

COLIFOAM

10% hydrocortisone acetate

FIRST CLASS TREATMENT WHICH TRAVELS TO WORK

- ☛ Colifoam is highly effective for distal ulcerative colitis.⁽¹⁾
- ☛ The retrograde spread of Colifoam increases with the extent of disease.⁽²⁾
- ☛ Colifoam is easier to retain than liquid enemas and causes less interference with social, sexual and occupational activities.^(1,3)

PRESCRIBED WITH CONFIDENCE FOR OVER 20 YEARS.



PRESCRIBING INFORMATION: **Presentation:** White odourless aerosol containing hydrocortisone acetate PhEur 10% w/w. **Uses:** Ulcerative colitis, proctosigmoiditis and granular proctitis. **Dosage and administration:** One applicatorful inserted into the rectum once or twice daily for two or three weeks and every second day thereafter. Shake can vigorously before use (illustrated instructions are enclosed with pack). **Contra-indications, warnings etc:** Local contra-indications to the use of intrarectal steroids include obstruction, abscess, perforation, peritonitis, fresh intestinal anastomoses and extensive fistulae. General precautions common to all corticosteroid therapy should be observed during treatment with Colifoam. Treatment should be administered with caution in patients with severe ulcerative disease because of their predisposition to perforation of the bowel wall. Safety

during pregnancy has not been fully established. Although uncommon at this dosage, local irritation may occur. **Pharmaceutical precautions:** Pressurised container containing flammable propellant. Protect from sunlight and do not expose to temperatures above 50°C. Keep away from sources of ignition. Do not pierce or burn even after use. Do not refrigerate, store below 25°C. Keep out of reach of children. For external use only. **Legal category:** POM. **Package quantity & basic NHS cost:** 20.8g canister plus applicator, £7.07. Provides approximately 14 doses. **Product Licence no:** 0036/0021. **References:** 1. Somerville KW *et al.* BMJ 1985;291:866. 2. Farthing MJG *et al.* BMJ 1979;2:822-824. 3. Ruddell WSJ *et al.* Gut 1980;21:885-889. Further information is available on request from Stafford-Miller Ltd., Broadwater Road, Welwyn Garden City, Herts. AL7 3SP. **Code:** DO2665.

CONSISTENT RESULTS IN A WORLD
OF CONSTANT CHANGE.



Gold Medal, Atlanta Olympics 1996 (Coress Pair)


OVER 90% H. PYLORI ERADICATION¹
OVER 90% PATIENT COMPLIANCE¹

Rx KLARICID 500MG B.D.

OMEPRAZOLE 20MG O.D.

AMOXYCILLIN 1G B.D. *for 10 days*

NEW TRIPLE THERAPY
KLARICID[®]500
Clarithromycin

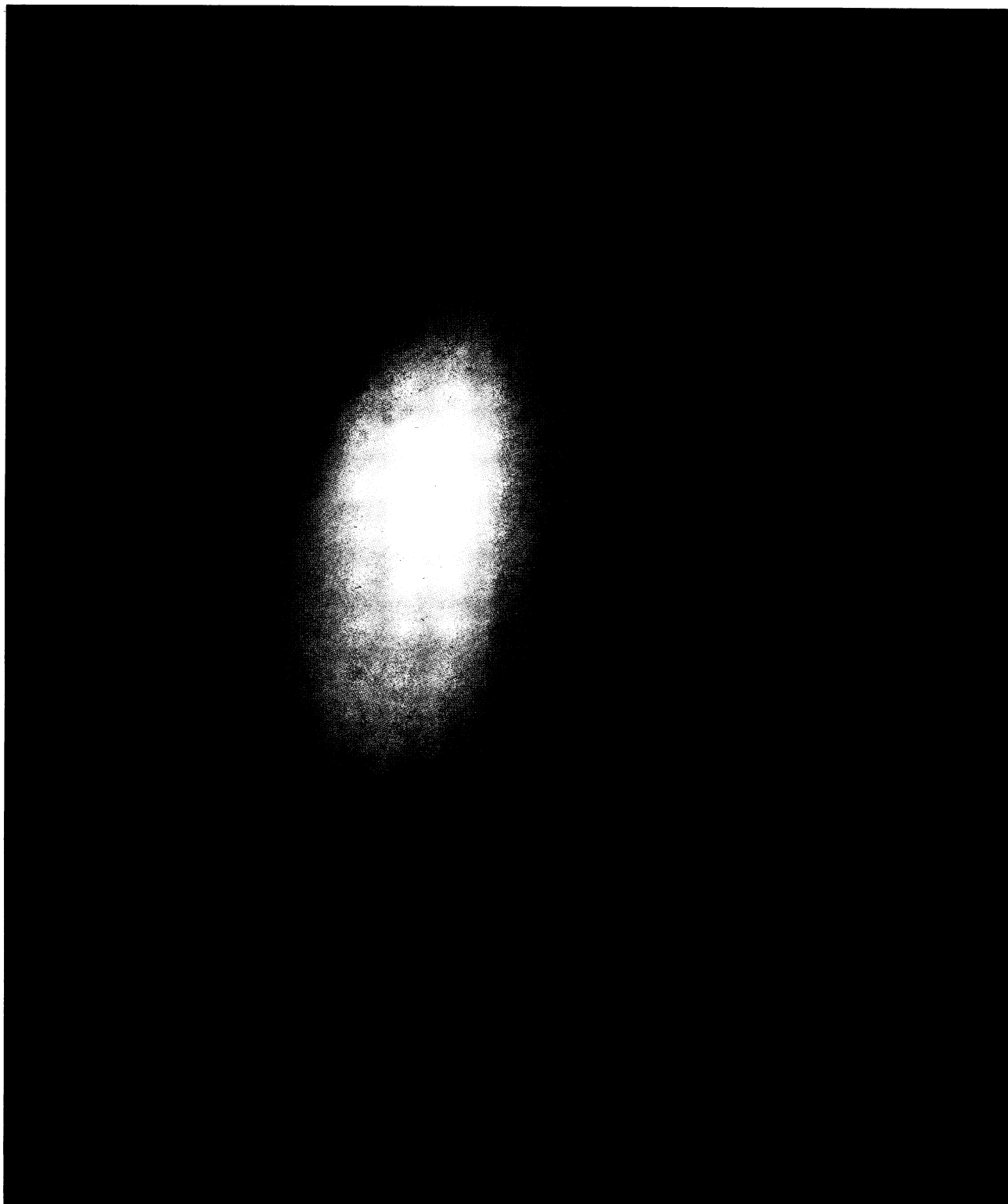
Prescribing information PI/1/4/002 **Klaricid 500**
Presentation: Yellow ovaloid film coated tablets containing 500mg of clarithromycin. Each tablet is engraved with  on one side. **Indications:** Klaricid in the presence of acid suppression effected by omeprazole is indicated for the eradication of H. pylori in patients with duodenal ulcers. **Dosage and Administration:** Adults: Dual therapy: clarithromycin 500mg t.d.s. for 14 days plus oral omeprazole 40mg o.d. The pivotal study was carried out with omeprazole 40mg o.d. for 28 days, whilst supportive studies were carried out with omeprazole 40mg o.d. for 14 days. Triple therapy: Klaricid (500mg) b.d. should be given with amoxicillin 1000mg twice daily and omeprazole 20mg daily for 10 days. See omeprazole and amoxicillin data sheet for further information on omeprazole dosing. **Contraindications, Warnings etc:** **Contraindications:** known hypersensitivity to macrolide drugs. Do not administer with any of the following: cisapride, pimozide, terfenadine, ergot derivatives. **Precautions:** Caution in adults with impaired hepatic

and renal function. Prolonged or repeated use of clarithromycin may result in an overgrowth of non-susceptible bacteria or fungi. If superinfection occurs, clarithromycin should be discontinued and appropriate therapy instituted. Caution in patients taking drugs metabolised by the cytochrome P450 system as there may be elevations in their serum levels. H. pylori organisms may develop resistance to clarithromycin in a small number of patients. **Interactions:** Potentiation of astemizole, theophylline, digoxin, warfarin and carbamazepine. Interaction of Klaricid tablets with simultaneously administered zidovudine in adults. No interaction with oral contraceptives. **Side-effects:** Klaricid is generally well tolerated. Side-effects include nausea, dyspepsia, vomiting, diarrhoea and rarely pseudomembranous colitis, abdominal pain, headache, taste perversion, reversible tongue discoloration, glossitis, stomatitis and oral monilia. Allergic reactions including anaphylaxis and Stevens-Johnson syndrome, and transient central nervous system side-effects have been reported. Hepatic dysfunction has

also been reported. There have been reports of hearing loss which is usually reversible on withdrawal of therapy. **Use in Pregnancy and Lactation:** The safety of Klaricid during pregnancy and breast feeding has not been established, and therefore if a patient becomes pregnant Klaricid should only be used if the benefits outweigh risks. Clarithromycin has been found in the milk of lactating animals and humans. **Overdose:** Should be treated with gastric lavage and supportive measures. **Legal Category:** POM. **Marketing Authorisation Number:** PL 0037/0254: 20 or 42 tablet calendar blister pack. **Basic NHS Price:** 500mg b.d. £3.21 per day; 500mg t.d.s. £4.82 per day.

Further information is available on request from Abbott Laboratories Ltd., Norden Road, Maidenhead, Berkshire SL6 4XE. Date of Preparation September 1996. Reference: 1. Data on file, Abbott Laboratories. PXXHP96227

 **ABBOTT**
ANTIBIOTICS



PRESCRIBING INFORMATION:

Indications Duodenal ulcer (including those associated with *H. pylori* infection), benign gastric ulcer, postoperative ulcer, oesophageal reflux disease, Zollinger Ellison Syndrome, prophylaxis of gastrointestinal haemorrhage from stress ulcer, recurrent haemorrhage from bleeding peptic ulcer, acid aspiration (Mendelson's Syndrome). Tablets, Syrup, Effervescent Tablets only, ulcers associated with non-steroidal anti-inflammatory drugs (NSAIDs), prevention of NSAID-associated duodenal ulcer, chronic episodic dyspepsia, severe oesophagitis, long-term management of healed oesophagitis. **Dosage Adults:** Duodenal ulceration and gastric ulceration: A single 300mg dose at bedtime or 150mg twice daily in the morning and evening for four weeks. In duodenal ulcer, 300mg twice daily produces higher healing rates at four weeks. Continued maintenance treatment of 150mg at bedtime is recommended for patients with a history of recurrent ulceration. Duodenal ulcers associated with *H. pylori*, 300mg at bedtime or 150mg twice daily with oral amoxicillin 750mg three times daily and metronidazole 500mg three times daily for 2 weeks. Zantac therapy then continued for a further 2 weeks. Ulcers following non-steroidal anti-inflammatory drugs: 150mg twice daily for up to eight weeks. Prevention of NSAID-associated duodenal ulcer: 150mg twice daily concomitantly with NSAID therapy. Chronic episodic

dyspepsia: 150mg twice daily for six weeks, investigate early relapsers and non-responders. Oesophageal reflux disease: 300mg at bedtime or 150mg twice daily for up to eight weeks. Moderate to severe oesophagitis: 150mg four times daily for up to twelve weeks. Long-term treatment of healed oesophagitis: 150mg twice daily. Obstetric patients at commencement of labour; oral dose of 150mg may be followed by 150mg at six-hourly intervals (see data sheet). Those at risk of acid aspiration syndrome; oral dose of 150mg two hours before induction of general anaesthesia with preferably 150mg the previous evening. Alternatively, Zantac Injection 50mg intramuscularly or by slow intravenous injection 45 to 60 minutes before general anaesthesia. Zantac Injection may be given every six to eight hours either as slow (over a period of at least two minutes) intravenous injection of 50mg, after dilution to a volume of 20ml per 50mg dose, or as intermittent intravenous infusion at a rate of 25mg per hour for two hours; alternatively, as intramuscular injection of 50mg (2ml) every six to eight hours. Prophylaxis of haemorrhage from stress ulceration or from bleeding peptic ulceration: parenteral administration may be continued until oral feeding commences. If still at risk, Zantac Tablets or Syrup 150mg may be given twice daily. Prophylaxis of haemorrhage from stress ulceration: priming dose of 50mg as a slow intravenous injection followed by continuous intravenous infusion of 0.125 to 0.250mg/kg/hr

cellent

Zantac

RANITIDINE HCl

*Zantac is now healing ulcers
in over 100 countries¹*

may be preferred. *Children:* Oral dose for peptic ulcer: 2mg/kg to 4mg/kg, twice daily to a maximum of 300mg per day. **Contra-indications** Patients with known hypersensitivity to ranitidine. **Precautions** Caution when using Effervescent Tablets in sodium-restricted patients. Exclude malignancy in gastric ulcer before instituting therapy, especially in middle-aged patients with new or recently changed dyspeptic symptoms. Regular supervision of patients taking NSAIDs concomitantly with Zantac recommended, especially if elderly. Protects against NSAID-associated ulceration in duodenum and not in stomach. Reduce dosage in the presence of severe renal failure (see data sheet). Avoid in patients with history of porphyria. Effervescent Tablets contain aspartame, use with caution in patients with phenylketonuria. Rapid administration of injection may rarely cause bradycardia; recommended rates of administration should not be exceeded. Like other drugs, use during pregnancy and lactation only if strictly necessary. **Side effects** Headache, dizziness, skin rash, occasional hepatitis, rarely arthralgia, myalgia, and with antibiotics, diarrhoea. Rarely, reversible mental confusion states, usually in very ill or elderly patients. Rare cases of leucopenia and thrombocytopenia, usually reversible, agranulocytosis and pancytopenia. Hypersensitivity reactions, anaphylactic shock, rare cases of breast symptoms in men. As with other H₂-receptor antagonists rare cases of bradycardia, A-V block

and asystole (see data sheet). **Presentations** Zantac 150 Tablets each containing 150mg ranitidine HCl, (Product licence number 10949/0042, 60 tablets £27.89); Zantac 300 Tablets each containing 300mg ranitidine HCl (Product licence number 10949/0043, 30 tablets £27.43); Zantac Effervescent Tablets each containing 150mg ranitidine HCl and 14.3mEq sodium, (Product licence number 10949/0137, 60 tablets £27.89); Zantac Effervescent Tablets each containing 300mg ranitidine HCl and 20.8mEq sodium (Product licence number 10949/0138, 30 tablets £27.43); Zantac Syrup each 10ml dose containing 150mg ranitidine HCl (Product licence number 10949/0108, 300ml bottle £22.32); Zantac Injection each 2ml dose containing 50mg ranitidine HCl (product licence number 10949/0109, 5 x 2ml £3.21). **Product licence holders** Glaxo Laboratories, Stockley Park West, Uxbridge, Middlesex UB11 1BT. **[POM]** Zantac is a trade mark of the Glaxo Wellcome Group of Companies. Further information is available on request from Glaxo Wellcome UK Limited, Stockley Park West, Uxbridge, Middlesex UB11 1BT. Telephone 0181-990 9444. Date of preparation: September 1995

Reference

1. Data on file. GlaxoWellcome UK Limited 1995.

GlaxoWellcome

STAY HEALED[†]



NOW APPROVED IN MAINTENANCE

15mg ZOTON[▼]

Lansoprazole

[†] Up to 87% remission rate in reflux oesophagitis after 1 year. (Range 69-87%)¹⁻⁵

ZOTON[▼] Abbreviated Prescribing Information

Presentation: Two tone lilac/purple capsules containing lansoprazole 30 mg. Opaque yellow capsules containing lansoprazole 15 mg. **Indications:** Healing and maintenance of gastro-oesophageal reflux disease (GORD) or duodenal ulcer. **Healing of benign gastric ulcer.** Effective for benign peptic lesions including reflux oesophagitis unresponsive to H₂ receptor antagonists. Eradication of *Helicobacter pylori* (*H. pylori*) in patients with duodenal ulcer or gastritis. **Dosage and Administration:** Duodenal ulcer: 30 mg for 4 weeks, then 15 mg for maintenance dose. GORD: 30 mg daily for 4-8 weeks, then 15 mg or 30 mg for maintenance dose. Benign gastric ulcer: 30 mg daily for 8 weeks. *H. pylori* eradication: 30 mg twice daily plus two of the following antibiotics for 7 days: clarithromycin 250 mg twice daily, amoxicillin 1 g twice daily or metronidazole 400 mg twice daily. Swallow capsules whole. No dosage adjustment is necessary in the elderly, or the renally or hepatically impaired. There is no experience with Zoton in children. **Contra-indications:** None known. **Precautions:** Exclude the possibility of malignancy when gastric ulcer is

suspected. When using in combination with antibiotics, refer to the prescribing information of the respective antibiotics. **Pregnancy and Lactation:** Avoid in pregnancy. Avoid during breast feeding unless essential. **Interactions:** Interactions with drugs metabolised by the liver are possible. Apply caution when used concomitantly with oral contraceptives, phenytoin, theophylline or warfarin. Antacids should not be taken within an hour of Zoton. **Side Effects:** Generally mild and transient, including gastro-intestinal disturbances, headache, dizziness, malaise, dry or sore mouth or throat, fatigue, rashes, urticaria, pruritis and alterations in liver function test values. A few cases of arthralgia, myalgia, peripheral oedema, depression, haematological changes, bruising, purpura, petechiae, jaundice, hepatitis, paraesthesia or blurred vision have been reported. **Legal Category:** POM **Package Quantities:** 30 mg capsules: Blister packs of 56, 28, 14 and 7 (hospital starter pack) capsules. 15 mg capsules: Blister packs of 56 and 28 capsules. **Product Licence Number:** 30 mg capsules: PL 0095/0264 15 mg capsules: PL 0095/0302 **Cost:** 30 mg capsules: 7 £9.09 (hospital starter pack) 14 £16.68 28 £33.36 56 £66.72 15 mg capsules: 28

£18.95 56 £37.90 Full prescribing information is available on request. **Name and Address of Licence Holder:** Cyanamid of Great Britain Ltd, Fareham Road, Gosport, Hampshire, PO13 0AS. **REFERENCES:** 1. Gough, A.L. et al, *Aliment Pharmacol Ther*, 1996, **10**, 529-539 2. Hatlebakk, J.G., and Berstad, A., *Gastroenterol*, 1995, **108** (4), A111 (102909) 3. Poynard, T. et al, *Gastroenterol*, 1995, **108** (4), A195 (102907) 4. Robinson, M., *Ann Intern Med*, 1996, **126**, 859-867 5. Baldi, F., *Gastroenterol*, 1996, **110** (4) Suppl A55 (107136), and Data on file, Lederle Laboratories (105806). * Trademark of Takeda Chemical Industries Ltd. Date of preparation: August 1996



Under Licence agreement with Takeda Chemical Industries Ltd, Japan.



Further information can be obtained from: Wyeth Laboratories, Huntercombe Lane South, Taplow, Maidenhead, Berks SL6 0PH



ZZ0T473/0896

EVERYDAY PEOPLE TAKE LOSEC.

Losec offers efficacy, flexibility, practicality and good tolerability. And with over 175 million prescriptions in 96 countries, it also inspires a high level of confidence. No wonder Losec is taking care of more people. Every day.

LOSEC® CAPSULES (omeprazole) PRESCRIBING INFORMATION

(refer to full data sheet before prescribing)

PRESENTATION: Losec Capsules containing 10mg, 20mg or 40mg omeprazole (O) as enteric coated granules with an aqueous based coating. **USES:** Oesophageal reflux disease. Duodenal and benign gastric ulcers (including NSAID-induced). *Helicobacter pylori* eradication: Relief of associated dyspeptic symptoms in combination treatment with antibiotics. Prophylaxis of acid aspiration. Zollinger-Ellison syndrome. **DOSAGE & ADMINISTRATION:** **Adults (including the elderly):** The usual dose in oesophageal reflux disease and peptic ulcer is 20mg once daily, increasing to 40mg once daily in severe or refractory cases, if required. **Oesophageal reflux disease:** Healing: 20mg daily for 4 weeks. Continue for further 4-8 weeks if required. **Maintenance in acid reflux disease:** Losec 10mg daily. Increase to 20mg daily if symptoms return. **Duodenal ulcer (DU):** Healing: 20mg daily for 4 weeks. **DU maintenance:** Losec 10mg daily increasing to 20mg daily if symptoms return. **Benign Gastric Ulcer:** 20mg daily for 8 weeks. **Helicobacter pylori eradication: DU disease: Triple therapies:** Losec 40mg daily with amoxicillin (A) 500mg and metronidazole (M) 400mg, both three times a day for 1 week. Or clarithromycin (C) 250mg and metronidazole 400mg (or tinidazole 500mg) both bd for 1 week. Or amoxicillin 1g and clarithromycin 500mg both bd for 1 week. **Dual therapies:** Losec 40mg daily with

amoxicillin 750mg to 1g bd or clarithromycin 500mg tid, both for 2 weeks. **GU disease:** Losec 40mg daily with amoxicillin 750mg to 1g bd for 2 weeks. **Prophylaxis of acid aspiration:** Losec 40mg on the evening before surgery followed by Losec 40mg on the morning of surgery. **Zollinger-Ellison Syndrome:** 60mg daily as long as clinically indicated. Individually adjust within range 20-120mg daily. If in excess of 80mg daily give in 2 equal divided doses. **Renal impairment:** No dose adjustment needed. **Hepatic impairment:** Adjust dose (maximum daily dose 20mg). **Children:** No experience of use. **CONTRA-INDICATIONS, WARNINGS, etc:** Known hypersensitivity to omeprazole. In gastric ulcer, exclude malignancy before starting therapy. Avoid in pregnancy unless no safer alternative. Discontinue breast feeding if Losec is considered essential. **Side effects:** Losec is well tolerated. Adverse reactions are generally mild and reversible (relationship to Losec not established in many cases). They include diarrhoea, headaches, skin disorders and in isolated cases, angioedema, musculoskeletal disorders, fatigue, insomnia, dizziness, blurred vision, dry mouth, vertigo, paraesthesia, anaphylaxis, liver enzyme and haematological changes. **Interactions:** The absorption of ketoconazole may be reduced. Losec can delay the elimination of diazepam, phenytoin and warfarin. Plasma concentrations of omeprazole and clarithromycin are increased when used concomitantly. Simultaneous treatment with omeprazole and digoxin may increase

the bioavailability of digoxin. **PHARMACEUTICAL PRECAUTIONS:** Store below 30°C. Bottles: Use within 3 months of opening. Replace cap firmly after use. Dispense in original container. **LEGAL CATEGORY:** POM. **FURTHER INFORMATION:** *Helicobacter pylori* (*Hp*) is associated with acid peptic and ulcer disease, contributing to gastritis and ulcer recurrence. Eradication of *Hp* with omeprazole and antibiotics gives rapid symptom relief, high healing rates and long-term remission of ulcer disease.

Quality of life. In recent clinical data, in patients with acute peptic ulcer disease, omeprazole *Hp* eradication therapy improved patients' quality of life. **PACKAGE QUANTITIES:** 10mg: bottles of 7* capsules £4.99, bottles of 28 capsules £19.95. 20mg: bottles of 7* capsules, £8.86, bottles of 28 capsules, £35.45; 40mg: blisters of 7 capsules £17.72. (*Hospital pack only). **MARKETING AUTHORIZATION NO:** PL 0017/0337 - Losec Capsules 10mg. PL 0017/0238 - Losec Capsules 20mg. PL 0017/0320 - Losec Capsules 40mg.

ASTRA
Astra Pharmaceuticals

For further information contact the **MARKETING AUTHORIZATION HOLDER:** Astra Pharmaceuticals Ltd, Home Park, Kings Langley, Herts WD4 8DH. Tel: (01923) 266191.

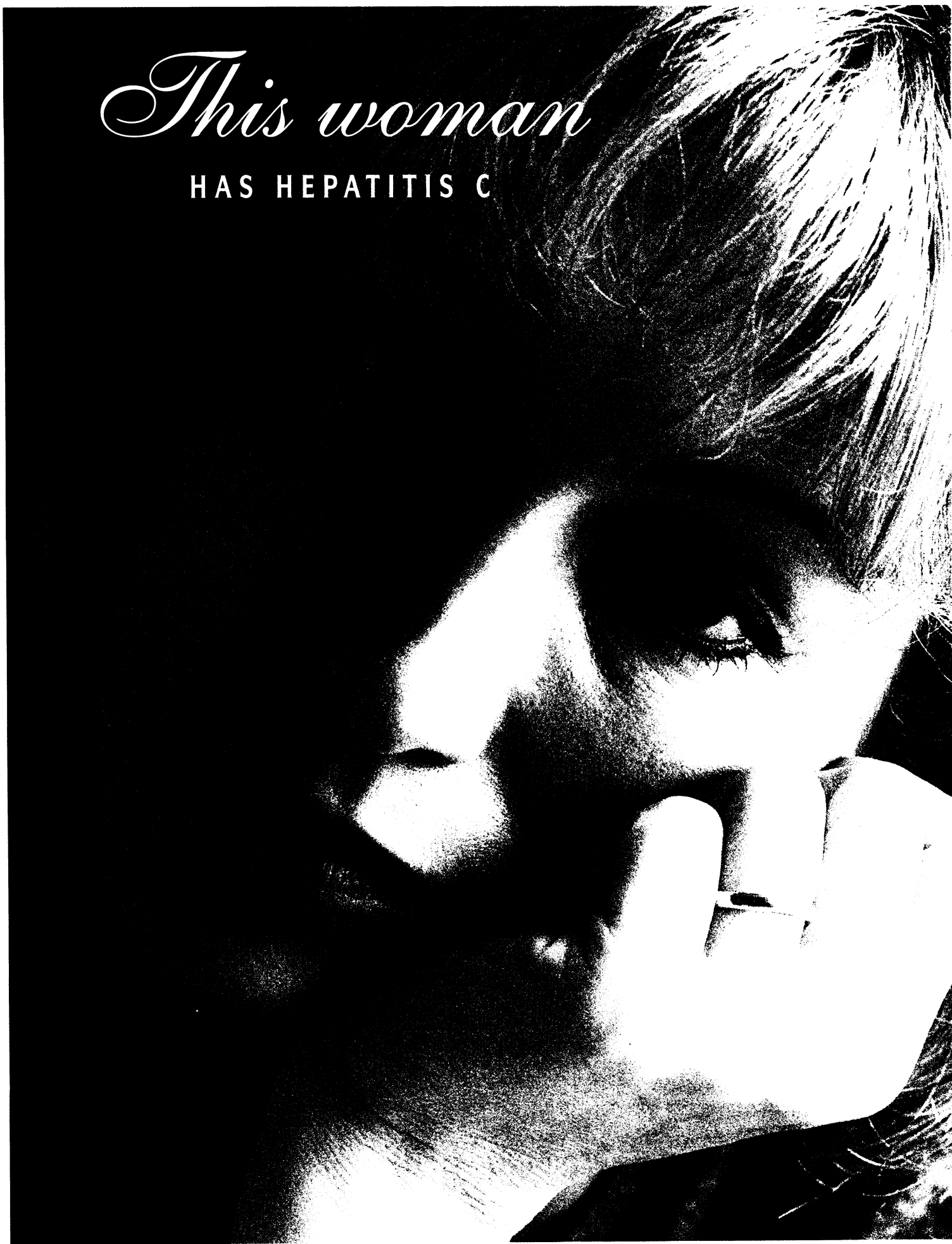
LOSEC is a registered trademark of Astra Pharmaceuticals Ltd.

Date of preparation: October 1996.

LOS/ADV 1327

This woman

HAS HEPATITIS C



Do you tell her IT COULD BE FATAL?

85,000 people in the UK
have chronic HCV*



50,000 of them will
develop cirrhosis*



10,000 will develop
liver cancer*



Many will die
prematurely



She could be one of them



Viraferon is not a vaccine,
nor a miracle cure.

But should this patient
develop chronic HCV
it could save her life.

VIRAFERON
INTERFERON ALFA-2B (rbe)



Today, for the future

* Estimates based on current incidence and epidemiology of hepatitis C ^{1,2}

PRESCRIBING INFORMATION

For full prescribing information, please refer to full Data Sheet for Viraferon. Viraferon is available as 5 million IU or 25 million IU/vial of recombinant interferon Alfa-2b (rbe) for intravenous use. **Uses:** Treatment of chronic hepatitis C (Non-A, Non-B). **Dosage and Administration:** The recommended dose is 3 million IU administered intravenously three times a week for a period of 18 months. **Chronic Hepatitis C/Non-A, Non-B:** The recommended dose is usually in the range of 2.5 million IU/m² of body surface area administered intravenously three times a week for a period of 18 months. **Chronic Hepatitis C/Non-A, Non-B:** The recommended dose is 3 million IU administered intravenously three times a week. Most patients who receive Viraferon show a moderate improvement in ALT levels within 4-6 weeks. In those patients, therapy should be continued for 18 months. **Contraindications, Warnings, Precautions:** **Contraindications:** A history of hypersensitivity to any component of Viraferon Injection contraindicates the use of Viraferon Injection. **Warnings and Precautions:** Patients with pre-existing cardiac disease, severe renal impairment, epilepsy and/or compromised respiratory system function, chronic hepatitis with advanced cirrhosis of the liver, chronic hepatitis who are being or have been treated with immunosuppressive agents, patients with a history of corticosteroid withdrawal, patients with a history of autoimmune disease not controlled by therapy, patients with a history of pulmonary disease, diabetes mellitus, coagulation disorders, depression. Moderate to severe depression may require reduction of dosage of Viraferon therapy. Patients with severe depression should be monitored as they may be at increased risk of clinical depression. Patients with a recent history of cardiac disease should be monitored as they may be at increased risk of hypotension, arrhythmias, myocardial infarction, or other cardiovascular events. Hypotension should be observed. Patients with a history of depression should be monitored. Patients with a history of pulmonary disease should be monitored. Pulmonary disease, including pneumonia, has been observed in patients receiving Viraferon. CNS effects have been observed in patients receiving Viraferon, manifested by symptoms such as fatigue, headache, depression, and irritability. These effects have been observed especially at high doses. Patients with chronic hepatitis B should be monitored for chronic hepatitis B development. Abnormalities of thyroid or pituitary function with Viraferon may occur. Patients with pre-existing disease. Ocular effects have been observed in patients taking concomitant narcotics. Patients taking xanthine derivatives should be monitored and dosage adjusted as necessary. No information is available on the use of Viraferon in human pregnancy or its effect on human lactation. Viraferon should only be given if the benefits clearly outweigh the potential hazard to the foetus or nursing infant. **Side Effects:** Elevated liver function test results, depression, and thrombocytopenia have been observed especially at higher doses. Other side effects include: haemorrhages, cotton wool spots and retinal changes, and vein obstruction have been observed rarely. The most common adverse effects are 'flu-like' symptoms, including fever, malaise, myalgia, headache, and fatigue, which are generally dose-related and reversible and can be ameliorated by dose adjustment. 'Flu-like' symptoms can be relieved by the use of paracetamol. **Package Quantities:** 10 Million IU (2ml) vials: 10 x 10M IU vial: £141.30. 25 Million IU (5ml) vials: 10 x 25M IU vial: £169.56. 25 Million IU (5ml) vials: 2 x 25M IU vials: £282.60. **Product Licence Numbers:** PL 00000000. **Further information is available from the Marketing Authoriser:** Schering-Plough Ltd, Kenilworth, New Jersey, NJ, USA. **Further information is available from the Marketing Authoriser:** Schering-Plough Ltd, Kenilworth, New Jersey, NJ, USA. **Further information is available from the Marketing Authoriser:** Schering-Plough Ltd, Kenilworth, New Jersey, NJ, USA. **Further information is available from the Marketing Authoriser:** Schering-Plough Ltd, Kenilworth, New Jersey, NJ, USA.

**Dosage in hepatitis C:
3M IU
three times a week
for up to 18 months**

Date of Preparation: January 1996

Schering-Plough

Medication or operation?

We have the figures to help you make the right decisions.

The causes of gastrointestinal diseases are often difficult to identify. Gastroesophageal reflux for example, can have a variety of causes, each of which may be most effectively treated with different therapies.

IDENTIFYING THE CAUSES OF GERD

Synectics' new Polygram for Windows™ software along with the Ditrappert™ Mk III and the PC Polygraf HR, give you the ability to look at the causes of GERD from every angle. Evaluate sphincter tone and function with the Polygraf HR and the Esophageal Manometry Analysis Module. Measure the severity of reflux while also looking for correlation between symptoms and reflux with the Ditrappert™ Mk III and the EsopHogram™ Reflux Analysis Module.

SIMPLE AND EFFICIENT

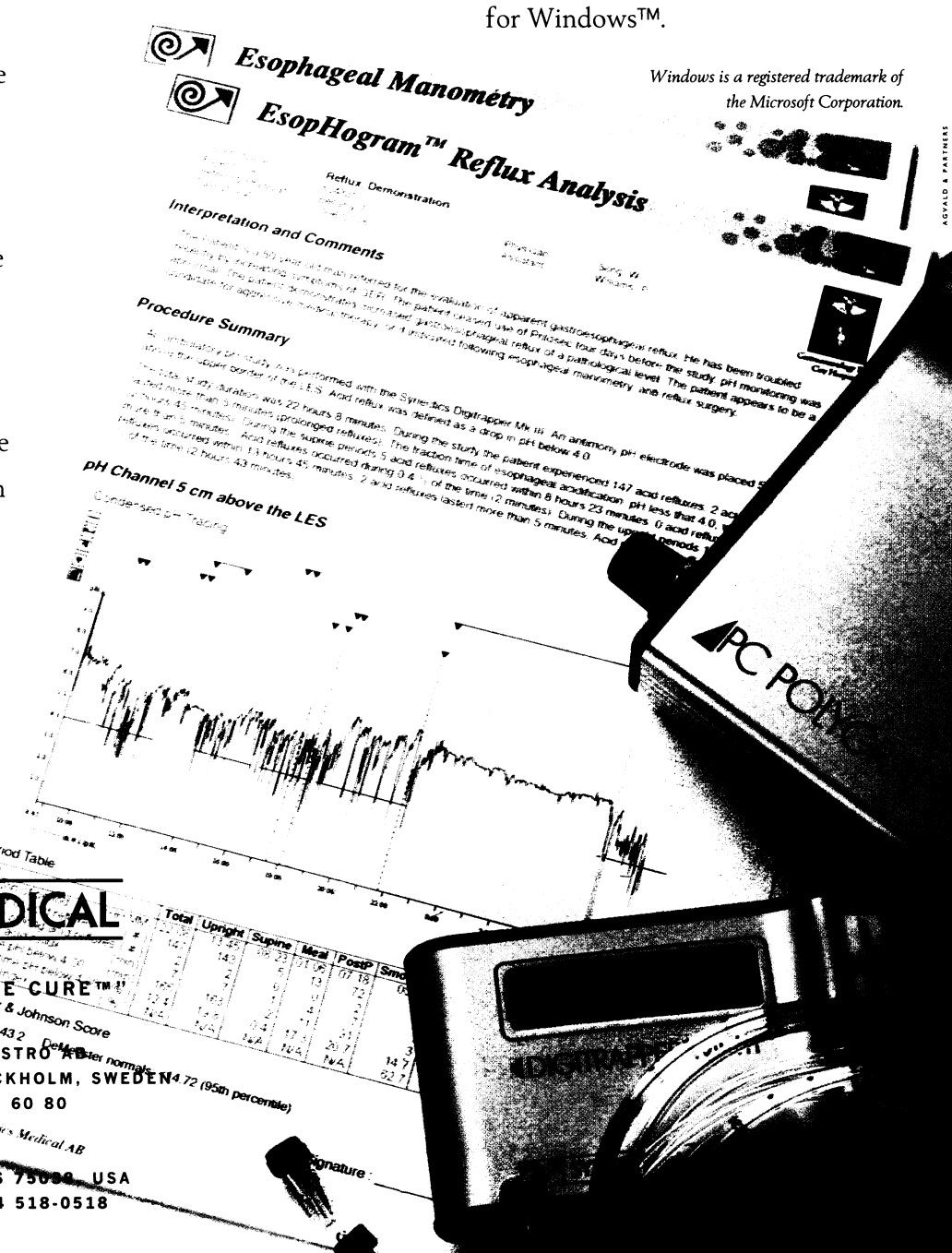
Extensive on-line help and performance checks will help you get through each procedure quickly and easily while also obtaining accurate results. Concise, one page, color reports make interpreting your studies and informing referring physicians both trouble free and efficient.

WITH MODULARITY COMES POWER

Select only the diagnostic tools you need today, adding more as your practice develops. We call this modular approach Synectics System Thinking.

Medication or operation? The answer is in your reach with Polygram for Windows™.

Windows is a registered trademark of the Microsoft Corporation.



"A GOOD DIAGNOSIS IS HALF THE CURE™"
 Copyright © 1998 Synectics Medical AB
INTERNATIONAL OFFICE: SYNECTICS GASTRO
 RENSTIERNAS GATA 12 • S-116 28 STOCKHOLM, SWEDEN
 TEL: +46 8 462 60 50 • FAX: +46 8 462 60 80
 E-MAIL: pw.info@synectics.se
USA OFFICE: SYNECTICS MEDICAL INC.
 1425 GREENWAY DRIVE • IRVING, TEXAS 75038 USA
 TEL: +1 800 798-3129 • OFFICE: +1 214 518-0518
 FAX: +1 214 518-0080

AGVALE & PARTNERS

Entocort® CR 3mg Capsules (budesonide)

PRESCRIBING INFORMATION (Refer to full Summary of Product Characteristics before prescribing). **Presentation:** Capsules containing 3mg budesonide **Use:** Induction of remission of mild to moderate Crohn's disease affecting the ileum and/or ascending colon. **Dosage and Administration: Adults:** 9mg once daily in the morning before breakfast for up to 8 weeks. When treatment is to be discontinued, the dose should normally be reduced for the last 2 to 4 weeks of therapy. **Children:** Not recommended. **Elderly:** No special dose adjustment, though limited experience in elderly. **Contra-Indications:** Bacterial, fungal or viral infections. Known hypersensitivity. **Precautions:** Treatment with Entocort CR Capsules results in lower systemic steroid levels than conventional oral

We've re-shaped steroid delivery for Crohn's -



steroid therapy. However, in common with all oral steroids use with caution in patients with tuberculosis, hypertension, diabetes mellitus, osteoporosis, peptic ulcer, glaucoma or cataracts, chicken pox and measles or patients with a family history of diabetes or glaucoma. Corticosteroids may cause suppression of HPA axis and reduce the stress response. As with any glucocorticoid steroids, blood levels may increase in patients with compromised liver function. **Interactions:** Cholestyramine may reduce uptake.

Pregnancy and lactation: Use during pregnancy should be avoided unless essential. It is not known whether budesonide passes into breast milk.

Adverse Events: Those characteristic of systemic corticosteroid therapy. Others include (most classed as mild to moderate) - dyspepsia, muscle cramps, tremor, palpitations, nervousness, blurred vision, skin reactions and menstrual disorders. In clinical studies adverse event incidence was similar to placebo. **Legal Category:** POM. **Packs and prices:** Polyethylene bottles of 100 capsules. Price: £90.00. **Pharmacological Properties:** Budesonide has a high local anti-inflammatory effect and a significantly lower effect on HPA axis and adrenal function than prednisolone.

Marketing Authorization No: PL0017/0359.

References: 1. Edsbäcker S, Wollmer P, Nilsson A, et al. *Abstract. Gastroenterol* 1993; **104** (4pt 2): A695. 2. Rutgeerts P, Löfberg R, Malchow H, et al. *N Engl J Med* 1994; **331**: 842-845. 3. Jewell DP, Campieri M, Järnerot G, et al. *Gastro Int* 1993; **6**: 1-4. 4. Campieri M, Ferguson A, Doe W, and The International Budesonide Study Group. *Abstract. Gastroenterol* 1995; **108** (4 suppl.): A790.

ASTRA Further information available from the **Marketing Authorization Holder:** Astra Pharmaceuticals Limited, Kings Langley, Herts, WD4 8DH. Tel: (01923) 266191.

Entocort is a registered trademark of Astra Pharmaceuticals Limited.

Date of preparation: May 1996

ENT/ADV 1047

and taken the edge off side effects.

Astra have developed Entocort® CR from an established steroid, budesonide, in a formulation that's designed specifically for Crohn's disease.

Entocort CR acts where it's needed - a unique delivery system targets the ileum and ileocaecal area,¹ achieving rapid results equivalent to prednisolone.² But

more importantly, the fact that 90% of the budesonide is metabolised on first pass through the liver,³ means that Entocort CR sets a low level of systemic steroid side effects.⁴

With efficacy and low steroid side effects,^{2,4} Entocort CR is tailor-made for Crohn's disease.

ENTOCORT® CR 3mg
budesonide **Capsules**

A tried and trusted steroid, adapted for Crohn's disease.

Entocort® CR 3mg Capsules (budesonide)

PRESCRIBING INFORMATION (Refer to full Summary of Product Characteristics before prescribing). **Presentation:** Capsules containing 3mg budesonide **Use:** Induction of remission of mild to moderate Crohn's disease affecting the ileum and/or ascending colon. **Dosage and Administration: Adults:** 9mg once daily in the morning before breakfast for up to 8 weeks. When treatment is to be discontinued, the dose should normally be reduced for the last 2 to 4 weeks of therapy. **Children:** Not recommended. **Elderly:** No special dose adjustment, though limited experience in elderly. **Contra-Indications:** Bacterial, fungal or viral infections. Known hypersensitivity. **Precautions:** Treatment with Entocort CR Capsules results in lower systemic steroid levels than conventional oral

We've re-shaped steroid delivery for Crohn's -



steroid therapy. However, in common with all oral steroids use with caution in patients with tuberculosis, hypertension, diabetes mellitus, osteoporosis, peptic ulcer, glaucoma or cataracts, chicken pox and measles or patients with a family history of diabetes or glaucoma. Corticosteroids may cause suppression of HPA axis and reduce the stress response. As with any glucocorticoid steroids, blood levels may increase in patients with compromised liver function. **Interactions:** Cholestyramine may reduce uptake. **Pregnancy and lactation:** Use during pregnancy should be avoided unless essential. It is not known whether budesonide passes into breast milk. **Adverse Events:** Those characteristic of systemic corticosteroid therapy. Others include (most classed as mild to moderate) - dyspepsia, muscle cramps, tremor, palpitations, nervousness, blurred vision, skin reactions and menstrual disorders. In clinical studies adverse event incidence was similar to placebo. **Legal Category:** POM. **Packs and prices:** Polyethylene bottles of 100 capsules. Price: £90.00. **Pharmacological Properties:** Budesonide has a high local anti-inflammatory effect and a significantly lower effect on HPA axis and adrenal function than prednisolone. **Marketing Authorization No:** PLO017/0359.

References: 1. Edsbäcker S, Wollmer P, Nilsson A, et al. Abstract. *Gastroenterol* 1993; **104** (4pt 2): A695. 2. Rutgeerts P, Löfberg R, Malchow H, et al. *N Engl J Med* 1994; **331**: 842-845. 3. Jewell DP, Campieri M, Järnerot G, et al. *Gastro Int* 1993; **6**: 1-4. 4. Campieri M, Ferguson A, Doe W, and The International Budesonide Study Group. Abstract. *Gastroenterol* 1995; **108** (4 suppl.): A790.

Further information available from the **Marketing Authorization Holder:** Astra Pharmaceuticals Limited, Kings Langley, Herts, WD4 8DH. Tel: (01923) 266191. Entocort is a registered trademark of Astra Pharmaceuticals Limited. **Date of preparation:** May 1996 ENT/ADV 1047

and taken the edge off side effects.

Astra have developed Entocort® CR from an established steroid, budesonide, in a formulation that's designed specifically for Crohn's disease.

Entocort CR acts where it's needed - a unique delivery system targets the ileum and ileocaecal area,¹ achieving rapid results equivalent to prednisolone.² But


more importantly, the fact that 90% of the budesonide is metabolised on first pass through the liver,³ means that Entocort CR sets a low level of systemic steroid side effects.⁴

With efficacy and low steroid side effects,^{2,4} Entocort CR is tailor-made for Crohn's disease.

ENTOCORT® CR 3mg Capsules
budesonide

A tried and trusted steroid, adapted for Crohn's disease.

NOW CROHN'S SUFFERERS
CAN EXPERIENCE THE
FREEDOM OF 'ASACOL'



Already the most widely prescribed 5-ASA in the treatment of inflammatory bowel disease, 'Asacol' is now indicated for the maintenance of remission of Crohn's ileo-colitis.

Now indicated for the maintenance of remission of Crohn's ileo-colitis.

SK **SmithKline Beecham**
Pharmaceuticals

Smith Kline & French Laboratories, Welwyn Garden City, Hertfordshire AL7 1EY.
Authorised user of the trade mark 'Asacol' in the UK. ©1996 Smith Kline & French Laboratories.
Prescribing information is available on reverse

ASACOL
MESALAZINE* (5-AMINOSALICYLIC ACID)
FREEDOM IN IBD



ALL THE
STRENGTH
OF 'ASACOL'
AVAILABLE
IN FOAM

Prescribing Information: Presentation 'Asacol' Tablets, PL 0002/0173, each containing 400 mg mesalazine (5-aminosalicylic acid) coated with a pH-dependent acrylic-based resin (Eudragit S) formulated to release the active ingredient in the terminal ileum and colon. Blister packs of 120 (12 x 10), £43.58. 'Asacol' Suppositories 250 mg, PL 0002/0158, each containing 250 mg mesalazine. 20, £7.15. 'Asacol' Suppositories 500 mg, PL 0002/0195 each containing 500 mg mesalazine. 10, £7.15. 'Asacol' Foam Enema, PL 0002/0222, 1 g mesalazine per metered dose. Carton containing can of 14 metered doses, 14 disposable applicators and 14 disposable plastic bags. £39.60. **Uses** *Ulcerative colitis*: Treatment of mild to moderate acute exacerbations. Suppositories and Foam Enema particularly appropriate for distal disease. Maintenance of remission, Tablets and Suppositories. *Crohn's ileo-colitis*: Maintenance of remission, Tablets. **Dosage and administration: Adults: Tablets: Acute disease**: 6 tablets a day, in divided doses, with concomitant corticosteroid therapy where clinically indicated. *Maintenance therapy*: 3 to 6 tablets a day, in divided doses. **Suppositories: 250 mg**: 3 to 6 a day, in divided doses, with the last dose at bedtime. **500 mg**: A maximum of 3 a day, in divided doses, with the last dose at bedtime. **Foam Enema**: 1 or 2 metered doses as single daily dose. **Children**: No dosage recommendation. **Contra-indications**: A history of sensitivity to salicylates or renal sensitivity to sulphasalazine. Severe renal impairment (GFR <20 ml/min). Children under 2 years of age. **Precautions**: Best avoided in patients with established renal impairment but, if necessary, use with caution. Very rare reports with mesalazine of serious blood dyscrasias, perform haematological investigations if patient develops unexplained bleeding, bruising, purpura, anaemia, fever or sore

throat. Stop treatment if suspicion or evidence of blood dyscrasia. Do not give tablets with lactulose or similar preparations which lower stool pH. Only use during pregnancy if benefits outweigh the risk. Avoid during lactation. Caution in elderly and only where renal function is normal. **Adverse reactions**: Nausea, diarrhoea, abdominal pain, headache. Exacerbation of symptoms of colitis. Rare reports of leucopenia, neutropenia, agranulocytosis, aplastic anaemia, thrombocytopenia, pancreatitis, hepatitis, allergic lung reactions, lupus erythematosus-like reactions, rash, interstitial nephritis and nephrotic syndrome, with oral mesalazine treatment, usually reversible on withdrawal. Renal failure has been reported. Suspect nephrotoxicity in patients developing renal dysfunction. **Legal category**: POM. 5.6.96. Smith Kline & French Laboratories, Welwyn Garden City, Hertfordshire AL7 1EY. Authorised user of the trade mark 'Asacol' in the UK. ©1995 Smith Kline & French Laboratories. *Mesalazine is the British approved name of 5-aminosalicylic acid.

ASACOL
MESALAZINE* (5-AMINOSALICYLIC ACID)

FIVE STAR, 5-ASA
COLITIS CONTROL

SB **SmithKline Beecham**
Pharmaceuticals
Healthy Alliance
partnership beyond prescription

NO

REFLUX

NO

BLOATING

NO

BELCHING

NO

PROBLEM



AFTER ANTACIDS
PrepulsidTM
 cisapride
 A PHYSIOLOGICAL APPROACH

ABBREVIATED PRESCRIBING INFORMATION Please refer to data sheet before prescribing. **Indications:** GASTRO-OESOPHAGEAL REFLUX DISEASE: Treatment of symptoms and healing of mucosal lesions; maintenance treatment of reflux oesophagitis. DYSPEPSIA: Treatment of symptoms such as epigastric pain, early satiety, bloating and belching where organic disease has been excluded. **IMPAIRED GASTRIC EMPTYING:** Relief of symptoms such as epigastric pain, early satiety, anorexia, bloating and nausea associated with delayed gastric emptying secondary to diabetes, systemic sclerosis and autonomic neuropathy. **Dosage and Administration:** Adults and children twelve years and over: Take 15 minutes before food. **REFLUX:** 20mg Prepulsid b.d. (before breakfast and at bedtime) or 10mg Prepulsid b.d. if necessary, night time symptoms can be treated with a fourth 10mg dose at bedtime for 12 weeks to heal oesophagitis. For long term maintenance therapy, 20mg once daily (at bedtime) or 10mg b.d. (before breakfast and at bedtime) increasing to 20mg b.d. in patients whose lesions were initially very severe. **DYSPEPSIA:** 10mg Prepulsid t.i.d. Usual course of treatment is 4 weeks. **IMPAIRED GASTRIC EMPTYING:** 10mg Prepulsid t.i.d. or q.i.d. An initial course of 6 weeks is recommended but longer treatment may be required. **CHILDREN:** Not recommended in children under 12. **Elderly:** As for adults, but monitor response. **ABNORMAL RENAL LIVER FUNCTION:** Initially dose should be halved. **Contra-indications:** Pregnancy, patients in whom gastrointestinal stimulation might be dangerous, concomitant oral or parenteral ketoconazole, itraconazole or miconazole, flucanazole.

erythromycin and clarithromycin, hypersensitivity to Prepulsid. **Warnings:** In view of reports of isolated cases of QT-interval prolongation and/or torsade de pointes (causal relationship not established), the recommended dose of Prepulsid should not be exceeded and it should be used with caution in patients with conditions leading to QT-interval prolongation, congenital QT-interval prolongation or in patients taking medication known to prolong QT-interval. Not recommended whilst breast feeding. **Drug Interactions:** Absorption from the stomach of concomitantly administered drugs may be diminished, whereas absorption of drugs from the small intestine may be accelerated. For drugs that require careful individual titration e.g. anti-coagulants, it may be useful to measure plasma concentrations. In patients receiving anti-coagulants, check prothrombin time as it may be increased. The sedative effects of benzodiazepines and alcohol may be accelerated when given with Prepulsid. The effects of Prepulsid are antagonised by anticholinergic drugs. Prepulsid is metabolised mainly via the cytochrome P450 3A4 enzyme. Oral ketoconazole significantly inhibits the metabolism of Prepulsid on the basis of *in vitro* data. Itraconazole and miconazole may also have this effect. Co-administration with oral ketoconazole can result in QT-interval prolongation, which can lead to ventricular arrhythmias (see warnings). Concomitant administration with oral or parenteral ketoconazole, itraconazole, miconazole, flucanazole, erythromycin and clarithromycin, is therefore contra-indicated. Concomitant administration with cimetidine increases peak plasma levels and the AUC of Prepulsid, while the absorption of cimetidine and

ranitidine is accelerated when co-administered with Prepulsid. The level of change is unlikely to be clinically significant. **Side Effects:** Abdominal cramps, borborygmi and loose stools are mainly transient. Should severe abdominal cramps occur with single administrations of 20mg Prepulsid halve the dose per administration and double the frequency of dosing. Less frequent side effects include headache and light-headedness. Reports of hyper-sensitivity, convulsions, extrapyramidal effects and increased urinary frequency have been received. Exceptionally, reversible liver function abnormalities have been reported - causal relationship not established. **Overdosage:** The most frequently reported symptoms are abdominal cramping and increased stool frequency. Treatment includes activated charcoal and close observation, and increased stool frequency. Treatment includes activated charcoal and close observation. Patients should also be evaluated for possible QT-interval prolongation and for factors that can predispose to the occurrence of torsade de pointes. **Preparation:** Prepulsid Tablets: pack of 120 tablets each containing 10mg cisapride. Prepulsid Suspension: 500ml bottles containing cisapride 5mg/5ml. **Pharmaceutical Precautions:** Prepulsid Tablets: store at room temperature in a dry place and protect from light. Prepulsid Suspension: store at room temperature (below 25°C). **PL Nos:** Prepulsid Tablets PL 0247/0136. Prepulsid Suspension PL 0242/0157. **Product Licence Holder:** Janssen-Cilag Ltd, Sanderson, High Wycombe Bucks, HP14 4HJ. **Basic NHS Cost:** 120 tablets £37.60; 500 ml bottle suspension £15.60. **Legal Category:** POM. **Date of Preparation:** April 1996. **Janssen-Cilag**

TM=Trademark. Copyright 00981 54