

A simple question...

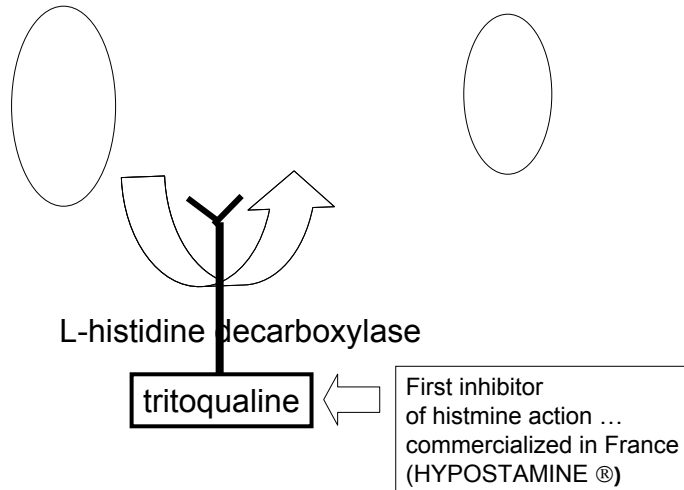
How was histamine discovered ?

- by chemical synthesis ...
- from the analysis of plant extracts (ergot fungus *Claviceps purpurea*)
- from the analysis of animal tissues extracts
- through none of these approaches

Histamine ...

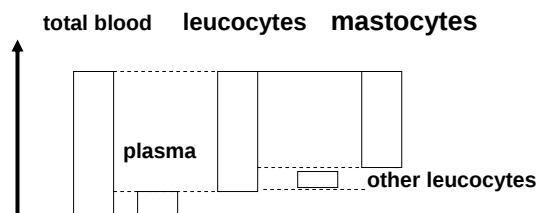
- ➔ obtained by synthetic chemist in 1907 ...as a chemical curiosity ...
 - detection of an identical compound in an extract from ergot fungus ... and shown to cause a marked vasodilatation
 - a similar effect is seen with tissues extracts
 - produces a similar picture as a very severe allergic reaction
- ➔ recognized as a "biological" molecule (and not a product from putrefaction in 1927 ...

From histidine to histamine ...



Localization of histamine

1. blood

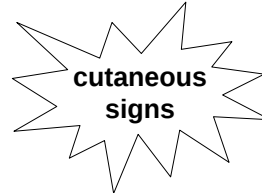


2. tissues ... the word comes from ιστος ("histos" = tissue !!)

- skin
- lung
- gastrointestinal tract
- central nervous system

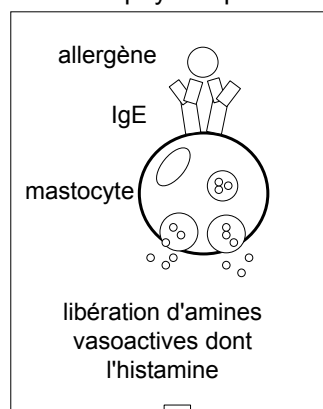
Actions of histamine

- **↑ of capillary permeability and vasodilatation**
 - rednesses
 - inflammation
- bronchoconstriction
important with the guinea-pig but under H_2 retrocontrol in man
- **↑ of HCl secretion**
(parietal cells of the stomach)
- **neurotransmission**
 - awakening reactions, tachycardia, hypertension
 - nausea, vomiting
 - migraines



Rappel: les 4 types de réactions d'hypersensibilité

Réaction de type I anaphylactique



- rhinite, conjonctivite, urticaire, asthme aigu, (bronchospasme), oedème
- délai: endéans les 30 min

Réaction de type II: cytotoxique

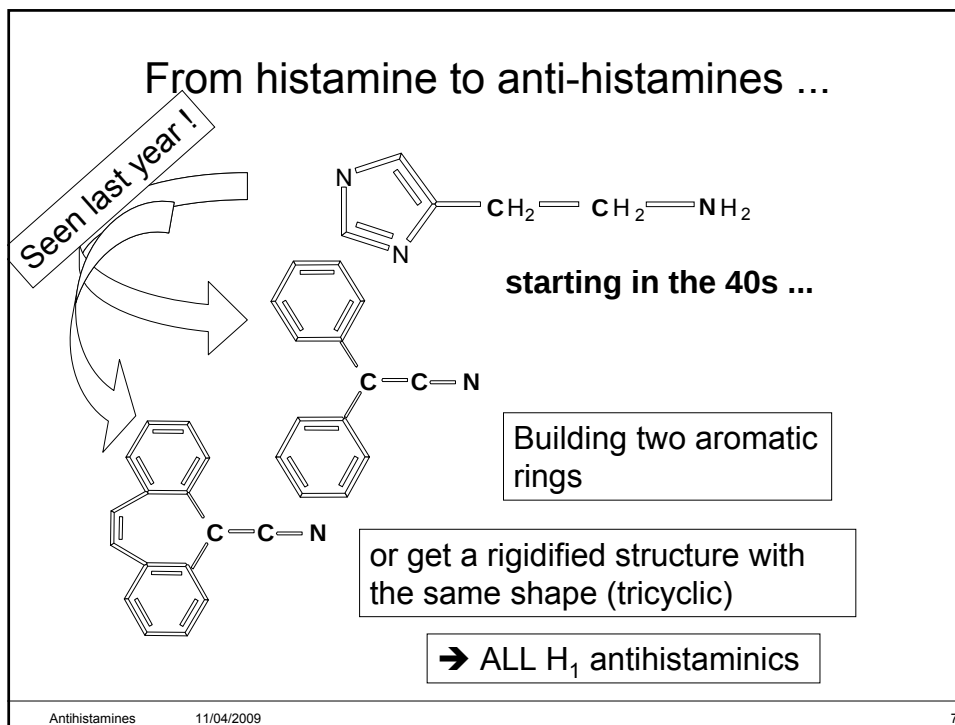
- médiée par les IgG et/ou les IgM
- action directe sur une cellule cible
- implique le complément
- lyse, phagocytose (anémie hémolytique, agranulocytose, thrombopénie)
- délai: 5-12h

Réaction de type III: formation de complexes immuns

- dépôts dans les tissus avec réaction inflammatoire disséminée
- activation du complément et libération de toxines des leucocytes
- agrégation plaquettaire, microthromboses...
- délai: 3-8h

Réaction de type IV : cellulaire

- activation directe des cellules T
- libération de cytokines et de $TNF\alpha$
- induit typiquement des manifestations cutanées (dermatite de contact, exanthèmes, eczéma, ...)
- délai: 24 à 48h



Rationalization through a deep understanding of the receptor

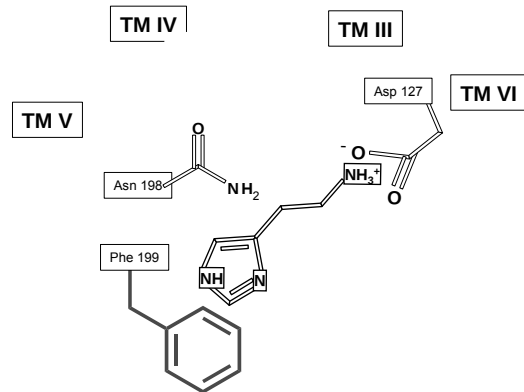
- H₁ receptor
 - CNS
 - périphery

action mediated by the phosphoinositides
- H₂ receptor
 - stomach
 - lung
 - CNS

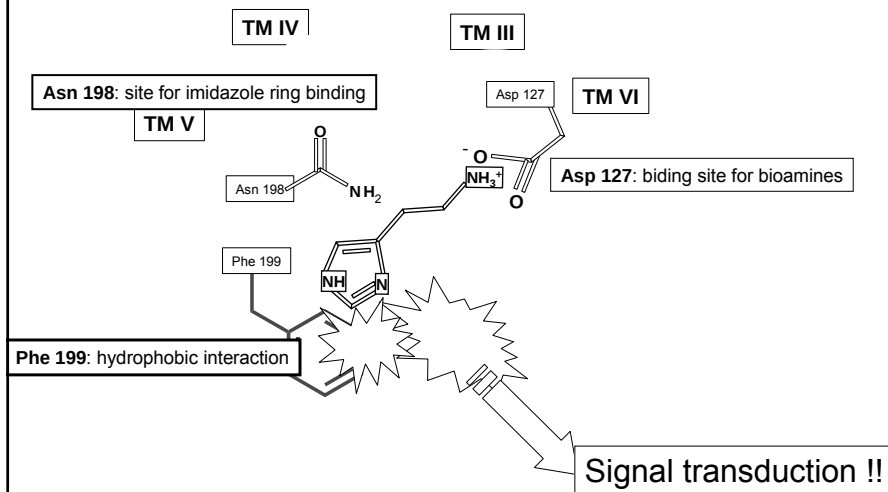
action mediated by cyclic AMP
- H₃ receptor
 - CNS

Antihistamines
11/04/2009
8

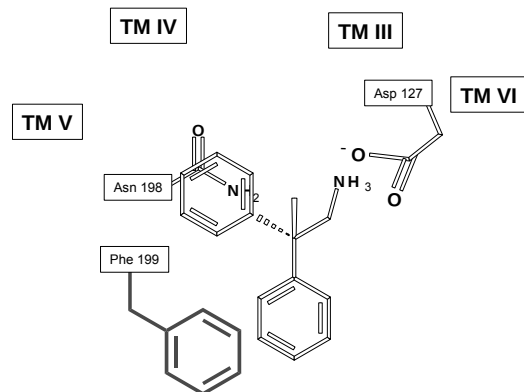
Binding of histamine to H1 receptor



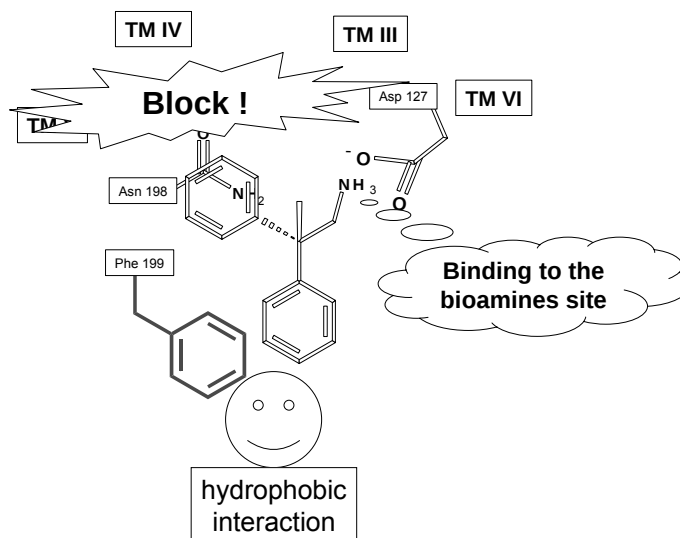
Binding of histamine to H1 receptor



Binding of an antagonist ...



Binding of an antagonist ...



Une famille d'antagonistes H1....

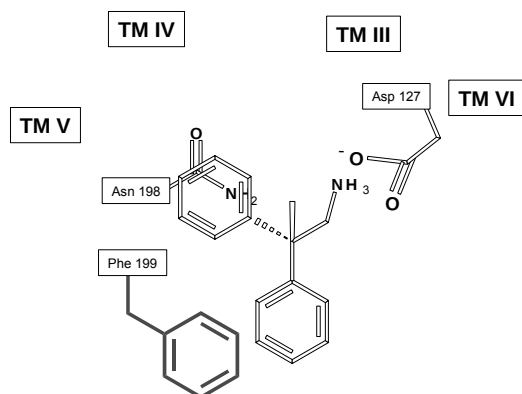
Nom DCI	nom commercial en Belgique *
• alimémazine	THERALENE
• prométhazine	PHENERGAN
• dimenhhydrinate	PARANAUSINE / VAGOMIN
• diphenhydramine	BENYLIN
• dexchlorphéniramine	POLARAMINE
• ciproheptadine	PERIACTIN
• dimétindène	FENISTIL
• méclozine	AGYRAX / POSTAFENE
• cetirizine	ZYRTEC / REACTINE /
• loratadine	CLARITINE / SANELOR
• fexofenadine	TELFAST

et plus récemment

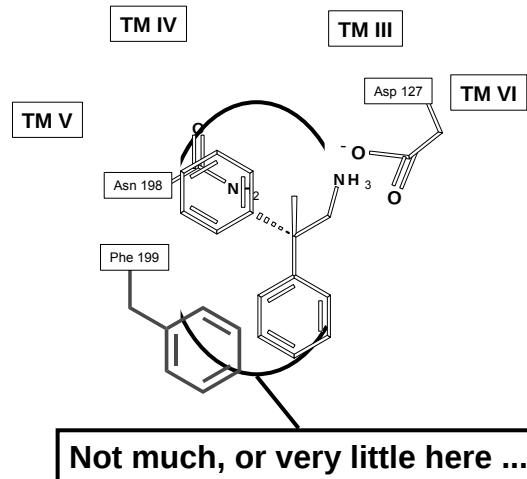
• lévocetirizine	XYZAL
• desloratadine	AERIUS

* liste non limitative...

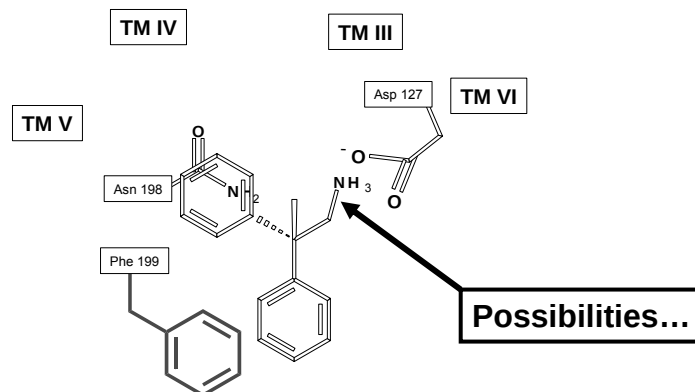
Binding of an antagonist: what can you modify ?



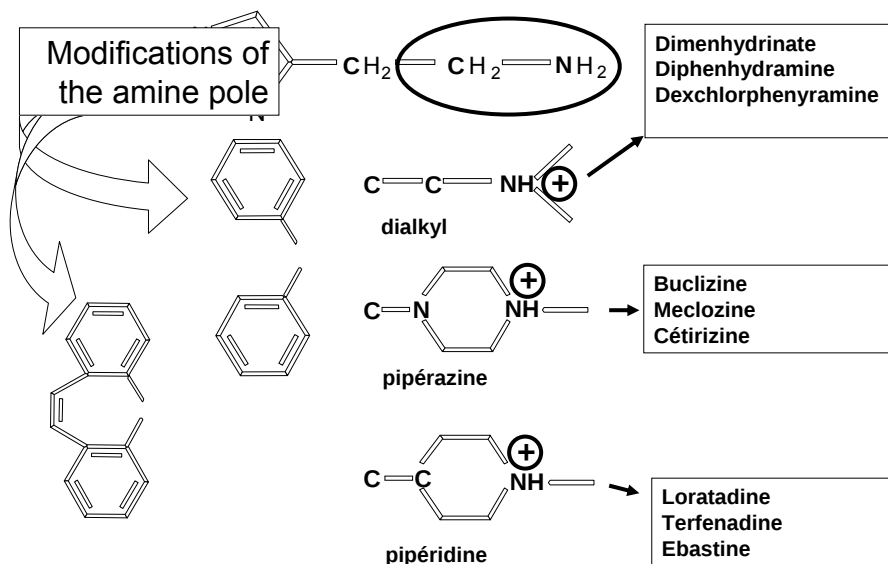
Binding of an antagonist: what can you modify ?



Binding of an antagonist: what can you modify ?



Variations among antihistamines....



Antihistamines

11/04/2009

17

The ideal antihistaminic drug for the treatment of allergy

What is your "wish list" ?

- Low sedation activity *
- No or little anticholinergic effects **
- Getting a rapid and prolonged action ***

* most "old" antihistamines make you to fall asleep...

** because their structure is reminiscent of atropine

*** I want a fast relief, and not needing taking pills every hour...

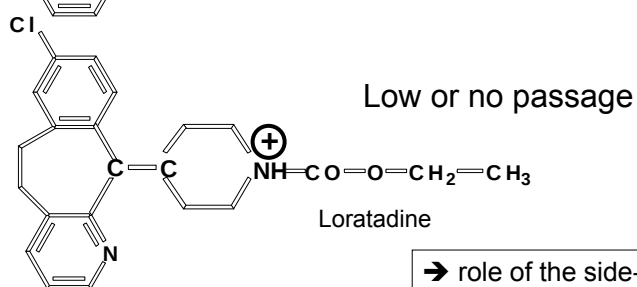
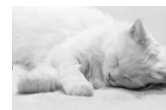
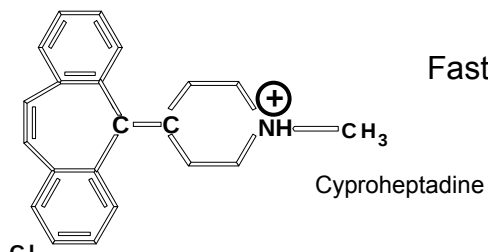
Antihistamines

11/04/2009

18

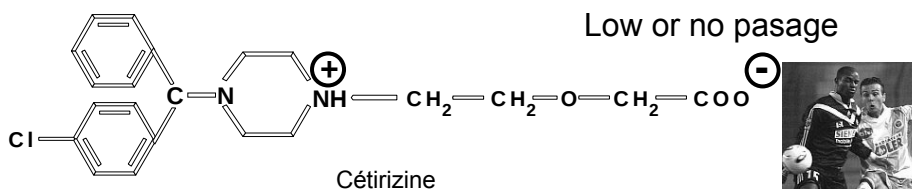
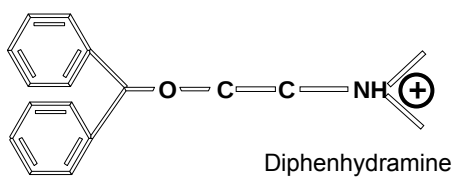
Low sedation activity ...

➔ Modulation of the hematoencephalic barrier passage...



Low sedation activity ...

Another example...



➔ role of the length and of the polarity of the side-chain

Molécules à passage hémato-méningé important et causant de la sédation ...

Nom DCI	sédation	OTC
alimémazine	+++	oui (partiel.)
prométhazine	+++	oui
dimenhydrinate	+++	oui
diphenhydramine	+++	oui
oxomémazine	++	non
dexchlorphéniramine	++	oui
ciproheptadine	++	oui
dimétindène	+	oui
méclozine	+	oui
	+	oui

Antihistamines

11/04/2009

21

The antihistaminic and the sedative actions of the "old" antihistaminics go side by side

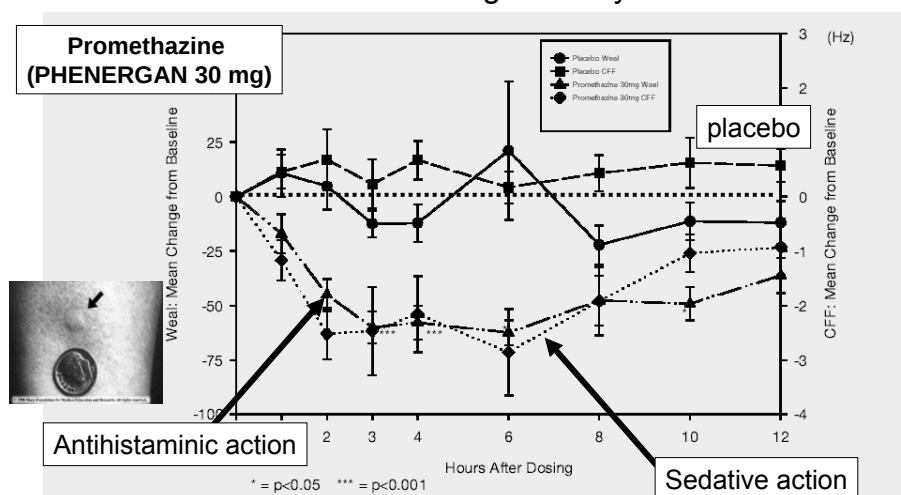


Figure 3. Change from baseline: peripheral antihistaminic suppression (weal) with respect to CFF threshold: acute dose promethazine 30 mg, day 1

Hindmarch et al., Curr. Med. Res. Opin., 17:241-255, 2001

Antihistamines

11/04/2009

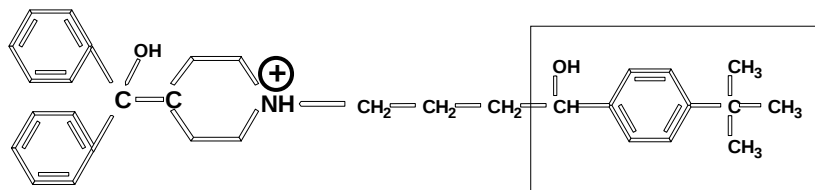
22

First molecules with low level of passage through the hemato-encephalic barrier

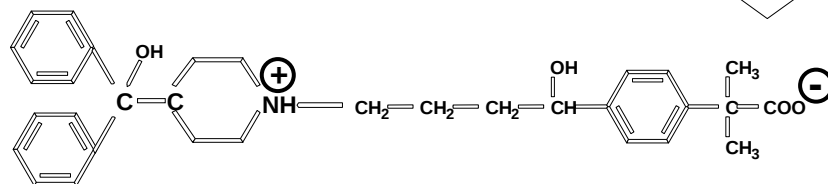
- astémisole
 - terfénadine
- withdrawn because of cardiac toxicity
Torsades de pointe !!!
- fexofénadine
- Active metabolite of terfenadine

What was terfenadine...

- terfenadine was a pro-drug

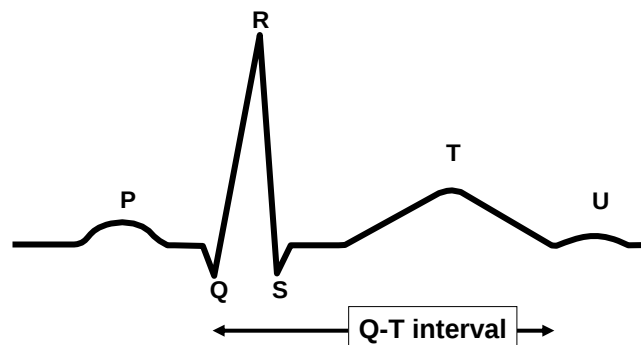


- which underwent a "first pass" liver metabolism that released fexofenadine, the active product



The main problem of terfenadine ...

- if terfenadine reaches the heart, it will block the K^+ canal, causing a delay in repolarization (that translate into a prolongation of Q-T interval [visible at the ECG] that may lead to **life-threatening** arrhythmia and "**Torsades de pointes**" ...

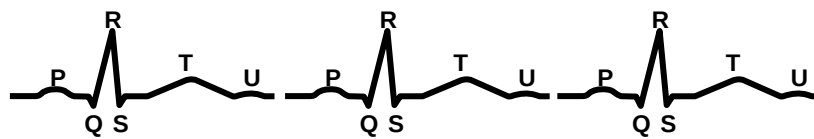


Antihistamines

11/04/2009

25

What is "Torsades de pointe" ?



J. Simkó et al. Proarrhythmic Potential of Antimicrobial Agents



Figure 1. Initiation of torsades de pointes. Note the prolonged QT interval of the last preceding beat, the twisting polarity and the changing amplitude of the QRS complexes during the arrhythmia.

Antihistamines

11/04/2009

26

Mechanism(s) of Torsade de Pointe

1. Role of ectopic beats

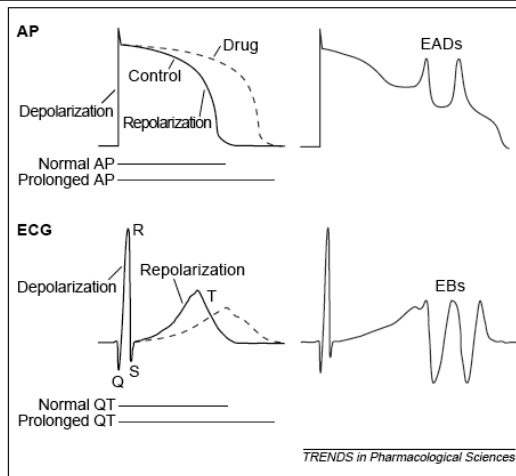


Figure 1. The relationship between ventricular transmembrane action potentials (APs) and the surface electrocardiogram (ECG). In the ECG traces the QRS and T waves denote the depolarization and repolarization, respectively, of the ventricles. The QT interval represents the time elapsed between the ventricular depolarization and repolarization. An increase in the duration of the AP (APD) is responsible for the prolongation of the QT interval. Shown on the left are the APs and ECG in the absence (control) and presence of a drug [e.g. a blocker of the rapid component of the delayed rectifier K^+ current (I_{Kr})] that prolongs the APD and consequently the QT interval. Shown on the right are two large early afterdepolarizations (EADs) occurring during the repolarization phase of a prolonged AP, giving rise to two ectopic beats (EBs) in the ECG trace.

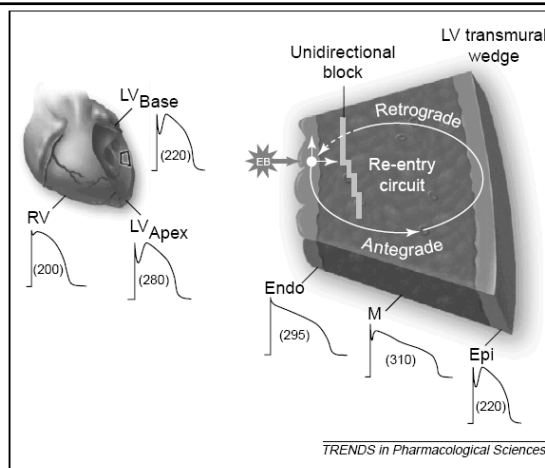
Balardinelli et al. TIPS (2003) 24:619-625

Antihistamines 11/04/2009

27

Mechanism(s) of Torsade de Pointes

2. role of dispersion of ventricular repolarization



Balardinelli et al. TIPS (2003) 24:619-625

Figure 3. Heterogeneity of ventricular repolarization caused by regional differences in action potential duration and the development of a re-entrant circuit. A heart and a transmural wedge of the left ventricle are shown with action potential (APs) and a re-entrant circuit. The APs shown are representative examples from the right and left (base and apex) ventricles (RV and LV, respectively), and the endocardial (Endo), mid-myocardial (M) and epicardial (Epi) regions across the LV wall. The numbers in parentheses denote the duration (in milliseconds) of the APs. The differences in the durations of the action potentials (APDs) are responsible for the heterogeneity of refractoriness, which in turn create the substrate for re-entrant arrhythmias, such as torsade de pointes. A re-entrant circuit that includes myocardial cells from the endocardial, mid-myocardial and epicardial regions of the LV transmural wedge is shown together with a region of functional unidirectional block (thick blue line) created by the long APDs of the mid-myocardium (long refractory period), and the reentrant excitation wave front (white line with arrows).

A simplified sequence of events that can initiate re-entry includes the following. (i) A premature beat [ectopic beat (EB)], presumably brought about by an early afterdepolarization, encounters a region of functional unidirectional block in its propagation path. (ii) The EB, however, propagates laterally through myocardium that is not refractory (cells with shorter APDs), resulting in excitation of the endocardium. (iii) This excitation wave front continues in an antegrade direction to excite various portions of the mid-myocardium, and the epicardium. (iv) The excitation wave front twists around and propagates in the opposite direction towards the endocardium (i.e. retrograde conduction) to activate the remainder of the mid-myocardium. (v) Finally, the wave front crosses the region of unidirectional block no longer refractory (dashed white line with arrow), to re-enter and thereby re-excite the endocardium. For a more detailed description of the electrophysiological basis for re-entry the reader is referred to Ref. [9].

Antihistamines 11/04/2009

28

Risk of Torsade de pointes and inhibitors of cyt P450 metabolism

Table 1

QT interval prolonging drugs metabolized by CYP 3A4, which may possibly interact both pharmacokinetically and pharmacodynamically with macrolides and imidazole antifungals.

Antiarrhythmics	Amiodarone (with roxithromycin [23]), quinidine (with erythromycin [116]), disopyramide (with clarithromycin [117, 118])
Antifungals	Fluconazole, ketoconazole, itraconazole, miconazole
Prokinetics	Cisapride (with clarithromycin, [119, 120], with erythromycin [121])
Antihistamines	Terfenadine (with erythromycin [122, 123], with troleandomycin [124]), astemizole (with erythromycin [125]), loratidine
Antipsychotics	Pimozide (with clarithromycin [126, 127]), chlorpromazine, haloperidol, ziprasidone, risperidone, clozapine, quetiapine
Immunosuppressive drugs	Tacrolimus
Opioid agonists	Methadone
Antimalarials	Quinine, chloroquine, halofantrine

Case reports on torsades de pointes or QT prolongation during coadministration of macrolide agents and other repolarization prolonging drugs are in brackets

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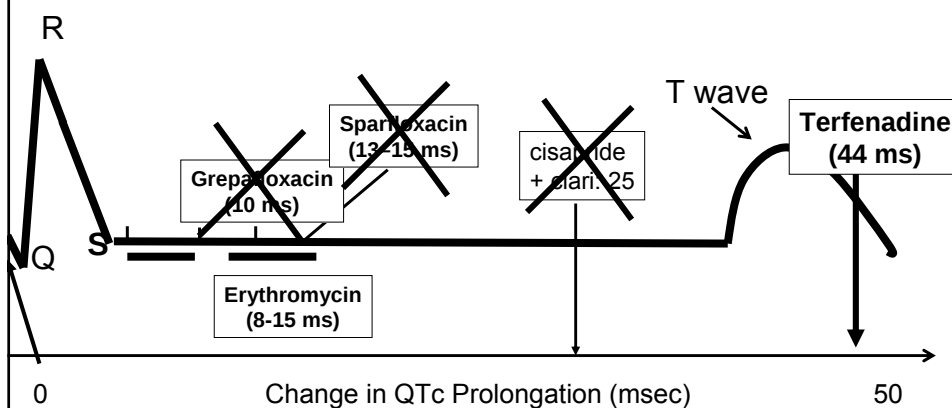
Simkó et al., Infection 2008;36:194-206

Antihistamines

11/04/2009

29

Molecules withdrawn because of risk of QT prolongation and *Torsades de pointe*



Adapted from Oberg and Bauman, 1995; Baker et al, 1997; van Haarst et al, 1998.

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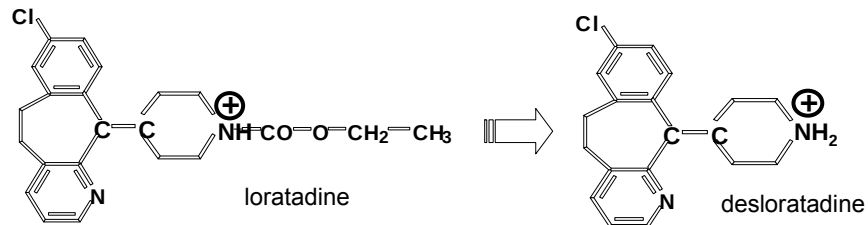
11/04/2009

30

Molecules with a weak hemato-encephalic passage ...

- loratadine

must be metabolized into desloratadine



- ebastin

- cetirizine

not very sedative and acting as such

Dissociation of the antiallergic and sedative activities

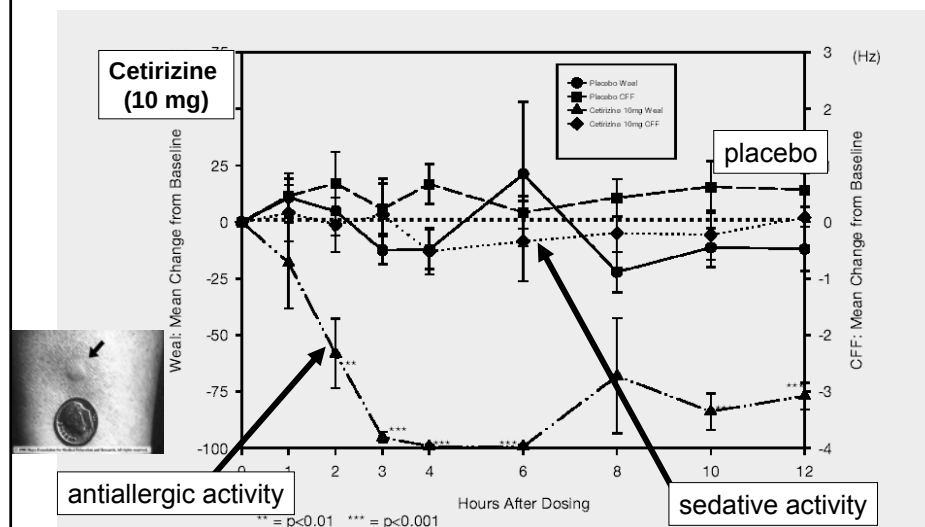


Figure 5. Change from baseline: peripheral antihistaminic suppression (weal) with respect to CFF threshold: acute dose cetirizine 10mg, day 1

Hindmarch et al., Curr. Med. Res. Opin., 17:241-255, 2001

Dissociation of anti-allergic and sédative activities...

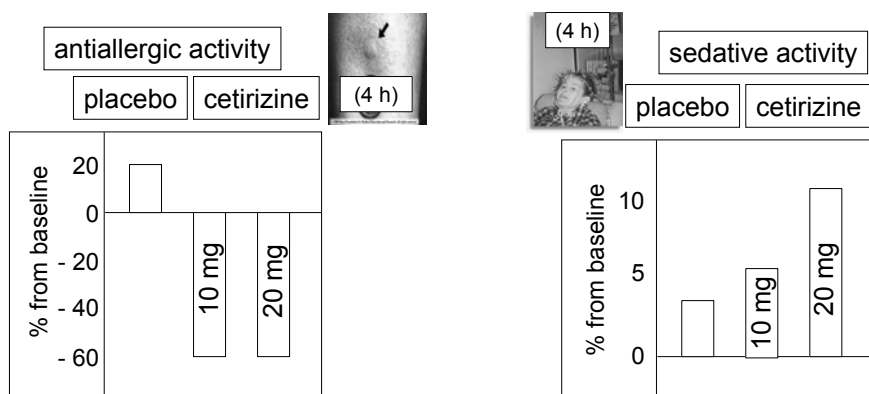
But, beware:



This is all related to dose...

Dissociation of anti-allergic and sedative activities ...

Everything is related to dose...



Sannita et al., Eur. J. Pharmacol. 300: 33-42, 1996

The ideal anti-H1 drug for treating allergy...

Specifications (*Cahier de charges*)

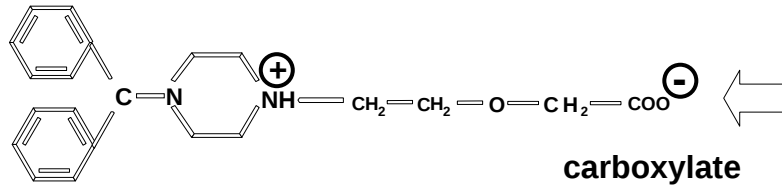
- Low sedative potential
- **Avoiding anti-cholinergic effects...**
 - important for old molecules
 - ➔ sight troubles, urinary retention ...
 - much improved for new ones
(loratadine, fexofénadine, cétirizine)
- Getting a rapid and sustained action

The ideal anti-H1 drug for treating allergy...

Specifications (*Cahier de charges*)

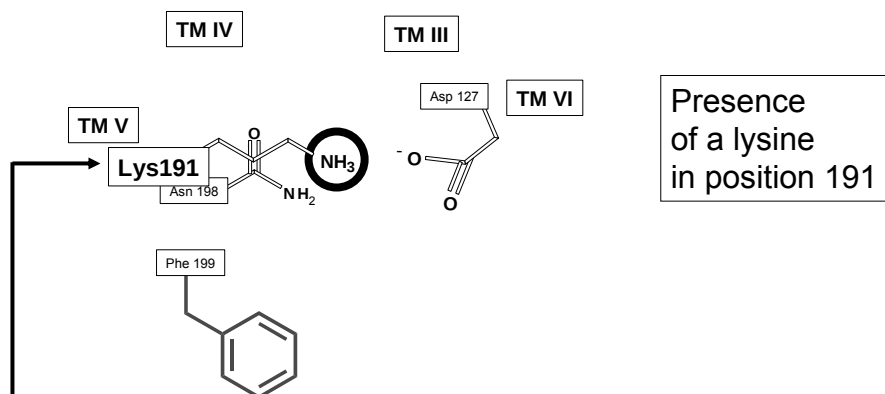
- Low sedative potential
- Avoiding anticholinergic effects
- **Getting a fast and sustained action**

Molecular properties of cetirizine



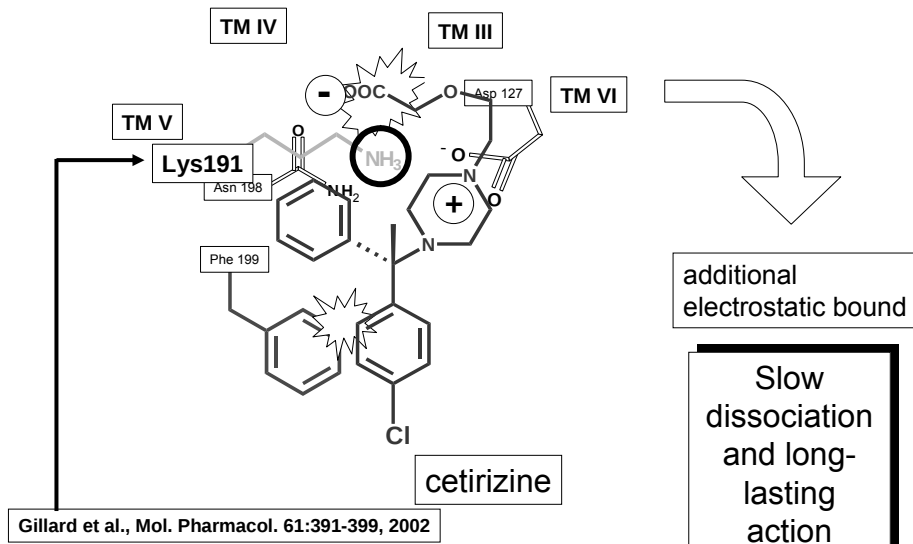
- fast action because no necessity of metabolic activation
(>< terfenadine, loratadine...)
- little or no penetration through the blood-brain barrier
- **long occupation of the receptor**

Binding of an H₁ antagonist to the receptor...



Gillard et al., Mol. Pharmacol. 61:391-399, 2002

Slow release of an antagonist ...



Prolonged action ...

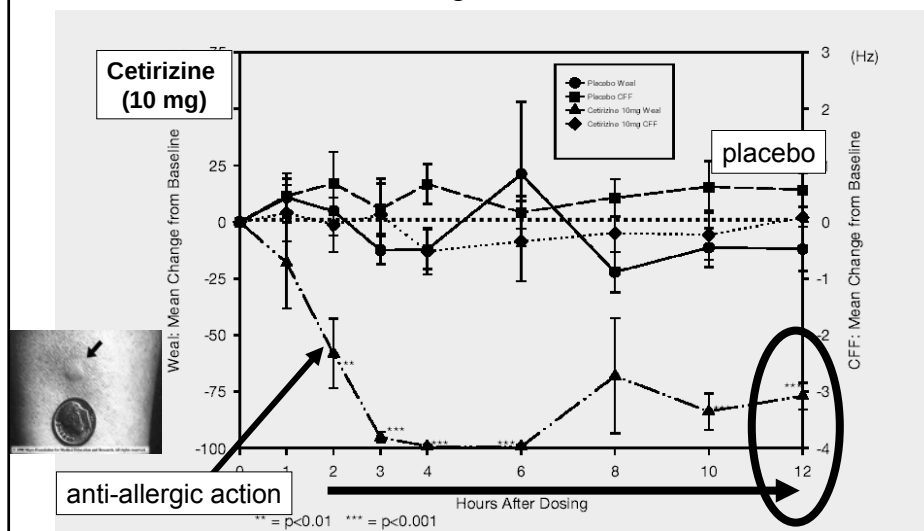
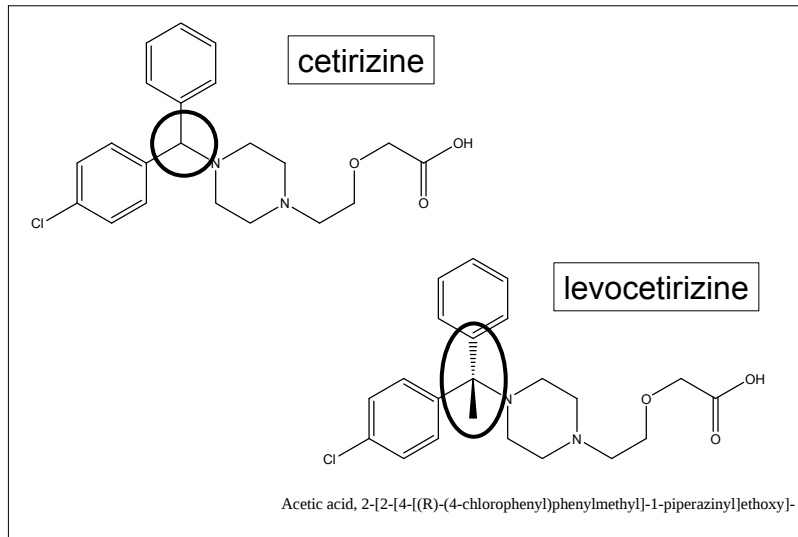


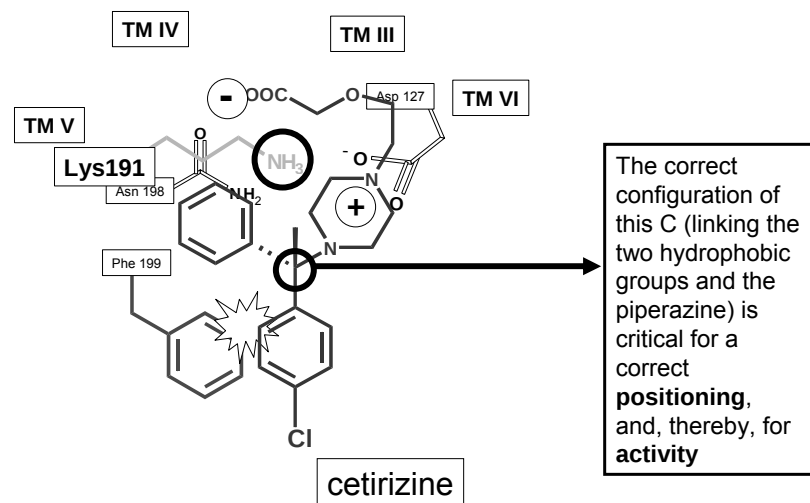
Figure 5. Change from baseline: peripheral antihistaminic suppression (weal) with respect to CFF threshold: acute dose cetirizine 10mg, day 1

Hindmarch et al., Curr. Med. Res. Opin., 17:241-255, 2001

Cetirizine and levocetirizine....



Optic isomers and binding to the receptor ...



Do you speak English in Braine l'Alleud or Gosselies ?



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Mol Pharmacol 61:391-399, 2002

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Binding Characteristics of Cetirizine and Levocetirizine to Human H₁ Histamine Receptors: Contribution of Lys¹⁹¹ and Thr¹⁹⁴

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This paper is available online at <http://molpharm.aspetjournals.org>

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Une longue histoire, mais ...

Sur base de ce que vous avez appris jusqu'ici, quel
est, à votre avis, le conseil le plus essentiel à
donner au patient lors de la délivrance d'un
antihistaminique de type cétérizine, loratadine ...

- faire attention aux autres médicaments
- ne pas abuser du produit
(ne pas reprendre trop rapidement)
- attention à l'alcool !
- respecter la posologie
- attention à la somnolence

Et pour la suite ...

Sur quel point, selon vous, le pharmacien doit-il être particulièrement bien informé par la firme XXX à propos de YYY qui est en vente sans prescription ?

- les propriétés de base du produit
- les indications
- la posologie
- les effets indésirables
- les interactions médicamenteuses

Et pour la vraie suite ...

Quelle est la différence entre A et B ... ?

Cétirizine UCB (UCB)
[cétirizine dichlorhydrate]
compr. (séc.)
20 x 10 mg € 7,00
40 x 10 mg € 12,00

Cétirizine-Ratiopharm (Ratiopharm)
[cétirizine dichlorhydrate]
compr. (séc.)
7 x 10 mg
20 x 10 mg

Zyrtec (UCB)
[cétirizine dichlorhydrate]
compr. (séc.)
7 x 10 mg € 5,21
20 x 10 mg € 9,99
40 x 10 mg € 16,25
gîtes
20 ml 10 mg/1 ml
(1 ml = 20 gîtes)
sol. (oral)
200 ml 5 mg/5 ml

A: Générique, donc sans supplément au ticket modérateur et en catégorie bon marché

B: Médicament original avec supplément au ticket modérateur, donc pas en catégorie bon marché

C: Médicament original avec supplément au ticket modérateur, donc pas en catégorie bon marché

Et pourquoi C est-il sous prescription ?

Et voyons les applications ...

