasis is an AIDS defining illness

Besides antiprotozoal treatment, additional measures are symptomatic treatment, rehydration and electrolyte management

besides antiprotozoal treatment, additional measures are symptomatic treatment, renydration and electrolyte management			
	Drug	Dose	Comments
Preferred therapy	TMP-SMX	320/1600 mg bid po or 160/800 mg bid po	Treat minimally 10 days, increase duration to 3-4 weeks if symptoms worsen or persist  Treat minimally 10 days, increase dose to 2 x 2 tablet/day, if symptoms worsen or persist
Alternative therapy, if TMP-SMX is not tolerated	pyrimethamine + folinic acid or ciprofloxacin	50-75 mg qd po 10-15 mg qd po 500 mg bid po	10 days Monitor for myelotoxicity, mostly neutro- penia, for <b>pyrimethamine</b> 7 days
Secondary prophylaxis / Maintenance th	erapy		
Stop: if CD4 count > 200 cells/µL and HIV-VL undetectable over 6 months and no signs of persistent cystoisosporiasis			
Preferred therapy	TMP-SMX	160/800 mg x 3/week po or 160/800 mg qd po or 320/1600 mg x 3/week po	
Alternative therapy, if TMP-SMX is not tolerated	pyrimethamine + folinic acid	25 mg qd po 10-15 mg qd po	Monitor for myelotoxicity, mostly neutropenia, for <b>pyrimethamine</b>

#### Visceral leishmaniasis

Treatment			
Diagnosis: microscopy or PCR in sme	ears, body fluids or tissue		
	Drug	Dose	Comments
Preferred treatment	liposomal amphotericin B	2-4 mg/kg qd iv for 10 consecutive days	Then secondary prophylaxis
	or liposomal amphotericin B	4 mg/kg qd iv on day 1-5, 10, 17, 24, 31 and 38	A recent trial suggested that addition of 50 mg bid of miltefosine for 14 days to a cumulative dose of 30 mg/kg of liposomal amphotericin B administered every other day (i.e. 5 mg/kg on days 1, 3, 5, 7, 9, and 11) is non-inferior to standard regimens
Alternative therapy	lipid complex amphotericin B	3 mg/kg qd iv	10 days
	or amphotericin B deoxycholate	0.5-1 mg/kg qd iv (total dose 1.5-2 g)	
	or pentavalent antimonium salt (Glucantime®)	20 mg/kg qd iv or im	4 weeks
	or miltefosine	1.5-2.5 mg/kg qd po	4 weeks
Secondary prophylaxis / Maintenan	ce therapy		
Consider stopping: if CD4 count > 20 or negative urinary antigen	00-350 cells/µL and HIV-VL undetectable	over 3 months, no relapse for at I	east 6 months and negative PCR in blood
Preferred treatment	liposomal amphotericin B	4 mg/kg every 2-4 weeks iv	
	or lipid complex amphotericin B	3 mg/kg every 3 weeks iv	
Alternative therapy	pentavalent antimonium salts (Glucantime®)	20 mg/kg every 4 weeks iv/im	
	or miltefosine	100 mg qd po	
	or pentamidine	300 mg every 3 to 4 weeks iv	



# **Diagnosis and Treatment of TB in Persons with HIV**

#### Treatment of TB in Persons with HIV

For standard treatment of TB in persons with HIV, including appropriate choice of ARVs, see table below and ART in TB/HIV Co-infection See online video lectures TB and HIV Co-infection-Part 1 and TB and HIV Co-infection Part 2 from the EACS online course Clinical Management of HIV

Disease	Drug	Dose <sup>(i)</sup>	Comments*
Susceptible Mycobacterium tuberculosis			
Initial phase	rifampicin + isoniazid (+ pyridoxine) + pyrazinamide + ethambutol	Weight based	Initial phase for 2 months. Possibility to omit ethambutol, if <i>M. tuberculosis</i> is known to be fully drug sensitive. Preventive steroid therapy may be considered to avoid IRIS, see IRIS section
Alternative Initial phase	rifabutin + isoniazid (+ pyridoxine) + pyrazinamide + ethambutol	Weight based	Initial phase for 2 months.  Possibility to omit ethambutol, if  M. tuberculosis is known to be fully drug sensitive
Continuation phase	rifampicin/rifabutin + isoniazid (+ pyridoxine)	Weight based	Total duration of therapy:  1. Pulmonary, drug susceptible TB: 6 months  2. Pulmonary TB & positive culture at 8 weeks of TB treatment: 9 months  3. Extrapulmonary TB with CNS involvement or disseminated TB: 9-12 months  4. Extrapulmonary TB with bone/joint in- volvement and in other sites: 6-9 months

An alternative shorter regimen of rifapentine, isoniazid, pyrazinamide and moxifloxacin for 2 months, followed by rifapentine, isoniazid and moxifloxacin for 2 months can be used, if rifapentine is available (see WHO communication, 2022)



<sup>\*</sup> Intermittent regimens (2 or 3 times per week) are contraindicated in persons with HIV. Missed doses can lead to treatment failure, relapse or acquired drug

i For dose details, please see separate table TB drug doses, page 138

#### Diagnosis of Multidrug Resistant TB (MDR-TB) / Extensively Drug-Resistant TB (XDR-TB)

MDR/XDR-TB should be suspected in case of:

- Previous or incomplete TB treatment
- Contact with MDR/XDR-TB index case
- Birth, travel or work in an area endemic for MDR-TB
- History of poor adherence
- No clinical improvement on standard therapy and/or sputum smear positive after 2 months of TB therapy or culture positive at 3 months
- Homelessness/hostel living and, in some countries, recent/current incarceration
- In areas with very high MDR/XDR-TB prevalence

MDR-TB: Resistance to isoniazid AND rifampicin

XDR-TB - since 2021: Resistance to isoniazid AND rifampicin AND fluoroquinolones AND at least one additional Group A drug, see below

#### Rapid detection

Gene Xpert or similar technology has the advantage of rapid detection of rifampicin resistance. Drug susceptibility testing is important for optimizing treatment

#### Treatment of resistant TB

#### Isoniazid-resistant TB

- rifampicin/rifabutin + pyrazinamide + ethambutol + levofloxacin or moxifloxacin for 6 months, (WHO 2020 recommendations)

#### Rifampicin-resistant (RR) and MDR-TB

- Treatment of MDR/XDR-TB is a specialist area. WHO has published new Guidelines (2020) and an additional more recent rapid communication (2022)

#### - Currently recommended all-oral regimen:

Can be used in persons with confirmed RR/MDR-TB who have not been exposed to bedaquiline, pretomanid, linezolid for > 1 month

6 months of bedaquiline, pretomanid, linezolid (600 mg gd) and moxifloxacin (BPaLM). This regimen may be used without moxifloxacin if resistance to fluoroquinolones (pre-XDR-TB) is documented (BPaL). In this case consider extension of 3 months. Data on the effectiveness of this regimen in extensive pulmonary TB disease or severe extra-pulmonary TB are currently not available (see WHO rapid communication)

#### - Alternative all oral regimen:

Can be used in persons with MDR/RR-TB without resistance to fluoroquinolones and without previous exposure to second-line drugs and without extensive pulmonary TB or severe extra-pulmonary TB

- 4-6 months bedaquiline (6 months) + levo-/ moxifloxacin + ethionamide\*\* + ethambutol + isoniazid (high-dose) + pyrazinamide + clofazimine followed by
- 5 months levo-/ moxifloxacin + ethambutol + pyrazinamide + clofazimine
- \*\*4 months of ethionamide can be replaced by 2 months of linezolid (600 mg qd)

#### Longer TB treatment regimens (>18 months)

Patients with XDR-TB and those not eligible to or failing all-oral short regimens may benefit from individualized longer treatment.

All three Group A drugs and at least one Group B drug should be included to ensure that treatment starts with at least four TB drugs likely to be effective, and that at least three agents are included for the rest of treatment if bedaquiline is stopped.

If only one or two Group A drugs are used, both Group B drugs are to be included.

If the regimen cannot be composed with drugs from Groups A and B alone, Group C drugs are added to complete it.

The duration of longer regimens must be individualized. For details, see WHO Guideline

Treatment compliance is crucial. If needed, each dose of medicines should be given as DOT throughout the whole treatment period Surgery

Surgical resection may be part of the management for selected persons with focal pulmonary MDR-TB

#### **Drug choices**

Each empiric regimen should be reassessed and modified if needed once drug sensitivity results become available

drug schsitivity results become available		
Group A: Include all three drugs	levofloxacin or moxifloxacin     bedaquiline     linezolid	
Group B: Add one or both drugs	• clofazimine • cycloserine or terizidone	
Group C: Add to complete the regimen and when drugs from Groups A and B cannot be used	ethambutol     delamanide     pyrazinamide     amikacin (or streptomycin – only if susceptible)     imipenem–cilastatin or meropenem     ethionamide or prothionamide     para-aminosalicylic acid	

Pretomanid is recommended but not yet included in Group A drugs

#### Drug interactions with ART and MDR/XDR regimens

When treating RR/MDR/XDR-TB careful review of DDIs and potential toxicities is mandatory before initiating ART, see Drug-drug Interactions between Anti-tuberculosis Drugs and ARVs



#### Latent tuberculosis

Indication: TST > 5 mm or positive IGRA or close contacts to persons with sputum smear positive tuberculosis. See Assessment at Initial & Subsequent Visits

Some national guidelines consider the ethnicity, CD4 count and ART usage to define indication for latent tuberculosis treatment

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- Other preventive regimens may be considered if high risk of latent infection with MDR/XDR-TB **Rifabutin** is not a WHO recommended regimen
- Rifapentine is not approved by EMA

# **TB Drug Doses**

Drug name	Dose	Comments	
First line drugs			
Isoniazid	5 mg/kg qd (usual dose 300 mg)	Max 375 mg qd Caution: neurotoxicity, add <b>pyridoxine</b> 20 mg qd	
Rifampicin	10 mg/kg qd (usual dose 600 mg)	Rifampicin is not recommended in persons receiving Pls, DOR, ETR, NVP, RPV, FTR, BIC, CAB, CAB/RPV LA, EVG/c. see Drug-drug Interactions between Anti-tuberculosis Drugs and ARVs and page 20	
Rifabutin without PIs, EFV, RPV with PIs with EFV with TAF or EVG/c	5 mg/kg qd (usual dose 300 mg) 150 mg qd 450-600 mg qd Not recommended		
<b>Pyrazinamide</b> 40-55 kg 56-75 kg 76-90 kg > 90 kg	1000 mg qd 1500 mg qd 2000 mg qd 2000 mg qd		
Ethambutol 40-55 kg 56-75 kg > 75 kg	800 mg qd 1200 mg qd 1200 mg qd	Max 1600 mg qd Caution: optic neuritis Baseline colour vision should be tested	
Other drugs			
Levofloxacin 30-45 kg > 46 kg	750 mg qd 1000 mg qd	Max 1500 mg qd	
Moxifloxacin	400 mg qd	Max 800 mg qd (used in the standardized shorter MDR-TB regimen) Monitor ECG in respect of QT prolongation	
Bedaquiline	400 mg qd for 2 weeks 200 mg qd three times weekly for 22 weeks	EFV, ETV: potential reduction of <b>bedaquiline</b> exposure and activity. Not recommended Boosted regimens: increase in <b>bedaquiline</b> exposure. Potential risk of QT interval prolongation, ECG monitoring recommended. Avoid coadministration > 14 days	
Linezolid	600 mg qd	Max 1200 mg qd Caution: hematological side effects and neurotoxicity, including optic neuropathy	
Clofazimine	100 mg qd	Alternative: 200 mg for 2 months then 100 mg qd Caution: skin toxicity Monitor ECG in respect of QT prolongation	
Cycloserine or terizidone 30-45 kg > 46 kg	500 mg qd 750 mg qd	Max 1000 mg qd Caution: neurotoxicity; add <b>pyridoxine</b> , up to 50 mg/250 mg <b>cycloserine</b>	
Delamanid	100 mg bid for 24 weeks	Monitor ECG in respect of QT prolongation	
Imipenem/cilastatin	1000/1000 mg bid iv		
Meropenem	1000 mg tid iv		
Amikacin 30-35 kg 36-45 kg 46-55 kg > 55 kg	625 mg qd iv 750 mg qd iv 750-1000 mg qd iv 1000 mg qd iv	After initial period can be reduced to trice weekly Baseline audiometry should be performed Caution: monitor renal function, audiometry and drug levels	
Streptomycin	12-18 mg/kg qd iv	Max 1000 mg qd iv	
Ethionamide or prothionamide 30-45 kg 46-70 kg > 70 kg	500 mg qd 750 mg qd 1000 mg qd	Caution: gastrointestinal toxicity; add <b>pyridoxine</b> , up to 50 mg/250 mg <b>prothionamide</b>	
Para-aminosalycilic acid	4000 mg bid	In weight > 70 kg can be increased to 4000-6000 mg bid Caution: gastrointestinal toxicity	
Pretomanid	200 mg qd	Use with <b>bedaquiline</b> and <b>linezolid</b> for 26 weeks Monitor ECG in respect of QT prolongation Peripheral neuropathy is common adverse effect	



### Management of COVID-19 in persons with HIV

#### Introduction

#### Epidemiology of SARS-CoV-2

- Incidence of SARS-CoV-2 infection in persons with HIV seems to be similar to that reported in the general population, although some studies reported higher SARS-CoV-2 incidence rate. Whether low CD4 count (< 200 cells/µL) and detectable HIV-VL are associated with increased rate of SARS-CoV-2 diagnosis is unclear
- Higher rates of SARS-CoV-2 breakthrough infections have been reported in fully vaccinated persons with HIV when compared to the general population

#### Risk factors for severe COVID-19 outcomes

- More adverse COVID-19 outcomes (hospitalization, disease severity, mortality) have been reported in persons with HIV and CD4 count < 200-350 cells/µL when compared to the general population and to persons with HIV with higher CD4 count, and in persons with untreated HIV infection and/or with detectable viremia when compared to those with controlled HIV infection, with a possible increasing association of COVID-19 severity with higher HIV-VL
- An increased incidence of severe COVID-19 has been described in persons with concomitant OIs (especially TB and PcP) and associated comorbidities. Among hospitalized COVID-19 patients, studies reported a younger age and higher rates of comorbidities in people with HIV when compared to the general population

#### Care during COVID-19 epidemic

- In case of lockdown or home isolation, it is important to ensure continuum of HIV-care
- It is recommended to develop local country-specific strategies to prevent disruption in HIV care, including teleconsultation and tele-pharmacy, and ensure continuous ART supply (consider providing at least 3 months of ART at a time)
- Telemedicine and phone visits can be used for chronically stable persons, not requiring changes in ART or co-medications. Retain in-person visits for persons recently diagnosed with HIV and initiating ART, or complaining of acute problems, adverse effects due to ART, virological failure, STIs or other complains/ co-morbidities requiring clinical evaluation
- New development or worsening of mental health problems (anxiety, depression, increased loneliness and stigma) have been very common following social distancing and lockdowns; psychological/psychiatric and social support should be actively offered
- Periodic updates to the position of EACS on SARS-CoV-2 risk and prevention in persons with HIV can be consulted here

#### **Management of COVID-19**

#### Diagnostic approach:

The same approach, as for general population, should be applied, according to the national or international recommendations. For details, see WHO

#### Differential diagnosis:

For persons with HIV, particularly for those with poor immune status, other respiratory diseases (e.g., PcP, and TB) should be considered as differential diagnosis. Appropriate diagnostic workup should follow current recommendations, see Opportunistic infections

- Treatment of COVID-19 should be the same as for general population. As treatment guidelines and prescription requirements for COVID-19 might vary between countries, refer to national guidelines. In absence of those, follow international recommendations: NIH; WHO
- Several early treatments with anti-SARS-CoV-2 directed agents, including antiviral drugs or monoclonal antibodies, are available to prevent COVID-19 progression to severe disease. People with HIV may be eligible for such treatments, according to local guidelines, and those with AIDS, poor immune responses to ART and/or ART-untreated HIV infection should be preferentially considered for early anti-SARS-CoV-2 treatment initiation.
- Check for drug-drug interactions and overlapping toxicities between COVID-19 treatments (particularly nirmatrelvir/ritonavir or other anti-SARS-CoV-2 directed agents, corticosteroids, and anticoagulants) and ARV drugs, see table Drug-drug Interactions between COVID-19 Therapies and ARVs, Drugdrug Interactions and Other Prescribing Issues, Drug-drug Interactions between Corticosteroids and ARVs
- Isolation precautions should be the same as for the general population, although persons with uncontrolled HIV infection may show long-term viral sheddina

#### Management of HIV infection during COVID-19 infection

- ART should neither be stopped nor modified due to recently diagnosed SARS-CoV-2 infection, unless strictly necessary (no ARV drug has proved to be clinically effective against SARS-CoV-2 infection)
- The ART regimen should be adapted in persons who are unable to swallow their ARV drugs (e.g., those on mechanical ventilation or ECMO therapy). see Administration of ARVs in Persons with Swallowing Difficulties
- Total lymphocytes, CD4 and CD8 subpopulations may decrease during acute COVID-19; in these cases, consider appropriate OI prophylaxis, see Primary Prophylaxis of Ols According to stage of Immunodeficiency
- HIV-RNA blips have been described during COVID-19, their clinical relevance is unknown
- Co-morbidities and co-infections should be managed as indicated in specific sections of the Guidelines, see Prevention and Management of Co-morbidities. Viral hepatitis co-infections. Opportunistic infections.
- Well-being general measures (e.g. diet/ exercise) should be recommended

#### Management of long-term symptoms and sequelae of COVID-19 (Post-acute COVID-19 syndrome, PACS)

- A substantial proportion of COVID-19 patients may show progressive or newly presenting symptoms, involving the lungs or other organs, weeks to months after COVID-19 (post-acute COVID-19 syndrome, PACS); studies addressing whether frequency of PACS is increased in persons with HIV are
- These conditions should be specifically addressed and evaluated; refer to the appropriate specialist consultations following local/national guidelines for persistent COVID-19 sequelae
- Check for drug-drug interactions if any treatment for post- COVID-19 complications is initiated, see Drug-drug Interactions and Other Prescribing Issues, Drug-drug Interactions between COVID-19 Therapies and ARVs, Drug-drug Interactions between Corticosteroids and ARVs



#### Prophylaxis of COVID-19 in persons with HIV

As vaccination guidelines and prescription requirements for anti-SARS-CoV-2 pharmacological prophylaxis might vary between countries, and the efficacy of specific antiviral agents may differ against different SARS-CoV-2 variants, please refer to national guidelines and local epidemiology. In absence of those, please follow international recommendations: NIH; WHO

#### SARS-CoV-2 vaccination:

- Several COVID-19 vaccines have been approved in Europe and other countries worldwide and numerous vaccine candidates are in development
- It is recommended for all persons with HIV to be vaccinated against SARS-CoV-2. There is no data to recommend a specific vaccine and the choice depends on the availability in individual countries. Priority should be given to those with immunosuppression (CD4 count < 350 cells/µL), if access to vaccines is limited
- Specific safety concerns with SARS-CoV-2 vaccines in people with have not been described so far. Serological testing before vaccination is not required, see Vaccination
- Persons with advanced disease (CD4<200 cells/mm³) and/or detectable VL have poorer humoral and cellular mediated immune responses to vaccination, and are candidates for booster doses following local guidelines (see also Vaccination)
- Other vaccines (particularly S pneumoniae and influenza) should be given as scheduled, see Vaccination
- Links to an overview of available vaccines and information regarding SARS-CoV-2 vaccination in persons with HIV: NIH; BHIVA; WHO; EACS

#### Pharmacological prevention of SARS-CoV-2 infection:

- Passive immunization with antibodies against SARS-CoV-2 can be considered as pre-exposure prophylaxis (PrEP) and post-exposure prophylaxis (PrEP) to prevent severe COVID-19 in unvaccinated persons or immunocompromised persons who do not respond adequately to vaccination
- These approaches can be considered in unvaccinated persons or in those with AIDS, CD4 count <200 cells/µL, or ART-untreated infection, following local guidelines</li>



# Part VII Paediatric HIV Treatment

#### Initiation of ART in Children and Adolescents

- We recommend the initiation of ART in all children and adolescents diagnosed with HIV irrespective of age, clinical stage, CD4 count and VL
- We emphasise the need for urgent diagnosis for infants born to women with HIV and prompt treatment of infants diagnosed with HIV infection
- We endorse the "U=U" campaign (undetectable (defined as VL < 200 copies/mL for > 6 months) = untransmissible) for sexual transmission of HIV, which is particularly relevant to sexually active adolescents and is potentially a motivational message to enhance adherence and prevent onward HIV transmission

# Initial Combination Regimen for ART-naïve Children and Adolescents, Table 1

- Where available, baseline resistance testing should be performed
- All first line regimens currently include 2 NRTIs together with a drug from a different class (third agent)
- DTG plus 2NRTI combination is the preferred option in all children over 4 weeks of age and 3 Kg
- Evidence for superiority of DTG compared to NNRTI or PI/b has been demonstrated by the ODYSSEY trial
- Whilst "preferred options" are recommended, "alternative options" are acceptable and remain important choices in settings where ART availability is limited or in individuals at particular risk of specific toxicity or DDIs
- Whenever possible first line 3rd agents with high barrier to resistance have been selected in view of potential difficulties with adherence in children and
- When choosing a regimen, potential transmitted resistance, including from maternal or infant ART exposure after failed prevention of vertical transmission should always be considered
- In infants under 4 weeks and/or under 3 kg, when NVP has been used in pregnancy or there is a risk of transmitted NVP resistance, non-NNRTI-based ART, including RAL from birth or LPV/r from 2 weeks are preferred

### **Additional Specific Paediatric Considerations**

- It should be noted that these Guidelines occasionally include recommendations for use of ARVs outside their European licence
- Local policy for the use of unlicensed medication in children and adolescents should be followed
- Apart from decisions on standard first line in high prevalence setting, options should be discussed within a multidisciplinary team (MDT)/paediatric virtual clinic (PVC)
- If local MDT or PVC are unavailable, an international PVC is accessible by contacting the Guideline Team.
- Adherence is key to achieving and maintaining viral suppression and adherence support and assessment should be provided at/prior to initiation of ART and at all subsequent visits
- The use of peer mentors, where available, is strongly recommended
- Although age cut offs are used in Table 1 it should be noted that weight as well as age are also included in the licensing of ARVs in children
- Detailed guidance on paediatric dosing is available from the Penta website, https://penta-id.org/hiv/treatment-guidelines/
- Drug formulations that are useful for paediatric dosing and administration are summarised in Table 2



Table 1. Preferred and Alternative First Line Options in Children and Adolescents

	Backbone		3 <sup>rd</sup> Agent (in alphabetical order)	
Age	Preferred	Alternative	Preferred	Alternative
0 - 4 weeks	ZDV <sup>(i)</sup> + 3TC	-	LPV/r <sup>(i, ii)</sup> NVP <sup>(iii)</sup> RAL <sup>(iii)</sup>	-
4 weeks - 3 years	ABC <sup>(v)</sup> + 3TC <sup>(v)</sup>	ZDV <sup>(i)</sup> + 3TC <sup>(vi)</sup> TDF <sup>(vii)</sup> + 3TC	DTG <sup>(viii)</sup>	LPV/r NVP RAL
3 - 6 years	ABC <sup>(v)</sup> + 3TC <sup>(v)</sup>		DTG <sup>(viii)</sup>	DRV/r EFV LPV/r NVP RAL
6 - 12 years	ABC(v) + 3TC(v) TAF(x) + XTC(tx)	TDF(vii) + XTC(ix)	DTG(viii)	DRV/r EFV EVG/c RAL
> 12 years	ABC(v) + 3TC(v) TAF(x) + XTC(ix)	TDF <sup>(vii)</sup> + XTC <sup>(ix)</sup>	BIC <sup>(xi)</sup> DTG <sup>(viii)</sup>	DRV/b EFV <sup>(xii)</sup> RAL <sup>(xii)</sup> RPV <sup>(xii)</sup>

#### Notes:

Toxicities as listed in the table on page 24 and 25 should be considered. Additional toxicity considerations specific to paediatrics are described in the footnotes below

- In view of potential long-term toxicity, any child on ZDV should be switched to ABC (preferred for younger children) or TAF/TDF (alternative for youngerer children, with renal/bone toxicity monitoring with TDF) once increase in age and/or weight makes licensed formulations available. When ABC is contraindicated in young children it is recommended that treatment options are discussed at MDT to decide between ZDV, TDF or TAF on a case by case basis
- LPV/r should not be administered to neonates before a postmenstrual age of 42 weeks and a postnatal age of at least 14 days although it may be considered if there is a risk of transmitted NVP resistance and appropriate INSTI formulations are unavailable. In these circumstances the neonate should be monitored closely for LPV/r related toxicity (e.g. metabolic, endocrine, cardiac)
- If starting a non-DTG 3rd agent in the neonatal period it is acceptable to continue this option. However, when over 4 weeks and 3 kg, a switch to DTG is recommended if and when an appropriate formulation is available
- ABC should NOT be prescribed to HLA-B\*57:01 positive individuals (where screening is available). ABC is not licensed under 3 months of age but dosing data for younger children are available from the WHO and DHHS
- At HIV-VL > 100,000 copies/mL ABC + 3TC should not be combined with EFV as 3rd agent
- If using NVP as a 3rd agent in children aged 2 weeks to 3 years, consider using 3 NRTI backbone (ABC + ZDV + 3TC) until VL consistently < 50 copies/mL
- TDF is only licensed from 2 years of age. In view of concerns about potential impact on bone development and renal toxicity TAF is recommended over TDF at all ages in settings where this is licensed and available
- viii DTG is licensed from 4 weeks and 3 kg. DTG has been associated with excessive weight gain in adults, especially in combination with TAF. This has not yet been demonstrated in paediatric and adolescent observational studies or trials, however the possibility of this should be considered when DTG is used. Families and young people should be counselled regarding this and weight should be monitored
- XTC indicates circumstances when FTC or 3TC may be used interchangeably
- TAF is only licensed in Europe for treatment of HIV in combination with FTC from 12 years of age and 35 kg in TAF/FTC and from 6 years of age and 25 kg in TAF/FTC/EVG/c. As TAF is licensed in younger ages and weights it can be included as a preferred option. TAF has been associated with excessive weight gain in adults, especially in combination with DTG. This has not yet been demonstrated in paediatric and adolescent observational studies or trials, however the possibility of this should be considered when TAF is used. Families and young people should be counselled regarding this and weight should be monitored
- BIC is a preferred first line option in adults. At time of writing it is not licensed under 18 years of age but may be considered in those aged less than 18 years following discussion at MDT/PVC
- Due to predicted poor adherence in adolescence, if preferred 3rd agents (BIC or DTG) are not available/appropriate then of the possible alternative 3rd line agents, DRV/b is favoured due to a higher barrier to resistance compared to EFV, RAL or RPV



Table 2. Antiretroviral Formulations Useful for Paediatric and Adolescent Dosing and Administration

NRTI	
ABC	tablet (300 mg) solution (20 mg/mL)
FTC	capsule (200 mg) solution (10 mg/mL)
3TC	tablet (300, 150 mg) solution (10 mg/mL)
TDF	tablet (245, 204, 163, 123 mg) granules (33 mg/g)
ZDV	capsule (250 mg, 100 mg) solution (10 mg/mL) iv infusion: 10 mg/mL (20 mL/vial)
TAF/FTC	tablet (25/200 mg and 10/200 mg)
TDF/FTC	tablet (300/200 mg)
ABC/3TC	tablet (600/300 mg)
ZDV/3TC	tablet (300/150 mg)
NNRTI	
EFV	tablet (600 mg) capsule (200, 100, 50 mg)
NVP	tablet (200 mg) prolonged release tablet (400, 100 mg) suspension (10 mg/mL)
RPV	tablet (25 mg)
TDF/FTC/EFV	tablet (300/200/600 mg)
TAF/FTC/RPV	tablet (25/200/25 mg)
TDF/FTC/RPV	tablet (300/200/25 mg)
Pl	
DRV	tablet (800, 600, 400, 150, 75 mg) solution (100 mg/mL)
DRV/c	tablet (800/150 mg)
LPV/r	tablet (200/50 mg and 100/25 mg) solution (80/20 mg/mL)
RTV	tablet (100 mg) powder for oral suspension (100 mg sachet)
TAF/FTC/DRV/c	tablet (10/200/800/150 mg)
INSTI	
DTG	tablet (50, 25, 10 mg) dispersible tablets (5 mg)
RAL	tablet (600 mg, 400 mg) chewable tablets (100, 25 mg) granules for oral suspension (100 mg)
ABC/3TC/DTG	tablet (600/300/50 mg)
TAF/FTC/BIC	tablet (25/200/50 mg)
TAF/FTC/EVG/c	tablet (10/200/150/150 mg)
TDF/FTC/EVG/c	tablet (300/200/150/150 mg)

#### Switch Strategies for virologically supressed children and adolescents

- · The general indications for switching when virologically suppressed are as for adults, see page 16 but with some additional considerations for children and adolescents relating to increasing age and weight, licensing, formulation availability, vulnerability to toxicity and predicted adherence issues in
- As children age and grow on suppressive ART, consideration should be given to simplification to robust once daily low pill burden regimens with optimal toxicity profiles and efficacy data. For example, in children aged less than 3 years commenced on liquid LPV/r, consider switching to once daily regimens when dispersible DTG is available or pill swallowing is achieved
- If "preferred" options become available for a child as they get older then a switch to this preferred option can be considered. However, if they are fully virologically suppressed on their current regimen with no toxicity or problems with convenience or adherence it is reasonable to remain on an alternative regimen
- Children and their carers should be involved in discussing the relative risk/benefit of switching when well and stable on an effective regimen
- Dual therapy is not recommended in first line or for simplification but can be considered on a case by case basis in adherent children and adolescents
- Simplification to monotherapy and treatment interruptions are not recommended and are discouraged

#### **Special Populations**

- Seek specialist expert advice e.g. through an MDT/PVC. If local MDT or PVC are unavailable, an international PVC is accessible by contacting the
- Adolescent girls of child bearing potential: First line options for adolescents of child bearing potential share the same considerations as discussed elsewhere in the EACS Guideline, see page 18, and should bear in mind contraceptive choices and DDIs with ARVs, see page 38, or whether the young person is trying to conceive
- HBV co-infection: requires an ART regimen that includes TAF or TDF in the NRTI backbone typically with 3TC or FTC, for recommendation in adults with HBV/HIV co-infection see pages 115-116
- HCV co-infection: DAAs against HCV are licensed and available in paediatric formulations down to 3 years of age. Seek specialist advice for consideration of curative HCV therapy for children and adolescents with HCV co-infection, for recommendation in adults with HCV/HIV co-infection see pages 115 and 117-120
- TB co-infection: From 3 years of age DTG bid, EFV (no dose modification), or double dose RAL can all be considered as 3rd agents for children when co-administered with rifampicin. Under 3 years of age, EFV is not recommended. DTG bid can be considered, acknowledging that there is limited data informing this approach in this age range. Super boosted LPV/r can also be considered when paediatric INSTI formulations are not available. Specialist advice should be sought with therapeutic drug monitoring recommended where available. For treatment recommendation in adults with TB/HIV co-infection see page 20



### Adherence, Virological Failure and Second Line ART

- Virological failure (defined as 2 consecutive VL >200 copies/mL at least 3 months apart with adherence support) is almost always due to suboptimal ART adherence, and always requires adherence assessment and support
- Resistance testing is recommended where possible. Choice of second line therapy is dependent both on ALL previous ART exposure and documented cumulative HIV resistance mutations at all times tested
- Second line options should ideally be discussed at a PVC/MDT including a virologist

#### Choosing a 3rd agent

#### Failed on first line NNRTI

- Switch to INSTI with a high barrier to resistance (i.e. DTG or BIC) or PI/b with optimised 2 NRTI
- · If high VL and extensive resistance impacting on NRTIs consider using regimen with at least 2 fully active drugs (e.g. INSTI with PI/b and 2 NRTI)

#### Failed on first line PI/b

- If no significant resistance to PIs, consider continuation of PI/b (consider switch to DRV/b) with optimised 2 NRTI or PI/b based STR to reduce pill burden
- Consider switch to INSTI with high barrier to resistance (i.e. DTG or BIC)
- Consider INSTI or PI based single tablet FDC with 2 NRTI to reduce pill burden (e.g. DRV/c (only in the absence of significant PI resistance), DTG or BIC where/when licensing allows)

#### Failed on first line INSTI

- If resistance testing demonstrates no INSTI resistance, consider switch to/continue INSTI with high barrier to resistance with optimised 2 NRTI
- Switch to PI/b with optimised 2 NRTI is also an option and required if INSTI resistance is demonstrated
- If INSTI resistance and substantial NRTI resistance, consider initial therapy with DTG (bid) + PI/b + optimised 2 NRTI. This should be discussed at MDT/PVC

#### **Optimising NRTI backbone**

- · If resistance testing available use results to guide choice of 2 NRTI
- · If NRTI resistance is demonstrated, XTC with either TAF or TDF are the preferred options, used according to license. If TAF or TDF are not available or contraindicated then ZDV can be considered but alternatives to ZDV should be regularly assessed in order to remove from the regimen as soon as possible due to risk of ZDV toxicity over time
- If resistance testing not available, switch to (or continue) TDF or TAF (or ZDV as per above) with 3TC or FTC (see below rationale)
- TDF or TAF are preferred in second line in combination with 3TC or FTC (even if failing on TDF or TAF)
- It is well established that M184V causes high level resistance to both FTC and 3TC. However ongoing use of either FTC or 3TC is still recommended in the presence of this mutation (especially if it minimises pill burden) as it is associated with an increased susceptibility to tenofovir and ZDV

## Virological Failure on Second Line Combination

- Subsequent virological failure on second line therapy requires further assessment of adherence and resistance testing, if available
- TDM may be useful if concerned about subtherapeutic drug levels
- Choice of subsequent regimens should be made through an MDT/PVC
- ART should continue despite virological failure (with a robust INSTI or PI/b based regimen including 3TC or FTC) to maintain CD4 count whilst additional adherence support is provided



# **References**

# **Video links**

EACS Guidelines	Video lectures	Link to video lecture
ART	When to Start ART Part 1	https://region-hovedstaden-ekstern.23vid- eo.com/secret/69954554/c401f8cf- 3bea2c6bb851a0886f523745
	When to Start ART Part 2	https://region-hovedstaden-ekstern.23vid- eo.com/secret/69954596/4dbab429a86e- ebc401f4c7cb6b66313f
	What ART to Start Part 1	https://region-hovedstaden-ekstern.23vid- eo.com/secret/68809298/066ed5598aa 3f94768fc5fba5b33ad2c
	What ART to Start Part 2	https://region-hovedstaden-ekstern.23vid- eo.com/secret/68809642/82861519cd6bd- cbec65c49924b013b92
Diagnostic Procedures for HCV in Persons with HCV/HIV Co-infection	Hepatitis C and HIV Co-infection Part 1	https://region-hovedsta- den-ekstern.23video.com/ secret/57391741/33aeca1d4a9baa8b- 9f7f408890f19f1f
Diagnosis and Treatment of TB in Persons with HIV	Tuberculosis and HIV Co-infection Part 1	https://region-hovedstaden-ekstern.23vid- eo.com/secret/69954460/7427cd7a76 ac33897ed905a5899278ba
	Tuberculosis and HIV Co-infection Part 2	https://region-hovedstaden-ekstern.23vid- eo.com/secret/69954502/e94125a138e- 7644680c22a589074a371



#### References to All Sections

#### Part I Assessment of Initial & Subsequent Visits

Please see references for Part IV

#### Part II ARV Treatment

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#### **Clinical Management and Treatment of Chronic Viral Hepatitis** Part V Co-infections

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WHO-Rapid communication: Key changes to the treatment of drug-resistant tuberculosis Geneva: World Health Organization; 2022. WHO Reference Number: WHO/UCN/TB/2022.2

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#### Part VII Paediatric HIV Treatment

PENTA Guidelines

https://penta-id.org/hiv/treatment-guidelines

WHO Guidelines

https://www.who.int/publications/i/item/9789240022232

Paediatric use of ABC

https://clinicalinfo.hiv.gov/en/guidelines/pediatric-arv/abacavir

