



ANTIVIRaux ACTIFS SUR LE VIRUS HIV ET PHARMACOTHERAPIE DU SIDA

Enseignant : F. Van Bambeke

FARM2129 – année 2007-2008

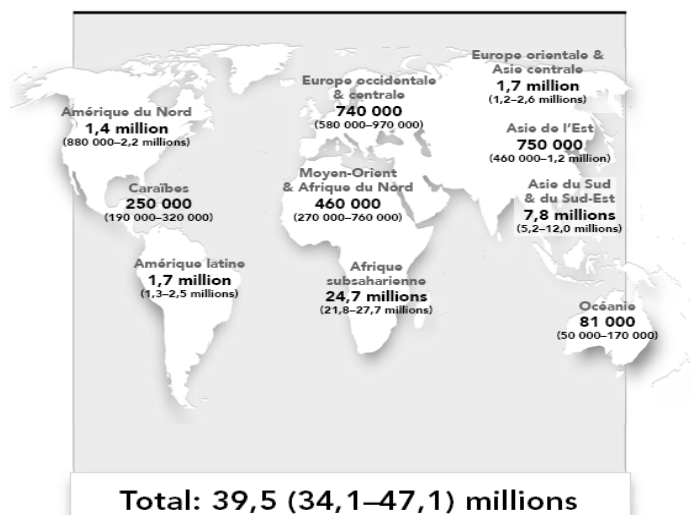
02/05/2008

SIDA

1

Le SIDA: données épidémiologiques

ADULTES ET ENFANTS VIVANT AVEC LE VIH ESTIMATIONS EN 2006



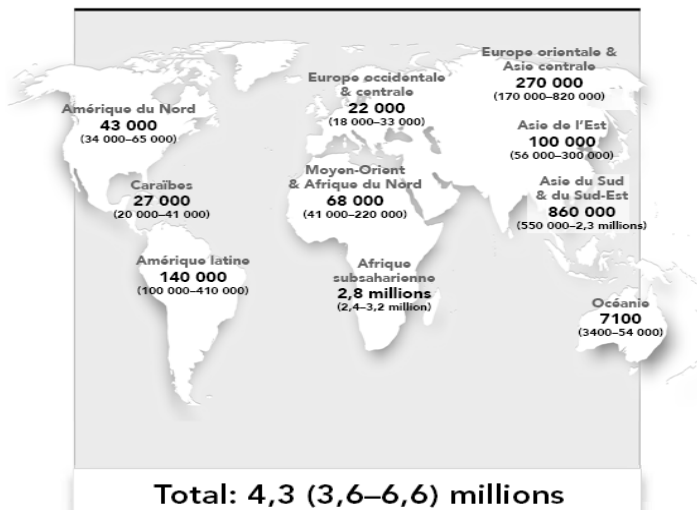
02/05/2008

SIDA

2

Le SIDA: données épidémiologiques

NOMBRE ESTIMATIF D'ADULTES ET D'ENFANTS NOUVELLEMENT INFECTÉS PAR LE VIH EN 2006



02/05/2008

SIDA

3

Le SIDA: mortalité en Afrique (2001)

Rank	% of total
• 1 HIV/AIDS	20.6
• 2 Acute lower respiratory infections	10.3
• 3 Malaria	9.1
• 4 Diarrhoeal diseases	7.3
• 5 Perinatal conditions	5.9
• 6 Measles	4.9
• 7 Tuberculosis	3.4
• 8 Cerebrovascular disease	3.2
• 9 Ischaemic heart disease	3.0
• 10 Maternal conditions	2.4

The World Health Report 2000, WHO

02/05/2008

SIDA

4

Le SIDA: données épidémiologiques



BELGIUM

I. DEMOGRAPHIC, SOCIAL AND ECONOMIC INDICATORS

Estimated Population	10 419 000
Population Growth Rate	0.2%
Life expectancy at birth	
Women	75
Men	75
Human Development Index	9
Human Poverty Index	
Rank	13 ¹
Value	12.4 ²
Percentage of people with less than US\$ 2 a day	-
Per Capita Gross National Income, ppp, Int'l dollar rate	31 360
Per Capita Government Expenditure on Health at Int'l dollar rate	1902

II. HIV AND AIDS ESTIMATES

Number of people living with HIV	14 000 [8100 – 22 000]
Adults aged 15 to 49 HIV prevalence rate	0.3 [0.2 – 0.5%]
Adults aged 15 and over living with HIV	14 000 [8100 – 22 000]
Women aged 15 and over living with HIV	5400 [2800 – 9500]
Deaths due to AIDS	<100 [<200]

02/05/2008

SIDA

5

Le SIDA: voies de transmission

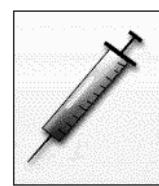


Unprotected
sexual intercourse
with an infected partner



Vertical
transmission
(from mother
to child)

- in utero
- during delivery
- breastmilk



Injection drug use
(rare: infected
blood/blood products)

HIV INFECTION

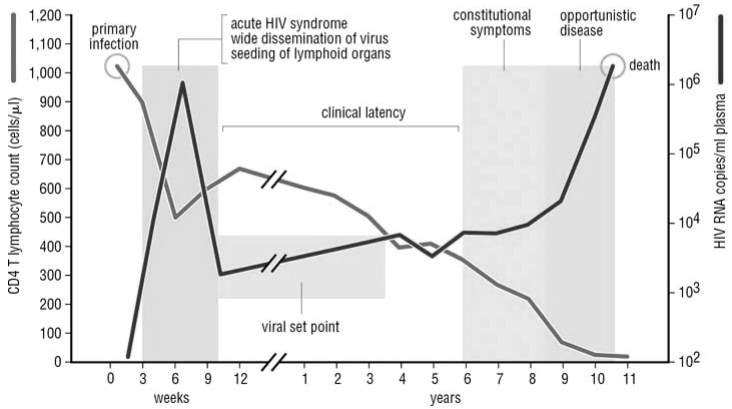
02/05/2008

SIDA

6

L'infection à HIV: histoire naturelle

From **Immunity: The Immune Response in Infectious and Inflammatory Disease**
by DeFranco, Locksley and Robertson

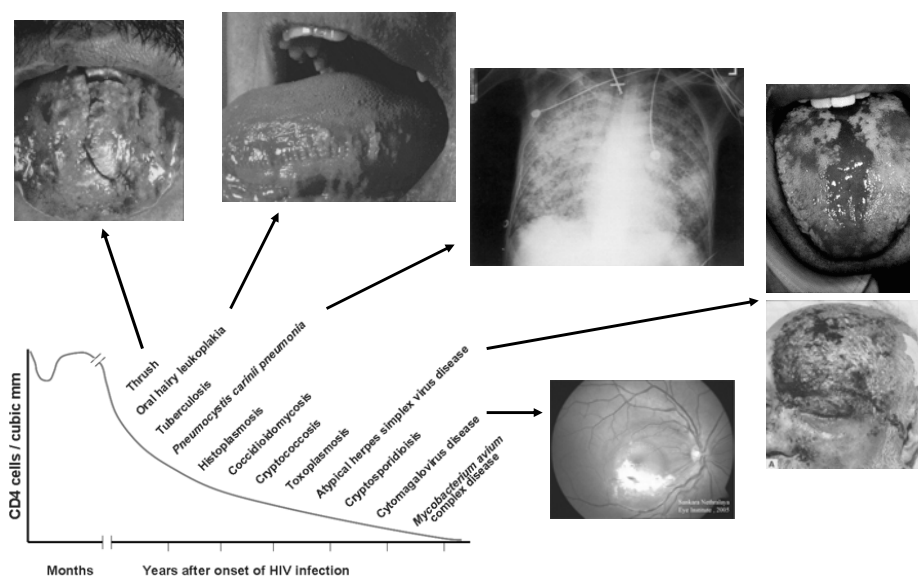


02/05/2008

SIDA

7

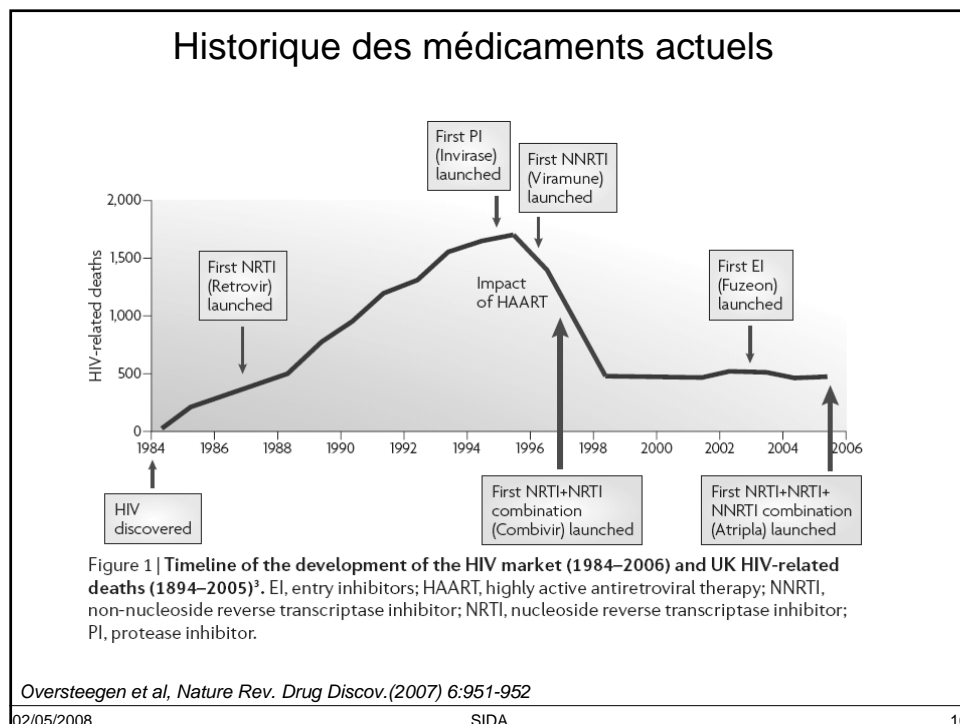
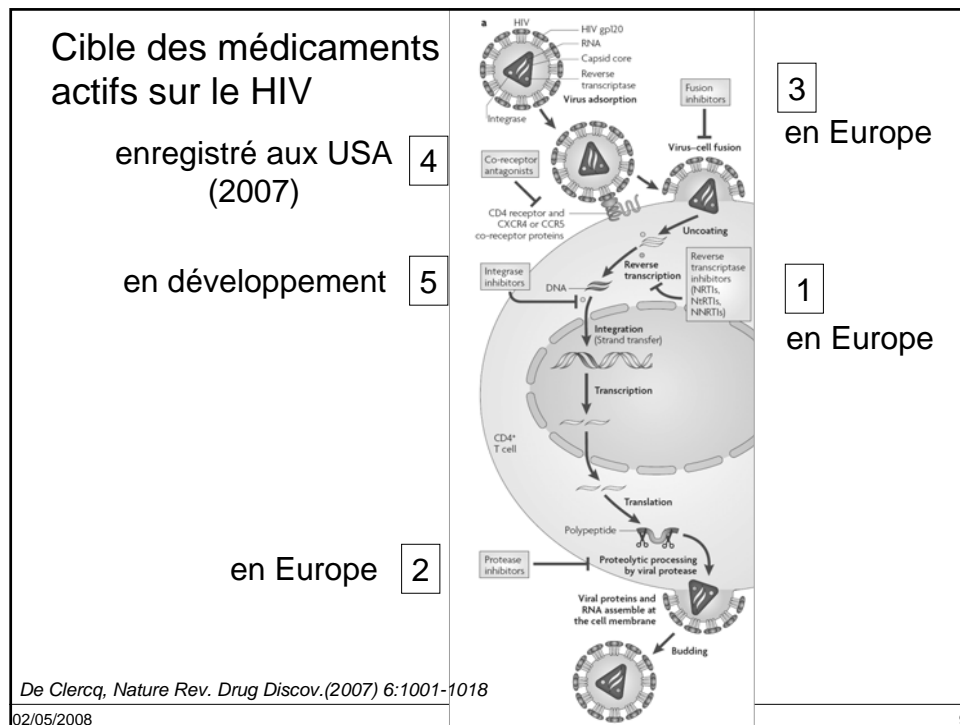
L'infection à HIV: infections opportunistes



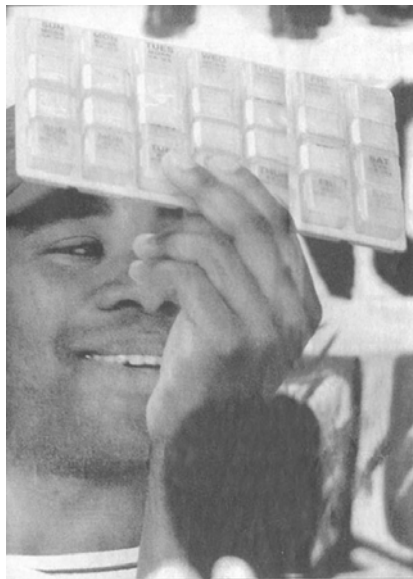
02/05/2008

SIDA

8



HAART : Highly Active Anti-Retroviral Therapy



02/05/2008

'Aids drugs made me well again'

LYNN ALLENBROOK AND JO-ANNE SMITHHAM

DOCTORS gave Matthew Damane, 25, five days to live after he was diagnosed with HIV, the virus that causes AIDS, in 1996.

At that time, life-saving AIDS medicines, widely available in the West, were not available for poor people in countries like South Africa.

Now he takes a single tablet, which cost R1 400 a month, even with discounts offered by drug companies, are still too expensive for him.

But Damane, 25, from Khayelitsha, has had access to his medicine since it was imported from Brazil, and he credits the drugs with restoring him to health.

"I am now well," he told a packed news conference in Cape Town yesterday. He held up a plastic pill box. It has one pill compartment for each day of the week, helping him take his AIDS medicines on schedule.

Damane, a nervous smile showing under his blue baseball cap, was one of many

activist groups announced it had imported the medicines from Brazil in violation of drug company patent rights but with the full support of the Medicines Control Council (MCC).

Citing preliminary results from a study in Khayelitsha, the activists said the AIDS drugs had reduced the presence of the virus in patients' blood streams to undetectable levels after less than one year of treatment.

"We are seeing people getting off their deathbeds and returning to productive work and living lives again."

"We literally resuscitated people," said Eric Goemans, who heads the Aids clinic run by Médecins Sans Frontières (MSF) in Khayelitsha.

The preliminary results of the study, which has reported findings for 85 patients taking the AIDS drugs, show that the prevalence of HIV in the community has dropped from a township clinic in South Africa that the AIDS drugs have had a dramatic long-term benefit and can have the same dramatic effect in improving health as they have had in industrialized countries.

ment Action Campaign (TAC), Oxfam and Cossat pointed to the findings yesterday to urge the government to set up pilot projects to import generic AIDS drugs to symptomatic AIDS patients in each province. They also referred to the findings to support their argument that the government should follow Brazil's lead and import generic and cost generic versions of the drugs.

"It is difficult, but it is feasible to do this without any additional cost," said Jackie Heywood, TAC secretary.

The government did not comment on the activists' calls. It said the MCC would check whether the legal import was legal.

The drug companies that own the patent rights to the AIDS drugs have refused to sue the activists. Peter Moore, medical director at GlaxoSmithKline, said the company would wait for the MCC to act.

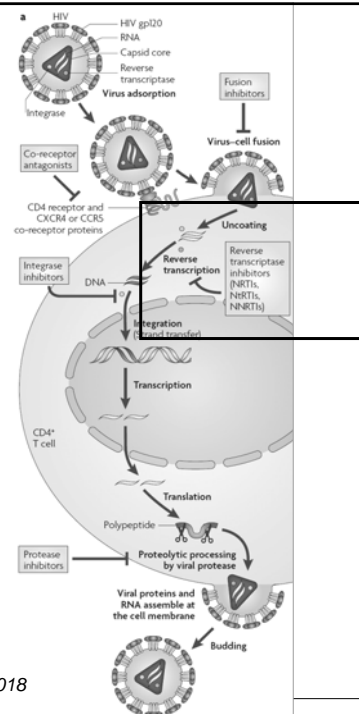
Boehringer-Ingelheim spokesman Michael Doherty said he was not surprised at the developments.

"I don't think we're falling off our chairs at the moment."

SIDA

11

Cible des médicaments actifs sur le HIV

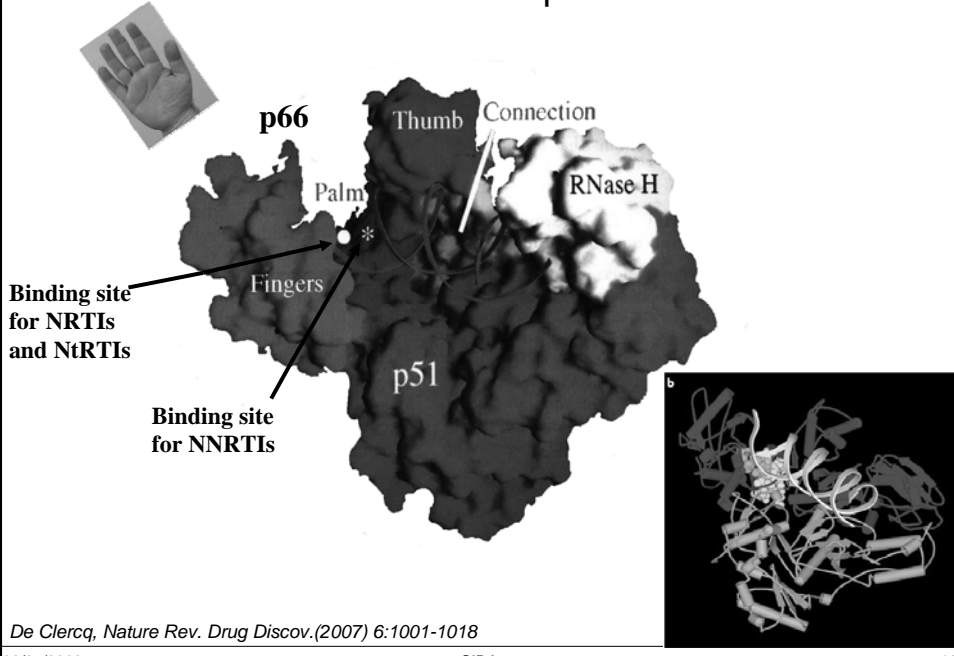


De Clercq, Nature Rev. Drug Discov. (2007) 6:1001-1018

02/05/2008

12

Inhibiteurs de transcriptase inverse

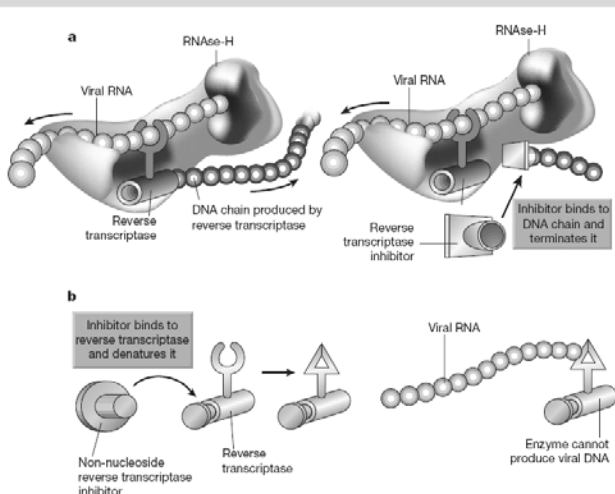


Inhibiteurs de transcriptase inverse

Figure 3 Mechanism of action of nucleoside and non-nucleoside reverse-transcriptase inhibitors. To enable HIV to be integrated into the host DNA and so use the cell's genetic machinery to make new virus, the single-stranded viral RNA must first be converted to double-stranded DNA by the viral enzyme reverse transcriptase, while the enzyme RNase-H hydrolyses the RNA after it has been copied. Nucleoside and non-nucleoside reverse-transcriptase inhibitors are two classes of antiretroviral drugs that suppress HIV replication by attacking reverse transcriptase.

a. Nucleoside reverse-transcriptase inhibitors are similar in structure to the building blocks that make up DNA. By incorporating themselves into the DNA nucleoside chain being produced by reverse transcriptase, they stop attachment of further nucleosides and so prevent ongoing viral DNA synthesis.

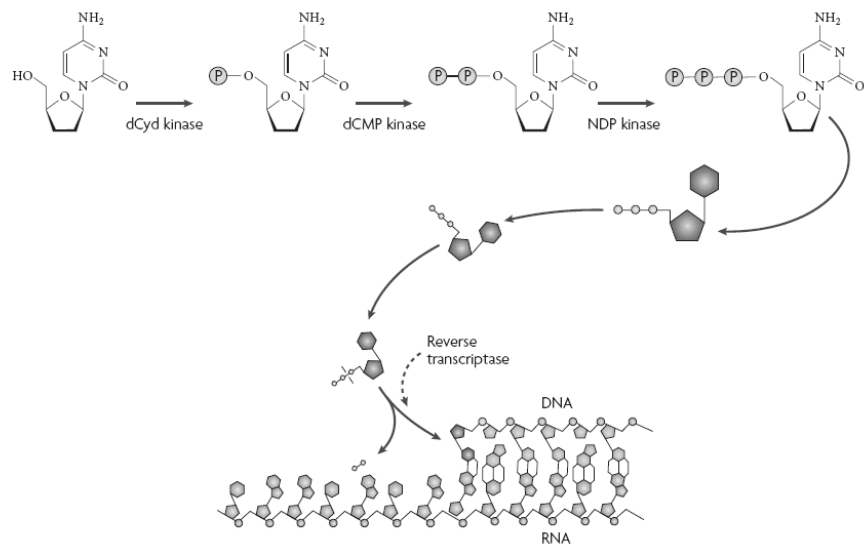
b. Non-nucleoside reverse transcriptase inhibitors attach to the reverse transcriptase and affect the activity of the enzyme by restricting its mobility and making it unable to function. (Adapted from ref. 108 with permission.)



Richman, Nature (2001) 410:995-1001

02/05/2008 SIDA 14

Mode d'action des inhibiteurs de transcriptase inverse analogues nucléosidiques/nucléotidiques



De Clercq, Nature Rev. Drug Discov. (2007) 6:1001-1018

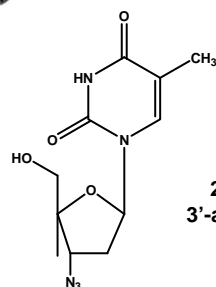
02/05/2008

SIDA

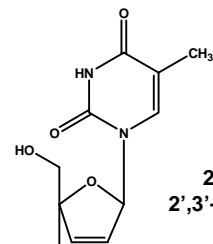
15

analogues nucléosidiques

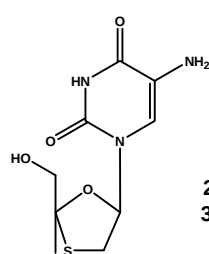
Analogues des bases pyrimidiques



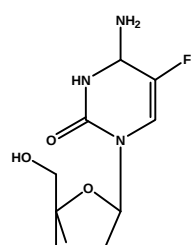
**2',3'-Dideoxy-
3'-azidothymidine
(AZT)
zidovudine**



**2',3'-Dihydro-
2',3'-dideoxythymidine
(D4T)
stavudine**



**2',3'-Dideoxy-
3'-thiacytidine
(3TC)
lamivudine**



**(FTC)
emtricitabine**

02/05/2008

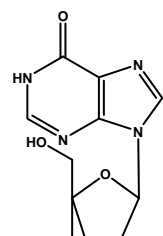
SIDA

16

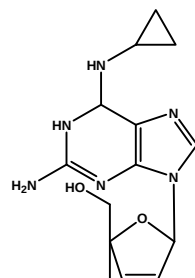


analogues nucléosidiques

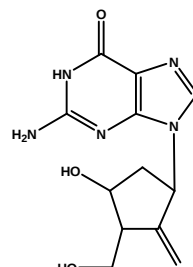
Analogues des bases puriques



2',3'-Dideoxy-inosine
(DDI)
didanosine



2',3'-Dideoxy-inosine
(ABC)
abacavir



entecavir

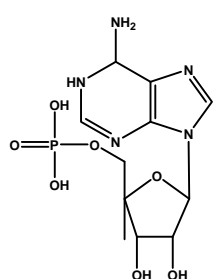
02/05/2008

SIDA

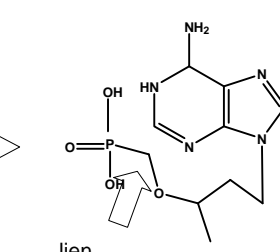
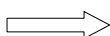
17



analogues nucléotidiques: tenofovir



adenosine-mono-P



lien
phosphonate
stable

tenofovir

02/05/2008

SIDA

18

Pharmacocinétique



- bonne biodisponibilité orale
 - didanosine : résorption réduite par l'acidité gastrique ou la prise de nourriture.
- distribution dans les liquides de l'organisme, y compris le LCR
- $T_{1/2}$ plasmatique courte mais fréquence d'administration dictée par $T_{1/2}$ cellulaire des formes triphosphorylées

Agent	Biodisponibilité orale (%)	$t_{1/2}$ sérique (h)	$t_{1/2}$ des formes triphosphate (h)	Voies d'élimination	Principal dosage (adulte)
Zidovudine	63	1.1	3-4	glucurono-conjugaison et élimination rénale	300 mg / 12 h
Didanosine	40 (à jeûn) 86	1.5 1.1	8-24 3	métabolisme cellulaire excrétion rénale	400 mg / 24 h 40 mg / 12 h
Stavudine	86	2.5	11-14	excrétion rénale	300 mg / 24 h
Lamivudine	86	1.5	3.3	glucurono-conjugaison et carboxylation	300 mg / 24 h
Abacavir	83	12-14	>12 *	excrétion rénale	300 mg / 24 h
Tenofovir	39 (avec un repas)	10	>24		200 mg / 24 h
Emtricitabine	93				

02/05/2008

SIDA

19

Combinaisons et compliance



Table 1 | Overview of currently launched fixed-dose combination products for the treatment of HIV*

Drug	Class	Pill volume	Total pills per day	Dosing schedule	Combination product	Total pills per day	Dosing schedule	2006 sales [‡]
Tenofovir	NRTI	300 mg	1	Once daily	Truvada	1	Once daily	1.125
Emtricitabine	NRTI	200 mg	1	Once daily	Atripla	1	Once daily	174
Efavirenz	NNRTI	600 mg	1	Once daily		-	-	-
Abacavir	NRTI	300 mg	2	Once daily	Epzicom	1	Once daily	396
Lamivudine	NRTI	300 mg	1	Once daily	Trizivir	2	Twice daily	478
Zidovudine	NRTI	300 mg	2	Twice daily	Combivir	2	Twice daily	789

Oversteegen et al, Nature Rev. Drug Discov. (2007) 6:951-952

02/05/2008

SIDA

20

Effets secondaires



Communs à la classe

- hyperlactacidémie (menant parfois à une acidose sévère)
- hépatomégalie et stéatose
(inhibition de la DNA-polymérase impliquée dans la réPLICATION du DNA mitochondrial (surtout pour didanosine, stavudine, et zidovudine)).

Particuliers à certaines molécules

molécule	Effet secondaire
zidovudine	Anémie neutropénique
didanosine	pancréatite, neuropathie périphérique
stavudine	neuropathie périphérique
abacavir	réactions d'hypersensibilité
tenofovir	toxicité rénale à long terme
emtricitabine	hyperpigmentation des mains et pieds

02/05/2008

SIDA

21

Interactions médicamenteuses



- excipient de la didanosine (sels de Mg²⁺ et d'Al³⁺) :
↓ absorption de nb médicaments:
(kétoconazole, dapsone, tétracyclines, fluoroquinolones)
- ranitidine: ↓ faible de l'absorption de la didanosine
- pentamidine : ↑ toxicité pancréatique (didanosine, stavudine et zalcitabine)
- probénécide, pyréméthamine/ sulfadiazine :
↓ glucuronoconjuguaison ou élimination rénale de l'azidothymidine
↑ sa toxicité
- ganciclovir (et autres médicaments myélotoxiques)
↑ risque de myélosuppression de l'azidothymidine

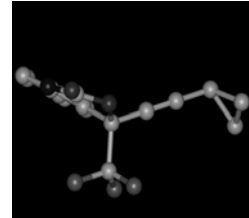
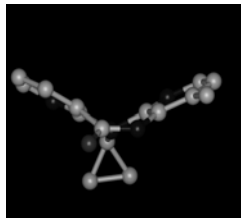
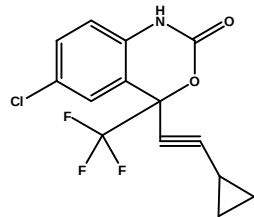
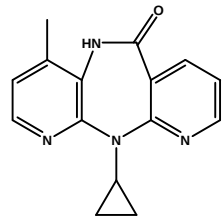
02/05/2008

SIDA

22



analogues non nucléosidiques



Inhibiteurs allostériques non compétitifs;
pas de résistance croisée avec les NRTI !



Pharmacocinétique

névirapine

- bonne résorption orale
- élimination par métabolisation hépatique; inducteur de son propre métabolisme
→ $t_{1/2} = 45$ h après une dose unique
= 25 h après administration répétée
→ augmentation des posologies après 15 jours de traitement

efavirenz

- forte liaison aux protéines et demi-vie prolongée (40 h)
→ administration 1X/jour
- inducteur et inhibiteur des cytochromes P450 (3A4 et 2B6), n'entraînant pas de modification importante de son propre métabolisme.

Effets secondaires



névirapine

- réactions cutanées fréquentes, parfois mortelles (syndrome de Stevens Johnson; nécrolyse cutanée).

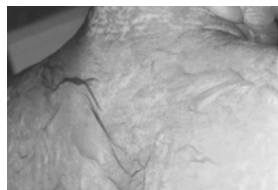


Figure 1. Typical Pattern of Toxic Epidermal Necrolysis.
Blisters and wrinkled areas result from full-thickness necrosis of the epidermis.

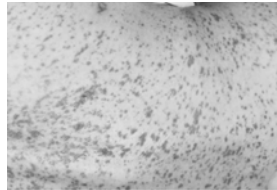


Figure 2. Typical Pattern of Stevens-Johnson Syndrome.
Blisters develop on widespread purpuric macules.

- Interrompre le traitement dès l'apparition de signes précurseurs (rash cutané, fièvre, lésions orales, conjonctivite, douleurs musculaires ou articulaires, malaise généralisé).
- toxicité hépatique (possibilité d'hépatites fulminantes).
- agranulocytose chez les enfants
- nausées, fièvre, maux de tête.

02/05/2008

SIDA

25

Effets secondaires



Efavirenz

- effets sur le système nerveux : étourdissements, vertiges, somnolence, maux de tête, dépression
- administration le soir
- rashes (ne demandent que rarement l'arrêt du traitement).

02/05/2008

SIDA

26

Interactions médicamenteuses



Inducteurs/inhibiteurs des CYP

Névirapine:

↓ taux sérique de rifabutine, rifampicine
kétoconazole
anticoagulants oraux

Efavirenz:

↓ taux sérique de inhibiteurs de la protéase du virus HIV
méthadone
rifabutine, clarithromycine.

↑ taux sérique de ritonavir

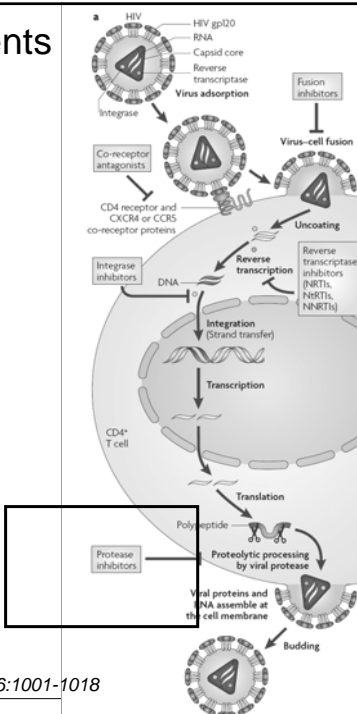
**Patients susceptibles
de développer
des infections
opportunistes !**

02/05/2008

SIDA

27

Cible des médicaments actifs sur le HIV



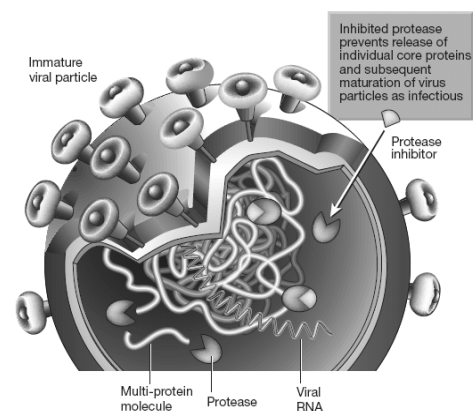
De Clercq, Nature Rev. Drug Discov. (2007) 6:1001-1018

02/05/2008

28

Protéase et inhibiteurs

Figure 4 Mechanism of action of protease inhibitors. After transcription in the nucleus, viral mRNA enters the cytoplasm and uses the host's cellular machinery to manufacture virus proteins. The viral components then gather at the cell membrane and immature viruses bud off the cell. Core proteins are produced as part of long polypeptides, which must be cut into smaller fragments by the enzyme protease in order to form mature, functional proteins. Protease inhibitors bind to the site where protein cutting occurs, and so prevent the enzyme from releasing the individual core proteins. In this way the new viral particles are unable to mature or become infectious. (Adapted from ref. 108 with permission.)



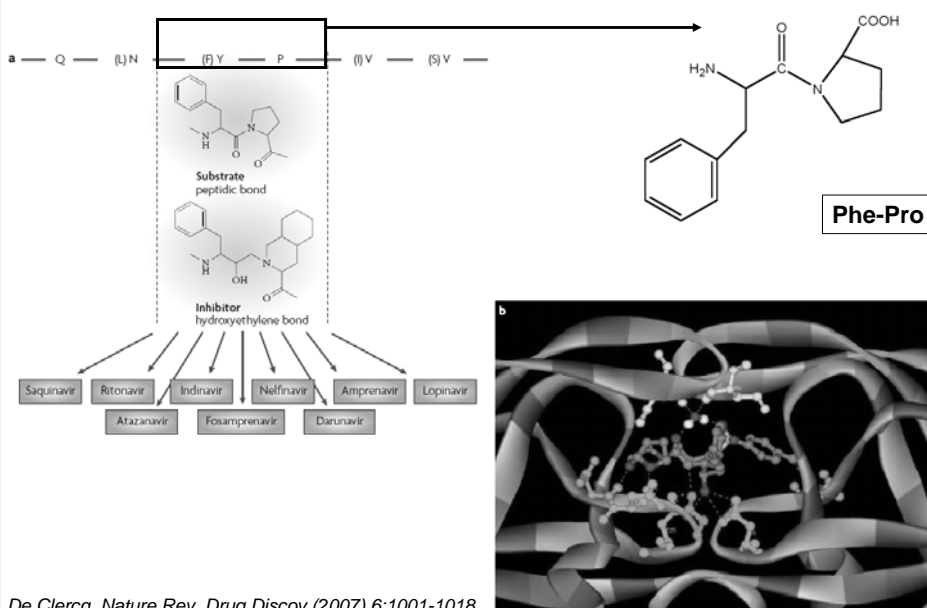
Richman, *Nature* (2001) 410:995-1001

02/05/2008

SIDA

29

Inhibiteurs de protéase HIV



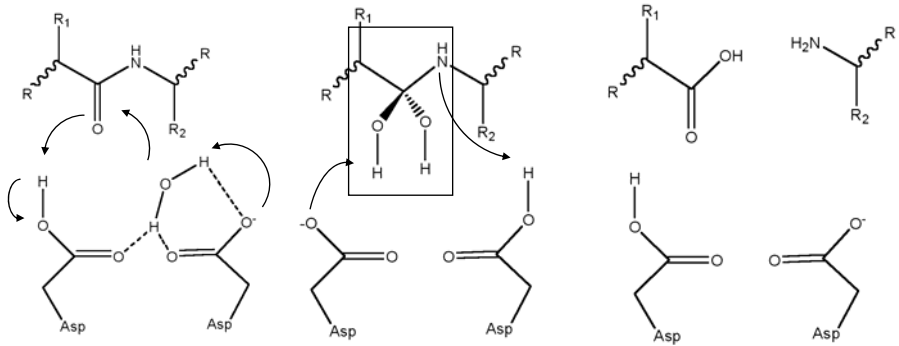
De Clercq, *Nature Rev. Drug Discov.* (2007) 6:1001-1018

02/05/2008

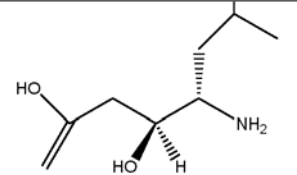
SIDA

30

La protéase HIV, une Aspartate- protease



Inhibiteur-type:
pepstatine...

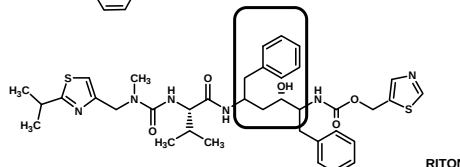
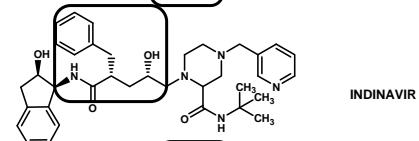
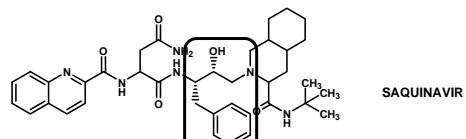
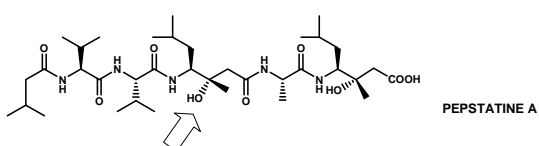


02/05/2008

SIDA

31

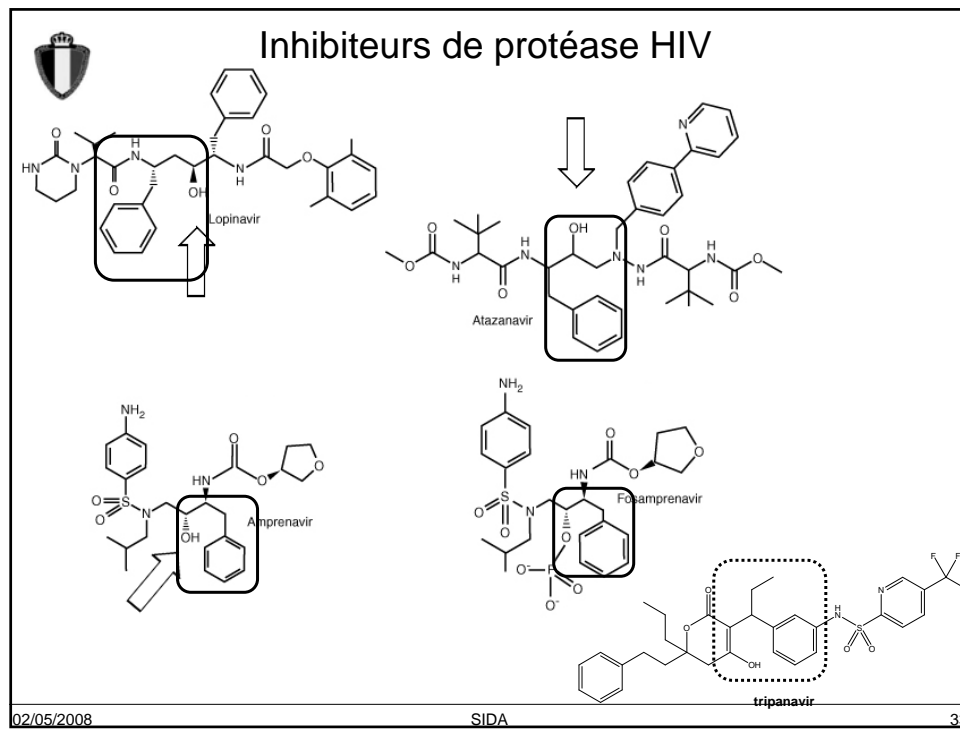
Inhibiteurs de protéase HIV



02/05/2008

SIDA

32



02/05/2008

SIDA

33

Résistance par mutation

MUTATIONS IN THE HIV PROTEASE GENE ASSOCIATED WITH REDUCED SUSCEPTIBILITY TO PROTEASE INHIBITORS (PIs)

Certaines mutations confèrent des résistances croisées !

http://www.iasusa.org/resistance_mutations/index.html

02/05/2008

SIDA

34

Pharmacocinétique



- faible biodisponibilité
(poids moléculaire élevé, mauvaise solubilité et instabilité)
- $T_{1/2}$ courte (quelques heures)
→ administrations 2 ou 3 X/jour
- métabolisation par les cytochrome P-450 hépatiques (principalement 3A4).
→ inhibiteurs ou activateurs du métabolisme de nb médicaments.

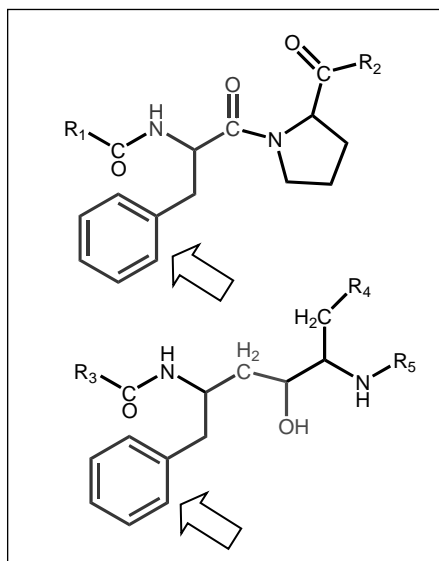
02/05/2008

SIDA

35

Inhibiteurs de protéase HIV et cytochromes

- la protéase doit scinder un lien Phe-Pro
- Les inhibiteurs miment donc tous une Phe...



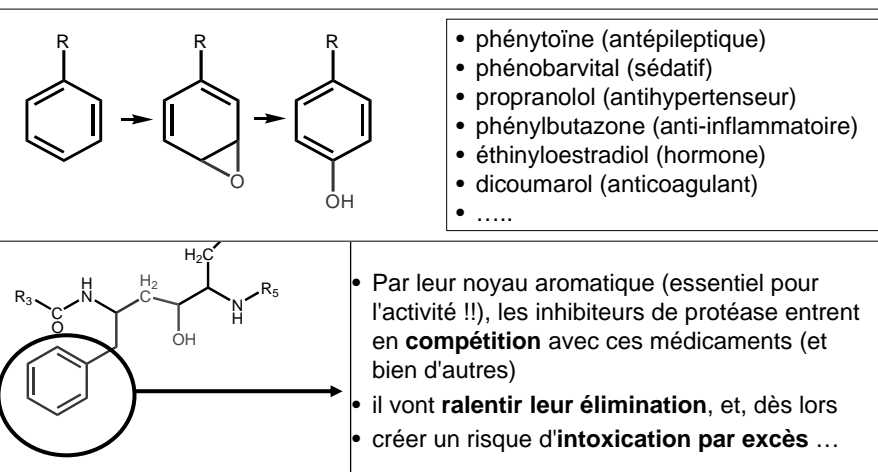
02/05/2008

SIDA

36

Inhibiteurs de protéase HIV et cytochromes

- La plupart des médicaments (et autres substances) à noyau aromatique sont métabolisées en dérivés hydroxylés, ce qui est essentiel pour leur élimination



02/05/2008

SIDA

37

Pharmacocinétique

- faible biodisponibilité
(poids moléculaire élevé, mauvaise solubilité et instabilité)
- $T_{1/2}$ courte (quelques heures)
→ administrations 2 ou 3 X/Jour
- métabolisation par les cytochrome P-450 hépatiques (principalement 3A4).
→ inhibiteurs ou activateurs du métabolisme de nb médicaments.

Très important pour le ritonavir; utilisé à faible dose comme inhibiteur du métabolisme des autres inhibiteurs de protéase.

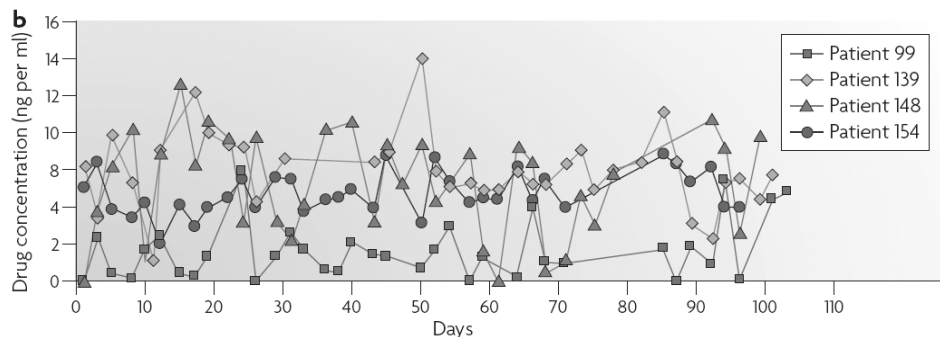
→ 200 mg lopinavir + 50 mg ritonavir (KALETRA®).
[tripranavir + ritonavir]

02/05/2008

SIDA

38

Lopinavir: variabilité pharmacocinétique



Measured intra-subject variability in concentrations of lopinavir dosed 400 mg every 12 hours. Participants had undetectable plasma HIV RNA on treatment for at least 3 months, and were seen in the clinic 3 times a week for up to 4 months. Blood for lopinavir concentration analysis was collected at approximately the same time of day at each visit

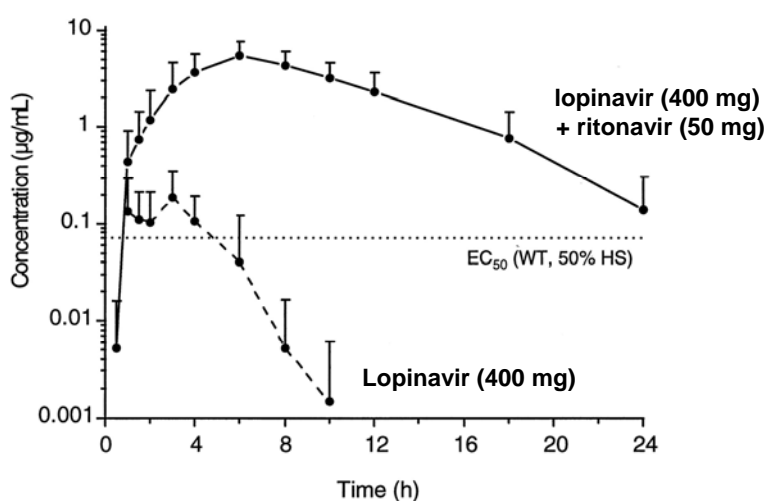
Flexner, *Nature Rev. Drug Discov.* (2007) 6:959-66

02/05/2008

SIDA

39

Lopinavir: influence du ritonavir sur le profil PK



Sham et al, AAC (1998) 42:3218-24.

02/05/2008

SIDA

40



Effets secondaires

"syndrome de lipodystrophie"

- bajoues et dépôts graisseux sur la face
- dépôts de graisse au niveau du cou ["bosse de bison"] et du tronc
- accumulation de graisse derrière les muscles abdominaux
- lipomes disséminés
- hyperplasie graisseuse des seins
- hyperglycémie, hyperinsulinémie
- augmentation des taux lipides sériques



risque de diabète non insulino-dépendant et de maladie cardiovasculaire.

02/05/2008

SIDA

41



Effets secondaires

Selon la molécule (orientation du choix!)

saquinavir	intolérance gastro-intestinale, diarrhée maux de tête ↗ taux sériques de triglycérides et de cholestérol
ritonavir:	intolérance digestive et diarrhée très importantes paresthesies ↗ transaminases hépatiques disgueusie
indinavir:	intolérance gastro-intestinale, diarrhée hyperbilirubinémie non conjuguée asymptomatique néphrolithiasies (peuvent être prévenues par hydratation) ↗ transaminases maux de tête, insomnie
nelfinavir	diarrhée et flatulence fréquentes altération de la formule sanguine ↗ transaminases

02/05/2008

SIDA

42

Effets secondaires



Selon la molécule (orientation du choix!)

amprenavir	maux de tête nausée et diarrhée fréquente rash
lopinavir	diarrhée et nausées ↗ importante des taux sériques de triglycérides et de cholestérol
tripanavir	diarrhées et nausées céphalées hépatotoxicité saignements éruptions cutanées
azatanavir	diarrhée et nausées hyperbilirubinémie non conjuguée asymptomatique

02/05/2008

SIDA

43

Interactions médicamenteuses



- inhibition des cytochromes : ritonavir > indinavir et nelfinavir > saquinavir
→ nb risques d'interactions à surveiller !
- modification des taux d'IP par d'autres médicaments
 - clarithromycine ↗ taux sérique du ritonavir et de l'indinavir
 - fluconazole ↗ taux sérique du ritonavir
 - kétoconazole ↗ ↗ taux sérique de saquinavir, d'indinavir et de nelfinavir
 - quinidine ↗ taux sérique de l'indinavir
 - rifampicine ↘ ↘ le taux sérique de saquinavir (et nelfinavir et ritonavir)
 - névirapine ↘ concentration-pic du saquinavir
- boissons acides ↘ taux sérique de l'indinavir et du nelfinavir
- substrats de P-glycoprotéine et inhibiteurs de MRP2 :
modulation de la pharmacocinétique et interaction avec d'autres médicaments

02/05/2008

SIDA

44

Interactions médicamenteuses



Interactions médicamenteuses importantes ou très dangereuses (! = contre indication) des inhibiteurs de protéase anti HIV (à l'exclusion des interactions entre anti-HIV).

Medicaments (classe)	Indinavir	Ritonavir	Saquinavir	Nelfinavir	Amprenavir ¹	Lopinavir (assoc. au ritonavir)	Azatanavir	Tripanavir (assoc. au ritonavir)
Antibiotiques		clarithromycine rifabutine			clarithromycine métronidazole (!) rifabutine	clarithromycine rifabutine	rifampicine (!)	rifampicine rifabutine clarithromycine
Analgesiques		opiacés mégépéridine (!) propoxyphène (!) poxicam (!)						opiacés et méthadone/ mégépéridine
Dérivés de l'ergot	tous (!)			tous (!)	tous (!)	tous (!)	tous (!)	
Antiarythmiques	tous (!)		quinidine		amiodarone quinidine		Amiodarone, bepridil, lidocaïne (systémique), et quinidine.	
Cardiotoniques		digoxine						
Anticoagulants		coumariniques					coumariniques	
Anticonvulsivants	carbamazépine phén妥toïne phénobarbital	tous	carbamazépine phén妥toïne phénobarbital	carbamazépine phén妥toïne phénobarbital		Carbamazépine, phénobarbital, phén妥toïne		
Antidépresseur		tous buspironne (!)						millepertuis desipramine
Antihistaminiques	terfénadine (!) astmineole (!) autres molécules (!)	terfénadine (!) astmineole (!) autres molécules (!)	terfénadine (!) astmineole (!) autres molécules (!)	terfénadine (!) astmineole (!) autres molécules (!)				
Antifongiques	kétoconazole	kétoconazole itraconazole			kétoconazole	Kétoconazole, itraconazole.		voriconazole (imprédictible)
Anticancéreux		étoposide alcaloïdes vinca tamoxifène						vinorelbane
Autres agents cardiovascul.	la plupart bepridil (!)	antagon. Ca ⁺⁺	antagon. Ca ⁺⁺		antagon. Ca ⁺⁺			

02/05/2008

SIDA

45

Interactions médicamenteuses



Voies de transmission

Table 135-2. Drug interactions between antiretrovirals and oral contraceptives. Recommended adjustments are listed. Data from CDC.²¹

Agent	Effect on oral contraceptive	Recommendation		
		No dose adjustment	No data	Use alternative agent or second method
Indinavir	Norethindrone ^R levels ↓26% ethynodiol levels ↓24%	X		
Ritonavir ^R	Ethinylestradiol levels ↓40%			X
Saquinavir ^R		X		
Nelfinavir	Norethindrone ^R levels ↓18% ethynodiol levels ↓47%			X
Amprenavir ^R	Potential for interaction	X		X
Lopinavir	Ethinylestradiol levels ↓42%			X
Nevirapine ^R	Ethinylestradiol levels ↓20%			X
Delavirdine		X		
Efavirenz ^R	Ethinylestradiol levels ↑37% no data on norethindrone ^R levels			X

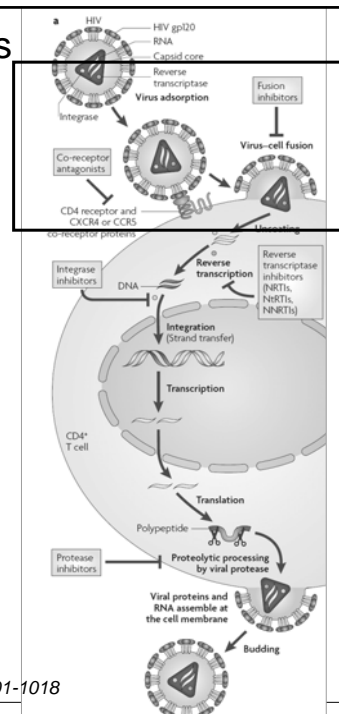
02/05/2008

SIDA

46

Cible des médicaments actifs sur le HIV

Inhibiteurs d'entrée



De Clercq, Nature Rev. Drug Discov. (2007) 6:1001-1018

02/05/2008

47

Récepteurs cellulaires au virus HIV

	CD4	CCR5	CXCR4
Structure	Four Ig-like domains	Seven transmembrane domains G-protein coupled receptor	Seven transmembrane domains G-protein coupled receptor
Function	Coreceptor for MHC class II during stimulation of T-helper cells	Receptor for CCL3 (MIP1- α) CCL4 (MIP- β) CCL5 (RANTES) Redundant system	Receptor for CXCL12 (SDF-1) Non-redundant system
Expression	CD4+ T cells Macrophages Microglia Dendritic cells	A subset of memory CD4+ cells Macrophages	Constitutive in many cell types, including CD4+ T cells and macrophages

Figure 2: Receptors for HIV-1 entry

Este & Telenti, Lancet (2007) 370:81-88

02/05/2008

SIDA

48

Fusion du virus avec la cellule hôte

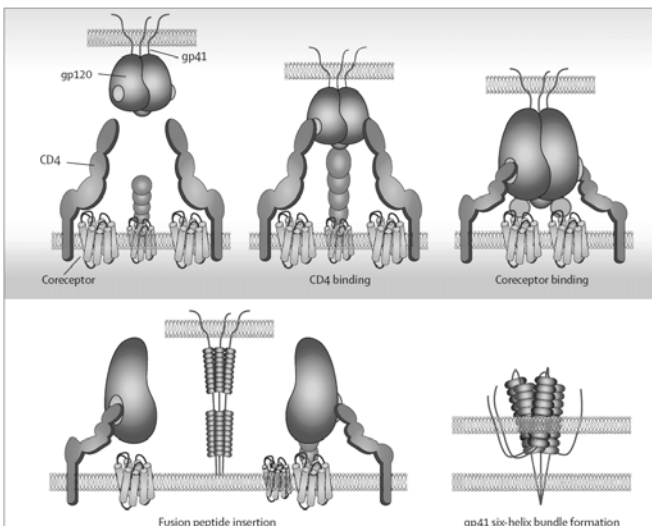


Figure 1: Mechanism of HIV entry
On CD4 binding (binding site for CD4 is shown in yellow), gp120 undergoes conformational changes. CD4-induced epitopes can then bind to chemokine receptors. Thereafter, gp41 is released or into a fusogenic conformation and its N-terminal (green) and C-terminal (red) helices form a hairpin structure, leading to the approximation of viral and cellular membranes, which results in membrane fusion.

Este & Telenti, Lancet (2007) 370:81-88

02/05/2008

SIDA

49

Inhibiteurs de fusion

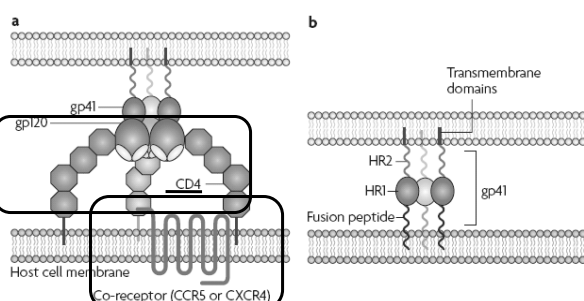
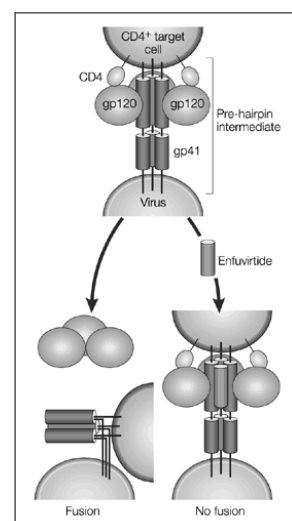


Figure 8 | Inhibiting human immunodeficiency virus (HIV) fusion. When HIV infects a CD4+ T cell (a), the viral glycoprotein gp120 first interacts with the CD4 receptor, then with the CCR5 or CXCR4 co-receptor, upon which the viral gp41 will bring the viral envelope in contact with the host cell membrane (b). The gp41 glycoprotein contains four major functional domains: starting from the N terminus towards the C terminus these are the fusion peptide, the heptad repeat 1 (HR1), the heptad repeat 2 (HR2) and the transmembrane domain that anchors gp41 into the viral lipid bilayer. Enfuvirtide is homologous to part of the HR2 region. When the N terminal fusion peptide of gp41 is inserted into the host cell membrane, the three HR2 domains of the gp41 trimer loop back in a triple hairpin and 'zip' themselves into three highly conserved hydrophobic grooves on the outer face of the HR1 trimeric bundle to form a six-helix bundle that pulls the outer membranes of the virus and the cell into close physical proximity, thus enabling the two membranes to fuse¹³. This process depends on an interaction of the heptad repeat HR2 with HR1. By being homologous to the HR2 domain, enfuvirtide blocks this interaction⁹⁰.



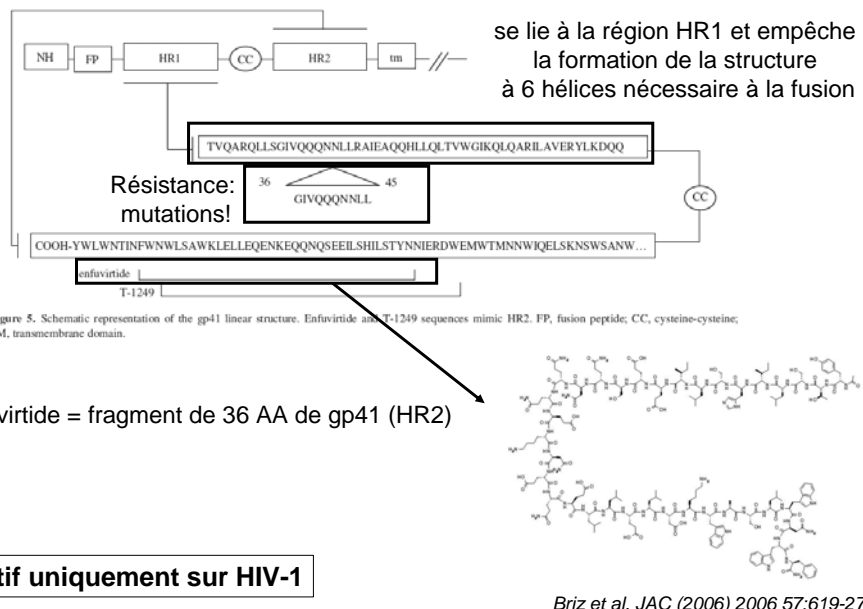
La Bonne et al (2003) & De Clercq (2007), Nature Rev. Drug Discov. 2: 345-346 & 6:1001-1018

02/05/2008

SIDA

50

Inhibiteurs de fusion

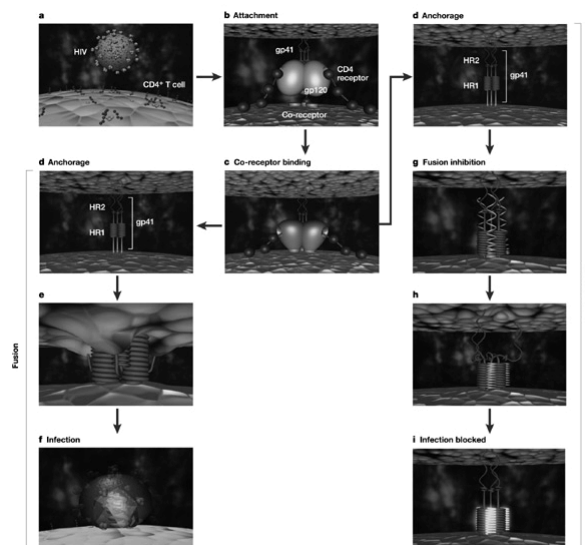


02/05/2008

SIDA

51

Inhibiteurs de fusion



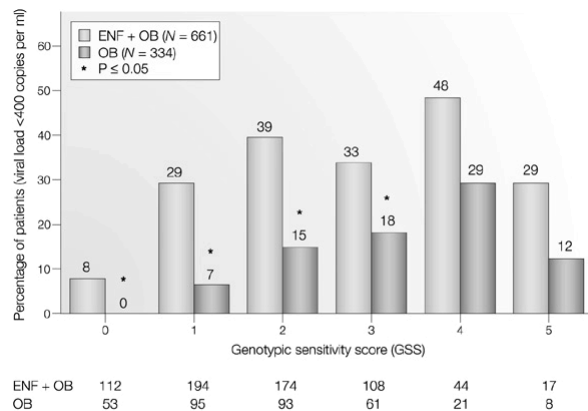
Nature Reviews | Drug Discovery
Matthews et al, Nature Rev. Drug Discov. (2004) 3:1215-25

02/05/2008

SIDA

52

Inhibiteurs de fusion: efficacité clinique



Percent responders with HIV-1 RNA <400 copies per ml, week 48 (intent-to-treat, discontinuation or virological failure = failure).

GSS is the actual number of antiretrovirals that the baseline virus is sensitive to as indicated by standard primary mutations that each virus possesses.

ENF, enfuvirtide; OB, optimized treatment background.

Matthews et al, Nature Rev. Drug Discov. (2004) 3:1215-25

02/05/2008

SIDA

53

Infuvirtide: propriétés pharmacologiques



Pharmacocinétique

- médicament peptidique:
 - administration par voie sous-cutanée
 - instable: préparation extemporanée



Risque de transmission par les aiguilles !

$t_{1/2}$: 3-4 heures (hydrolyse); administration 2 X / jour

Effets secondaires:

- réactions cutanées au site d'injection
- réactions d'hypersensibilité pouvant imposer l'arrêt du traitement
- augmentation du risque de pneumonie en début de traitement (raison peu claire)



Patient à risque d'infection opportuniste !

Usage clinique:

- en association avec d'autres antiviraux; patients phase avancée (souches multirésistantes)

02/05/2008

SIDA

54

Infuvirtide: conseils d'auto-administration



2 Injection Sites and Syringe Information

Injection Sites

Changing where you inject FUZEON on your body each time is an important way to lessen how bad your injection site reactions get. For more detailed information about each injection site, see *Your Guide to Taking FUZEON*.



Abdomen



Upper Thighs

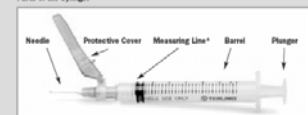


Upper Arms

About the Safety Syringes

- * There are two different-sized safety syringes, a 3-cc/mL (large) syringe and a 1-cc/mL (small) syringe.
- * Before using the safety syringe, be sure the clear plastic capped needle is tight by pushing it down gently while twisting it clockwise [2A].
- * The safety syringes have a colored piece of plastic that covers the needle after use, lowering the risk of needlestick injuries [2B, 2C].
- * Your healthcare provider may recommend other types of syringes for use with FUZEON.
- * Never throw your used syringes into the trash. Put them in the sharps container.

Parts of the Syringe:



* The measuring line of the syringe is the edge line of the plunger closest to the needle.



02/05/2008

SIDA

55

Infuvirtide: conseils d'auto-administration



4 Mixing FUZEON

Draw Up Sterile Water

- * Gently tap the FUZEON vial to loosen the powder
- * Pick up the 3-cc/mL (large) syringe
- * Using your index finger, pull the colored protection device away from the capped needle
- * To ensure that the needle is secure, hold the clear plastic cap and tighten the needle with a gentle clockwise twist. Do not use too much force as the needle may loosen [4A].
- * Pull the clear plastic cap off
- * Pull the plunger back to get 1.1 cc/mL of air [4A].
- * Before turning the sterile water vial upside down, slowly inject the air into the vial—and keep the needle in the vial
- * Turn the vial upside down. Make sure the tip of the needle is always below the surface of the water to help keep air bubbles from entering the syringe [4B].
- * Slowly pull the plunger back to get 1.1 cc/mL of sterile water into the syringe. *Tip!* Gently tap or flick the barrel and push and pull the plunger to remove extra air and bubbles. To be sure you end up with 1.1 cc/mL of sterile water in the syringe, you may need to pull the plunger past the 1.1 cc/mL mark. [4C]
- * Carefully remove the needle and syringe from the vial



- * Remove the needle from the vial. Using one hand, gently press the colored protective cover against a flat surface until you hear a click and the needle is re-covered. *Never use your hand to re-cover the needle* [4E, 4F].

* Put the used syringe in the sharps container [4G].

Gently Mix FUZEON

- * Gently tap the FUZEON vial with your fingertip for 10 seconds to start dissolving the powder. Then gently roll the FUZEON vial between your hands to reduce the mixing time. [4H] Make sure no FUZEON is stuck to the vial walls. After tapping, it could take up to 45 minutes to dissolve. *Important!* Never shake the FUZEON vial. Shaking will make the medicine foam and it will take much longer to dissolve.
- * Once the powder starts to dissolve, just set it aside and it will completely dissolve
- * When completely mixed, the liquid FUZEON should be clear and without foam. [4I] If the FUZEON is foamy [4H] or jelled, allow more time for it to dissolve.
- * If you see bubbles, gently tap the vial until they disappear
- * If you see any particles in the FUZEON once it is completely mixed, do not use that vial. Contact the pharmacy that provided it
- * Mixed FUZEON must be used right away or stored in the vial in the refrigerator and used within 24 hours. Do not store mixed FUZEON in the syringe [4J].



02/05/2008

SIDA

56

Infuvirtide: conseils d'auto-administration



5 Giving the Injection

Choose the injection Site

- Clean your FUZEON Plunger to help you, choose a site different from the one you used for your last injection
- Important!** With the tips of your fingers, feel for any hard bumps. Do not inject in or near bumps or any other types of reactions from past injections. Also, do not inject into moles, scars, bruises, your belly button or areas that could be irritated by a belt or waistband. [5A]
- Clean the injection site with a new alcohol pad. Start in the center, apply pressure and clean in a circular motion, working outward. Allow the site to air-dry [5B]



- Make sure the tip of the needle is always below the surface of the FUZEON to help keep air bubbles from entering the syringe. Slowly pull the plunger to get 1 cc/mL of FUZEON [5F]

Tip! Gently tap or flick the barrel and push and pull the plunger to remove extra air and bubbles. To be sure you end up with 1 cc/mL of FUZEON in the syringe, you may need to pull the plunger past the 1 cc/mL mark. [5E]



- Carefully remove the needle and syringe from the vial

inject FUZEON

- Pinch and hold a fold of skin around the injection site [5F]
- Pierce the skin at a 45-degree angle. The needle should be inserted most of the way in [5G]
- Tip!* Your healthcare provider may teach you to inject in a different way.
- Slowly push the plunger all the way to inject FUZEON
- Remove the needle from your skin
- Using one hand, gently press the colored protective cover against a flat surface until you hear a click and the needle is re-covered. Never use your hand to re-cover the needle [5B, 5C]
- Put the used syringe in the sharps container [5H]
- Cover the site with a small bandage if you see any blood or medicine

For additional details on FUZEON, please see the accompanying patient package insert.

02/05/2008

SIDA

57

Inhibiteurs des co-récepteurs CCR5

a

gp120
gp41
CD4
Host cell membrane
Co-receptor (CCR5 or CXCR4)

b

Co-receptor binding site

c

d

Fusion peptide

e

UK-427857
Maraviroc
Selzentry

SCHD (SCH-417690)
Vicriviroc

873140 (GW-873140, ONO-4128, AK-602)
Aplaviroc

Figure 9 | Human immunodeficiency virus (HIV) co-receptor antagonists. When the HIV glycoprotein gp120 binds to CD4 (a), it induces a conformational change in gp120 that exposes the co-receptor binding site (b); this is a complex domain comprising the V3 loop and specific amino-acid residues in CD4, collectively termed the 'bridging sheet'. Exposure of the co-receptor binding site permits binding of gp120 to the co-receptor (c). Co-receptor antagonists inhibit this step by binding to the co-receptor and changing its shape so that gp120 cannot recognize it. Co-receptor binding induces conformational changes in gp41 and insertion of the fusion peptide into the host cell membrane (d), ultimately resulting in fusion of the viral envelope with the host cell membrane⁹¹. (e) Structural formulae of selected CCR5 antagonists.

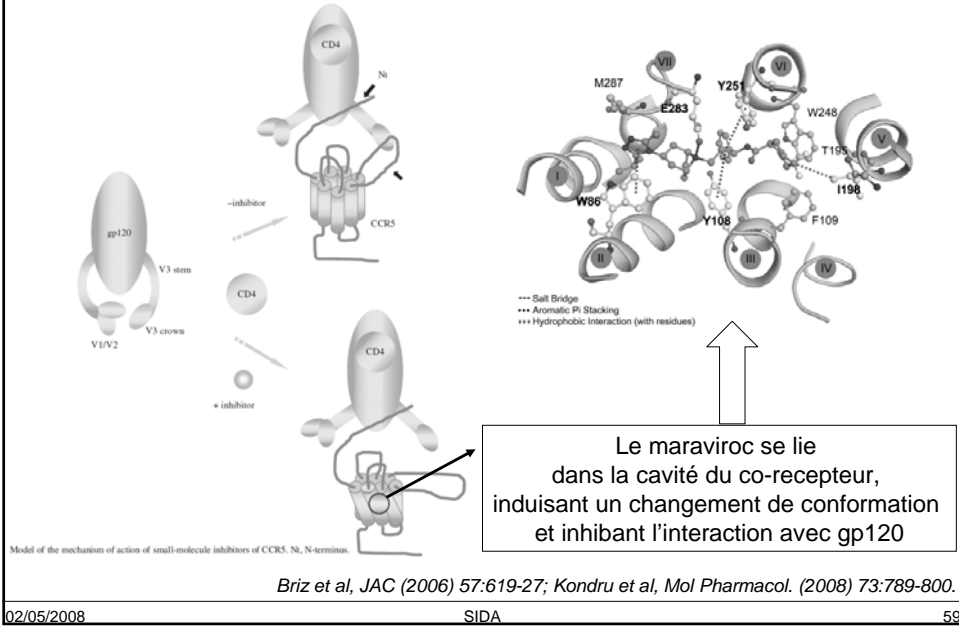
De Clercq, Nature Rev. Drug Discov. (2007) 6:1001-1018

02/05/2008

SIDA

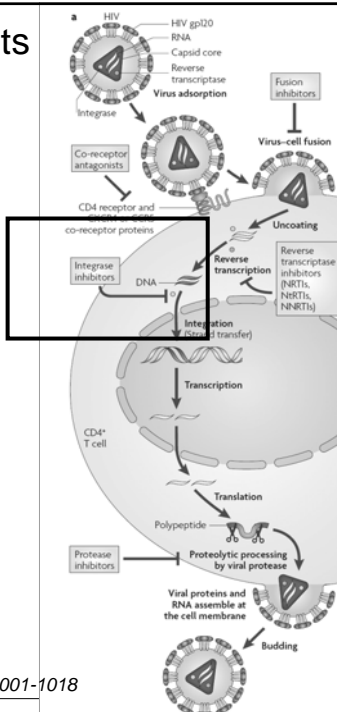
58

Antagonistes de CCR5



Cible des médicaments actifs sur le HIV

en développement



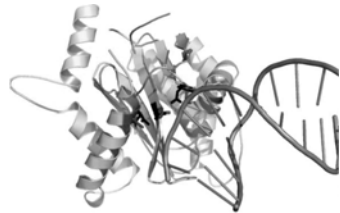
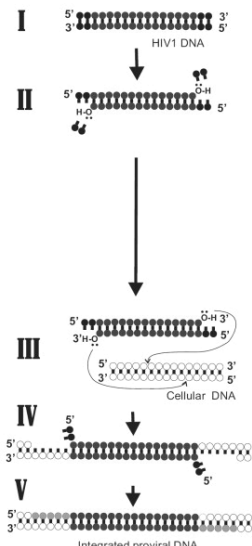
De Clercq, Nature Rev. Drug Discov. (2007) 6:1001-1018

02/05/2008

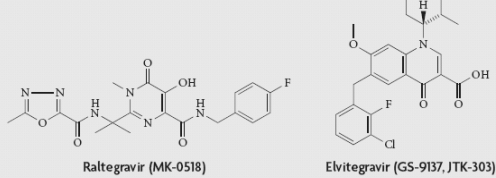
60

Inhibiteurs d'intégrase

INTEGRASE



a



Savarino, Retrovirology. (2007) 4:21

02/05/2008

SIDA

61

PHARMACOTHERAPIE DU SIDA



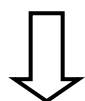
02/05/2008

SIDA

62

Buts du traitement

- ↓ charge virale
0.5-0.75 log₁₀ en 4 semaines ou 1 log₁₀ en 8 semaines
- charge virale non détectable à 4-6 mois
(< 50 - 20 copies)
- restaurer ou préserver la fonction immunitaire
- réduire la morbidité et la mortalité



Trithérapie pour éviter la sélection de résistance

02/05/2008

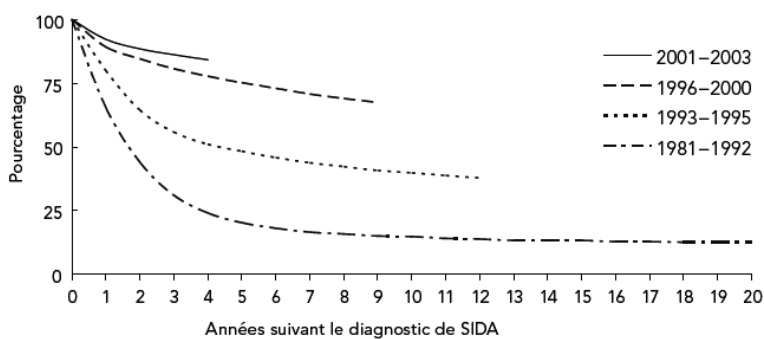
SIDA

63

Grâce au HAART la survie des patients s'améliore



Pourcentage de personnes encore en vie en juin 2006,
par cohortes selon les années suivant le diagnostic de SIDA
entre 1981 et 2003 et par année de diagnostic



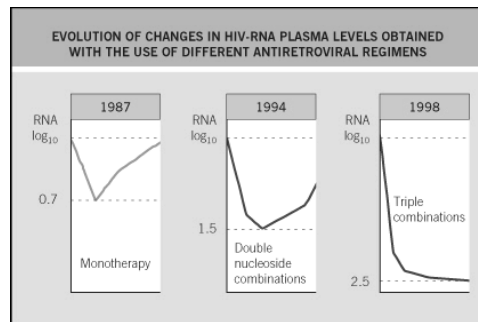
Source : CDC Twenty-five years of HIV/AIDS – Etats-Unis, 1981-2006. MMWR 2006.

02/05/2008

SIDA

64

Grâce au HAART la survie des patients s'améliore



© Elsevier 2004. Infectious Diseases 2e - www.idreference.com

02/05/2008

SIDA

65

Algorithme de traitement proposé par l'OMS (1/3)

TABLE 6. RECOMMENDATIONS FOR INITIATING ART IN PLHIV

WHO clinical stage ^a	CD4 cell count	Recommendation
1	<200/mm ³	Treat
	200–350/mm ³	Consider treatment ^b
2	<200/mm ³	Treat
	200–350/mm ³	Consider treatment ^b
3	200–350/mm ³	Treat
4	Regardless of CD4 count	Treat

1. asymptomatique, adénopathie
2. Candidose, infections respiratoire, herpes
3. Candidose récurrente, ulcération de la bouche, infections pulmonaires sévères, diarrhée inexpliquée
4. Infections opportunistes

www.euro.who.int/document/e90840.pdf

02/05/2008

SIDA

66

Algorithme de traitement proposé par l'OMS (2/3)

TABLE 7. RECOMMENDED FIRST-LINE HAART

ARV drug classes	HAART regimens
2 NRTIs + 1 NNRTI	ZDV + 3TC + (EFV ^a or NVP) or TDF + FTC + (EFV ^a or NVP) or ABC + 3TC + (EFV ^a or NVP)

^a EFV is highlighted as the preferred NNRTI.



TABLE 8. CRITERIA FOR TREATMENT SUCCESS

	Virological		Immunological	Clinical
Marker	Viral Load		CD4 cell count	Clinical stage
Time ^a	24 weeks		24–48 weeks	By 12 weeks of treatment initiation should be asymptomatic or have few symptoms
Suggested ranges ^a	<400 copies/ml	<50 copies/ml	Increase from baseline by at least 50–100 cells/mm ³	Stage 1 or 2 ^b

www.euro.who.int/document/e90840.pdf

02/05/2008

SIDA

67

Algorithme de traitement proposé par l'OMS (3/3)



TABLE 9. RECOMMENDED SECOND-LINE HAART FOR ADULTS AND ADOLESCENTS

First-line HAART regimens	Second-line HAART regimens after treatment failure
ZDV + 3TC + (EFV or NVP)	LPV/r ^a (or ATV/r, SQV/r, FPV/r, IDV/r) + ddI + ABC or LPV/r ^a (or ATV/r, SQV/r, FPV/r, IDV/r) + TDF + ABC or LPV/r ^a (or ATV/r, SQV/r, FPV/r, IDV/r) + TDF + (ZDV + 3TC) ^b
TDF + FTC + (EFV or NVP)	LPV/r ^a (or ATV/r, SQV/r, FPV/r, IDV/r) + ddI + ABC or LPV/r ^a (or ATV/r, SQV/r, FPV/r, IDV/r) + ddI + ZDV
ABC + 3TC + (EFV or NVP)	LPV/r ^a (or ATV/r, SQV/r, FPV/r, IDV/r) + ddI + ZDV or LPV/r ^a (or ATV/r, SQV/r, FPV/r, IDV/r) + ZDV + TDF (+ 3TC) ^b

www.euro.who.int/document/e90840.pdf

02/05/2008

SIDA

68

Suivi des patients



TABLE 10.

FREQUENCY OF LABORATORY TESTING, GENERALLY AND WITH SPECIFIC ARV USE

	Baseline	Week 2	Week 4	Week 8	Week 16	Week 24	Week 36	Week 48
Viral load	X			X		X	X	X
CD4 count	X			X		X	(X)	X
Complete blood count	X		X	X	X (ZDV)	X	(X)	X
Liver Function Test (LFT)	X	X (NVP)	X	X (NVP, ZDV, PIs)	X (NVP, PIs)	X	(X)	X
Cholesterol triglycerides	X (PIs)				X (PIs)			X (PIs)
Renal function test	X	X (TDF)	X (TDF, IDV)			X	(X)	X

X: laboratory tests to be performed irrespective of the ARVs being administered; X (ARV): laboratory tests to be performed if an ARV in parentheses is being administered; (X): optional test.

www.euro.who.int/document/e90840.pdf

02/05/2008

SIDA

69

Prévention de la transmission foeto-maternelle

REVERSE TRANSCRIPTASE INHIBITORS FDA approved					
Agent	Transmission to fetus prevented*	Neonates	Children	FDA pregnancy category ^[dagger]	Placental transfer (%)
Zidovudine ^{Rx}	Yes	Yes	Yes	C	85
Didanosine ^{Rx}	No	Yes	Yes	B	50
Lamivudine ^{Rx}	Yes	No	≥3 months	C	100
Stavudine ^{Rx}	No	No	≥1 months	C	76 (rhesus monkeys)
Zalcitabine ^{Rx}	No	No	No	C	30-50 (rhesus monkeys)
Abacavir	No	No	≥3 months	C	Yes (rats)
Nevirapine ^{Rx}	Yes	No	≥2 months	C	100
Delavirdine	No	No	No	C	?
Efavirenz ^{Rx}	No	No	≥3 years	C	100 (rhesus monkeys)
Tenofovir	No	No	No	B	Yes (rat, monkey)

PROTEASE INHIBITORS FDA approved					
Agent	Transmission to fetus prevented*	Neonates	Children	FDA pregnancy category ^[dagger]	Placental transfer
Nelfinavir	No	No	≥2 years	B	Minimal
Indinavir	No	No	No	C	Minimal
Ritonavir	No	No	≥2 years	B	Minimal
Saquinavir ^{Rx}	No	No	No	B	Minimal
Amprenavir ^{Rx}	No	No	≥4 years	C	?
Lopinavir/ritonavir	No	No	≥6 months	C	?

Ne passent pas
la barrière placentaire



02/05/2008

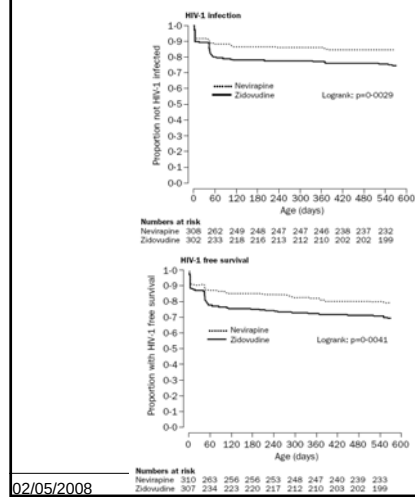
SIDA

70

Prévention de la transmission foeto-maternelle

→ Traitement suggéré:

- AZT or AZT/3TC – pendant la gestation et continuer pendant l'accouchement
- Nevirapine – 1 dose à la mère et à l'enfant (zones défavorisées)



Methods From November, 1997, to April, 1999, HIV-1 infected pregnant women in Kampala, Uganda, were randomly assigned nevirapine (200 mg at labour onset and 2 mg/kg for babies within 72 h of birth; regimen A) or zidovudine (600 mg orally at labour onset and 300 mg every 3 h until delivery, and 4 mg/kg orally twice daily for babies for 7 days, regimen B). Infant HIV-1 testing was done at birth, age 6–8 and 14–16 weeks, and age 12 months by HIV-1 RNA PCR, and by HIV-1 antibody at 18 months.

Traitement court:
la nevirapine
est plus efficace !



Jackson et al, Lancet (2003) 362:859-68.

02/05/2008

71

Prévention lors d'une exposition accidentelle à un matériel contaminé



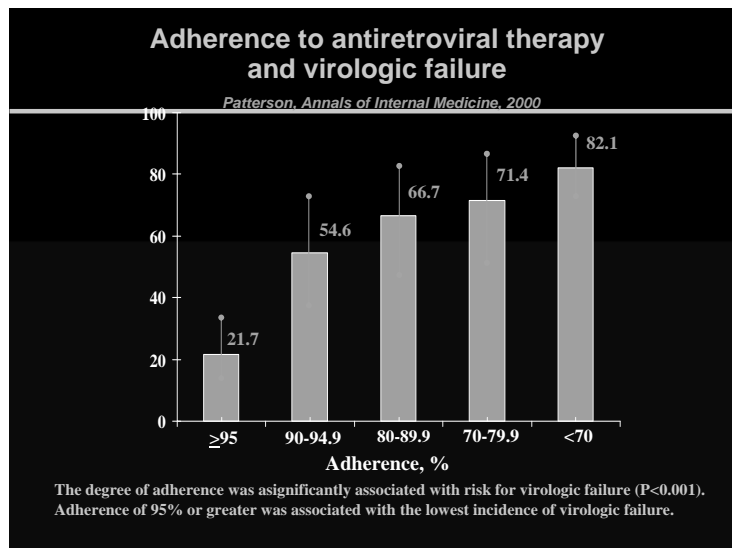
- traitement administré le plus rapidement possible ; 4 semaines
- association puissante : 2 NRTI et 1 IP (zidovudine-lamivudine-indinavir)
[bonne tolérance et interactions médicamenteuses limitées]
- surveillance clinique et biologique à maintenir plus longtemps.

02/05/2008

SIDA

72

Importance de l'adhérence au traitement



J. Nacheega, 2006

02/05/2008

SIDA

73

Comment améliorer la compliance ?



02/05/2008

SIDA

74

Comment améliorer la compliance ?



Simplification des régimes thérapeutiques: comparaison entre 1996 et 2004

1996:

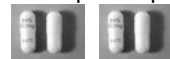
ddl + d4T + SQV

-24 gélules/jour:

-SQV: 6 gel 3 X/jour avec la nourriture



-ddl: 2 gel 2 X jour ½ hr avant
ou 2 h après repas



-d4T: 1 co 2 X /jour



2004:

TDF/FTC or ABC/3TC + EFV

- 1 co 2 X/ jour + 1 co 1X/jour



pas restriction par rapport au repas

02/05/2008

SIDA

75

Comment améliorer la compliance ?



Reasons for Missing Doses of Antiretroviral Therapy

US

Chesney

- Simply forgot
- Slept through dose
- Away from home
- Change in routine
- Busy with other things
- Too sick
- Depressed

Africa

Weidle, Orrell, Nachege, Brown,

- Forgot
- Away from home
- Schedule difficulties
- Ran out of pills
- Cost
- Home language
- Fear of stigmatization by sexual partner

J. Nachege, 2006

02/05/2008

SIDA

76

.. Un rôle de choix pour le pharmacien !



http://www.ascp.com/public/pubs/tcp/1998/nov/hiv aids.shtml

AMERICAN SOCIETY OF CONSULTANT PHARMACISTS

Membership Practice Resources
Meetings & Education Government Affairs
Publications & Products ConsultNet™
Students & New Practitioners ASCP Calendar
ASCP Foundation News

Quick jump to... ▾

Current Concepts in

HIV/AIDS Pharmacotherapy

Pharmacists have assumed an increasingly important role in monitoring and fine-tuning HIV drug therapy for maximal effectiveness....

http://www.fip.org/activities/activities_working_groupmember.htm

The International Pharmaceutical Federation (FIP) and World Health Organisation (WHO) Working Group on AIDS and Drug Addiction

PHARMACISTS AS KEY FOR PREVENTION AND PHARMACEUTICAL CARE PROVIDERS FOR PEOPLE LIVING WITH HIV

COMPOSITION OF THE WORKING GROUP

BELGIUM

M. Laurent RAVEZ - Conseiller Ethique
Association Chrétienne des Institutions Sociales et de Santé,

M. F. DE BRABANTER - Directeur du Secrétariat National
Ordre des Pharmaciens Belges

M. HANOT - President
Conseil National de l'Ordre des pharmaciens