Cheatography

Anti-Epileptic Agents 4				
1. Carboxamide derivatives	2. Benzodiazepines	3. Barbiturates	4. Hydantoin derivative	5. Fatty acid derivative
Cabamazepine	Clonazepam	Phenobarbital	Phenytoin	Valproate
	Diazepam			
	Lorazepam			
	Midazolam			

1.Cabamazepine

Trade	Tegretol, Degranol and
names:	Sandoz-Carbamazepine
Oral dosage	sustained release tablet
form tablet:	and suspension

Primary drug for partial and generalized tonic clonic seizures

Not effective in absence seizure

Mechanism of action 4

 In seizure dysfunctional sodium channels allow for too much influx of Sodium ion which prolongs neuronal excitability (depolarization)

 Carbamazepine limits repetitive firing of action potential

(limits too much influx of sodium ion)

This is due to prolonging the Sodium channel inactivation

Side effects and Toxicity

Cardiac arrhythmias with or without hypertension

Behaviroal change

Gastroinstinal symptoms

Hirudism

Megaloblastic anemia

Pharmacokinetics 4

• Phenytoin distributes into the body tissues, including the brain,

within 30 to 60 minutes after reaching the systemic circulation

Effective for up to 24 hours

Metabolized by hepatic cytochrome P450

By SmeNdlela cheatography.com/smendlela/

Drug interactions:

Reduces the effect of oral contraceptives	Reduces the effect of warfarin		
Could result in unplanned pregnancy	Could result in deep vain thrombosis		
Cytochrome P450 inducer			

 5.Valproate

 Trade
 Epilium liquid, Epilium CR,

 Names:
 Epilium IV, Epilium crushable,
Convulox, Navalpro, Eprolep

 Oral and
 Tablet, capsule, suspension,
parantral

 IV
 IV

 dosage
 Form:

Effective against absence, myoclonic partial and tonic clonic seizures, suitable for **HIV positive children with epilepsy**

Mechanism of action 4

- Mechanism of action is not fully understood
- It is believed that valproic acid leads to increased production of GABA

 In addition to this, valproate is also thought to enhance the effect of GABA that

- already exists in the area on the receptors.
- Valproate mimics the action of GABA

Pharmacokinetics

• Absorbed rapidly and completely after oral administration with 81-89% bioavailability. Active for around 9-16 hours

Metabolized by hepatic cytochrome P450

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Side Effects and Toxicity 4

Anorexia
Vomiting
Nausea
Sedation atexia
Tremor
Rash
Alopecia
Increased appetite

Drug interactions

All benzodiazepines cause excessive sedation when combined with other medications that slow the brain's processes. (i.e, alcohol, barbiturates narcotics, and tranquilizers)

For Benzodiazepines

2. BENZODIAZ	ZEPINES 5	
Drug type:	Trade names:	Indica- tion:
Clonazepam	Rivotril, Clonam	Drops, tablet, IV
Diazepam	Pax, Valium, A- Lenon, Transjet, Betapam, Doval	Tablet, IV
Lorazepam	Ativan, Tranqipam	Tablet, IV
Midazolam	Accord, Sabax, Midaium, Dormicum, Midazoject	IV, tablet, INF

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Cheatography

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Mechanism of	action 4			Benzodiaze	pines (cont)			Benzodiaze	epines (cont)
G Vig Na" E		BA Ca ²⁴	49	Effective in Status Epilep- ticus	Rarely used for long term	Rarely used for long term	used fo	ortolerance	g term use - develops metabolized in liver and urine
Indication of E Prolonged s minutes betwee Can last up Emiliation of the minutes betwee Emiliation of the minutes betwee Can last up EML recommission	eizure activit een seizures to 20 minutes mends which	republic network action problem proble	a ^{2*} de	Absence seizure	Good oral absorption	Better than Diazepam	Epilep- ticus th has no improv followin other	ve Side Effect Drowsiness at Blurred visi eSurred spe duack of coor Difficulty br ents Coma For Benzoor	s and Toxicity f s and confusion on eech ordination and weakness eathing diazepines
In Status Epile Mechanism of	-			Given in	Completely	Good oral		t ⊅fade ≎ hames oral	Phenobarb Vital, Propain Forte, Donatal Elexir, Lethyl Sedabarb
	pines cause a tor and openi	-	ie	combin- ation	metabo- lized in liver and excreted in	absorption	absorp	dosage form	Tablet, syrup
receptor • The neurona	al membrane	is hyperpola-			urine			Low toxicity young child	 inexpensive, widely used for lren
rized and is le	ss likely to fir excitatory glu		ors	Good oral absorption		Completely metabo- lized in liver and			
Clonazepam	Diazepam	Lorazepam	Midaz	olam		excreted in urine			
Used for broad spectrum of seizures	First line treatment in Status Epilep- ticus	First line treatment in Status Epilepticus	First I treatn in Sta Epilep	nent tus					



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3. Phenobarbital (cont)

Non-selective to the type of seizure. Mainly used for generalized tonic clonic and partial seizures and status epilepticus 20mg/kg, crushed and given by nasogastric tube (if one dose of midazolam or two doses of diazepam fail to show response)

Mechanism of action 4

Through its action on GABA receptors, phenobarbital increases flux of chlorine ions into the neuron which decreases excitability.

□ Direct blockade of excitatory glutamate signaling is also believed to contribute to the hypnotic/anticonvulsant effect

Pharmacokinetics

- · Oral absorption is complete but slow
- Peak concentration in plasma is seen after several hours (8-12 hours) after oral administration
- Remains in the body for a long time (2-7 days)
- Metabolized by liver (cytochrome p450)

Side effects and Toxicity

Sedation (tolerance develops after continues use)

Driving and use of heavy machinery must be avoided

Irritability and hyperactivity in children

Agitation and confusion in the elderly

Some skin allergies in rare cases

Drug interactions:

Reduces the effect of	
oral contraceptives	

By SmeNdlela

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Reduces the effect of warfarin

Drug interactions: (cont)

Could result unplanned pregnancy		ould result in deep in thrombosis
Cytochrome	P450 inducer	
4. Phenytoin		
Trade Names:		rte suspension, atabs, Epanutin d Parantral
Oral and parantral dosage	Tablet, caps IV	ule, suspension,

form:

Active against all types of partial and tonic clonic seizures but not absence seizures

Causes antiseizure activity without causing general CNS depression

Mechanism of action 4

 In seizure dysfunctional sodium channels allow for too much influx of Sodium ion which prolongs neuronal excitability (depolarization)

 Phenytoin Limits repetitive firing of action potential (too much influx of sodium ion)

This is due to slowing the rate of recovery

of Sodium channels from inactivation.

Pharmacokinetics 4

Phenytoin distributes into the body

tissues, including the brain,

within 30 to 60 minutes after reaching the systemic circulation

- · Effective for up to 24 hours
- Metabolized by hepatic cytochrome P450

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Side effects and Toxicity

Cardiac arrhythmias with or without hypert-
ension
Behaviroal change
Gastroinstinal symptoms
Hirudism
Megaloblastic anemia