

Antidepressants

Pharmacology and Toxicology Central Nervous System Module Third Year Medical Students Tareq Saleh Faculty of Medicine

The Hashemite University



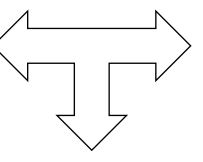


Mood Disorders

Major depressive disorder

- <u>2 weeks</u> of at least **5** of the following symptoms:
- Depressed mood
- Anhedonia (diminished interest or loss of pleasure in almost all activities)
- Weight change or appetite disturbance
- Sleep disturbance (insomnia or hypersomnia)
- Psychomotor agitation
- Fatigue or loss of energy,
- Feelings of worthlessness, diminished ability to think or concentrate;
- suicidal ideation or a suicide attempt

Mood disorders



<u>Others</u>

Bipolar disorder

- periods of prolonged depression that alternate with periods of an excessively elevated mood (mania)
- Manic episodes: <u>1 week</u> of at least **3** of the following symptoms:
- Grandiosity
- Diminished need for sleepexcessive talking or pressured speech
- Racing thoughts or flight of ideas-distractibility
- Increased level of goal-focused activity at home, at work, or sexually
- excessive pleasurable activities





Pathophysiology of Depression

• NOT fully understood.

Monoamine Theory of Depression:

- norepinephrine (NE), dopamine (DA) & serotonin (5-HT) are neurotransmitters responsible for mood.
- Depression is due to a deficiency in monoamines such as NE and 5-HT.
 Very simplistic----fails to explain the long time course of most antidepressants.





Treatment of Depression





Classes of antidepressants

- Selective serotonin reuptake inhibitors (SSRIs)
- Serotonin/norepinephrine reuptake inhibitors (SNRIs)
- Tricyclic antidepressants (TCAs)
- Atypical antidepressants
- Monoamine oxidase inhibitors (MAOIs)
- Serotonin-Dopamine Activity Modulators (SDAMs)





SELECTIVE SEROTONIN REUPTAKE INHIBITORS (SSRIs)

Citalopram CELEXA Escitalopram LEXAPRO Fluoxetine PROZAC Fluvoxamine LUVOX CR Paroxetine PAXIL Sertraline ZOLOFT



Selective Serotonin Reuptake Inhibitors (SSRIs)

Mechanism of action

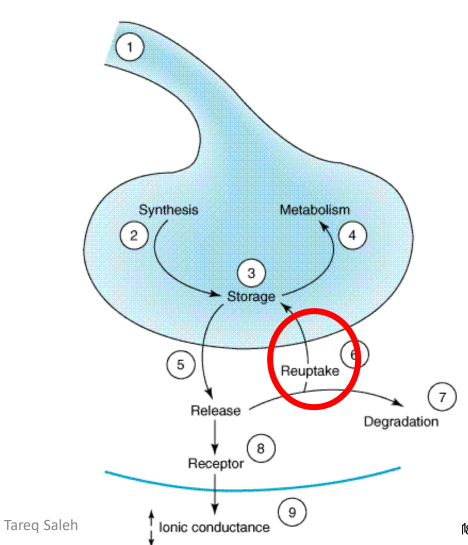
• SSRIs <u>block the reuptake of</u> <u>serotonin</u> →increase its concentrations in the synaptic cleft.

DRUG	UPTAKE INHIBITION			
	Nor- epinephrine	<mark>Serotonin</mark>		
Selective serotonin reuptake inhibitor <i>Fluoxetine</i>	0	++++		
Selective serotonin/ norepinephrine reuptake inhibitors <i>Venlafaxine</i>	++*	++++		
Duloxetine	++++	++++		
Tricyclic antidepressants				
Imipramine	++++	+++		
Nortriptyline	++++	++		





Sites and Mechanisms of CNS Drug Action



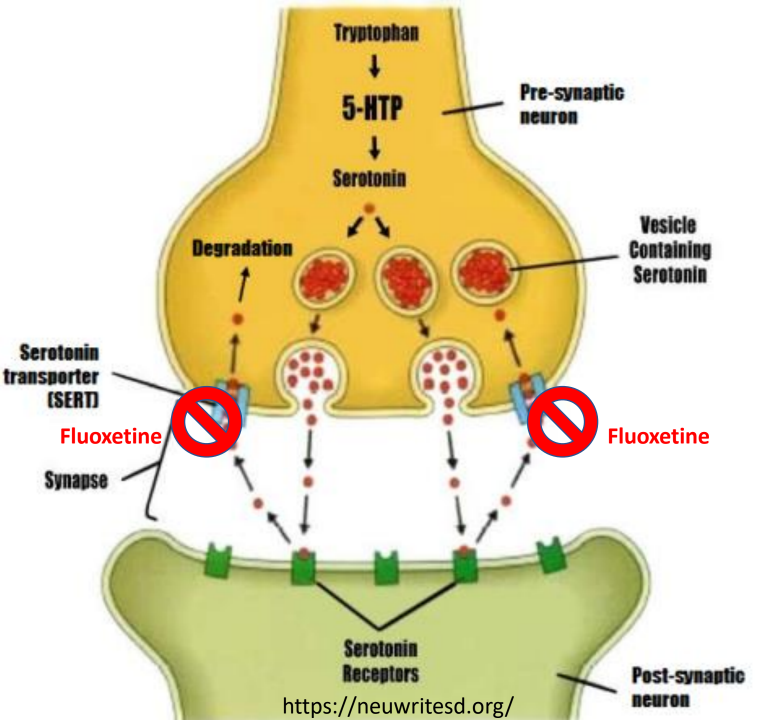
NT reuptake:

•Antidepressants



Selective Serotonin Reuptake Inhibitors (SSRIs)

- Fluoxetine inhibits SERT and interferes with serotonin reuptake.
- This results in increased serotonin availability in the synaptic cleft.









Therapeutic uses

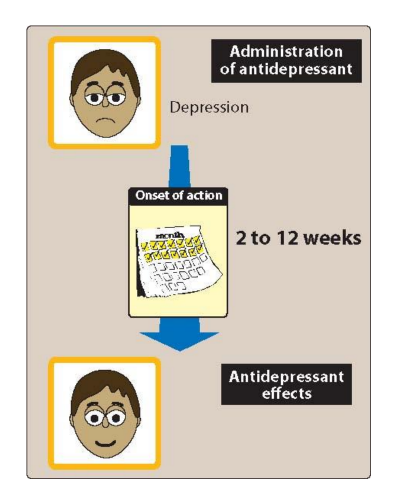
- **Depression** (The primary indication)
- Obsessive Compulsive disorder (OCD)
- Panic disorder
- Generalized anxiety disorder
- Social anxiety disorder
- Post-traumatic stress disorder
- Premenstrual dysphoric disorder
- Bulimia Nervosa (*Only fluoxetine*)





Therapeutic Effect

• SSRIs require 2 weeks to establish a significant alteration in mood (up to 12 weeks and more).







Pharmacokinetics

• Oral.

- Food has little impact on their absorption (*except for sertraline, for which food increases its absorption*).

• Metabolized by CYP450 enzyme family

- fluoxetine differs from the other members of the family in that it has a much longer half life (~50 hours), and the half life of its metabolite can be longer than 10 days.

- fluoxetine and paroxetine are a potent inhibitors of CYP2D6



Selective Serotonin Reuptake Inhibitors (SSRIs)

Adverse effects

- Headache, sweating, nausea, vomiting and diarrhea.
- Sleep disturbances:
 - -Paroxetine and fluvoxamine are sedative
 - -Fluoxetine and sertraline are more activating.
- Sexual dysfunction: loss of libido, delayed ejaculation, anorgasmia.
 - Very common
 - Could require switching to another family of antidepressants









Anxiety

Drowsiness

Nausea



Sexual dysfunction

Drua



Adverse effects

- Overdose: "serotonin syndrome" especially when used with another MAOi (includes seizures, hyperthermia, muscle rigidity, sweating, myoclonus, ...)
- **Discontinuation syndrome**: occurs due to abrupt withdrawal (includes headache, malaise, flu-like symptoms, irritability, nervousness, sleep disturbances).

Particularly by the agents with the shorter half-lives.

SSRIs should be discontinued gradually.





Use of SSRIs in Children/Adolescents

• Used with caution.

[reports of suicidal ideation]

- Fluoxetine and escitalopram are approved to treat childhood depression.
- Fluoxetine, fluvoxamine and sertraline are approved to treat OCD in children



Serotonin/Norepinephrine Reuptake Inhibitors (SNRIs)



SEROTONIN/NOREPINEPHRINE REUPTAKE INHIBITORS (SNRIs) Desveniafaxine PRISTIQ

Duloxetine CYMBALTA Levomilnacipran FETZIMA Venlafaxine EFFEXOR

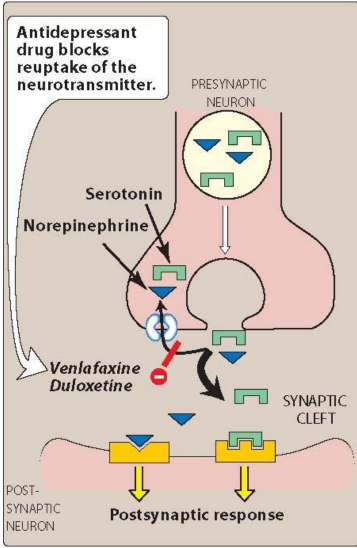


Serotonin/Norepinephrine Reuptake Inhibitors (SNRIs)



Mechanism of action

• SNRIs inhibit the <u>reuptake of</u> <u>BOTH serotonin and</u> <u>norepinephrine</u>





Serotonin/Norepinephrine Reuptake Inhibitors (SNRIs)

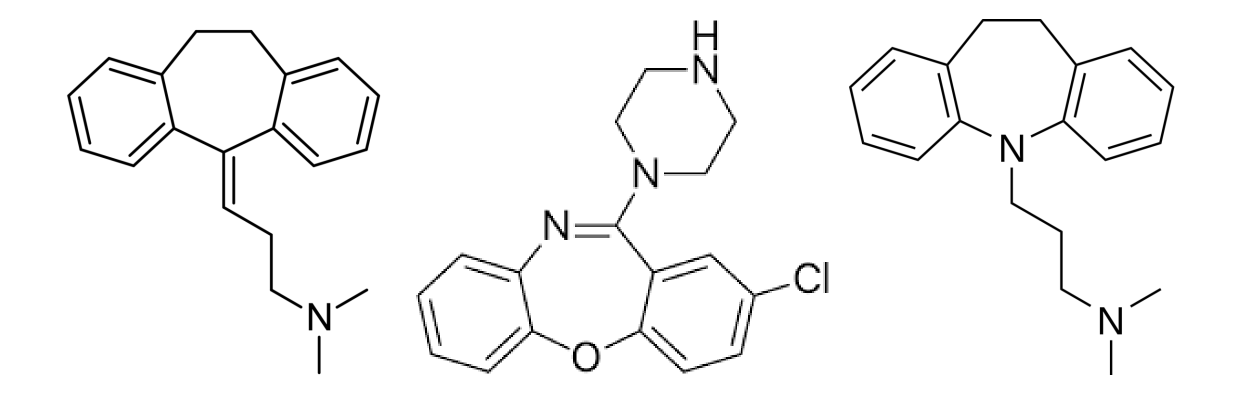


Therapeutic uses

- **1. Depression** (when SSRIs are ineffective).
- 2. Depression accompanied by a chronic painful condition.
- **3. Neuropathic Pain** (diabetic neuropathy, postherpetic neuralgia, fibromyalgia, etc....)

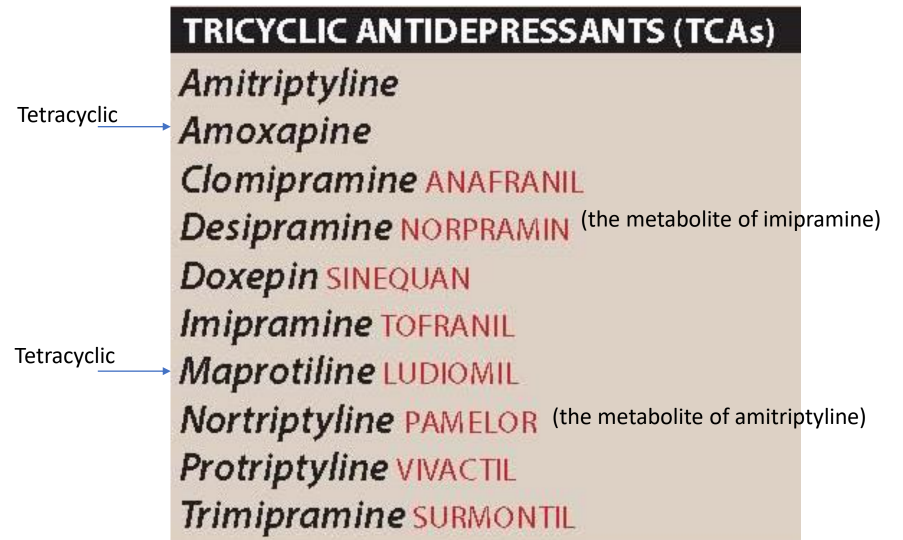
















Mechanism of action

- Inhibition of neurotransmitter (NE and 5-HT) reuptake:
- Receptor antagonism:

- TCAs also block <u>serotonergic</u>, α-adrenergic, histaminic and <u>muscarinic receptors</u>.

- Amoxapine also blocks <u>5-HT₂</u> and dopamine <u>D₂ receptors</u>.

many of the side effects of TCAs result from this non-selective receptor antagonism.

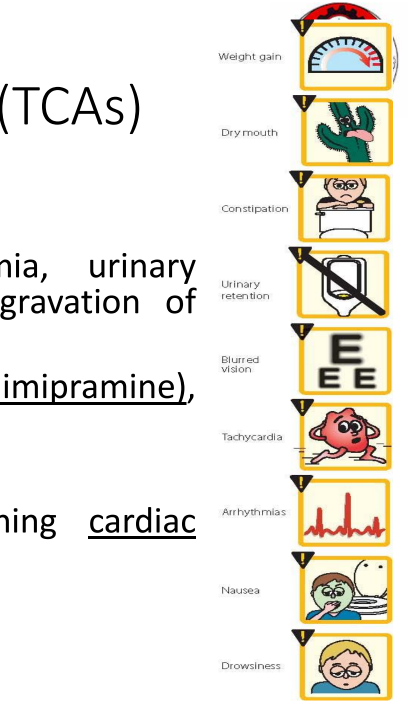




Therapeutic uses

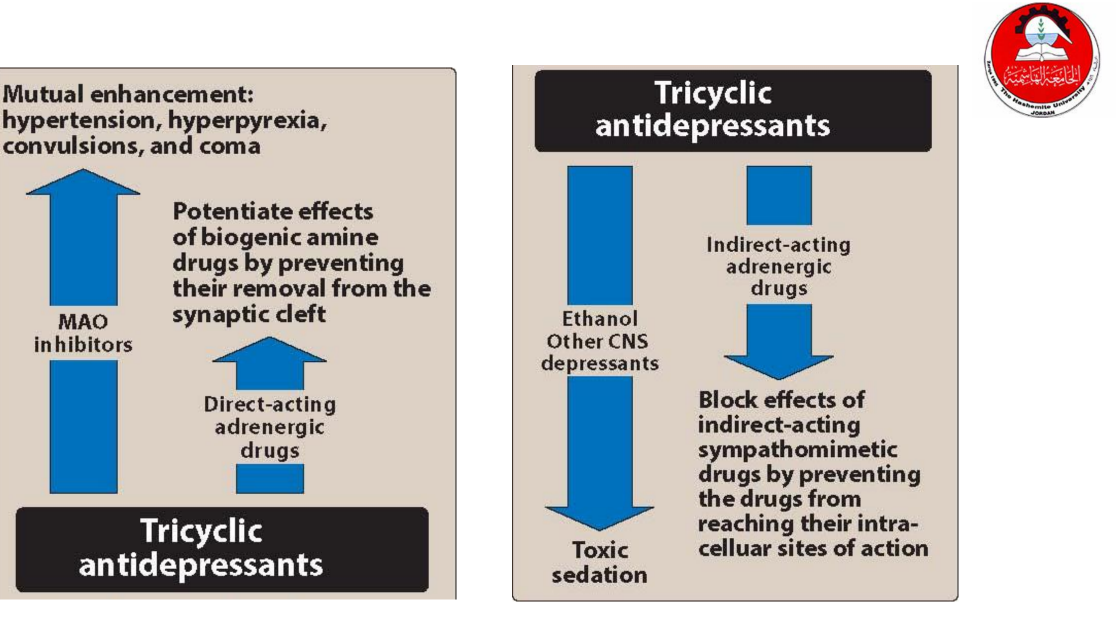
- Moderate to severe depression
- Panic disorder
- Nocturnal enuresis (bedwetting): Imipramine (largely replaced by desmopressin).
- Migraine and chronic pain conditions: amitriptyline.
- Insomnia: doxepin.





Adverse effects

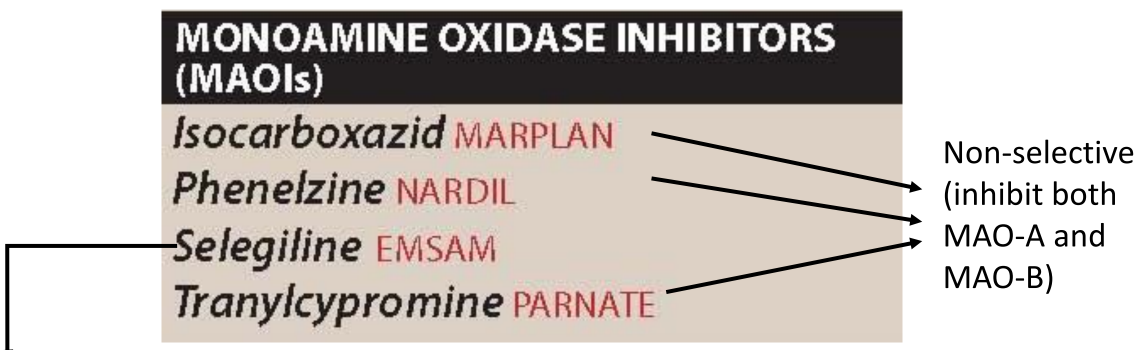
- **Muscarinic blockade:** blurred vision, xerostomia, urinary retention, sinus tachycardia, constipation and aggravation of angle-closure glaucoma.
- α-adrenergic blockade: <u>orthostatic hypotension (imipramine)</u>, dizziness and reflex tachycardia.
- H1 histamine blockade: sedation.
- Overdose: can be associated with life-threatening <u>cardiac</u> <u>arrhythmias</u>.
- Sexual dysfunction: less than SSRIs.



MAO







-Selective for MAO-B

-also used for the treatment of Parkinson's disease.





The use of MAOi is limited (last line) due to the dietary restrictions required while taking these agents, toxicity and drug-drug interaction.

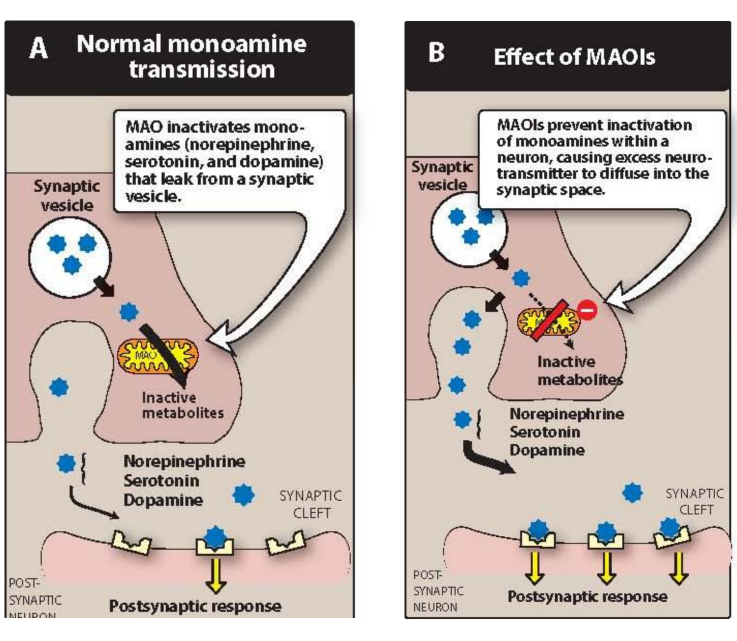




Mechanism of action:

- MAO enzyme exists in 2 forms:
- 1- MAO-A: responsible for metabolism of NE and 5-HT.
- 2- MAO-B: more selective for dopamine (DA) metabolism.
- Most MAOis form stable complexes with the enzyme causing <u>irreversible inactivation</u>.
- Inhibition of MAO results in 个 NE + 5-HT + DA











Mechanism of action:

• The action of MAOi is delayed for several weeks.

MAOi also interfere with hepatic and intestinal isoforms of the enzyme which accounts for their high drug-drug and food-drug interactions.





Therapeutic uses:

- Last line for the treatment of depression: for patients who are unresponsive to SSRIs or TCAs.
- Atypical depression.





Adverse effects:

- Orthostatic hypotension, insomnia and convulsions.
- Hepatotoxicity (Phenelzine).
- Serious food (tyramine-rich) and drug interactions.



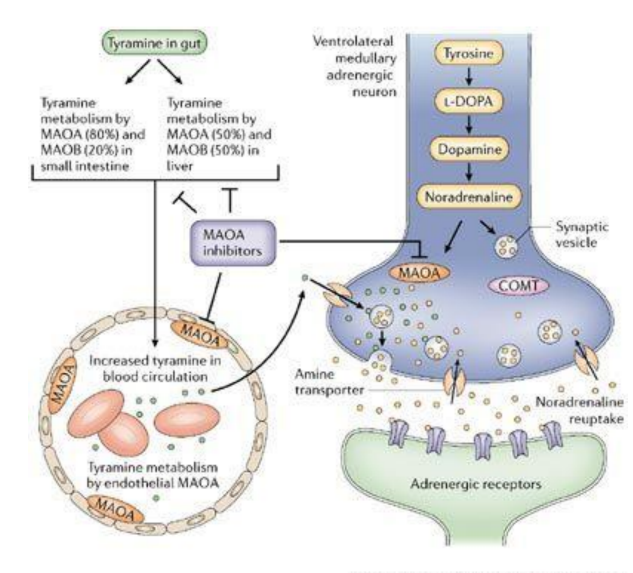


Tyramine-rich diet and MAOi

- Tyramine is contained in foods such as aged cheese, meats, chicken liver, smoked fish and red wine.
- Tyramine is inactivated by hepatic and intestinal MAOs.
- MAOi interfere with the degradation of dietary tyramine.
- Tyramine accumulation results in the release of large amounts of stored catecholamines → Hypertensive crisis!!!







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> Youdim et al, 2006 Wolters Kluwer

MAOIs +					
1. Tyramine -	2. Cold Remedies	3. TCAs	4. Pethidine	5. SSRIs	
rich food	(sympathomimetic)	(† CA)		(† 5HT)	
t	↓ ↓	Ļ	↓ ↓	ł	
Hypertensive crisis	Hypertensive	-Hypertension	-Respiratory	"Serotonin	
(Cheese reaction)	Crisis.	-Hyperthermia	depression	syndrome":	
Tyramine in food is metabolized in GIT by MAO-A & MAO-B. MAOIs allow tyramine in tyramine-rich food (old cheese, chicken liver, chocolate) to escape metabolism & release ↑ ↑ ↑ amounts of catecholamines from neurons → hypertensive crisis.		-Convulsions	-Hyperthermia -Convulsions	-Hyperthermia -Convulsions	





Precautions with MAOi

- Patients on nonselective MAOIs should be warned against serious drug interactions and should be given a list of the foods they should avoid.
- Patients on MAOIs should not receive TCAs or SSRIs except after 2 weeks from stopping MAOIs (effect persists for 2 weeks or 6 for fluoxetine).
- Avoid in the elderly because of postural hypotension.





Atypical antidepressants

ATYPICAL ANTIDEPRESSANTS

Bupropion WELLBUTRIN, ZYBAN Mirtazapine REMERON Nefazodone Trazodone DESYREL Vilazodone VIIBRYD Vortioxetine BRINTELLIX





Bupropion

- Mechanism of action: Weak DA and NE reuptake inhibitor
- **Therapeutic uses:** <u>Depression</u> and <u>smoking cessation</u> (reduces cravings and attenuates nicotine withdrawal symptoms.
- Adverse effects: associated with a dose-dependent increased risk for seizures.

-----it has a very <u>low</u> incidence of sexual dysfunction.



Mirtazapine

 Mechanism of action: presynaptic α₂ antagonist and partially due to 5-HT₂ antagonism (enhances serotonin and norepinephrine neurotransmission)



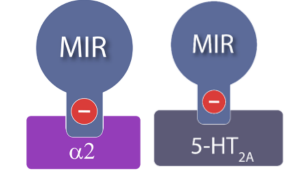
-patients intolerant to TCAs or SSRIs.

-sedating antidepressant improve insomnia

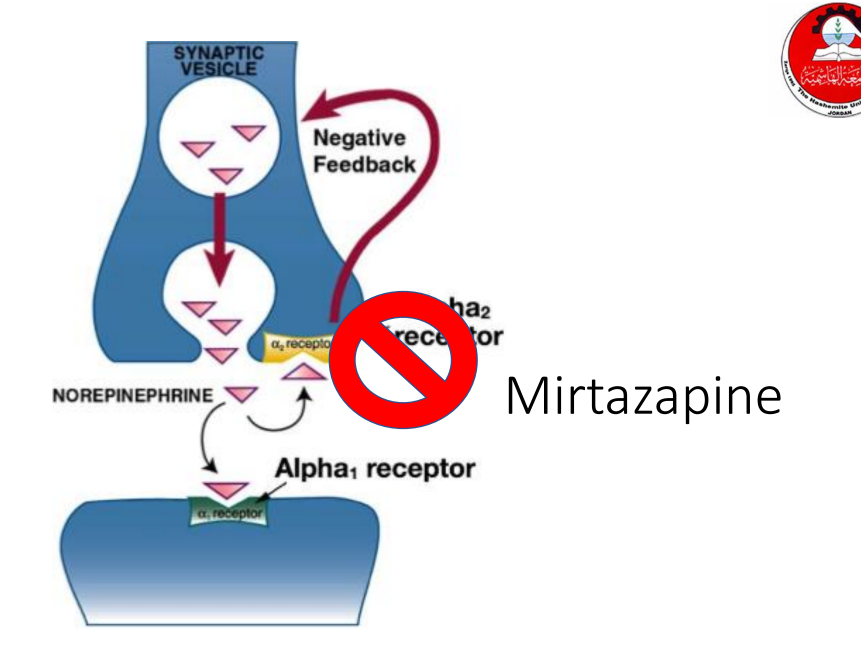
• Advantages: <u>No</u> sexual dysfunction, nausea, anxiety of SSRIs.















Other atypical antidepressants

- Nefazodone and trazodone: weak serotonin reuptake inhibitors + 5-HT_{2a} antagonists + H₁-blocking + α_1 antagonism
- Vilazodone: serotonin reuptake inhibitor + 5-HT_{1a} partial agonism
- **Vortioxetine**: serotonin reuptake inhibitor + 5-HT_{1a} agonism + 5-HT₃ and 5-HT₇ antagonism





Novel therapies

- Brexpiprazole.
- Serotonin-dopamine activity modulator.
- Reading assignment:
- https://www.ncbi.nlm.nih.gov/pubmed/26849053





Good news?

NMDA receptor antagonists

• Esketamine







Summary of antidepressants mechanisms of action

Amine Pump Inhibitors		MAO Inhibitors	<u>Presynaptic</u>
Inhibit uptake-I of biogenic		Inhibit metabolism of	<u>α2</u> Blockers
amines into neurons resulting in		biogenic amines by	↑ NA release into
their accumulation in synaptic		MAO enzyme inside	synaptic cleft by
cleft, potentiating their action at		nerve endings $\rightarrow \uparrow$ stores	preventing α_2
post synaptic receptors.		available for release.	auto-inhibition.
Members		Members	<u>Members</u>
1. TCAs	2. TTAD	Tranylcypromine	Mirtazapine
3. SSRI	4. NSRI	Phenelzine	
5. Bupropion		Moclobemide	

Mechanisms of Increase of Biogenic Amines by Antidepressants

 TCAs: Tricyclic antidepressants
 NSRI: Norepinephrine Serotonin Reuptake Inhibitor

 TTADs: Tetracyclic antidepressants
 SSRIs: Selective Serotonin Reuptake Inhibitor.





Drug class used as first-line therapy of major depressive disorder is <u>SSRIS</u>

Consuming aged cheese and meat is contraindicated while on MAOIS for the treatment of depression

How can you manage major depression in patients on SSRI that are suffering from persistent sexual dysfunction? Switch to atypical antidepressants





Overall Therapeutic Strategy

- The **goal** of initial treatment for depression is symptom remission and restoring baseline functioning.
- The treatment strategy includes *combination of pharmacotherapy and psychotherapy* (based upon randomized trials that found combination treatment was more effective than either of these treatments alone).
- First line treatment: SSRIs
- Alternatives: second generation antidepressants: SNRIs, atypical antidepressants and serotonin modulators.
- TCAs and MAOis are typically **not** used as initial treatment because of concerns about safety and adverse effects.





Treatment of Bipolar Disorder





Drugs Used to Manage Bipolar Disorder

DRUGS USED TO TREAT MANIA and BIPOLAR DISORDER

Carbamazepine TEGRETOL, EQUETRO, CARBATROL

Lamotrigine LAMICTAL Lithium Valproic acid DEPAKENE, DEPAKOTE





Lithium

- Used acutely and prophylactically for managing bipolar patients. (effective in 60-80% of patients).
- Mechanism of action: Unknown.
- Pharmacokinetics:
 - very narrow therapeutic window (highly toxic).

- entirely eliminated by <u>renal clearance</u> (best choice in patients with hepatic dysfunction)

- Adverse effects: headache, xerostomia, polyuria, polydipsia, polyphagia, dermatologic reactions and <u>sedation</u>.
- Toxicity: <u>ataxia</u>, <u>slurred speech</u>, <u>confusion</u>, <u>seizures and thyroid</u> <u>dysfunction</u>.





Treatment of Bipolar Disorder

Other drugs

- Antiepileptics: Carbamazepine, valproic acid and lamotrigine.
- Antipsychotics: Chlorpromazine, haloperidol, risperidone, olanzapine, aripiprazole.





- Thank you
- Questions?

Please contact me tareq@hu.edu.jo

