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1.3.1 SUMMARY OF PRODUCT CHARACTERISTICS

1. NAME OF THE MEDICINAL PRODUCT

Spectromax 250

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each film coated tablets contains Clarithromycin 250 mg

Sr. No	Ingredients	Quantity / tablet in mg	Function of Ingredients
1	Clarithromycin	250	Active Ingredient
2	Sodium Starch Glycolate	25.0	Disintegrant
3	Calcium Phosphate Anhydrous	74.5	Filler
4	Maize starch	78.2	Filler
5	Hydroxypropyl Cellulose (Klucel EF)	10.3	Binder
6	Sodium Starch Glycolate	25.0	Disintegrant
7	Purified talc	15.0	Lubricant
8	Colloidal Anhydrous Silica (Aerosil 200)	7.0	Glidant

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9	Magnesium Stearate	10.0	Lubricant
10	Opadry-OY 8487 Yellow	20.0	Coating
11	White Bees wax (Pellets)	q.s	Polishing agent
12	Purified water	q.s	Vehicle

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3. PHARMACEUTICAL FORM

Yellow circular, shallow biconvex film coated tablets with "609" embossed on one side and plain on other side.

4. THERAPEUTIC INDICATIONS

Clarithromycin is indicated for the treatment of infections due to susceptible or ganisms. Such infections include:

- 1. Lower respiratory tract infections (e.g. bronchitis, pneumonia).
- 2. Upper respiratory tract infections (e.g. pharyngitis, sinusitis).
- 3. Skin and soft tissue infections (e.g. folliculitis, cellulitis, erysipalis).
- 4. Disseminated or lo calised mycobacterial infections due to *Mycobacterium avium* or *Mycobacterium intracellulare*. Localised i nfections due t o *Mycobacterium chelonae*, *Mycobacterium fortuitum* or *Mycobacterium kansasii*.
- 5. Clarithromycin is indicated for the prevention of disseminated *Mycobacterium avium* complex infection in HIV infected patients with CD4 lymphocyte counts less than or equal to 100/mm³.
- 6. Clarithromycin in the presence of acid suppression is indicated for the eradication of *H.pylori*, resulting in decreased recurrence of duodenal ulcer. (See further information).

Clarithromycin tablets are indicated in adults and children 12 years and older.

Consideration should be given to official guidance on the appropriate use of antibacterial agents.

Further Information: *H. pylori* is strongly associated with peptic ulcer disease. Ninety to 100% of patients with duode nal ulcers are infected with this a gent. E radication of *H.pylori* has be en shown to markedly reduce the rate of duode nal ulcer recurrence, thereby reducing the need for maintenance anti-secretory therapy.

In a well controlled double-blind study, *H.pylori* infected patients with duodenal ul cer received clarithromycin 500mg TID for 14 days with omeprazole 40mg daily for 28 days.

Clarithromycin has been used in other treatment regimens for the eradication of *H.pylori*. These regimens i nclude: c larithryomycin pl us t inidazole a nd om eprazole; a nd c larithromycin pl us tetracycline, bismuth subsalicylate, and ranitidine.

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4.1 POSOLOGY AND METHOD OF ADMINISTRATION

Adults: The usual recommended dos age of clarithromycin in a dults is one 250m g tablet twice daily. In more severe infections, the dos age can be increased to 500m g twice daily. The usual duration of therapy is 6 to 14 days.

Children under 12 years: Use of clarithromycin tablets is not recommended for children under 12 years. Clinical trials have been conducted using clarithromycin paediatric suspension in children 6 months to 12 years of age. Therefore, children under 12 years of age should use clarithromycin paediatric suspension (granules for oral suspension).

Children over 12 years: As for adults.

In patients with renal impairment with creatinine clearance less than 30 ml/min, the dos age of clarithromycin should be reduced by one-half, i.e. 250mg once daily or 250mg twice daily in more severe infections. Treatments should not be continued beyond 14 days in these patients.

Dosage in patients with mycobacterial infections: The recommended starting dose is 500mg twice daily. If no c linical or bacteriologic r esponse is observed in 3 t o 4 w eeks, the dose may be increased to 1000mg twice daily. Treatment of disseminated MAC infections in A IDS patients should be continued, as long as clinical microbiological benefit is demonstrated. Clarithromycin should be used in conjunction with other antimycobacterial agents.

Treatment of other non-tuberculous mycobacterial infections should continue at the discretion of the physician.

Dosage for MAC prophylaxis: The recommended dos age of clarithromycin in a dults is 500m g twice daily.

Eradication of H.pylori:

Dual Therapy (14 days): The recommended dose of clarithromycin is 500mg three times daily for 14 days. (see Further Information).

Triple Therapy (7 days): Clarithromycin (500mg) twice daily and a proton pump inhibitor (at the approved daily dose)* should be given with amoxycillin 1000mg twice daily for 7 days.

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Triple Therapy (7 days): Clarithromycin (500mg) twice daily and a proton pump inhibitor (at the approved daily dose)* should be given with metronidazole 400mg twice daily for 7 days.

Triple Therapy (7-10 days): Clarithromycin (500mg) twice daily should be given with amoxycillin 1000mg twice daily and omeprazole 20mg daily for 7-10 days.

* see individual data sheets/SPCs for the dose recommendations for *H. pylori* eradication.

4.2 CONTRAINDICATIONS

Clarithromycin is contraindicated in patients with known hypersensitivity to macrolide antibiotic drugs or any of its excipients.

Concomitant administration of clarithromycin and any of the following drugs is contraindicated: astemizole, cisapride, pimozide or terfenadine as this may result in QT prolongation and cardiac arrhythmias, including v entricular ta chycardia, ventricular fibrillation, and torsades de pointe (see s ection 4.5) . C oncomitant a dministration of c larithromycin and e rgotamine or dihydroergotamine is contraindicated, as this may result in ergot toxicity.

Clarithromycin should not be given to patients with history of QT prolongation or ventricular cardiac arrhythmia, including torsades de pointe (see sections 4.4 and 4.5).

Clarithromycin should not be used concomitantly with HMG-CoA reductase inhibitors (statins), lovastatin or simvastatin, due to the risk of rhabdomyolysis. Treatment with these agents should be discontinued during clarithromycin treatment (see section 4.4).

Clarithromycin should not be given to patients with hypokalaemia (risk of prolongation of QT-time).

Clarithromycin's hould not be us ed i n pa tients who suffer from severe he patic failure i n combination with renal impairment.

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4.3 SPECIAL WARNINGS AND PRECAUTIONS FOR USE

The physician should not prescribe clarithromycin to pregnant women without carefully weighing the benefits against risk, particularly during the first three months of pregnancy (see section 4.6).

Caution is advised in patients with severe renal insufficiency (see section 4.2).

Clarithromycin is principally ex creted by the liver. Therefore caution should be exercised in administering the antibiotic to patients with impaired he patic function. Caution should also be exercised when administering clarithromycin to patients with moderate to severe renal impairment.

Cases of fatal hepatic failure (see section 4.8) have been reported. Some patients may have had pre-existing he patic di sease o r m ay have be en t aking ot her he patotoxic m edicinal p roducts. Patients should be a dvised to s top treatment and c ontact their doc tor if signs and s ymptoms of hepatic disease develop, such as anorexia, jaundice, dark urine, pruritus, or tender abdomen.

Pseudomembranous c olitis ha s be en r eported with ne arly a ll a ntibacterial a gents, i ncluding macrolides, and ma y r ange in severity f rom mild to life-threatening. *Clostridium difficile* associated diarrhoea (CDAD) ha s b een reported with use of ne arly all ant ibacterial agents including clarithromycin, and may range in severity from mild diarrhoea to fatal colitis. Treatment with antibacterial agents alters the normal flora of the colon, which may lead to overgrowth of *C. difficile*. CDAD must be considered in all patients who present with diarrhoea following antibiotic use. Careful medical history is necessary since CDAD has been reported to occur over two months after the a dministration of a ntibacterial a gents. T herefore, di scontinuation of c larithromycin therapy should be considered regardless of the indication. Microbial testing should be performed and adequate treatment initiated. Drugs inhibiting peristalsis should be avoided.

Exacerbation of s ymptoms of m yasthenia gravis has be en r eported in pa tients r eceiving clarithromycin therapy.

There have been post-marketing reports of colchicine toxicity with concomitant use of clarithromycin and colchicine, especially in the elderly, some of which occurred in patients with renal insufficiency. Deaths have been reported in some such patients (see section 4.5). If concomitant administration of colchicine and clarithromycin is necessary, patients should be monitored for clinical symptoms of colchicine toxicity.

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Caution i s a dvised r egarding concomitant administration of c larithromycin and triazolobenzodiazepines, such as triazolam, and midazolam (see section 4.5).

Caution is a dvised regarding concomitant a dministration of clarithromycin with other ototoxic drugs, especially with aminoglycosides. Monitoring of vestibular and auditory function should be carried out during and after treatment.

Due to the risk for QT prolongation, clarithromycin should be used with caution in patients with coronary artery disease, severe cardiac insufficiency, hypomagnesemia, bradycardia (<50 bpm), or when co-administered with other medicinal products associated with QT prolongation (see section 4.5). C larithromycin m ust not be us ed in patients with c ongenital or do cumented a cquired QT prolongation or history of ventricular arrhythmia (see section 4.3).

<u>Pneumonia</u>: In view of the emerging resistance of *Streptococcus pneumoniae* to macrolides, it is important that sensitivity testing be performed when prescribing clarithromycin for community-acquired pn eumonia. In hos pital-acquired pn eumonia, c larithromycin s hould be us ed in combination with additional appropriate antibiotics.

Skin and soft tissue infections of mild to moderate severity: These infections are most often caused by *Staphylococcus aureus* and *Streptococcus pyogenes*, bot h of w hich m ay be r esistant t o macrolides. Therefore, it is important that sensitivity testing be performed. In cases where *beta*—lactam antibiotics cannot be used (e.g. allergy), other antibiotics, such as clindamycin, may be the drug of first choice. Currently, macrolides are only considered to play a role in some skin and soft tissue i nfections, s uch as t hose c aused by *Corynebacterium minutissimum* (erythrasma), acne vulgaris, and erysipelas and in situations where penicillin treatment cannot be used.

In the eve nt of s evere acute h ypersensitivity r eactions, such as an phylaxis, Stevens-Johnson Syndrome, a nd t oxic e pidermal ne crolysis, c larithromycin t herapy s hould be di scontinued immediately and appropriate treatment should be urgently initiated.

Clarithromycin should be used with caution when administered concurrently with medications that induce the cytochrome CYP3A4 enzyme (see section 4.5).

<u>HMG-CoA reductase inhibitors:</u> Concomitant use of clarithromycin with lovastatin or simvastatin is contraindicated (see section 4.3). As with other macrolides, clarithromycin has been reported to increase conc entrations of H MG-CoA r eductase i nhibitors (see s ection 4.5). R are reports of rhabdomyolysis have been reported in patients taking these drugs concomitantly. Patients should be monitored for signs and symptoms of myopathy. Rare reports of rhabdomyolysis have also been

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reported in patients taking atorvastatin or rosuvastatin concomitantly with clarithromycin. When used with clarithromycin, a torvastatin or rosuvastatin should be a dministered in the lowest possible doses. Adjustment of the statin dose or use of a statin that is not dependent on C YP3A metabolism (e.g. fluvastatin or pravastatin) should be considered.

Oral hypoglycemic agents/Insulin: The concomitant use of clarithromycin and oral hypoglycemic agents a nd/or i nsulin c an r esult in significant hypoglycemia. With certain hypoglycemic drugs such as nateglinide, pi oglitazone, repaglinide and rosiglitazone, inhibition of CYP3A enzyme by clarithromycin may be involved and could cause hypolgycemia when used concomitantly. Careful monitoring of glucose is recommended.

<u>Oral a nticoagulants</u>: T here is a risk of serious he morrhage and significant e levations in International N ormalized Ratio (INR) and prothrombin time when clarithromycin is coadministered with warfarin (see section 4.5). INR and prothrombin times should be frequently monitored while patients are receiving clarithromycin and oral anticoagulants concurrently.

Use of any antimicrobial therapy, such as clarithromycin, to treat *H. pylori infection* may select for drug-resistant organisms.

Long-term us e m ay, as with other antibiotics, result in colonization with increased numbers of non-susceptible bacteria and fungi. If superinfections oc cur, a ppropriate therapy should be instituted.

Attention should also be paid to the possibility of cross resistance between clarithromycin and other macrolide drugs, as well as lincomycin and clindamycin.

4.4 DRUG INTERACTIONS:

The use of the following drugs is strictly contraindicated due to the potential for severe drug interaction effects:

Cisapride, pimozide, astemizole and terfenadine

Elevated cisapride levels have been reported in patients receiving clarithromycin and cisapride concomitantly. This may result in QT prolongation and cardiac arrhythmias including ventricular tachycardia, ventricular fibrillation and torsades de pointes. Similar effects have been observed in patients taking clarithromycin and pimozide concomitantly (see section 4.3, Contraindications).

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Macrolides have been reported to alter the metabolism of terfenadine resulting in increased levels of t erfenadine w hich has oc casionally been as sociated with cardiac arrhythmias, such as Q T prolongation, ventricular tachycardia, ventricular fibrillation and torsades de pointes (see section 4.3, Contraindications). In one study in 14 he althy volunteers, the concomitant administration of clarithromycin and terfenadine resulted in a 2 t o 3-fold increase in the serum level of the acid metabolite of t erfenadine and in prolongation of the Q T interval w hich did not 1 ead t o a ny clinically detectable effect. Similar effects have been observed with concomitant administration of astemizole and other macrolides.

Ergotamine/dihydroergotamine

Post-marketing r eports indi cate that c o-administration of c larithromycin with ergotamine or dihydroergotamine has been associated with acute ergot toxicity characterized by vasospasm, and ischemia of the extremities and other tissues including the central nervous system. Concomitant administration of clarithromycin and these medicinal products is contraindicated (see section 4.3, Contraindications).

Effects of Other Medicinal Products on Clarithromycin

Drugs that are inducers of CYP3A (e.g. rifampicin, phenytoin, carbamazepine, phenobarbital, St John's wort) may induce the metabolism of clarithromycin. This may result in sub-therapeutic levels of clarithromycin leading to reduced efficacy. Furthermore, it might be necessary to monitor the plasmalevels of the CYP3A inducer, which could be increased owing to the inhibition of CYP3A by clarithromycin (see also the relevant product information for the CYP3A4 inhibitor administered). Concomitant administration of rifabutin and clarithromycin resulted in an increase in rifabutin, and decrease in clarithromycin serum levels together with an increased risk of uveitis.

The following drugs are known or suspected to affect circulating concentrations of clarithromycin; clarithromycin dosage adjustment or consideration of alternative treatments may be required.

Efavirenz, nevirapine, rifampicin, rifabutin and rifapentine

Strong i nducers of the cytochrome P 450 m etabolism system such as e favirenz, ne virapine, rifampicin, rifabutin, and rifapentine may accelerate the metabolism of clarithromycin and thus lower the plasmal evels of clarithromycin, while i ncreasing those of 14-OH-clarithromycin, a metabolite that is a lso microbiologically active. Since the mic robiological activities of clarithromycin and 14-OH-clarithromycin are different for different bacteria, the intended therapeutic effect could be impaired during concomitant administration of clarithromycin and enzyme inducers.

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Fluconazole

Concomitant administration of fluconazole 200 mg daily and clarithromycin 500 mg twice daily to 21 healthy vol unteers led to increases in the mean steady-state minimum clarithromycin concentration (Cmin) and area under the curve (AUC) of 33% and 18% respectively. Steady state concentrations of the active metabolite 14-OH-clarithromycin were not significantly affected by concomitant administration of fluconazole. No clarithromycin dose adjustment is necessary.

Ritonavir

A pharmacokinetic study demonstrated that the concomitant administration of ritonavir 200 m g every eight hours and clarithromycin 500 mg every 12 hours resulted in a marked inhibition of the metabolism of clarithromycin. The clarithromycin C_{max} increased by 31%, C_{min} increased 182% and AUC increased by 77% with concomitant administration of ritonavir. An essentially complete inhibition of the formation of 14-OH-clarithromycin was noted. Because of the large therapeutic window for clarithromycin, no dosage reduction should be necessary in patients with normal renal function. However, for patients with renal impairment, the following dosage adjustments should be considered: For patients with CL_{CR} 30 to 60 mL/min the dose of clarithromycin should be decreased by 50%. For patients with CL_{CR} <30 mL/min the dose of clarithromycin should be decreased by 75%. Doses of clarithromycin greater than 1 gm/day should not be coadministered with ritonavir.

Similar dos e a djustments s hould be c onsidered i n pa tients w ith r educed r enal f unction w hen ritonavir i s us ed as a p harmacokinetic e nhancer w ith ot her H IV pr otease i nhibitors i ncluding atazanavir and saquinavir (see section below, Bi-directional drug interactions).

Effect of Clarithromycin on Other Medicinal Products

CY3A4-based interactions

Co-administration of clarithromycin, known to inhibit CYP3A, and a drug primarily metabolized by CYP3A may be associated with elevations in drug concentrations that could increase or prolong both therapeutic and adverse effects of the concomitant drug. Clarithromycin should be used with caution in patients receiving treatment with other drugs known to be CYP3A enzyme substrates, especially if the CYP3A substrate has a n arrow safety margin (e.g., carbamazepine) and/or the substrate is extensively metabolized by this enzyme.

Dosage a djustments m ay be c onsidered, a nd when pos sible, s erum c oncentrations o f dr ugs primarily metabolized by CYP3A should be monitored closely in patients concurrently receiving clarithromycin.

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The following dr ugs or dr ug c lasses are k nown or s uspected to be m etabolized by the same CYP3A is ozyme: a lprazolam, astemizole, carbamazepine, cilostazol, cisapride, cyclosporine, disopyramide, ergot a lkaloids, lovastatin, m ethylprednisolone, m idazolam, om eprazole, or al anticoagulants (e.g. w arfarin), pi mozide, qui nidine, r ifabutin, s ildenafil, s imvastatin, s irolimus, tacrolimus, terfenadine, tr iazolam a nd vinblastine. Drugs int eracting by s imilar me chanisms through other isozymes within the cytochrome P450 system include phenytoin, theophylline and valproate.

Antiarrhythmics

There have been postmarketed reports of torsade de points occurring with the concurrent use of clarithromycin and quinidine or disopyramide. Electrocardiograms should be monitored for QT prolongation during c o-administration of c larithromycin with these d rugs. S erum le vels of quinidine and disopyramide should be monitored during clarithromycin therapy.

Omeprazole

Clarithromycin (500 mg every 8 hours) was given in combination with omeprazole (40 mg daily) to healthy adult subjects. The steady-state plasma concentrations of omeprazole were increased (C_{max} , AUC_{0-24} , and t $_{1/2}$ increased by 30%, 89%, and 34%, respectively), by the concomitant administration of clarithromycin. The mean 24-hour gastric pH value was 5.2 when omeprazole was administered alone and 5.7 when omeprazole was co-administered with clarithromycin.

Sildenafil, tadalafil and vardenafil

Each of these phosphodiesterase inhibitors is metabolized, at least in part, by CYP3A, and CYP3A may be inhibited by concomitantly administered clarithromycin. Co-administration of clarithromycin with sildenafil, tadalafil or vardenafil would likely result in increased phosphodiesterase inhibitor exposure. Reduction of sildenafil, tadalafil and vardenafil dos ages should be considered when these drugs are co-administered with clarithromycin.

Theophylline, carbamazepine

Results of clinical studies indicate that there was a modest but statistically significant ($p \le 0.05$) increase of circulating theophylline or carbamazepine levels when either of these drugs were administered concomitantly with clarithromycin. Dose reduction may need to be considered.

Tolterodine

The pr imary route of m etabolism f or tolterodine is vi a the 2D 6 i soform of c ytochrome P 450 (CYP2D6). However, in a subset of the population devoid of CYP2D6, the identified pathway of metabolism is via CYP3A. In this population subset, inhibition of CYP3A results in significantly higher serum concentrations of tolterodine. A reduction in tolterodine dosage may be necessary in the pr esence of C YP3A i nhibitors, s uch a s c larithromycin i n the C YP2D6 poor m etabolizer population.

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<u>Triazolobenzodiazepines (e.g., alprazolam, midazolam, triazolam)</u>

When midazolam w as co -administered with clarithromycin tablets (500 mg tw ice daily), midazolam AUC was increased 2.7-fold after intravenous administration of midazolam and 7-fold after oral administration. Concomitant administration of oral midazolam and clarithromycin should be avoided. If intravenous midazolam is co-administered with clarithromycin, the patient must be closely monitored to a llow dose a djustment. The same precautions should a lso a pply to other benzodiazepines that a remetabolized by C YP3A, including triazolam and a lprazolam. For benzodiazepines which a renot dependent on CYP3A for the ire limination (temazepam, nitrazepam, lorazepam), a clinically important interaction with clarithromycin is unlikely.

There have been post-marketing reports of drug interactions and central nervous system (CNS) effects (e.g., somnolence and confusion) with the concomitant use of clarithromycin and triazolam. Monitoring the patient for increased CNS pharmacological effects is suggested.

Other drug interactions

Colchicine

Colchicine i s a s ubstrate f or bot h C YP3A and t he e fflux t ransporter, P -glycoprotein (Pgp). Clarithromycin and other macrolides are known to inhibit CYP3A and Pgp. When clarithromycin and colchicine are administered together, inhibition of Pgp and/or CYP3A by clarithromycin may lead to increased exposure to colchicine. Patients should be monitored for clinical symptoms of colchicine toxicity (see section 4.4 Warnings and Precautions).

Digoxin

Digoxin i s t hought t o be a s ubstrate f or t he e fflux t ransporter, P-glycoprotein (Pgp). Clarithromycin i s kno wn t o i nhibit P gp. W hen c larithromycin and di goxin a re a dministered together, inhibition of Pgp by clarithromycin may lead to increased exposure to digoxin. Elevated digoxin serum concentrations in patients receiving clarithromycin and digoxin concomitantly have also been reported i n pos t m arketing s urveillance. S ome pa tients ha ve s hown c linical s igns consistent w ith digoxin toxicity, including potentially f atal arrhythmias. Serum di goxin concentrations s hould be c arefully m onitored w hile pa tients a re r eceiving di goxin a nd clarithromycin simultaneously.

Zidovudine

Simultaneous or all administration of clarithromycin tablets and zidovudine to HIV-infected adult patients may result in decreased steady-state zidovudine concentrations. Because clarithromycin appears to interfere with the absorption of simultaneously administered or alzidovudine, this interaction can be largely avoided by staggering the doses of clarithromycin and zidovudine to all for a 4-hour interval between each medication.

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This interaction does not appear to occur in paediatric HIV-infected patients taking clarithromycin suspension with zidovudine or dideoxyinosine. This interaction is unlikely when clarithromycin is administered via intravenous infusion.

Phenytoin and Valproate

There have been spontaneous or published reports of interactions of CYP3A inhibitors, including clarithromycin w ith dr ugs not t hought t o be m etabolized b y C YP3A (e.g. phe nytoin a nd valproate). S erum l evel de terminations a re r ecommended f or t hese dr ugs w hen a dministered concomitantly with clarithromycin. Increased serum levels have been reported.

Bi-directional drug interactions

Atazanavir

Both clarithromycin and atazanavir are substrates and inhibitors of CYP3A, and there is evidence of a bi-directional drug interaction. Co-administration of clarithromycin (500 mg twice daily) with atazanavir (400 mg once daily) resulted in a 2-fold increase in exposure to clarithromycin and a 70% de crease i n e xposure to 14 -OH-clarithromycin, w ith a 28% i ncrease i n t he A UC of atazanavir. B ecause of t he l arge t herapeutic w indow f or c larithromycin, no dos age r eduction should be ne cessary in patients w ith normal r enal function. For p atients w ith moderate renal function (creatinine clearance 30 to 60 m L/min), the dose of clarithromycin should be decreased by 50%. For patients with creatinine clearance <30 mL/min, the dose of clarithromycin should be decreased by 75 % us ing a n a ppropriate c larithromycin f ormulation. D oses of c larithromycin greater than 1000 mg per day should not be co-administered with protease inhibitors.

Itraconazole

Both clarithromycin and itraconazole are substrates and inhibitors of C YP3A, I eading to a bidirectional drug interaction. Clarithromycin may increase the plasma levels of itraconazole, while itraconazole may increase the plasma levels of clarithromycin. Patients taking itraconazole and clarithromycin concomitantly should be monitored closely for signs or symptoms of increased or prolonged pharmacologic effect.

Saquinavir

Both clarithromycin and saquinavir are substrates and inhibitors of CYP3A, and there is evidence of a bi-directional drug interaction. Concomitant administration of clarithromycin (500 mg twice daily) and saquinavir (soft gelatin capsules, 1200 mg three times daily) to 12 he althy volunteers resulted in steady-state AUC and C_{max} values of saquinavir which were 177% and 187% higher than those seen with saquinavir alone. Clarithromycin AUC and Cmax values were approximately 40% higher than those seen with clarithromycin alone. No dose adjustment is required when the two drugs are co-administered for a limited time at the doses/formulations studied. Observations from drug interaction

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studies using the soft gelatin capsule formulation may not be representative of the effects seen using the saquinavir hard gelatin capsule. Observations from drug interaction studies performed with saquinavir a lone may not be representative of the effects seen with saquinavir/ritonavir therpy. When saquinavir is co-administered with ritonavir, consideration should be given to the potential effects of ritonavir on clarithromycin.

Verapamil

Hypotension, br adyarrhythmias a nd l actic a cidosis ha ve be en obs erved i n pa tients t aking clarithromycin and verapamil concomitantly.

4.5 USE IN PREGNANCY AND LACTATION

The safety of clarithromycin for use in pregnancy and breast feeding of infants has not been established. Based on variable results obtained from studies in mice, rats, rabbits and monkeys, the possibility of adverse effects on embryofoetal development cannot be excluded. Therefore, use during pregnancy is not a dvised without carefully weighing the benefits a gainst risk. Clarithromycin is excreted into human breast milk.

4.6 EFFECTS ON ABILITY TO DRIVE AND USE MACHINES

There are no da ta on the effect of clarithromycin on the ability to drive or use machines. The potential for di zziness, vertigo, confusion and disorientation, which may occur with the medication, should be taken into account before patients drive or use machines.

4.7 UNDESIRABLE EFFECTS

a. Summary of the safety profile

The most frequent and common adverse reactions related to clarithromycin therapy for both adult and pa ediatric populations are a bdominal p ain, diarrhea, na usea, vomiting and taste perversion. These a dverse reactions are usually mild in intensity and are consistent with the known safety profile of macrolide antibiotics. (see section b of section 4.8)

There was no significant difference in the incidence of these gastrointestinal adverse reactions during clinical trials be tween the patient population with or without preexisting mycobacterial infections.

b. Tabulated summary of adverse reactions

The following table displays adverse reactions reported in clinical trials and from post-marketing experience with clarithromycin immediate-release tablets, granules for oral suspension, powder for solution for injection, extended-release tablets and modified-release tablets.

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The reactions considered at least possibly related to clarithromycin are displayed by system organ class and frequency using the following convention: very common ($\geq 1/10$), common ($\geq 1/100$) to < 1/10), uncommon ($\geq 1/1,000$ to < 1/100) and not known (adverse reactions from post-marketing experience; cannot be estimated from the available data). Within each frequency grouping, adverse reactions are presented in order of decreasing seriousness when the seriousness could be assessed

System Organ Class	Very common	Common	Uncommon	Not Known
	(≥1/10	≥ 1/100 to < 1/10	≥ 1/1,000 to < 1/100	(cannot be estimated from the available data)
Infections and infestations			Cellulitis ¹ , candidiasis, gastroenteritis ² ,infection ³ , vaginal infection	Pseudomembranous colitis, erysipelas, erythrasma
Blood and lymphatic system			Leukopenia, neutropenia ⁴ , thrombocythemia ³ , eosinophilia ⁴	Agranulocytosis, thrombocytopenia
Immune system disorders ⁵			Anaphylactoid reaction ¹ , hypersensitivity	Anaphylactic reaction
Metabolism and nutrition disorders			Anorexia, decreased appetite	Hypoglycaemia ⁶
Psychiatric disorders		Insomnia	Anxiety, nervousness ³ , screaming ³	Psychotic disorder, confusional state, depersonalisation, depression, disorientation, hallucination, abnormal dreams
Nervous system disorders		Dysgeusia, headache, taste perversion	Loss of consciousness ¹ , dyskinesia ¹ , dizziness, somnolence ⁷ , tremor	Convulsion, ageusia, parosmia, anosmia
Ear and labyrinth disorders			Vertigo, hearing impaired, tinnitus	Deafness
Cardiac disorders			Cardiac arrest ¹ , atrial fibrillation ¹ , electrocardiogram QT prolonged ⁸ , extrasystoles ¹ , palpitations	Torsade de pointes ⁸ , ventricular tachycardia ⁸

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Vascular disorders		Vasodilation ¹		Haemorrhage ⁹
Respiratory, thoracic and mediastinal disorder			Asthma ¹ , epistaxis ² , pulmonary embolism ¹	
Gastrointestinal disorders		Diarrhoea ¹⁰ , vomiting, dyspepsia, nausea, abdominal pain	Oesophagitis ¹ , gastrooesophageal reflux disease ² , gastritis, proctalgia ² , stomatitis, glossitis, abdominal distension ⁴ , constipation, dry mouth, eructation, flatulence,	Pancreatitis acute, tongue discolouration, tooth discoloration
Hepatobiliary disorders		Liver function test abnormal	Cholestasis ⁴ , hepatitis ⁴ , alanine aminotransferase increased, aspartate aminotransferase increased, gamma-glutamyltransferase increased ⁴	Hepatic failure ¹¹ , jaundice hepatocellular
Skin and subcutaneous tissue disorders		Rash, hyperhidrosis	Dermatitis bullous ¹ , pruritus, urticaria, rash maculo-papular ³	Stevens-Johnson syndrome ⁵ , toxic epidermal necrolysis ⁵ , drug rash with eosinophilia and systemic symptoms (DRESS), acne
Musculoskeletal and connective tissue disorders			Muscle spasms ³ , musculoskeletal stiffness ¹ , myalgia ²	Rhabdomyolysis ^{2,12} , myopathy
Renal and urinary disorders			Blood creatinine increased ¹ , blood urea increased ¹	Renal failure, nephritis interstitial
General disorders and administration site conditions	Injection site phlebitis ¹	Injection site pain ¹ , injection site inflammation ¹	Malaise ⁴ , pyrexia ³ , asthenia, chest pain ⁴ , chills ⁴ , fatigue ⁴	
Investigations			Albumin globulin ratio abnormal ¹ , blood alkaline phosphatase increased ⁴ , blood lactate dehydrogenase increased ⁴	International normalised ratio increased ⁹ , prothrombin time prolonged ⁹ , urine color abnormal

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¹ ADRs reported only for the Powder for Solution for Injection formulation

c. Description of selected adverse reactions

Injection site phl ebitis, inj ection site pa in, v essel punc ture s ite pa in, and injection site inflammation are specific to the clarithromycin intravenous formulation.

In very rare instances, hepatic failure with fatal outcome has been reported and generally has been associated with serious underlying diseases and/or concomitant medications (see section 4.4).

A s pecial attention to di arrhoea s hould be pa id a s *Clostridium difficile*-associated di arrhoea (CDAD) has been reported with use of nearly all antibacterial agents including clarithromycin, and may range in severity from mild diarrhoea to fatal colitis. (see section 4.4)

In the event of s evere acute hypersensitivity r eactions, s uch a s a naphylaxis, S tevens-Johnson Syndrome and t oxic e pidermal ne crolysis, c larithromycin t herapy s hould be di scontinued immediately and appropriate treatment should be urgently initiated (see section 4.4).

As with other macrolides, QT prolongation, ventricular tachycardia, and *torsade de pointes* have rarely been reported with clarithromycin (see section 4.4 and 4.5).

Pseudomembranous c olitis has be en reported with nearly all antibacterial agents, including clarithromycin, and may range in severity from mild to life threatening. Therefore, it is important to consider this diagnosis in patients who present with diarrhoea subsequent to the administration of antibacterial agents (see section 4.4).

In some of the reports of rhabdomyolysis, clarithromycin was a dministered concomitantly with statins, fibrates, colchicine or allopurinol (see section 4.3 and 4.4).

There have be en post-marketing reports of colchicine tox icity with concomitant use of clarithromycin and colchicine, especially in elderly and/or patients with renal insufficiency, some with a fatal outcome (see sections 4.4 and 4.5).

There h ave b een rare r eports of h ypoglycemia, some of w hich have o ccurred in patients on concomitant oral hypoglycemic agents or insulin (see section 4.4 and 4.5).

There have been post-marketing reports of drug interactions and central nervous system (CNS) effects (e.g. somnolence and confusion) with the concomitant use of clarithromycin and triazolam. Monitoring the patient for increased CNS pharmacological effects is suggested (see section 4.5).

There is a risk of serious haemorrhage and significant elevations in INR and prothrombin time when c larithromycin is co-administered with warfarin. INR and prothrombin times should be frequently

²ADRs reported only for the Extended-Release Tablets formulation

³ ADRs reported only for the Granules for Oral Suspension formulation

⁴ ADRs reported only for the Immediate-Release Tablets formulation ^{5,8,10,11,12}See section a)

^{6,7,9}See section c)

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monitored while patients are receiving clarithromycin and or all anticoagulants concurrently (see section 4.4 and 4.5).

There have been rare reports of clarithromycin ER tablets in the stool, many of which have occurred in patients with a natomic (including i leostomy or colostomy) or functional gastrointestinal disorders with shortened GI transit times. In several reports, tablet residues have occurred in the context of diarrhoea. It is recommended that patients who experience tablet residue in the stool and no improvement in their condition should be switched to a different clarithromycin formulation (e.g. suspension) or another antibiotic.

Special population: Adverse Reactions in Immunocompromised Patients (see section e)

d. Paediatric populations

Clinical t rials ha ve be en c onducted us ing c larithromycin p aediatric s uspension i n c hildren 6 months to 12 years of age. Therefore, children under 12 years of age should use clarithromycin paediatric suspension.

Frequency, type and severity of adverse reactions in children are expected to be the same as in adults.

e. Other special populations

Immunocompromised patients

In A IDS and other immunocomprimised patients treated with the higher doses of clarithromycin over long periods of time for mycobacterial infections, it was often difficult to distinguish adverse events possibly associated with clarithromycin administration from underlying signs of Human Immunodeficiency Virus (HIV) disease or intercurrent illness.

In adult patients, the most frequently reported adverse reactions by patients treated with total daily doses of 1000 m g a nd 2000m g of c larithromycin w ere: n ausea, vo miting, t aste pe rversion, abdominal pa in, di arrhoea, rash, flatulence, he adache, c onstipation, he aring di sturbance, S erum Glutamic Oxaloacetic Transaminase (SGOT) and Serum Glutamic Pyruvate Transaminase (SGTP) elevations. A dditional low-frequency events i ncluded d yspnoea, i nsomnia a nd dr y m outh. T he incidences w ere c omparable f or p atients t reated w ith 1000m g a nd 2000 mg, but w ere generally about 3 t o 4 t imes a s frequent f or those pa tients w ho r eceived t otal daily dos es of 4000m g of clarithromycin.

In these immunocompromised patients evaluations of laboratory values were made by analysing those values out side the seriously a bnormal level (i.e. the extreme high or low limit) for the specified test. On the basis of these criteria, about 2% to 3% of those patients who received 1000mg or 2000mg of clarithromycin daily had seriously abnormal elevated levels of SGOT and SGPT, and abnormally low white blood cell and platelet counts. A lower percentage of patients in these two dos age groups also had elevated Blood U rea N itrogen (BUN) levels. S lightly higher incidences of abnormal values were not ed for patients who received 4000mg daily for a ll parameters except White Blood Cell.

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5. PHARMACOLOGICAL PROPERTIES

5. 1 Pharmacodynamic Properties

Clarithromycin is a semi-synthetic derivative of erythromycin A. It exerts its antibacterial action by binding to the 50s ribosomal sub-unit of susceptible bacteria and suppresses protein synthesis. It is highly pot ent a gainst a wide variety of a erobic and anaerobic gram-positive and gram-negative or ganisms. The minimum inhibitory concentrations (MICs) of clarithromycin are generally two-fold lower than the MICs of erythromycin.

The 14-hydroxy metabolite of clarithromycin also has antimicrobial activity. The MICs of this metabolite are equal or two-fold higher than the MICs of the parent compound, except for H. influenzae where the 14-hydroxy metabolite is two-fold more active than the parent compound.

Klacid is usually active against the following organisms in vitro. Please see below for table of MIC breakpoints.

Gram-positive Bacteria: Staphylococcus aureus (methicillin susceptible); Streptococcus pyogenes (Group A be ta-hemolytic s treptococci); al pha-hemolytic s treptococci (viridans group); Streptococcus (Diplococcus) pneumoniae; Streptococcus agalactiae; Listeria monocytogenes.

Gram-negative Bacteria: H aemophilus inf luenzae; H aemophilus pa rainfluenzae; M oraxella (Branhamella) catarrhalis; Neisseria gonorrhoeae; Legionella pneumophila; Bordetella pertussis; Helicobacter pylori; Campylobacter jejuni.

Mycoplasma: Mycoplasma pneumoniae; Ureaplasma urealyticum.

Other Organisms: Chlamydia trachomatis; Mycobacterium avium; Mycobacterium leprae.

Anaerobes: M acrolide-susceptible Bacteroides fragilis; C lostridium pe rfringens; P eptococcus species; Peptostreptococcus species; Propionibacterium acnes.

Clarithromycin has bactericidal activity against several bacterial strains. The organisms include Haemophilus i nfluenzae, S treptococcus pne umoniae, S treptococcus p yogenes, S treptococcus agalactiae, Moraxella (Branhamella) cat arrhalis, Neisseria gonorrhoeae, H. p ylori and Campylobacter spp.

H. pylori is associated with acid peptic disease including duodenal ulcer and gastric ulcer in which about 95% and 80% of pa tients r espectively are infected with the agent. H. pylori is also implicated as a major contribution factor in the development of gastric and ulcer recurrence in such patients.

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Clarithromycin has been used in small numbers of patients in other treatment regimens. Possible kinetic interactions have not been fully investigated. These regimens include:

Clarithromycin pl us t inidazole a nd om eprazole; c larithromycin plus te tracycline, bismuth subsalicylate and ranitidine; clarithromycin plus ranitidine alone.

Clinical studies using various different H. pylori eradication regimens have shown that eradication of H. pylori prevents ulcer recurrence.

Breakpoints

The following breakpoints for clarithromycin, separating susceptible organisms from resistant organisms, have been established by the European Committee for Antimicrobial Susceptibility Testing (EUCAST).

Breakpoints (MIC, µg/ml)				
Microorganism	Susceptible (≤)	Resistant (>)		
Streptococcus spp.	0.25 μg/ml	0.5 μg/ml		
Staphylococcus spp.	1 μg/ml	2 μg/ml		
Haemophilus spp.	1 μg/ml	32 μg/ml		
Moraxella catarrhalis	0.25 μg/ml	0.5 μg/ml		

Clarithromycin is used for the eradication of *H. pylori*; minimum inhibitory concentration (MIC) $\leq 0.25 \,\mu\text{g/ml}$ which has been established as the susceptible breakpoint by the Clinical and Laboratory Standards Institute (CLSI).

5. 2 Pharmacokinetic Properties

H. p ylori is a ssociated with a cid pe ptic di sease i ncluding duode nal ulcer and gastric ul cer i n which about 95% and 80% of patients respectively are infected with the agent. H. pylori is also implicated as a major contribution factor in the development of gastric and ulcer recurrence in such patients.

Clarithromycin has been used in small numbers of patients in other treatment regimens. Possible kinetic interactions have not been fully investigated. These regimens include:

Clarithromycin plus tini dazole a nd omeprazole; c larithromycin plus tetracycline, bismuth subsalicylate and ranitidine; clarithromycin plus ranitidine alone.

Clinical s tudies us ing various di fferent H . p ylori e radication r egimens ha ve s hown t hat eradication of H.pylori prevents ulcer recurrence.

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Clarithromycin i s r apidly and w ell a bsorbed f rom t he g astro-intestinal tr act a fter or al administration of K lacid tablets. The microbiologically active me tabolite 14 hydroxyclarithromycin is formed by first pass metabolism. Klacid may be given without regard to meals as food does not affect the extent of bioavailability of Klacid tablets. Food does slightly delay the onset of absorption of clarithromycin and formation of the 14-hydroxymetabolite. The pharmacokinetics of clarithromycin are non linear; however, steady-state is attained within 2 days of dosing. At 250 mg b.i.d. 15-20% of unchanged drug is excreted in the urine. With 500 mg b.i.d. da ily dos ing ur inary excretion i s g reater (approximately 36%). T he 14 hydroxyclarithromycin is the major ur inary metabolite and a counts for 10-15% of the dose. Most of the remainder of the dose is eliminated in the faeces, primarily via the bile. 5-10% of the parent drug is recovered from the faeces.

When clarithromycin 500 mg is given three time s d aily, the c larithromycin pl asma concentrations are increased with respect to the 500 mg twice daily dosage.

Klacid provides t issue concentrations that are several times higher than the circulating drug levels. Increased levels have been found in both tonsillar and lung tissue. Clarithromycin is 80% bound to plasma proteins at therapeutic levels.

Klacid also penetrates the gastric mucus. Levels of clarithromycin in gastric mucus and gastric tissue a re hi gher w hen c larithromycin i s c o-administered w ith om eprazole t han w hen clarithromycin is administered alone.

5. 3 Preclincal safety data

In acute mouse and rat studies, the median lethal dose was greater than the highest feasible dose for administration (5g/kg).

In repeated dose studies, toxicity was related to dose, duration of treatment and species. Dogs were more sensitive than primates or rats. The major clinical signs at toxic doses included emesis, weakness, reduced food consumption and weight gain, salivation, dehydration and hyperactivity.

In a ll s pecies the live r was the p rimary ta rget organ at tox ic dos es. Hepatotoxicity was detectable by e arly elevations of liver function tests. Discontinuation of the drug generally resulted in a r eturn to or t oward nor mal r esults. O ther t issues less commonly a ffected included the stomach, thymus and other lymphoid tissues and the kidneys. At near therapeutic doses, c onjunctival i njection and l acrimation oc curred only in dogs. At a massive dose of 400 mg/kg/day, some dogs and monkeys developed corneal opacities and/or oedema.

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Fertility and reproduction s tudies in r ats ha ve s hown no adverse e ffects. T eratogenicity studies in rats (Wistar (p.o.) and Spraque-Dawley (p.o. and i.v.)), New Zealand White rabbits and c ynomolgous m onkeys failed to demonstrate a ny te ratogenicity from c larithromycin. However, a further similar study in Sprague-Dawley rats indicated a low (6%) incidence of cardiovascular abnormalities which appeared to be due to spontaneous expression of genetic changes. Two mouses tudies r evealed a variable incidence (3-30%) of cleft p alate and embryonic loss was seen in monkeys but only at dose levels which were clearly toxic to the mothers.

6. PHARMACEUTICAL PARTICULARS

6. 1 List of excipient (s)

Sodium Starch Glycolate

Calcium Phosphate Anhydrous

Maize starch

Hydroxy propyl Cellulose (Klucel EF)

Sodium Starch Glycolate

Purified talc

Colloidal Anhydrous Silica (Aerosil 200)

Magnesium Stearate

Opadry yellow OY 8487

White Bees wax (Pellets)

Purified water

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6. 2 Incompatibilities

Not applicable

6. 3 Shelf life

3 years

6. 4 Special precautions for storage

Store below 30°C. Protect from light and moisture.

6.5 NATURE AND CONTENTS OF CONTAINER

Spectromax 250 is packed as 7 Tablets/blister of PVC film with printed laminated Aluminum backing foil.

6. MARKETING AUTHORISATION HOLDER

National Pharmaceutical Industries Co. (SAOG)

Road No. 15,

P.O Box 120

Postal Code 124

Rusayl, Sultanate of Oman