

QUALITATIVE AND QUANTITATIVE COMPOSITION

ent to respectively 4, 8 and 12 mg galanta

base.

REMINYL oral solution contains galantamine hydrobromide, equivalent to 4 mg/ml galantamine base For excipients, see List of Excipients.

- PHARMACEUTICAL FORM
 Film-Coated Tablets for Oral Use
 4 mg galantamine as off-white, circular, biconvex tablets with the inscription "JANSSEN" on one side and
 "G4" on the other side;
 8 mg galantamine as pink, circular, biconvex tablets with the inscription "JANSSEN" on one side and "G8" on the other side;
- 12 mg galantamine as orange-brown, circular, biconvex tablets with the inscription "JANSSEN" on one side and "G12" on the other side.

Oral Solution 4 mg/ml

CLINICAL PARTICULARS

Therapeutic Indications
REMINYL is indicated for the treatment of mild to moderately severe dementia of the Alzheimer type
Posology and Method of Administration

Adults
REMINIVL should be administered twice a day, preferably with morning and evening meals. Ensure adequate fluid intake during treatment.
Starting Dose
The recommended starting dose is 8 mg/day (4 mg twice a day) for 4 weeks.
Maintenance Dose

- g Dose

 commended starting dose is 8 mg/day (4 mg twice a day) for 4 weeks.

 rearize Dose
 initial maintenance dose is 16 mg/day (8 mg twice a day) and patients should be maintained on
- The missi manifestance cose is to inguisty to mig wave a cary and patients should be manifested of fill migdaly for all sasts 4 weeks. An increase to the maximum recommended maintenance dose of 24 mg/day (12 mg twice a day) should be considered after appropriate assessment including evaluation of clinical benefit and tolerability. There is no rebound effect after about discontinuation of treatment (e.g., in preparation for surgery).
- Children
 Use of REMINYL in children is not recommended. No data on the use of REMINYL in pediatric patients are

available. Hepatic and Renal Impairment

respace and Henal Impairment
Galantamine plasma levels may be increased in patients with moderate to severe hepatic or renal impairment.
In patients with moderately impaired hepatic function, based on pharmacokinetic modeling, dosing could begin with 4 mg once daily, preferably taken in the morning for at least one week. Thereafter, patients should proceed with 4 mg b.i.d. for at least four weeks. In these patients, daily doses should not exceed 8 mg b.i.d. in patients with severe hepatic impairment, the use of REMINYL is not recommended.

For patients with a creatinine clearance greater than 9 ml/min, no dosage adjustment is required. In patients with severe renal impairment (creatinine clearance less than 9 ml/min), the use of REMINYL is not recommended since no data are available.

Concomitant Treatment

In patients treated with potent CYP2D6 or CYP3A4 inhibitors, dose reductions can be considered (see: Inter actions with Other Medicinal Products and Other Forms of Interaction).

Contraindications
REMINYL should not be administered to patients with a known hypersensitivity to galantamine hydrobromide

or to any excipients used in the formulations.

Special Warnings and Special Precautions for Use

REMINUT is includated for patients with mild to moderately severe dementia of the Alzheimer's type. The benefit of ERMINUT in patients with other types of dementia or other types of memory impairment has not been demonstrated.

It of REMINYL in patients with other types of dementa or other types of memory impatitivents has not occur
demonstrated. Patients with Alzheimer's disease lose weight. Treatment with cholinesterase inhibitors, including galantamine,
has been associated with weight loss in these patients. During therapy, patient's weight should be monitored.
As with other cholinomimetics, ReMINYL should be given with caution in the following conditions:
Cardiovascular Conditions: because of their pharmacological action, cholinomimeties may have vagotoric
effects on heart rate (e.g. bradycardia). The potential for this action may be particularly important to patients
with sick sinus syndrome' or other supraventricular cardiac conduction disturbances or who use drugs that
significantly reduce heart rate concomitantly, such as digoxin and beta-blockers, in clinical trials, use of
REMINYL has been associated with syncope and rarely with severe bradycardia.
Gastrointestinal Conditions; patients at increased risk of developing peptic ulcers, e.g., those with a history of
ulcer disease or those predisposed to these conditions, including those receiving concurrent nonsteroidal
anti-inflammatory drugs (NSAIDS), should be monitored for symptoms. However, clinical studies with
REMINYL showed no increase, relative to placebo, in the incidence of either peptic ulcers disease or gastrointestinal beleding. The use of REMINYL is not recommended in patients with gastrointestinal bodies under
recovering from gastrointestinal surgery.

intestinal bleeding. The use of REMINYL is not recommended in patients with gastrointestinal obstruction or recovering from gastrointestinal surgery. Neurological Conditions: Although cholinomimetics are believed to have some potential to cause seizures, seizure activity may also be a manifestation of Alzheimer's disease. Pulmonary Conditions: because of their cholinomimetic actions, cholinomimetics should be prescribed with care for patients with a history of severe asthma or obstructive pulmonary disease. Genitiourinary: the use of PEMINIYL is not recommended in patients with urinary outflow obstruction or recovering from bladder surgery. Satety in Subjects With Mild Cognitive Impairment (MCI) REMINIYL is not indicated for individuals with mild cognitive impairment (MCI), i.e., those who demonstrate isolated memory impairment greater than expected for their age and education, but do not meet criteria for Alzheimer's disease.

Alzheimer's dis

Alzheimer's disease.

Two, 2-year controlled trials in subjects with MCI did not meet dual primary efficacy outcomes. Although mortality in both realment arms was low, more deaths were initially recorded in subjects randomized to galantamine than to placebo, but the incidence of serious adverse events was identical between freatment groups. The deaths were due to various causes that are not unexpected in an elderly population. When data retrieved from the large proportion of patients who discontinued prior to completion of the double-blind period was included, there was no evidence of an increasing insk of death in FEMINVI-treated subjects over time. More subjects from the placebo than the galantamine group discontinued prior to death, which may account for the difference in mortality initially recorded.

The MCI study results are discrepant from those observed in studies of Alzheimer's disease. In pooled studies in Alzheimer's disease (n=4614), the mortality rate was numerically higher in the placebo than the REMINVI group.

REMINYL group.
Interactions With Other Medicinal Products and Other Forms of Interaction

Pharmacodynamic Interactions Because of its mechanic

Pharmacodynamic Interactions
Because of its mechanism of action, galantamine should not be given concomitantly with other cholinomimetics. Galantamine antagonises the effect of anticholinergic medication. As expected with cholinomimetics, a pharmacodynamic interaction is possible with drugs that significantly reduce the heart rate (e.g. digoxin and beta blockers).

Galantamine, as a cholinomimetic, is likely to exaggerate succinylcholine-type muscle relaxation during

Galantamine, as a cholinomimetic, is likely to exaggerate succinylcholine-type muscle relaxation during anaesthesia.

Multiple metabolic pathways and renal excretion are involved in the elimination of galantamine. Based on in virto studies, CYP2D6 and CYP3A4 were the major enzymes involved in the metabolism of galantamine. Inhibition of gastric acid secretion will not impair the absorption of galantamine. Inhibition of gastric acid secretion will not impair the absorption of galantamine. Other Drugs Affecting the Metabolism of Galantamine Drugs that are potent inhibitors for CYP2D6 or CYP3A4 may increase the AUC of galantamine. Multiple dose pharmacokinetic studies demonstrated that the AUC of galantamine increased 30% and 40%, respectively, during oc-administration of ketocoracide and paroxetine. As co-administred with erythromycin, another CYP3A4 inhibitor, the galantamine AUC only increased approximately 10%. Population PX analysis for patients with Azhemier's Gasease showed that the clearance of galantamine was decreased about 25-36% by concurrent administration of amétripyline, fluoxetine, fluoxoxamine, paroxetine and quinidine, known inhibitors of CYP2D6. Therefore, during initiation of treatment with potent inhibitors of CYP2D6 or CYP3A4 patients may experience an increased incidence of cholinergic side effects, predominantly nausea and vomiting. Under these circumstances, based on tilevalation, an N-methyt-D-aspartate (NMDA) receptor antagonist, at a dose of 10 mg/daily for 2 days followed by 10 mg BID for 12 days had no effect on the pharmacokinetics of galantamine 16 mg/day at steady state.

state.

Effect of Galantamine on the Metabolism of Other Drugs

Therapeutic obses of galantamine (12 mg b.i.d.) had no effect on the kinetics of digoxin and warfarin. Galantamine did not affect the increased prothrombin time induced by warfarin.

In vitro studies indicated that the inhibition potential of galantamine with respect to the major forms of human in rivide outsets indicated that the influence potential or gradient with respect to the major forms or indirat-cytochrome P250 is very low.

Pregnancy and Lactation

Use During Pregnancy

Reproduction studies conducted in pregnant rats at doses up to 16 mg/kg (or about 25 times the human ther-

Reproduction studies conducted in pregnant rats at doses up to 16 mg/kg (or about 25 times the human therapeutic dose) and in pregnant rabbits up to 40 mg/kg (or about 63 times the human therapeutic dose) did not show any evidence of a teratogenic potential. A non-significant increase in the incidence of minor skeletal abnormalities was noted at a dose of 16 mg/kg in rats.

No studies are available on the use of FEMINY1 in pregnant women. REMINY1 should be used during pregnancy only if the potential benefit justifies the potential risk to the foetus.

Use During Lactation
It is not known whether REMINY1. Is excreted in human breast milk and there are no studies in lactating women. Therefore, women on REMINY1 should not breast-feed.

Effects on Ability to Drive and Use Machines
Alzheimer's disease may cause gradual impairment of driving performance or compromise the ability to use machinery. Furthermory, Furthermory, is other cholinomimetics, REMINY1 may cause disziness and somnolence, which could affect the step into of the remains and the step of the step o

AIZHEIMER'S unsease may cause gradual impairment of uniting personnels of uniting machinery. Furthermore, like other cholinominieties, REMINIVE, may cause dizziness and somnolence, could affect the ability to drive or use machines, especially during the first weeks after initiation of treatments.

System/Organ Class

Undesirable Effects Clinical Trial Data

Clinical Irial Data — Adverse Drug Reactions Reported at ≥1% Frequency Druble-Blind Data – Adverse Drug Reactions Reported at ≥1% Frequency The safety of REMINYL was evaluated in 4457 subjects with mild to moderately severe dementia of the Alzheimer's type who participated in 7 placebo-controlled, double-blind clinical trials. The information pre-sented in this section was derived from pooled data. Adverse Drug Reactions (ADRs) reported by ≥1% of REMINYL-treated subjects in these trials are shown in

rse Drug Reactions Reported by ≥1% of REMINYL-Treated Subjects in 7 Placebo-Controlled,

Adverse Reaction	%	%
Metabolism and Nutrition Disorders		
Decreased appetite	5.2	1.4
Anorexia	3.8	1.0
Psychiatric Disorders		
Depression	4.2	2.9
Nervous System Disorders		
Dizziness	8.9	4.6
Headache	7.6	5.4
Tremor	2.0	0.8
Syncope	1.8	0.7
Lethargy	1.7	0.7
Somnolence	1.7	0.8
Cardiac Disorders		
Bradycardia	1.2	0.3
Gastrointestinal Disorders		
Nausea	25.0	7.6
Vomiting	12.8	3.1
Diarrhea	9.0	6.3
Abdominal pain	2.4	0.9
Abdominal pain upper	2.0	1.4
Dyspepsia	1.8	1.3
Stomach discomfort	1.6	0.6
Abdominal discomfort	1.0	0.4
Skin and Subcutaneous Tissue Disorders		
Hyperhidrosis	1.2	0.7
Musculoskeletal and Connective Tissue Disorders		
Muscle spasms	1.5	0.8
General Disorders and Administration Site Conditions		
Fatigue	4.0	2.2
Asthenia	2.3	1.7
Malaise	1.4	0.7
Investigations		
Weight decreased	5.1	1.4

In a randomized, double-blind, placebo-controlled clinical trial, the safety profile of once-daily treatment with

In a randomized, double-blind, placebo-controlled clinical trial, the safety profile of once-daily treatment with REMINTV. prolonged release capsules was similar in frequency and nature to that seem with tablets. Nausea and vomiting, the most frequent adverse drug reactions, occurred mainly during tittation periods, lasted less than a week in most cases and the majority of patients had one episode. Prescription of anti-emetics and ensuring adequate fluid intake may be usually in these instances. Open-Label Data – Adverse Drug Reactions Reported at ≥1% Frequency. The safety of REMINTVL was evaluated in 1454 subjects with mild to moderately severe dementia of the Alzheimer's type who participated in 5 open-label clinical trials. The information presented in this section was derived from pooled data. Adverse Drug Reactions (ADRs) reported by ≥1% of REMINTVL-treated subjects in these trials and not listed in Table 1 included Fall, which occurred at a rate of 6.5% in open-label trials.

Double Blind and Open-Label Data – Adverse Drug Reactions Reported at <1% Frequency. Additional ADRs that occurred in <1% of REMINTVL-treated subjects in the double-blind and open-label clinical datasets are listed in Table 2.

Table 2. Adverse Drug Reactions Reported by <1% of REMINYL-Treated Subjects in Either Double-Blind or Open-Label Clinical Trials Metabolism and Nutrition Disorders

Dehydration Nervous System Disorders

Dysgeusia, Hypersomnia, Paresthesia Eye Disorders

Atrioventricular block first degree, Palpitations, Sinus bradycardia, Supraventricular extrasyst

Vascular Disorders

Flushing, Hypotension Gastrointestinal Disorders

Gastion Research Parking Retching Rusculoskeletal and Connective Tissue Disorders

In Table 3, ADRs are presented by frequency category based on spontaneous reporting rates. In Table 4, ADRs are presented by frequency category based on incidence in clinical trials, wh

Table 3. Adverse Drug Reactions Identified During Postmarketing Experience with REMINYL by Frequency Category Estimated from Spontaneous Reporting Rates Psychiatric Disorders Linearcae, Lethicipation Individual Hallucipation visual Hallucipation auditory.

/ery rare – Hallucination, I r and Labyrinth Disord /ery rare – Tinnitus scular Disorders

Very rate - Huminos
Vascular Disorders
Very rare - Hypertension
Hepatobiliary Disorders
Very rare - Hepatitis
Investigations

Very rare - Hepatic enzyme increased

Table 4. Adverse Drug Reactions Identified During Postmarketing Experience with REMINYL by Frequency Category Estimated from Clinical Trials

Psychiatric Disorders Common – Hallucination

Uncommon - Hallucination vis Ear and Labyrinth Disorders Uncommon - Tinnitus fascular disorders Common - Hypertension

Common - Hypertension

Hepatobiliary Disorders

Rare - Hepatitis

nvestigations

Uncommon - Hepatic enzyme increased

Overdose

Overdose
Symptoms of significant overdosing of galantamine are predicted to be similar to those of overdosing of other cholinomimetics. These effects generally involve the central nervous system, the parasympathetic nervous system, and the neuromuscular junction. In addition to muscle weakness or fasciculations, some of other signs of a cholinergic crisis may develop: severe nausea, vomiting, gastiro-intestinal cramping, salivation, lacirnation, urination, defectation, sweeting, bradycardia, hypotension, collapse and convulsions. Increasing muscle weakness together with tracheal hypersecretions and bronchospasm, may lead to vital air-way compromise.

There have been post-marketing reports of Torsade de Pointes, QT prolongation, bradycardia, ventricular tachycardia and brief loss of consciousness in association with inadvertent overdoses of galantamine. In one case

where the dose was known, eight 4 mg tablets (32 mg total) were ingested on a single day. Two additional cases of accidental ingestion of 32 mg (nausea, womling, and dry mouth; nausea, womling, and substemat chest pain) and one of 40 mg (womling) resulted in brief hospitalizations for observation with full recovery. One patient, who was prescribed 24 mg place and a history of hallucinations over the previous two years, mistakenily received 24 mg place daily for 34 days and developed hallucinations requiring hospitalization. Another patient, who was prescribed 16 mg (34 mg) or of as lobuluin, inadvertenity ingested 160 mg (34 mg). And experienced sweeting, vomining, bradycardia, and near-syncope one hour later, which necessitated hospital treatment. His symptoms resolved within 24 hours. prescribed 16 mg/day of oing, bradycardia, and ne resolved within 24 hours.

Treatment
As in any case of overdose, general supportive measures should be used. In severe cases, anticholinergics such as atropine can be used as a general antidote for cholinomimetics. An initial dose of 0.5 to 1.0 mg i.v. is recommended, with subsequent doses based on the clinical response.

Because strategies for the management of overdose are continually veolving, it is advisable to contact a poison control centre to determine the latest recommendations for the management of an overdose.

PharmacoLugical Properties

Pharmacodynamic Properties

Galantamine, a tertiary alkaloid is a selective, competitive and reversible inhibitor of acety/cholinesterase. In addition, galantamine enhances the intrinsic action of acety/choline on nicotinic receptors, probably through binding to an allosteric site of the receptor. As a consequence, an increased activity in the cholinergic system associated with improved cognitive function can be achieved in patients with dementia of the Alzheimer type. Clinical Studies

The dosages of REMINYL shown to be effective in controlled efficient and the studies of the studies o

Clinical Studies

The disages of REMINYL shown to be effective in controlled clinical trials in Alzheimer's disease were 16, 24 and 32 mg/day, Of these doses, 16 and 24 mg/day were determined to have the best benefithrisk relationship and are the recommended doses. Galantamine's efficacy has been studied using four specific outcom measures: the ADAS-cog (a performance based measure of cognition), the CIBIC-plus (a global assessmen by an independent physician based on a clinical interview with the patient and caregiver), several measure ments of the activities of daily living and the Neuropsychiatric Inventory (NPI, a scale that measures behavioured feithrogeness.

ments of the activities of daily living and the Neuropsychiatric Inventory (NPI, a scale that measures behavioural disturbances).
In clinical studies, performance of galantamine treated patients on the ADAS-Cog (see Figure) and CIBIC-plus was consistently statistically significantly better than that of patients who were on placebo. Patients who were treated for 6 months with galantamine had ADAS-cog scores that were significantly improved compared to their baseline scores. Compared to the untreated patients there was a substantial and sustained benefit in cognitive functioning. Galantamine treatment also significantly preserved the activities of daily living, such a dressing, hygiene, meal preparation. These were assessed using the Disabliny Assessment in Demental (the DAD) and the Alzheimer's Disease Cooperative Study (ADCS)-ADL-Inventory, caregiver-rated assessments. Galantamine doses of 16 and 24 mg daily maintained the NPI score throughout the observation period whereas the score of the placebo patients clearly deteriorated, as a result of the emergence of behavioural disturbances.

See figure 1 and 2.

Long-term treatment (combination of 6 months double-blind followed by 6 months open treatment) suggested

2.
ent (combination of 6 months double-blind followed by 6 months open treatment) sugg
nitive and functional performance was maintained for a full year.

Long-term treatment (combination of 6 months double-blind followed by 6 months open treatment) suggested that patients' cognitive and functional performance was maintained for a full year. Alzheimer's Disease With Cerebrovascular Disease (AD+CVD)

The efficacy and sately of galantamine in subjects with Alzheimer's disease and significant cerebrovascular diseases (AD+CVD) was investigated in a double-blind, placebro-controlled study. There were 282 subjects, 48% of the total study population (N=592), who met criteria for AD+CVD. Although the clinical trial was not

48% of the total study population (N=592), who met criteria for AD+CVD. Although the clinical trial was not powered for subgroup analyses, galantamine-treated subjects experienced a statistically significant improvement, compared to placeb-treated subjects to both primary outcomes [(cognition: ADAS-cogn't 1]-p-0.001]; global clinical assessment: CIBIC-plus [p-0.001] and on a measure of activities of daily living ((ADA) [p-0.001]). Overall, the safety and tolerability of galantamine in subjects with AD+CVD was similar to that seen in previous studies of galantamine and 11% of placebo subjects). Other events, occurring in >5% of AD+CVD subjects and reported adverse event in subjects was nausea (19% of galantamine and 11% of placebo subjects). Other events, occurring in >5% of AD+CVD subjects and reported more frequently in the galantamine than the placebo group, were duzieness, vomiting, abdominal pain, diarrhea, and fatigue. The incidence of 'cerebrovascular disorders' (e.g., stocke) was higher in the placebo group (placebo, 596 [5%] subjects; galantamine, 2/186 [1%] subjects with Alzheimer's disease.

Mild Cognitive Impairment (MCI)
Two, 2-year controlled trials in subjects with MCI did not meet dual primary efficacy outcomes. Although mortality

Overall, the safety profile in AD-CVD was consistent with that observed in studies of galantamine in subjects with Alzheimer's disease.

Midi Cagnitive Impairment (MCI)

Tivo, 2-year controlled trials in subjects with MCI did not meet dual primary efficacy outcomes. Although mortality was low (077%), more deaths were initially recorded in subjects randomized to galantamine (13/1026) than to placebo (1/1022), but the incidence of serious adverse events was identical (19%) between treatment groups. When data retrieved from the large proportion of patients in both treatment groups who docontinued prior to completion of the double-blind period (GAL-OCG-3002) were included, a total of 102 deaths were identified, 56 in the galantamine group and 46 in the placebo group (relative risk) (5%) Cl = 1.24 (0.241, 183); p = 0.274). The 24-month intent-to-treat analysis recorded 20 deaths among subjects randomised to placebo compared to 34 deaths recorded among subjects randomised for EREMINVI, (relative risk) (5%) Cl = 1.70; 11.00, 2.90); p = 0.051). Of subjects who died within the protocot-specified period of 30 days of discontinuing double-blind subdy medication, there were 14 in the galantamine group and 3 in the placebo group relative risk (5%) Cl = 4.08 (1.5.7, 10.375) p = 0.004). More placebo-treated than galantamine-treated subjects discontinued prior to death, which may account for the difference in mortality mitially recorded. Thirteen deaths in the placebo group and 20 deaths in the galantamine group were found to be directly related to adverse events that occurred while the subjects were exposed to double-blind study drug (relative risk) (5%) Cl = 1.54 (0.78, 3.04); p = 0.218.

The deaths were due to various causes that are not unexpected in an elderly population. About half of the deaths in both placebo and achieve retainment group serve found to be directly related to adverse events that occurred while the subjects were exposed to double-blind study drug (relative risk) (5%) Cl = 1.54 (0.78, 3.04); p = 0.218.

T

tration). The disposition of galantamine was studied in young subjects with varying degrees of renal function. Elimination of galantamine decreased with decreasing creatinine clearance. Plasma concentrations of galantamine increased in subjects with impaired renal function by 38% in moderate ($C|_{CB}$ =52-104 ml/min) or 67% in severe renal impairment ($C(C_{CB}$ =515 ml/min), compared to age and weight-matched healthy subjects ($C(C_{CB}$ =512 ml/min). A population or pharmacoknetic enalpsis and simulations indicate that no dose-adjustments are needed in Alzheimer patients with renal impairment provided that the $C|_{CB}$ is at least 9 ml/min (see: Posology and Method of Administration) as the galantamine clearance is lower in the Alzheimer population. Plasma arminin hindrina The aliasma crotein bindring of galantamine is (C_{CB}) and C_{CB} in the Alzheimer population.

Plasma protein binding: The plasma protein binding of galantamine is low: 17.7 ± 0.8%. In whole blood, galantamine is mainly distributed to blood cells (52.7%) and plasma water (39.0%), whereas the fraction of galantamine bound to plasma proteins in only 8.4%. The blood-to-plasma concentration ratio of galantamine is 1.17.

All other preclinical safety data relevant to the prescriber have been included in the appropriate sec

All other preclinical satery oata relevant to the prescriber handles made appropriate state of preclinical states of the preclinical states of preclinical states of preclinical states. Film-Coater Tablet Core:

The inactive ingredients are colloidal silicon dioxide, crospovidone, lactose monohydrate, magnesium stearate and microcrystalline cellulose.

Film-Coating:
The inactive ingredients are hypromeliose [hydroxypropyl methylcellulose], propylene glycol, talc and titanium dioxide. The 4 mg tablets also contain yellow ferric oxide. The 8 mg tablets contain red ferric oxide. The 12 mg tablets contain red ferric oxide and orange yellow S aluminium lake.

Oral Solution The inactive ingredients are methyl parahydroxybenzoate, propyl parahydroxybenzoate, sodium saccharin, sodium hydroxide and purified wate

Incompatibilities Not applicable. Shelf Life

Stell Line

Observe expiry date on the outer pack.

Special Precautions for Storage

REMINYL tablets store between 15° and 30°C.

REMINYL or all solution: store between 15° and 30°C, protect from freezing, use within 3 months of first open-

e and Contents of Container The tablets are packaged in a PVC-PE-PVDC/Alu blister that holds 14 tablets. Blisters are packed in a card-

Oral Solution

The oral solution is packaged in a 100 ml amber glass bottle with a LDPE insert, a PP/LDPE child resistant closure and a HDPE/LDPE/PS pipette, calibrated in millilitres. The pipette has a minimum volume of 0.5 ml

Istructions to soer efigure 3
open the bottle and use the pipette:
The bottle comes with a child-resistant cap, and should be opened as follows:
- Push the plastic screw cap down while turning it counter clockwise.
- Remove the unscrewed cap.
Insert the pipette into the bottle.
While holding the bottom ring, pull the top ring up to the mark corresponding to the number of millilitres

you need to give books in mile, you are up mile up to the man corresponding to the intended or mile you need to give Holding the bottom ring, remove the entire pipette from the bottle. Empty the pipette into any non-alcoholic drink by sliding the upper ring down and drink it immediately. Close the bottle. Rinse the pipette with some water.

DATE OF REVISION OF THE TEXT

JANSSEN-CILAG

Manufactured by: see outer pack for Janssen Pharmaceutica N.V., Turnhoutseweg 30, B-2340 Beerse, Belgium

