1.Name of the Medicinal Product Pantobex (Pantoprazole) Delayed Release Tablet 20 mg Pantobex (Pantoprazole) Delaved Release Tablet 40 mg

2. Qualitative and Quantitative Composition

Pantobex Delayed Release Tablet 20 mg

1 Delayed-Release (enteric coated) tablet contains 20 mg pantoprazole (as pantoprazole sodium) for oral use. Active ingredient: 20 mg pantoprazole (as pantoprazole sodium sesquihydrate 22.6 mg)

Pantobex Delayed Release Tablet 40 mg

1 Delayed-Release (enteric coated) tablet contains 40 mg pantoprazole (as pantoprazole sodium) for oral use. Active ingredient: 40 mg pantoprazole (as pantoprazole sodium sesquihydrate 45.1 mg)

3 Pharmaceutical Form

Pantobex Delayed Release Tablet 20 mg Delayed-Release tablets, yellow, oval, biconvex, enteric coated tablets, plain on both the sides, pack size of 30. Pantobex Delayed Release Tablet 40 mg

Delayed-Release tablets, yellow, oval, biconvex, enteric coated tablets, plain on both the sides, pack size of 30.

4. Clinical Particulars

4.1. Therapeutic indications Pantobex Delayed Release Tablet 20 mg

• Mild reflux disease and associated symptoms (e.g. heartburn, acid regurgitation, pain on swallowing).

· Long-term management and prevention of relapse in reflux oesophagitis.

 Prevention of gastroduodenal ulcers induced by non-selective non-steroidal anti- inflammatory drugs (NSAIDs) in patients at risk with a need for continuous NSAID treatment

Pantobex Delayed Release Tablet 40 mg

• In combination with two appropriate antibiotics (see Dosage) for the eradication of H.pylori in patients with peptic ulcers with the 4.4 Special warnings and precautions for use objective of reducing the recurrence of duodenal and gastric ulcers caused by this microorganism Duodenal ulcer

Gastric ulcer

· Moderate and severe reflux oesophagitis

4.2. Posology and method of administration

Pantobex Delayed Release Tablet 20 mg

Pantobex 20 mg delayed-release tablets should not be chewed or crushed, and should be swallowed whole with water

Therapeutic Indication	Posology and method of oral administration
Mild reflux disease and associated	The recommended oral dosage is one gastro- resistant tablet Pantobex 20 mg per
Symptoms (e.g. heartburn, acid	day. Symptom relief is generally accomplished within 2-4 weeks, and a 4-week
regurgitation, pain on swallowing)	treatment period is usually required for healing of associated oesophagitis. If this is
	not sufficient, healing will normally be achieved within a further
	4 weeks.
Long-term management and	For long-term management, a maintenance dose of one delayed-release tablet
prevention of relapse in reflux	Pantobex 20 mg per day is recommended, increasing to 40 mg Pantoprazole per
oesophagitis	day if a relapse occurs.
	Pantobex 40 mg is available for this case. After healing of the relapse the dosage
	can be reduced again to 20 mg Pantoprazole.
Prevention of gastroduodenal ulcers	The recommended oral dosage is one delayed-release coated tablet Pantobex
induced by non-selective	20 mg per day. The use of Pantoprazole 20 mg as a preventive of gastroduodenal
non-steroidal anti-inflammatory drugs	ulcers induced by non-selective nonsteroidal anti-inflammatory drugs (NSAIDs)
(NSAIDs) in patients at risk with a	should be restricted to patients who require continued NSAID treatment and have
need for continuous NSAID treatment	an increased risk to develop gastrointestinal complications. The increased risk
	should be assessed according to individual risk factors, e.g. high age (>65 years),
	history of gastric or duodenal ulcer or upper gastrointestinal bleeding

Pantobex Delayed Release Tablet 40 mg

Pantobex 40 mg delayed-release tablets should not be chewed or crushed, and should be swallowed whole with water

Therapeutic Indication	Posology and method of oral administration
Therapeutic Indication Eradication of H.pylori in combination with appropriate antibiotics	Posology and method of oral administration In Helicobacter pylori positive patients with gastric and duodenal ulcers, eradication of the germ by a combination therapy should be achieved. Depending upon the resistance pattern, the following combinations can be recommended for the eradication of H. pylori: a) twice daily one tablet Pantobex 40 mg + twice daily 1000 mg amoxycillin + twice daily 500 mg clarithromycin b) twice daily one tablet Pantobex 40 mg + twice daily 500 mg netronidazole + twice daily 500 mg metronidazole + twice daily 1000 mg amoxycillin + twice daily 500 mg metronidazole In combination therapy for eradication of Helicobacter pylori infection, the second Pantobex 40mg tablet should be taken before the evening meal. The combination therapy is implemented for 7 days in general and can be prolonged to up to two weeks maximum. If, to ensure healing of the ulcers, further treatment with pantoprazole is indicated, the dosage recommendations for duodenal and gastric ulcers should be considered. A duodenal ulcer generally heals within 2 weeks. If a 2-week period of treatment is not sufficient, healing will be achieved in almost all cases within a further 2 weeks. A 4-week period is usually required for the treatment of gastric ulcers and reflux esonbardits. If this is not sufficient healing will usually usually be achieved within a further
	not sufficient, healing will be achieved in almost all cases within a further 2 weeks. A 4-week period is usually required for the treatment of gastric ulcers and reflux esophagitis. If this is not sufficient, healing will usually be achieved within a further 4 weeks.
Treatment of gastric ulcer and duodenal ulcer	4 weeks. If combination therapy is not an option, e.g. if the patient has tested negative for Helicobacter pylori, take one 40 mg tablet daily. In individual cases, the dose may be doubled (increased to two 40 mg tablet daily) especially when there has been no response to other medicines.

Special Patient Populations

Pediatric patients:

The experience in children is limited. Pantoprazole 20 mg and 40 mg tablet is not recommended for use in children below 12 ye of age due to limited data on safety and efficacy in this age group. Pantoprazole 40 mg powder for solution for injection is recommended for use in patients below 18 years of age.

Impaired hepatic function

A daily dose of Pantoprazole 20 mg should not be exceeded in patients with severe liver impairment (See section 4.4)

In addition. Pantoprazole 40 mg tablet must not be used in combination treatment (e.g. amoxicillin, clarithromycin.) for eradicat of H. pylori in patients with moderate to severe hepatic dysfunction since currently no data are available on the efficacy and safe of pantoprazole in combination treatment of these patients (See section 4.3).

Impaired renal function:

No dose adjustment is necessary in those with impaired renal function.

In addition, pantoprazole 40 mg tablet must not be used in combination treatment (e.g. amoxicillin, clarithromycin) for eradicati of H. pylori in patients with impaired renal function, since currently no data are available on the efficacy and safety of pantopraze in combination treatment for these patients.

Elderly patients:

Generally, no dose adjustment is necessary in elderly patients. However, the daily dose of 40mg should not be exceeded treatment of gastric or duodenal ulcer.

4.3 Contraindications

Pantobex should generally not be used in cases of known hypersensitivity to one of the other constituents of Pantobex 20 Pantobex 40 mg tablet or of the combination partners.

Pantobex 40 mg tablet must not be used in combination treatment for eradication of H.pylori in patients with moderate to seve liver or kidney function disturbances since currently no clinical data are available on the efficacy and safety of Pantoprazole 40 n in combination treatment of these patients

In presence of alarm symptoms In the presence of any alarm symptom (e. g. significant unintentional weight loss, recurrent vomiting, dysphagia, haematemet anaemia or melaena) and when gastric ulcer is suspected or present, malignancy should be excluded, as treatment pantoprazole may alleviate

symptoms and delay diagnosis. Further investigation is to be considered if symptoms persist despite adequate treatment. Hepatic Impairment

In patients with severe liver impairment the daily dose has to be reduced to 20 mg pantoprazole. In patients with severe liver in the daily dose has to be reduced to 20 mg pantoprazole. impairment, liver enzymes should be monitored during therapy regularly. In the case of a rise of the liver enzymes, the treatme should be discontinued. (See Dosage and Method of Administration).

Gastrointestinal infections caused by bacteria, including Clostridium difficile

Pantoprazole, like all proton pump inhibitors (PPIs), might be expected to increase the counts of bacteria normally present in the upper gastrointestinal tract. Treatment with Pantobex may lead to a slightly increased risk of gastrointestinal infections caused bacteria such as Salmonella and Campylobacter. In hospitalized patients, PPI therapy may be associated with an increased risk Clostridium difficile infection.

Published observational studies suggest that proton pump inhibitor (PPI) therapy like pantoprazole may be associated with increased risk of Clostridium difficile-associated diarrhoea (CDAD), especially in hospitalised patients. This diagnosis should I considered for diarrhoea that does not improve (see 4.8 Undesirable Effects). Patients should use the lowest dose and short duration of PPI therapy appropriate to the condition being treated.

Clostridium difficile-associated diarrhea (CDAD) has been reported with use of nearly all antibacterial agents. For more informati specific to antibacterial agents (clarithromycin and amoxicillin) indicated for use in combination with Pantobex, refer to Warning and Precautions sections of those package inserts.

<u>Hypomagnesaemia</u>

Hypomagnesaemia, symptomatic and asymptomatic, has been reported in patients treated with PPIs for at least three months. most cases after a year of therapy. Serious manifestations of hypomagnesamia such as fatigue, tetany, delirium, convulsio dizziness and ventricular arrhythmias can occur but they may begin insidiously and be overlooked. Hypomagnesemia may lead hypocalcemia and/or hypokalemia. In most patients, treatment of hypomagnesemia (and hypomagnesemia associate hypocalcemia and/or hypokalemia) required magnesium replacement and discontinuation of the PPI.

For patients expected to be on prolonged treatment or who take PPIs with medications such as digoxin or drugs that may cau hypomagnesaemia (e.g., diuretics), health care professionals may consider monitoring magnesium levels prior to initiation of treatment and periodically. [See Undesirable effects]. Bone Fractures

Several published observational studies suggest that proton pump inhibitor (PPI) therapy may be associated with an increased r for osteoporosis-related fractures of the hip, wrist, or spine. The risk of fracture was increased in patients who received high-do defined as multiple daily doses, and long term PPI therapy (a year or longer). Patients should use the lowest dose and short duration of PPI therapy appropriate to the condition being treated. Patients at risk for osteoporosis- related fractures should managed according to established treatment guidelines (see Dosage and Administration and Undesirable effects). Concomitant use with Methotrexate

Literature suggests that concomitant use of PPIs with methotrexate (primarily at high dose; see methotrexate prescribin information) may elevate and prolong serum levels of methotrexate and/or its metabolite, possibly leading to methotrex toxicities. In high-dose methotrexate administration a temporary withdrawal of the PPI may be considered in some patients (s Drug Interactions).

HIV protease inhibitors

Co-administration of pantoprazole is not recommended with HIV protease inhibitors for which absorption is dependent on aci intragastric pH such as atazanavir, nelfinavir; due to significant reduction in their bioavailability.

Influence on vitamin B12 absorption

Daily treatment with any acid-suppressing medication over a prolonged period of time (several years) may lead to malabsorpti of cyanocobalamin (vitamin B₁₂) caused by hypo- or achlorhydria. Cyanocobalamin deficiency should be considered in patie with Zollinger-Ellison syndrome and other pathological hypersecretory conditions requiring long-term treatment, individuals with reduced body stores or risk factors for reduced vitamin B₁₂ absorption (such as the elderly) on long- term therapy or if relevant clinical symptoms are observed

Interference with Laboratory Tests

Increased Chromogranin A (CgA) level may interfere with investigations for neuroendocrine tumours. To avoid this interference proton pump inhibitor treatment should be stopped 14 days before CgA measurements

Cutaneous lupus ervthematosus (CLE) and systemic lupus ervthematosus (SLE)

Cutaneous lupus erythematosus (CLE) and systemic lupus erythematosus (SLE) have been reported in patients taking PP These events have occurred as both new onset and an exacerbation of existing autoimmune disease. The majority of PPI-induc lupus ervthematosus cases were CLE.

The most common form of CLE reported in patients treated with PPIs was subacute CLE (SCLE) and occurred within weeks years after continuous drug therapy in patients ranging from infants to the elderly. Generally, histological findings were observed without organ involvement.

Systemic lupus erythematosus (SLE) is less commonly reported than CLE in patients receiving PPIs. PPI associated SLE usually milder than non-drug induced SLE. Onset of SLE typically occurred within days to years after initiating treatment prima in patients ranging from young adults to the elderly. The majority of patients presented with rash; however, arthralgia and cytopenia

	were also reported.					
	Avoid administration	of PPIs for longer that	n medically indicated	. If signs or symptom	is consistent with CLE	E or SLE are noted in
s	patients receiving pan	toprazole, discontinue	e the drug and refer the	e patient to the approp	priate specialist for eva	aluation. Most patients
ot	improve with discontir	nuation of the PPI alor	ne in 4 to 12 weeks. Se	erological testing (e.g	., Antinuclear antibody	may be positive and
	elevated serological to	est results may take le	onger to resolve than	clinical manifestations	S.	
	Co-administration with	h NSAIDs (Applies to	Pantobex 20 mg only)		
	The use of Pantobex	20 mg as a preventi	ve of gastroduodenal	l ulcers induced by n	on-selective nonstero	idal anti-inflammatory
	drugs (NSAIDs) shou	ld be restricted to par	tients who require cor	ntinued NSAID treatm	ent and have an incr	eased risk to develop
	gastrointestinal comp	lications. The increas	sed risk should be as	ssessed according to	individual risk factor	s, e.g. high age (>65
	years), history of gast	ric or duodenal ulcer	or upper gastro-intest	inal bleeding.		
	Combination therapy	(Applies to Pantobex	40 mg only) acribian information fo	u tha raanaatiya duya	a must be abaamiad	
	In the case of combin	Anon merapy, me pre	Scribing information ic	n the respective drug	s must be observed.	
	In long-term treatmen	t, especially when exe	ceeding a treatment p	eriod of 1 year, patier	nts should be kept und	ler regular surveillance
	4.5 Interaction with	other medicinal proc	lucts and other form	is of interaction		
	Drugs with pH-Depen	dent Absorption Phar	macokinetics:			
	Pantoprazole may rec	duce or increase the a	bsorption of drugs wh	nose bioavailability is	pH-dependent (e.g. k	etoconazole)
	HIV medications (e.g.	atazanavir)				
	Co-administration of I	Pantoprazole is not re	commended with HIV	/ protease inhibitors f	or which absorption is	s dependent on acidic
	intragastric pH such a	is atazanavir, nelfinav	ir; due to significant re	eduction in their bloav	allability.	
	It has been shown that	at co-administration of	atazanavir 300 mg/ri	tonavir 100 mg with o	meprazole (40 mg on	ce dally) or atazanavir
	atazanavir. The absor	azole (60 mg single (nH dependent	inteers resulted in a s		In the bloavallibility of
	Coumarin anticoacula	ants (phenprocourson	or warfarin)			
	Although no interact	tion during concomit	ant administration of	f phenprocourron or	r warfarin has heen	observed in clinical
	pharmacokinetic stud	ies, a few isolated or	ases of changes in In	ternational Normalise	ed Ratio (INR) and pr	othrombin have been
	reported during conco	mitant treatment in th	e post-marketing peri	od. Increases in INR	and prothrombin time	may lead to abnormal
	bleeding, and even	death. Therefore. in	patients treated with	h coumarin anticoad	ulants (e.g. phenpro	coumon or warfarin).
	monitoring of increase	e prothrombin time / II	NR is recommended a	after initiation, termina	tion or during irregula	r use of pantoprazole.
	Methotrexate			,	<u> </u>	1 1 1 1 1 1
	Case reports, publish	ed population pharma	cokinetic studies, and	d retrospective analys	ses suggest that conc	omitant administration
	of PPIs and methotrex	kate (primarily at high	dose; see methotrexa	te prescribing information	ation) may elevate and	l prolong serum levels
	of methotrexate and/c	or its metabolite hydro	xymethotrexate. Howe	ever, no formal drug ir	nteraction studies of m	nethotrexate with PPIs
	have been conducted	(see Warnings and F	recautions).			
	Other interactions stu	dies				
	Pantoprazole is metal	bolized in the liver via	the cytochrome P450	U enzyme system. Th	e main metabolic pati	nway is demethylation
	by CYP2C19 and othe	er metabolic pathway	s include oxidation by	UYP3A4.	using the server	mo ovotom or
	An interaction of pant		ings of compounds w	mich are metabolized	using the same enzy	me system cannot be
	nathways like carbon	o cimically significant	nteractions were obse	ne phenytoin and an	oral contracentivo cor	netabolized with these
	and ethinvi pestradio	iazepine, uiazepaili, (nioenolamide, miedipi	no, phonytoin anu an	oral contraceptive COI	itaning evolutiongestien
	Results from a range	of interaction studies	demonstrate that na	intoprazole does not	effect the metabolism	of active substances
	metabolised by CYP1	A2 (such as caffeine	theophylline), CYP20	C9 (such as piroxican	n, diclofenac, naproxe	en), CYP2D6 (such as
	metoprolol), CYP2E1	(such as ethanol), or	does not interfere wit	h p-glycoprotein relate	ed absorption of digox	kin.
	There were also no in	teractions with conco	mitantly administered	antacids.	,	
	Interaction studies h	ave also been perfo	ormed by concomitar	ntly administering pa	ntoprazole with the	respective antibiotics
	(clarithromycin, metro	nidazole, amoxicillin)	No clinically relevant	interactions were fou	ind.	
	<u>Clopidogrel</u>					
	Concomitant administ	tration of pantoprazol	e and clopidogrel in h	ealthy subjects had r	no clinically important	effect on exposure to
	ine active metabolite	u ciopiaogrei or clopia	uogrei-induced platele	a innibition. No dose a	aujustment of clopidog	rel is necessary when
	Druge that Inhibit or li	approved dose of par	itoprazole.			
	Inhibitors of CVP2C1	9 such as fluvovamin	e would likely increase	se the systemic expos	sure of pantoprazole	Inducers of CVP2C19
	may decrease the sys	stemic exposure to pa	ntoprazole.			
	4.6 Pregnancy and l	actation				
	Pregnancy					
	The limited data from	the use of pantopra	cole in preanant wom	en does not indicate	foetal /neonatal toxici	ty. Studies in animals
	have shown reproduc	tive toxicity. The poter	itial risk for humans is	unknown. Pantobex s	should not be used dur	ing pregnancy. unless
	clearly necessary.	.,				5, 5
	Lactation					
	Animal studies have	shown excretion of pa	antoprazole in breast	milk. Excretion into h	uman milk has been i	reported. Therefore, a
	decision on whether	to continue/discontinu	ue breast-feeding or	to continue/discontinu	ue therapy with Panto	bex should be made
	taking into account the	e benefit of breastfee	ding to the child, and	the benefit of Pantobe	ex therapy to women.	
	4.7 Effects on the ab	pility to drive and us	e of machines			
	Pantoprazole is not e	xpected to adversely	attect the ability to dr	ive or use machines.	Adverse drug reactio	ns, such as dizziness
	and visual disturbance	es may occur (see se	ction 4.8). If affected,	patients should not di	rive or operate machir	nes.
	4.8 Undesirable offe	cte				
	Approximately 5 % of	natients can be even	cted to experience ad	verse drug reactions	(ADBs)	
	The most commonly r	reported ADRs are dia	urrhoea and headache	both occurring in an	proximately 1% of pa	tients.
	Table 1 lists adverse	drug reactions report	ed with pantoprazole	in clinical studies and	post- marketing exp	erience. The following
	convention is used for	or the classification o	f the frequency of an	adverse drug reacti	on (ADR) and is bas	ed on the Council for
	International Organiza	ations of Medical Scie	nces (CIOMS) guideli	ines:	. ,	
	Very common (≥1/10); common (≥1/100 t	o <1/10); uncommon	n (≥1/1,000 to <1/100); rare (≥1/10,000 to	<1/1,000); very rare
	(<1/10,000); not know	n (cannot be estimate	ed from the available	data).		
	For all adverse reaction	ons reported from pos	t-marketing experience	e, it is not possible to	apply any Adverse Re	eaction frequency and
	therefore they are me	ntioned with a "not kn	own" frequency.	d in order of dearers	na poriouonoco	
	within each frequency	y grouping, adverse ri	eactions are presente	u in order of decreasi	ng senousness.	
	Table 1. Adverse read	tions with pantoprazo	le in clinical trials and	post-marketing expe	rience	
	Frequency	Common	Uncommon	Rare	Verv Rare	Not known
	Organ class	(≥1/100 to <1/10)	(≥1/1,000 to	(≥1/10,000 to	(<1/10,000)	
		,	<1/100)	<1/1,000)		
	Blood and			Agranulocytosis	Leukopenia;	

a: Pancvtopenia

Frequency Organ class	Common (≥1/100 to <1/10)	Uncommon (≥1/1,000 to <1/100)	Rare (≥1/10,000 to <1/1 000)	Very Rare (<1/10,000)	Not known
Immune system disorders			Hypersensitivity (including anaphylactic reactions including anaphylactic shock)	Leukopenia; Thrombocytopeni a; Pancytopenia	Systemic lupus erythematosus
Metabolism and nutrition disorders			Hyperlipidaemia; Weight changes		Hyponatraemia; Hypomagnesaen ia; Hypocalcemia*; Hypokalemia*
Psychiatric disorders		Sleep disorders	Depression	Disorientation	Hallucination; Confusion
Nervous system		Headache,	Taste		
disorders		Dizziness	Disorders		
Eye disorders			Disturbances in vision/ blurred vision		
Gastrointestinal disorders		Diarrhoea Nausea; vomiting; Abdominal distension and bloating; Constipation; Dry mouth; abdominal pain and discomfort			
Hepatobiliary disorders		Liver enzymes increased	Bilirubin increased		Hepatocellular injury, jaundice, hepatocellular failure
Skin and subcutaneous tissue disorders		Rash/ Exanthema; eruption; Pruritus	Urticaria Angioedema;		Stevens-Johnsor syndrome; Lyell syndrome; Erythema multiforme; Photosensitivy, Cutaneous lupus erythematosus; DRESS^
Musculoskeletal, connective tissue disorders			Arthralgia; Myalgia		Fracture of wrist, hip and spine
Renal and urinary disorders					Tubulointerstitial nephritis (TIN) (with possible progression to renal failure)
Reproductive system and breast disorders			Gynecomastia		
General disorders and administration site conditions		Asthenia; Fatigue and Malaise	Body temperature increased; Peripheral oedema		

* Hypocalcemia and/or hypokalemia may be related to the occurrence of hypomagnesemia (see Special Warnings and Special Precautions for use, 4.4)

^DRESS: Drug Rash with Eosinophilia and Systemic Symptoms

4.9 Overdosage

Systemic exposure with doses up to 240 mg administered intravenously over 2 minutes was well tolerated. As pantoprazole is extensively protein bound, it is not readily dialyzable.

In the case of overdose with clinical signs of intoxication, the usual rules of intoxication therapy apply.

4.10 Drug abuse and dependence

Drug dependence has not been observed

5. Pharmacological Properties

Pantobex: contains Pantoprazole, a proton pump inhibitors which inhibits the gastric H⁺K⁺ ATPase which is responsible for acid secretion in the parietal cells of the stomach. Pantoprazole is white to off-white powder with a molecular weight of 432.4. Pantoprazole is freely soluble in water, very slightly soluble in phosphate buffer at pH 7.4, and practically insoluble in n-hexane. Pantoprazole is a racemic mixture with a melting point of 138° C.

The chemical name for Pantoprazole is sodium-5-(difluoromethoxy)-2-[[(3,4-dimethoxy-2- pyridinyl)methyl] sulfinyl]-1 H-benzimidazole sesquihydrate and is represented by the following chemical structure :



Empirical chemical structure: C₁₆H₁₄F₂N₃NaO₄Sx1.5H₂O

5.1 Pharmacodynamic properties Pharmacotherapeutic / indication group / action mechanism

Selective proton pump inhibitor, substituted benzimidazole

(1) Mechanism of Action

Pantoprazole is a substituted benzimidazole which inhibits the secretion of hydrochloric acid in the stomach by specific action the proton pumps of the parietal cells. Pantoprazole is converted to its active form in the acidic canaliculi in the parietal cells which inhibits the H*. K* - ATPase enzyme, i.e. the final stage in the production of hydrochloric acid in the stomach.

The inhibition is dose-dependent and affects both basal and stimulated acid secretion. In most patients, freedom from sympti is achieved in 2 weeks. As with other proton pump inhibitors and H2 receptor inhibitors, treatment with pantoprazole cause reduced acidity in the stomach and thereby an increase in gastrin in proportion to the reduction in acidity. The increase in gas is reversible.

Since Pantoprazole binds to the enzyme distal to the cell receptor level, the substance can affect hydrochloric acid secret independently of stimulation by other substances (acetylcholine, histamine, gastrin). The effect is the same whether the ac substance is

given orally or intravenously. The fasting gastrin values increase under pantoprazole. On short-term use, in most cases they do exceed the normal upper limit. During long-term treatment, gastrin levels double in most cases. An excessive increase, howe occurs only in isolated cases. As a result, a mild to moderate increase in the number of specific endocrine (ECL) cells in stomach is observed in a minority of cases during long-term treatment (simple to adenomatoid hyperplasia). However, accord to the studies conducted so far, the formation of carcinoid precursors (atypical hyperplasia) or gastric carcinoids as were foun animal experiments have not been observed in humans.

An influence of a long term treatment with Pantoprazole exceeding one year cannot be completely ruled out on endor parameters of the thyroid and liver enzymes according to results in animal studies.

5.2 Pharmacokinetic properties

(1) Absorption

After ingestion, Pantoprazole is rapidly absorbed into the bloodstream. On average the maximum serum concentrations (Cma: 1 to 1.5 µg/mL (pantoprazole 20 mg tablet) or 2 to 3 µg/mL (pantoprazole 40 mg tablet) are achieved at about 2 to 2.5 hours a administration. After single and repeated administration of Pantoprazole, the pharmacokinetic characteristics of Pantoprazole very similar.

Both oral and I.V. administration of Pantoprazole in the dose range of 10 mg to 80 mg result in linear serum pharmacokinetics. absolute bioavailability from the tablet was found to be about 77%.

Concomitant intake of food had no relevant influence either on the AUC or on the Cmax and, thus, bioavailability. Only variability of the lao-time will be increased by concomitant food intake.

(2) Distribution

Pantoprazole's serum protein binding is about 98%, and in keeping with this, Pantoprazole has a low volume of distribution (a 0.15 l/kg) and limited tissue distribution.

(3) Metabolism

Pantoprazole is rapidly eliminated from the circulation and extensively metabolized in the liver. Metabolism occurs via oxidatio the CYP enzyme system, predominantly by CYP2C19 and CYP3A4 (Phase I metabolism, which is saturable). Pantopra undergoes further biotransformation by conjugation with sulphate, which involves the cytoplasmic enzyme sulphotransfer (phase II metabolism, which is not saturable), and which presents the main metabolism of Pantoprazole.

(4) Excretion and elimination

About 80% of the metabolites of pantoprazole are eliminated via the renal route, the rest via the feces. None of the metabolites considered as biologically active. The main metabolite in both the serum and urine is desmethylpantoprazole, which is conjug with sulphate.

T1/2 of the main metabolite is about 1.5 hour (which is not much longer than that of pantoprazole, 1 hour).

(5) Special populations

Impaired renal function

In patients with impaired renal function (including dialysis), pantoprazole showed no prolonged elimination half-life and accumulation when compared with healthy subjects. No dose reduction is requested when pantoprazole is administered to patie with restricted kidney function (incl. dialysis patients). As with healthy subjects, pantoprazole's half-life is short. Only very s amounts of Pantoprazole can be dialyzed. Although the main metabolite has a moderately delayed half-life (2-3h), excretion is rapid and thus accumulation does not occur.

Impaired hepatic function

Although for patients with liver cirrhosis (classes A and B according to Child) the half-life values increased to between 3 and (pantoprazole 20 mg tablet) and the AUC values increased by a factor of 3 - 5 (pantoprazole 20 mg tablet), the maximum set concentration only increased slightly by a factor of 1.3 (after oral administration) compared with healthy subjects.

A slight increase in AUC and Cmax in elderly volunteers compared with younger counterparts is also not clinically relevant. Age, Gender, Race

As with other clinically used PPIs, a small percentage of the population (about 3% Caucasians, 20% Asians) shows slo elimination of pantoprazole (T1/2 being up to 10 hours as compared with 1hour).

Such persons are known as poor metabolizers as a result of a deficiency of the CYP2C19 enzyme. In these individuals metabolism of Pantoprazole is probably mainly catalyzed by CYP3A4. After a single-dose administration of 40 mg Pantopraz the mean area under the plasma concentration- time curve was approximately 6 times higher in poor metabolizers than in subj having a functional CYP2C19 enzyme (extensive metabolizers). Mean peak plasma concentrations were increased by al 60%. These findings have no implications for the posology of Pantoprazole.

Compared with younger subjects, slight increases in AUC and Cmax were noted after single and repeated oral administration Pantoprazole to healthy elderly subjects (age >65 years). However, no dose adjustment is generally necessary in elderly patients

(6) Drug Interactions

Pantoprazole is metabolized in the liver via the CYP enzyme system. An interaction of pantoprazole with other drugs compounds, which are metabolized using the same enzyme system, cannot be ruled out.

Nevertheless, in specific tests pantoprazole did not affect the clearance of several compounds metabolized by CYP enzyr Vice-versa, all drugs that were tested regarding their potential influence on the pharmacokinetics of pantoprazole had no rele effect.

No detectable interactions between Pantoprazole and any other commonly prescribed co- medication tested so far were foun Metabolism of Pantoprazole occurs via oxidation by the CYP enzyme system, predominantly by CYP2C19 and CYP2 Interaction studies with drugs also metabolized by these pathways, like carbamazepine, diazepam, glibenclamide, nifedig phenytoin, and an oral contraceptive containing levonorgestrel and ethinyl estradiol did not reveal clinically significant interactio. Results from a range of interaction studies demonstrate that pantoprazole does not affect the metabolism of active substar metabolized by CYP1A2 (such as caffeine, theophylline), CYP2C9 (such as piroxicam, diclofenac, naproxen), CYP2D6 (such metoprolol), or CYP2E1 (such as ethanol), or does not interfere with pglycoprotein related absorption of digoxin. There were interactions with concomitantly administered antacids. Interaction studies have also been performed administering pantopra concomitantly with the respective antibiotics (clarithromycin, metronidazole, amoxicillin). No clinically relevant interactions vertice.

(See Interaction with other medicinal products and other forms of interaction).

	5.3. Preclinical safety data	
	Carringgenesis Mutagenesis Impairment of Fertility	
	Cardinogenesis, waagenesis, impairment of returny	
	Non-clinical data reveal no special hazard to humans based on conventional studies of safety pharmacology, repeated dose	
on on	toxicity and genotoxicity.	
here	In the two-year carcinogenicity studies in rats, neuroendocrine neoplasms were found. In addition, squamous cell papillomas were	
	found in the forestomach of rats. The mechanism leading to the formation of gastric carcinoids by substituted benzimidazoles has	
toms	been carefully investigated and it can be concluded that it is a secondary reaction to the massively elevated serum castrin levels	
	occurring in the rat during chronic high dose treatment. In the two-year rodent studies, an increased number of liver tumors was	
otrin	abanya in the ratio and in simple more interpreted as being due to particular, an included in the ratio as the interpreted as being due to particular the stability and use in the liver	
ISUIII	observed in rats and in remark mice and was interpreted as being due to partoprazole's high metabolic rate in the liver.	
	Animai Toxicology and/or Pharmacology	
etion	A slight increase of neoplastic changes of the thyroid was observed in the group of rats receiving the highest dose (200 mg/kg).	
ctive	The occurrence of these neoplasms is associated with the Pantoprazole- induced changes in the breakdown of thyroxine in the rat	
	liver. As the therapeutic dose in man is low, no side effects to the thyroid glands are expected.	
o not	In animal reproduction studies, signs of fetotoxicity were observed at doses above 3 mg/kg	
ovor	Investigations revealed no evidence of imprint protocological attractions offente. Crossing of the placents was investigated in the rat	
	investigations revealed no evidence of imparted remainly on teradigence energies. Or ossing of the placenta was investigated in the rate	
i the	and was found to increase with advanced gestation. As a result, the concentration of partoprazole in the letus is increased shortly	
rding	before birth.	
nd in		
	6. Pharmaceutical Particulars	
crine	6.1. List of excipients	
	Pantobex 20 mg and Pantobex 40 mg	
	Mannital Sadium Carbanata Anburraus, Craspovidana, Bavidana K.30, Calaium Staarata, Burifiad watar	
	Marinio, Soulin Carbonale Annyorous, Crospovidone, Fovidone K-so, Calcium Stearate, Funned Water	
\ -		
ax) of	Hydroxypropylmetnyl Gellulose, Polyethylene Glycol-6000, Purified Talc, Purified Water	
after		
e are	Enteric Coating	
	Methacrylic Acid- ethyl acrylate Copolymer Dispersion, Triethyl Citrate, Purified Talc, Purified Water	
The		
	Film Coating	
(the	. m. course	
y ule	Opacity Anno 601052172, Fullineu watel, Fullineu watel q.S. Solvent Ph. Eur Carnauda Wax (POWder)	
	6.2 Storage conditions and shelf life	
about	Storage Conditions:	
	Store below 30° C. Protect from light.	
	Shelf life:	
	2 years	
n hv		
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	Manufactured by	
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