

TECHNICAL UNIVERSITY OF MOMBASA

FACULTY OF APPLIED AND HEALTH SCIENCES

DEPARTMENT OF PURE & APPLIED SCIENCES

UNIVERSITY EXAMINATION FOR:

BTAC

ACH 4402: MEDICINAL CHEMISTRY II

END OF SEMESTER EXAMINATION

SERIES: APRIL 2016

TIME: 2HOURS

DATE: 9th May 2016

Instructions to Candidates

You should have the following for this examination

-Answer Booklet, examination pass and student ID This paper consists of **FIVE** questions. Attempt question ONE (Compulsory) and any other TWO questions.

Do not write on the question paper.

Question ONE:

- a) Define the following terms as used in medicinalchemistry.
 - i. Anticonvulsant and antiepileptic drugs(AEDs)
 - ii. Cholinergic and anticholinergicdrugs.
 - iii. Sympathomimetics and sympatholytics,
 - iv. sedative-hypnoticsandantipsychotics

(4marks)

b) Sevoflurane is a general anesthetic that reacts with desiccated carbon dioxide adsorbents, to produce compounds (A and B) with knowntoxicity.

$$F H H H$$

$$F - C C O C F \longrightarrow A + B$$

$$F CF_3 H$$

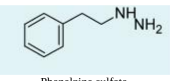
Sevoflurane

- i. Draw the structures of compounds AandB (2mks)
- ii. The degree to which sevoflurane breaks down is influenced by three factors. Name the threefactors

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(3 mks)

- c) Antidepressant therapy usually implies therapy directed against major depressive disorders of the unipolar type and is centered on three groups of chemical agents. Namethesegroups (3mks)
- **d**) Phenelzine sulfate, an effective antidepressant agent irreversibly inactivates the enzyme or its cofactor via irreversibleinhibition.

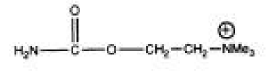


Phenelzine sulfate

- i. Why do we say that phenelzineisirreversible (2mks)
- ii. Discuss the biotransformation of phenelzine to the active species in irreversibleinhibition

(5 mks)

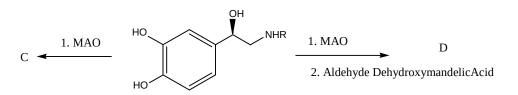
e) Carbachol is a long acting cholinergic agent. It is certainly stable to hydrolysis and is the right size to fit the cholinergicreceptor.



Carbachol

- i. Explain why carbachol is resistant to hydrolysis morethanacetylcholine (3mks)
- ii. Explain why even after a hydrophobic methyl group has been replaced with a polar NH₂ group, carbachol does fit the cholinergic receptor andisactive (2mks)
- iii. Give reasons as to why Carbachol though a long acting cholinergic agent is used locally and cannot be ingested (2mks)
- f) In the metabolism of catecholamines (CAs), monoamine oxidase (MAOs) oxidatively deaminate CAs to their corresponding aldehydes, which are rapidly oxidized to the corresponding acid by the enzyme aldehyde dehydrogenase (AD). In some circumstances, the aldehyde is reduced to the glycol by aldehyde reductase (AR). Propose structures of compounds C and D formed from the metabolism of CAs

(3 mks)



Question TWO:

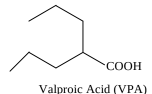
a) In the structure activity relationship (SAR) of anesthetics, the chemical structure is related to the activity of the drugmolecule.

i.	Discuss the potency of alkanes, cycloalkanes, and aromatic hydrocarbons in relation to the anesthetic			
	activity		(6mks)	
ii.	Explain t	n the reason for the reduced activity of the anesthetic compounds beyond their cut-offnumber		
			(2 mks)	
	iii.	Discuss the effect of halogenationonanesthetics	(3mks)	

b) At cellular level, three basic mechanisms are believed to contribute to the antiepileptic action of the currently marketed anticonvulsants. State the three mechanisms of action of anticonvulsants

c)	State four possible causes of epilepsyandconvulsions	(4mks)

4. Valproic Acid (VPA) is an established AED with a simple chemical structure but an unusually broad spectrum of action. It is generally well tolerated, but its use is limited by two rare but significant toxic side effects (hepatotoxicity and teratogenicity).



Atropine and the cholinergic receptor that makesthispossible.

Identify two possible reactive metabolites of VPA that are responsible for this observation

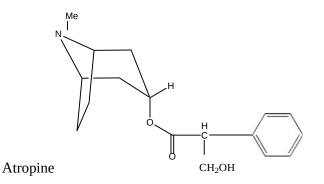
(2 mks)

(3 mks)

Question THREE

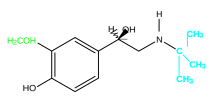
a) Differentiate between nucleotidesandnucleosides (2mks.
 b) Explainindetailsthebiosynthesisofacetylcholineatsynapsesinvolvingacetylcholineastheneurotransmitter (7 mks).
 c) Explain why it is not feasible to give a patient more acetylcholine if there is lack of acetylcholine acting at a certain part ofthebody (3mks).
 d) Antidepressant therapy usually implies therapy directed against major depressive disorders of the unipolar type and is centered on three groups of chemical agents. Name these threegroups
 e) Atropine is an antagonist of the cholinergic receptor. It can bind to and block the receptor. Explain the SAR of

(5mks)



Question FOUR

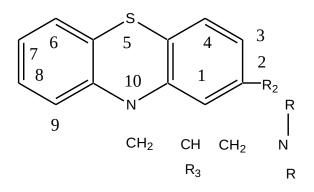
- a) To which class of compounds do adrenaline, noradrenaline anddopaminebelong? (1mk)
- **b**) Name the natural chemical messenger for the following receptors
 - i. Adrenalinereceptor (1mk)
 - ii. GABAreceptors (1mk)
- c) Salbutamol is an important clinical agent. The coloured groups are modifications that distinguish the molecule from adrenaline ornoradrenaline.



Salbutamol

- i. What role does the t-butylgroupplay? (2mks)
- ii. Why was the hydroxymethylenegroupintroduced? (3mks)
- **d**) Phenothiazine is an antipsychotic agent. The best position for substitution is the 2-position. Activity increases (with some exceptions) as electron-withdrawing ability of the 2-substituent increases. Explainwhy.

(4 mks).



e) 2-Thioimidazole derivatives are antithyroid agents whose mechanism of action is by prevention of the iodination of the precursors of thyroxine andtriiodothyronine.



These compounds are absorbed well after oral administration and excreted in the urine. However, they are tautomeric compounds. Draw three tautomeric compounds of 2-thioimidazole

		(3 mks)
f)	The mechanisms of action of all antithyroid a	gents are similar. State the mechanism ofaction
		(1mk)
g)	Discuss the four stages of general anesthesia	(4mks)

Question FIVE

a)	Define	e the following terms as used in medicinalchemistry	
	i.	Antihyperlipidermicagents	(1mk)
	ii.	Hyperlipidemia	(1mk)
	iii.	Hyperthyroidism	(1mk)
	iv.	Nucleotidesand Nucleosides	(2mks)
b)	What	is the disadvantage of using adrenaline as anadrenergicagonist	(2mks)
c)	Lipids pathw	are transported in the body by both exogenous and endogenous pathways. I ays	Differentiate between the two (4mks)
c) d)	pathw		
->	pathw	ays	
->	pathw There	ays are two mechanism of action of inhaledanesthetics.	(4mks)