1 PRODUCT NAME

Modafinil 100 mg tablets

2 QUALITATIVE AND QUANTITATIVE COMPOSITION

Each tablet contains 100 mg of modafinil.
Excipients with known effect: Lactose
For the full list of excipients, see section 6.1.

3 PHARMACEUTICAL FORM

Tablet
A white to off white capsule shape tablet embossed with ‘100’.

4 CLINICAL PARTICULARS

4.1 Therapeutic indications

Modafinil is indicated:
• to improve wakefulness in patients with excessive daytime sleepiness associated with narcolepsy;
• to treat excessive sleepiness associated with moderate to severe chronic shift work sleep disorder where nonpharmacological interventions are unsuccessful or inappropriate;
• as an adjunct to continuous positive airways pressure (CPAP) in obstructive sleep apnoea/hypopnoea syndrome in order to improve wakefulness.

4.2 Dose and method of administration

Modafinil should be used only in patients who have had a complete evaluation of their excessive sleepiness, and in whom a diagnosis of either narcolepsy, OSAHS, and/or SWSD has been made in accordance with ICSD or DSM diagnostic criteria. Such an evaluation usually consists of a complete history and physical examination, and testing in a laboratory setting. Some patients may have more than one sleep disorder contributing to their excessive sleepiness (e.g., OSAHS and SWSD coincident in the same patient).

Treatment with modafinil should be initiated and supervised by physicians with appropriate experience in the treatment of sleep disorders who have access to sleep laboratory diagnostic facilities.

Narcolepsy

The dose of modafinil is 200 to 400 mg/day, given as a single dose in the morning, or as two divided doses, in the morning and at noon. Tablets should be swallowed whole.

Doses of 400 mg/day have been well tolerated, but there is no statistically significant evidence that this dose confers additional benefit beyond that of the 200 mg dose.

For patients who require more than 200 mg/day, the dose should be increased, to a maximum of 400 mg/day, in increments of 100 mg as needed and tolerated.
Obstructive Sleep Apnoea/Hypopnoea Syndrome

The dose of modafinil is 200 to 400 mg/day, given as a single dose in the morning, or as two divided doses, in the morning and at noon. Tablets should be swallowed whole.

Doses of 400 mg/day have been well tolerated, but there is no statistically significant evidence that this dose confers additional benefit beyond that of the 200 mg dose.

For patients who require more than 200 mg/day, the dose should be increased, to a maximum of 400 mg/day, in increments of 100 mg as needed and tolerated.

For patients with obstructive sleep apnoea/hypopnoea syndrome, modafinil treats the symptoms of excessive daytime sleepiness associated with the condition. In addition to this symptomatic treatment, disease-modifying interventions (e.g., Continuous Positive Airway Pressure) should be commenced or continued.

Moderate to Severe Chronic Shift Work Sleep Disorder

The recommended daily dose is 200 mg. modafinil should be taken as a single dose approximately 1 hour prior to the start of the work shift. Tablets should be swallowed whole.

Dosing in Special Populations

In patients with severe hepatic impairment, the dose of modafinil should be reduced to one-half of that recommended for patients with normal hepatic function (see section 4.4 “Special warnings and precautions for use”).

There is inadequate information to determine safety and efficacy of modafinil dosing in patients with severe renal impairment (see section 4.4 “Special warnings and precautions for use”).

In elderly patients, elimination of modafinil and its metabolites may be reduced as a consequence of aging. Therefore, consideration should be given to the use of lower doses in this population (see section 4.4 “Special warnings and precautions for use”).

Paediatric population

Modafinil should not be used in children aged less than 18 years old because of safety and efficacy concerns (see section 4.4).

4.3 Contraindications

- Hypersensitivity to modafinil or any other component of the product.
- Use in pregnancy.

4.4 Special warnings and precautions for use

Serious Rash, including Stevens-Johnson Syndrome

Serious Skin Rash, including Stevens-Johnson Syndrome

Serious rash requiring hospitalization and discontinuation of treatment has been reported in adults and children in association with the use of modafinil.

Modafinil is not approved for use in pediatric patients for any indication.

In clinical trials of modafinil, the incidence of rash resulting in discontinuation was approximately 0.8% (13 per 1,585) in pediatric patients (age <17 years); these rashes included 1 case of possible Stevens-Johnson Syndrome (SJS) and 1 case of apparent multiorgan hypersensitivity reaction.
Several of the cases were associated with fever and other abnormalities (e.g., vomiting, leukopenia). The median time to rash that resulted in discontinuation was 13 days. No such cases were observed among 380 pediatric patients who received placebo. No serious skin rashes have been reported in adult clinical trials (0 per 4,264) of modafinil.

Rare cases of serious or life-threatening rash, including SJS, Toxic Epidermal Necrolysis (TEN) and Drug Rash with Eosinophilia and Systemic Symptoms (DRESS) have been reported in adults and children in world-wide post-marketing experience (see section 4.8 Undesirable effects “Post Marketing Experience”). The reporting rate of TEN and SJS associated with modafinil use which is generally accepted to be an underestimate due to underreporting, exceeds the background incidence rate. Estimates of the background incidence rate for these serious skin reactions in the general population range between 1 to 2 cases per million-person years.

While little is known about factors that can predict the risk of occurrence or the severity of rash associated with modafinil, the risk may increase with higher doses. Nearly all cases of serious rash associated with modafinil occurred within 1 to 5 weeks after treatment initiation. Isolated cases have been reported after prolonged treatment (e.g. 3 months). Duration of therapy cannot be relied upon as a means to predict the potential risk heralded by the first appearance of a rash.

Although benign rashes also occur with modafinil, it is not possible to reliably predict which rashes will prove to be serious. Accordingly, modafinil should ordinarily be discontinued at the first sign of rash, unless the rash is clearly not drug-related. Discontinuation of treatment may not prevent a rash from becoming life-threatening or permanently disabling or disfiguring.

**Multi-organ Hypersensitivity Reactions**

Multi-organ hypersensitivity reactions have occurred in close temporal association to the initiation of modafinil. Although there have only been a limited number of reports, multi-organ hypersensitivity reactions may result in hospitalization or be life-threatening. There are no factors that are known to predict the risk of occurrence or the severity of multi-organ hypersensitivity reactions associated with modafinil. Signs and symptoms of these reactions were diverse; however, patients typically, although not exclusively, presented with fever and rash associated with other organ system involvement. Other associated manifestations included myocarditis, hepatitis, liver function test abnormalities, haematological abnormalities (e.g., eosinophilia, leukopenia, thrombocytopenia), pruritus, and asthenia. Because multi-organ hypersensitivity is variable in its expression, other organ system symptoms and signs, not noted here, may occur.

If a multi-organ hypersensitivity reaction is suspected, modafinil should be discontinued and not restarted. Although there are no case reports to indicate cross-sensitivity with other medicines that produce this syndrome, the experience with medicines associated with multi-organ hypersensitivity would indicate this to be a possibility.

**Psychiatric Symptoms and Disorders**

Psychiatric adverse experiences have been reported in patients treated with modafinil in clinical trials and from post-marketing experience. Patients should be monitored for the development of de novo psychiatric disorders or exacerbation of pre-existing psychiatric disorders at every adjustment of dose and regularly during treatment. If psychiatric symptoms develop in association with modafinil treatment, discontinuation of modafinil may be required. Caution should be exercised in giving modafinil to patients with a history of psychiatric disorders including psychosis, depression, mania, major anxiety, agitation, insomnia or substance abuse.
Aggressive or hostile behaviour

The onset or worsening of aggressive or hostile behaviour has been reported in patients treated with modafinil. Patients treated with modafinil should be carefully monitored for the appearance or worsening of aggressive or hostile behaviour. If symptoms occur, modafinil should be discontinued.

Suicidal ideation and suicide-related behaviour

Suicidal ideation and suicide-related behaviour (including suicide attempts) have been reported in patients treated with modafinil. Patients treated with modafinil should be carefully monitored for the appearance or worsening of suicidal thinking and/or suicide-related behaviour. If suicide-related symptoms develop in association with modafinil, treatment should be discontinued.

Psychotic or manic symptoms

The onset or worsening of psychotic symptoms or manic symptoms (including hallucinations, delusions, agitation or mania) has been reported in patients treated with modafinil. Patients treated with modafinil should be carefully monitored for the appearance or worsening of psychotic or manic symptoms. If psychotic or manic symptoms occur, modafinil should be discontinued.

Bipolar disorders

Care should be taken in using modafinil in patients with co-morbid bipolar disorder because of concern for possible precipitation of a mixed/manic episode in such patients.

Depression

The onset of depression or the aggravation of underlying depressive disorder has been reported in patients treated with modafinil. Patients treated with modafinil should be carefully monitored for the appearance of or worsening of depression.

Anxiety

The onset or worsening of anxiety has been reported in patients treated with modafinil. Anxiety and nervousness are adverse events that appear to be closely dose related.

General

Although modafinil has not been shown to produce functional impairment, any medicine affecting the CNS may alter judgment, thinking or motor skills. Patients with major anxiety should only receive treatment with modafinil in a specialist unit.

If a hypersensitivity reaction is suspected, modafinil treatment should be discontinued.

In patients with obstructive sleep apnoea/hypopnoea syndrome, the underlying condition and any associated cardiovascular pathology should be monitored.

Patients should be advised that modafinil is not a replacement for sleep and good sleep hygiene should be maintained.

Cardiovascular System

In hypertensive patients, blood pressure should be adequately controlled before initiating treatment with modafinil and monitored regularly during treatment. Blood pressure, heart rate and general cardiovascular status should be monitored in all patients during treatment with modafinil.

In clinical studies of modafinil, signs and symptoms including chest pain, palpitations, dyspnoea and transient ischemic T-wave changes on ECG were observed in three subjects in association with mitral valve prolapse or left ventricular hypertrophy. It is recommended that modafinil not be used in
patients with a history of left ventricular hypertrophy or ischaemic ECG changes, chest pain, arrhythmia or other clinically significant manifestations of mitral valve prolapse in association with CNS stimulant use.

The safety of modafinil has not been established in patients with coronary artery disease, a recent history of myocardial infarction or unstable angina. Patients with these conditions were not included in the controlled clinical trials. Post marketing adverse events of ischaemic heart disease have been reported in patients with and without a history of cardiovascular disease while being treated with modafinil. The risks of using modafinil in patients with coronary artery disease, a recent history of myocardial infarction or unstable angina should be carefully weighed against the potential therapeutic benefit. It is recommended that such patients receive further specialist evaluation before modafinil treatment is considered.

Postmarketing adverse events of cardiac arrhythmia, such as atrial fibrillation and premature ventricular contractions, have been reported in patients treated with modafinil. In some of these cases there was a close temporal association to the use of modafinil, a resolution of the arrhythmia upon medicine discontinuation and, in a few cases, a recurrence of arrhythmia after modafinil rechallenge. It is recommended that patients have an ECG before modafinil is initiated. Patients with abnormal findings should receive further specialist evaluation before modafinil treatment is considered.

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**Dose Dependency and adverse effects**

The development of skin and hypersensitivity reactions, central nervous system, psychiatric and cardiovascular system adverse reactions appear to be related to higher doses of modafinil. Cardiovascular and central nervous system adverse reactions increase significantly after a total daily dose of more than 400 mg. Always start at the lowest recommended dose (see section 4.2 “Dose and method of administration”).

**Patients (Women) Using Contraception**

Sexually active women of child-bearing potential should be established on a contraceptive program before taking modafinil. The effectiveness of oral contraceptives may be impaired due to enzyme induction activity of modafinil. Alternative or concomitant methods of contraception are recommended for patients treated with modafinil, and for one month after discontinuation of treatment (see section 4.5 “Interaction with other medicines and other forms of interaction”).

**Carcinogenicity and Mutagenicity**

Carcinogenicity studies were conducted in which modafinil was administered in the diet to mice for 78 weeks and to rats for 104 weeks at doses up to 60 mg/kg/day. The highest dose studied in these studies would have achieved systemic exposure levels less than human exposure at the maximum recommended dose. There was no evidence of tumourigenesis associated with modafinil administration in these studies; however, the carcinogenic potential of modafinil has not been fully evaluated.

There was no consistent evidence for genotoxic activity of modafinil in *in vitro* assays of gene mutation (reverse mutation in *S. typhimurium* and *E. coli*, forward mutation in Chinese hamster V79 fibroblasts) or in the chromosomal damage assay (human lymphocytes *in vitro*, Chinese hamster bone marrow cells *in vivo*, mouse micronucleus assay). Modafinil did not increase unscheduled DNA synthesis in rat hepatocytes. In a cell transformation assay in BALB/3T3 mouse embryo cells,
modafinil did not cause an increase in the frequency of transformed foci in the presence or absence of metabolic activation.

**Abuse and Dependence Potential**

In addition to its wakefulness-promoting effect and increased locomotor activity in animals, in humans, modafinil may produce psychoactive and euphoric effects, alterations in mood, perception, thinking and feelings. In in vitro binding studies, modafinil binds with low affinity to the dopamine reuptake site and causes an increase in extracellular dopamine, but no increase in dopamine release. Modafinil is reinforcing, as evidenced by its self-administration in monkeys previously trained to self-administer cocaine. In some studies, modafinil was also partially discriminated as stimulant-like. Caution should be exercised in administering modafinil to patients with history of alcohol, medicine or illicit substance abuse. Patients with such history should be monitored for signs of misuse or abuse (e.g. increasing the recommended dosage).

**Withdrawal:** In one US Phase 3 clinical trial of nine weeks of modafinil use, the effects of modafinil cessation were monitored for 14 days. No specific symptoms of withdrawal were observed during the 14 days; however, sleepiness returned in patients with narcolepsy.

**Use in Children and Adolescents (<18 years of age)**

The efficacy and safety of modafinil in this age group has not been established. modafinil is not approved for use in paediatric patients for any indication. The use of modafinil in this age group is not recommended. Neuropsychiatric and serious skin reactions have been reported in children and adolescents treated with modafinil.

**Special Populations**

**Use in the Elderly:** There are no satisfactory data on the safety and efficacy of modafinil in patients ≥ 65 years of age. The clearance of modafinil may be reduced in the elderly (see section 4.2 “Dose and method of administration”).

**Renal Impairment:** In a single-dose 200 mg modafinil study, severe chronic renal failure (creatinine clearance ≤ 20 mL/min) did not significantly influence the pharmacokinetics of modafinil, but exposure to modafinil acid (an inactive metabolite) was increased 9-fold (see section 4.2 “Dose and method of administration”).

**Hepatic Impairment:** The dose of modafinil should be reduced by half in patients with severe hepatic impairment (see section 4.2 “Dose and method of administration”).

4.5 **Interaction with other medicines and other forms of interaction**

**CNS Active Drugs**

Methylphenidate – The absorption of modafinil may be delayed by approximately one hour when co-administered with methylphenidate.

Clomipramine – The coadministration of a single dose of clomipramine (50 mg) on the first three days of treatment with modafinil (200 mg/day) in healthy volunteers did not show an effect on the pharmacokinetics of either medicine. However, one incident of increased levels of clomipramine and its active metabolite desmethylclomipramine has been reported in a CYP2D6 poor metabolizer with narcolepsy during treatment with modafinil. (See below “Potential Interactions with Drugs That Inhibit or are Metabolised by Cytochrome P-450 Isoenzymes and Other Hepatic Enzymes”).
Triazolam – In healthy, female volunteers, who were receiving long-term treatment with ethinyl estradiol, the co-administration of two single doses of 0.125 mg of triazolam (one administered before and the other at the end of treatment) with modafinil (200 mg for seven days, followed by 400 mg for 21 days) indicated that, for triazolam, the $C_{\text{max}}$ and $AUC_{0-\infty}$ were reduced by 59% and 42% respectively, and the elimination rate was increased by approximately 50%. Therefore, dosage adjustment of triazolam may be necessary when co-administered with modafinil.

Monoamine Oxidase (MAO) Inhibitors – Interaction studies with monoamine oxidase inhibitors have not been performed. Therefore, caution should be used when concomitantly administering MAO inhibitors and modafinil.

**Potential Interactions with Drugs That Inhibit, Induce, or are Metabolised by Cytochrome P-450 Isoenzymes and Other Hepatic Enzymes**

Diazepam, Phenytoin, Propranolol, Tricyclic Antidepressants, Selective Serotonin Reuptake Inhibitors – Because modafinil is a reversible inhibitor of the drug-metabolising enzyme CYP2C19, co-administration of modafinil with medicines such as diazepam, phenytoin, and propranolol, which are largely eliminated via that pathway, may increase the circulating levels of those compounds. In addition, in individuals deficient in the enzyme CYP2D6, the levels of CYP2D6 substrates such as tricyclic antidepressants and selective serotonin reuptake inhibitors, which have ancillary routes of elimination through CYP2C19, may be increased by co-administration of modafinil. Dose adjustments may be necessary for patients being treated with these and similar medications.

Steroidal Contraceptives, Cyclosporin, Theophylline – Chronic administration of modafinil also causes modest induction of the metabolising enzyme CYP3A4, thus reducing the levels of co-administered substrates for that enzyme system, such as steroidal contraceptives, cyclosporin and to a lesser degree, theophylline. Dose adjustments may be necessary for patients being treated with these and similar medications.

Inducers or Inhibitors of CYP3A4 – Co-administration of potent inducers of CYP3A4 (e.g., carbamazepine, phenobarbital, rifampicin) or inhibitors of CYP3A4 (e.g., ketoconazole, itraconazole) could alter the levels of modafinil due to the partial involvement of that enzyme in the metabolic elimination of the compound (see section 4.4 Special warnings and precautions for use: “Patients (Women) Using Contraception”).

Warfarin, Phenytoin – The exposure of human hepatocytes to modafinil in vitro produced an apparent concentration-related suppression of expression of CYP2C9 activity. The clinical relevance of this finding is unclear, since no other indication of CYP2C9 suppression has been observed. However, monitoring of prothrombin times is suggested as a precaution for the first several months of co-administration of modafinil and warfarin, a CYP2C9 substrate, and thereafter whenever modafinil dosing is changed. In addition, patients receiving modafinil and phenytoin, a CYP2C9 substrate, concomitantly should be monitored for signs of phenytoin toxicity.

It should be noted that evaluation of medicine interactions based on in vitro systems might not necessarily reflect those seen in vivo situations. This information should be used as a guide to assess the risks associated with the use of concomitant medications.

### 4.6 Fertility, pregnancy and lactation

**Impairment of Fertility**

No effects on fertility were observed in male or female rats treated with modafinil prior and throughout mating and gestation at oral doses up to 100 mg/kg/day (the highest dose investigated would have achieved systemic exposure levels less than human exposure at the maximum
recommended dose). However, sufficiently high enough doses or large enough sample sizes to adequately assess effects on fertility were not used in the study.

Use in Pregnancy (Category B3)
Animal studies to assess the effects of modafinil on reproduction and the developing foetus were not conducted at adequately high doses or according to guidelines which would have been able to provide a comprehensive evaluation of the potential of modafinil to adversely affect fertility, or cause embryolethality or teratogenicity.

Embryotoxicity, in the absence of maternal toxicity, was observed in rats receiving oral modafinil throughout the period of organogenesis. At a dose of 200 mg/kg/day (less than human exposure at the maximum recommended daily clinical dose of 400 mg), there was an increase in resorption, hydronephrosis and skeletal variations. The no effect dose for these effects was 100 mg/kg/day. Embryotoxicity was not observed in rabbits receiving oral modafinil throughout organogenesis at doses up to 100 mg/kg/day (0.6 times the human exposure at the maximum recommended daily dose of 400 mg, based on AUC). However, neither of these studies used optimal doses for the evaluation of embryotoxicity. Although a threshold dose for embryotoxicity has been identified, the full spectrum of potential toxic effects on the foetus has not been characterised. Modafinil was embryotoxic in rats dosed during late gestation and lactation, or prior to and throughout mating and gestation, at oral doses ≥ 50 mg/kg/day; the no effect dose was 20 mg/kg/day (less than human exposure at the maximum recommended clinical daily dose of 400 mg).

As there are no adequate and well-controlled trials with modafinil in pregnant women, it should be contraindicated during pregnancy.

Patients should be cautioned regarding the potential increased risk of pregnancy when using steroidal contraceptives with modafinil and for one month after discontinuation of therapy (see section “4.5 Interaction with other medicines and other forms of interaction”).

Use in Lactation
No developmental toxicity was noted postnatally in the offspring of rats given oral modafinil up to 100 mg/kg/day during late gestation and throughout lactation. The highest dose studied in these studies would have achieved systemic exposure levels less than human exposure at the maximum recommended dose.

Modafinil and/or its metabolites have been found in the milk of lactating rats. It is not known whether modafinil or its metabolites are excreted in human milk. Therefore, breastfeeding is not recommended during administration of modafinil.

4.7 Effects on ability to drive and use machines
Patients should be cautioned about operating an automobile or other hazardous machinery until they are reasonably certain that modafinil therapy will not adversely affect their ability to engage in such activities.

4.8 Undesirable effects
Modafinil has been evaluated for safety in over 3500 patients, of whom more than 2000 patients with excessive sleepiness associated with primary disorders of sleep and wakefulness were given at least one dose of modafinil. In clinical trials, modafinil has been found to be generally well tolerated and most adverse experiences were mild to moderate.
The most commonly observed adverse events (≥5%) associated with the use of modafinil more frequently than placebo-treated patients in the placebo-controlled clinical studies in primary disorders of sleep and wakefulness were headache, nausea, nervousness, rhinitis, diarrhoea, back pain, anxiety, insomnia, dizziness, and dyspepsia. The adverse event profile was similar across these studies.

In the placebo-controlled clinical trials, 74 of the 934 patients (8%) who received modafinil discontinued due to an adverse experience compared to 3% of patients that received placebo. The most frequent reasons for discontinuation that occurred at a higher rate for modafinil than placebo patients were headache (2%), nausea, anxiety, dizziness, insomnia, chest pain and nervousness (each <1%). In a Canadian clinical trial, a 35 year old obese narcoleptic male with a prior history of syncopal episodes experienced a 9-second episode of asystole after 27 days of modafinil treatment (300 mg/day in divided doses).

**Incidence in Controlled Trials**

The following table (Table 1) presents the adverse experiences that occurred at a rate of 1% or more and were more frequent in patients treated with modafinil than in placebo patients in the principal, placebo-controlled clinical trials.

The prescriber should be aware that the figures provided below cannot be used to predict the frequency of adverse experiences in the course of usual medical practice, where patient characteristics and other factors may differ from those occurring during clinical studies. Similarly, the cited frequencies cannot be directly compared with figures obtained from other clinical investigations involving different treatments, uses, or investigators. Review of these frequencies, however, provides prescribers with a basis to estimate the relative contribution of drug and non-drug factors to the incidence of adverse events in the population studied.

**Table 1: Incidence of Treatment-Emergent Adverse Experiences in Parallel-Group, Placebo-Controlled Clinical Trials in Narcolepsy, OSAHS, and SWSD with modafinil (200mg. 300mg. 400mg).**

<table>
<thead>
<tr>
<th>Body System</th>
<th>Preferred Term</th>
<th>Modafinil (n=934, %)</th>
<th>Placebo n=567, %</th>
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<tr>
<td>Body as a whole</td>
<td>Headache</td>
<td>34</td>
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<td>Back pain</td>
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<td>Flu syndrome</td>
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<td></td>
<td>Chills</td>
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<td>Cardiovascular system</td>
<td>Hypertension</td>
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<td>Tachycardia</td>
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<td></td>
<td>Palpitation</td>
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<td>Vasodilatation</td>
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<td>Digestive system</td>
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<td></td>
<td>Diarrhoea</td>
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<td></td>
<td>Dyspepsia</td>
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<tr>
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<td>Dry mouth</td>
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<td>Anorexia</td>
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<tr>
<td></td>
<td>Constipation</td>
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<td>Abnormal liver function²</td>
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<tr>
<td></td>
<td>Flatulence</td>
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<td>Mouth ulceration</td>
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NEW ZEALAND DATA SHEET

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<tr>
<th>System</th>
<th>Conditions</th>
<th>Incidence</th>
<th>Placebo</th>
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<td>Haemic/Lymphatic system</td>
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<td>Insomnia</td>
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<td>Dizziness</td>
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<td>Paraesthesia</td>
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<td>Hypertonia</td>
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<tr>
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<td>Dyskinesia(^1)</td>
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<td>Hyperkinesia</td>
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<tr>
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<td>Agitation</td>
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</tr>
<tr>
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<td>Confusion</td>
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<td>Epistaxis</td>
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<td>Herpes simplex</td>
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</tr>
<tr>
<td>Special Senses</td>
<td>Amblyopia</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Abnormal vision</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Taste perversion</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Eye pain</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Urogenital</td>
<td>Urine abnormality</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Haematuria</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>Pyuria</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

*Six double-blind, placebo controlled clinical studies in narcolepsy, OSAHS and SWSD

1. Events reported by at least 1% of patients treated with modafinil that were more frequent than in the placebo group are included; incidence is rounded to the nearest 1%. The adverse experience terminology is coded using a standard modified COSTART Dictionary.

Events for which the modafinil incidence was at least 1%, but equal to or less than placebo are not listed in the table. These events included the following: infection, pain, accidental injury, abdominal pain, hypothermia, allergic reaction, asthenia, fever, viral infection, neck pain, migraine, abnormal electrocardiogram, hypotension, tooth disorder, vomiting, periodontal abscess, increased appetite, ecchymosis, hyperglycaemia, peripheral oedema, weight loss, weight gain, myalgia, leg cramps, arthritis, cataplexy, thinking abnormally, sleep disorder, increased cough, sinusitis, dyspepsia, bronchitis, rash, conjunctivitis, ear pain, dysmenorrhea\(^4\), urinary tract infection

2. Elevated liver enzymes
3. Oro-facial dyskinesias
4. Incidence adjusted for gender
Post Marketing Experience
Post Marketing Experience for modafinil, principally from spontaneous reporting based on reporting rates and not incidence rates, has documented the following adverse events:

Common 1/100 to <1/10
Uncommon 1/1,000 to <1/100
Rare 1/10,000 to <1/1,000
Very rare <1/10,000

Cardiac disorders
Rare Palpitations
Very rare Ischaemic heart disease, cardiac arrhythmias

Gastrointestinal disorders
Rare Dry mouth, nausea, diarrhoea, vomiting, abdominal pain

General disorders
Rare Tolerance, chest pain, lack of efficacy, condition aggravated, malaise, fatigue
Very rare Oedema

Immune system disorders
Very rare Multi-organ system hypersensitivity reactions, urticaria (hives), angioedema, anaphylaxis

Investigations
Rare Increased hepatic enzymes, increased gamma-GT, weight increase, weight decrease, blood pressure increased
Very Rare Abnormal ECG

Musculoskeletal and connective tissue disorders
Rare Muscle weakness

Nervous system disorders
Uncommon Headache
Rare Dizziness, tremor, paraesthesia, dyskinesia
Very rare Dyskinesias, including reports of tardive dyskinesia; convulsions

Psychiatric disorders
Rare Nervousness, agitation, irritability, psychomotor hyperactivity, depression, anxiety, confusion, insomnia, suicide attempt, aggravated depression, psychosis, mania, delusions, hallucinations, suicidal ideation, thinking abnormal and aggression

Renal and urinary disorders
Rare Foul urine odour

Skin and subcutaneous tissue disorders
Rare Rash, acne, eczema, pruritus
Very rare Serious or life threatening rash, including erythema multiforme, Stevens Johnson Syndrome (SJS), Toxic Epidermal Necrolysis (TEN), and Drug Rash with Eosinophilia and System Symptoms (DRESS), and hyperhidrosis

Vascular disorders
Rare Hypertension
Reporting suspected adverse reactions after authorisation of the medicine is important. It allows continued monitoring of the benefit/risk balance of the medicine. Healthcare professionals are asked to report any suspected adverse reactions [https://nzphvc.otago.ac.nz/reporting/](https://nzphvc.otago.ac.nz/reporting/)

4.9 Overdose

**Symptoms**

A small number of individuals have each taken modafinil at doses of 1000 mg/day (2.5 times the maximum recommended daily dose of 400 mg) or more. The adverse experiences observed were limited, expected and non-life threatening, and the patients recovered fully by the following day. The adverse experiences included excitation or agitation, insomnia and slight or moderate elevations in haemodynamic parameters. No specific organ toxicities were observed. Other observed high dose effects in clinical studies have included anxiety, irritability, aggressiveness, confusion, nervousness, tremor, palpitations, sleep disturbances, nausea, diarrhoea and decreased prothrombin time.

Death has occurred with modafinil overdose alone or in combination with other medicines. Symptoms accompanying modafinil overdose, alone or in combination with other medicines, have included: insomnia; central nervous system symptoms such as restlessness, disorientation, confusion, agitation, anxiety, excitation and hallucination; digestive changes such as nausea and diarrhea; and cardiovascular changes such as tachycardia, bradycardia, hypertension and chest pain.

**Management**

Management of overdosage is primarily symptomatic, as no specific antidote to the toxic effects of modafinil overdose has been identified. Overdoses should be managed empirically, with supportive care, including cardiovascular monitoring. As for any overdose, the physician should consider contacting a Poison-control centre regarding treatment.

For advice on the management of overdose please contact the National Poisons Centre on 0800 POISON (0800 764766).

5 PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Psychoanaleptics, centrally acting sympathomimetics

ATC code: N06BA07

The precise mechanism(s) through which modafinil promotes wakefulness is unknown. Modafinil has wake-promoting actions but a pharmacological profile that is distinct from sympathomimetic amines, which increase wakefulness by other mechanisms.

Modafinil does not bind to most of the potentially relevant receptors for sleep/wake regulation, including those for noradrenaline, serotonin, dopamine, GABA, adenosine, histamine-3 and benzodiazepines. Modafinil is not a direct- or indirect-acting dopamine receptor agonist and is inactive in several *in vivo* preclinical models capable of detecting enhanced dopaminergic activity. *In vitro*, modafinil binds to the dopamine reuptake site with low affinity and causes an increase in extracellular dopamine, but no increase in dopamine release. Modafinil does not appear to be a direct or indirect α1-adrenergic agonist. Although modafinil-induced wakefulness can be attenuated by the α1-adrenergic receptor antagonist prazosin, modafinil has no activity in assay systems known to be responsive to the α-adrenergic agonists.
In rats, the wakefulness induced by amphetamine, but not modafinil, was antagonised by the dopamine receptor antagonist haloperidol. In cats, modafinil evoked neuronal activation in brain regions different from methylphenidate and amphetamine. Modafinil served as a positive reinforcer for cocaine in monkeys and was partially discriminated as stimulant-like in rats (see section 4.4 Special warnings and precautions for use: “Abuse and Dependence Potential”).

The optical enantiomers of modafinil have similar pharmacological actions in mice, but have not been studied individually in humans. The two major metabolites of modafinil, modafinil acid and modafinil sulfone, showed little CNS-activating activity in animal studies.

Modafinil in humans restores and/or improves the level of wakefulness. Changes are found in electrophysiological parameters reflecting alertness (ratio of power of alpha rhythm to power of theta rhythm), starting from a dose of 100 mg in the morning. An increase is seen in latency periods in the multiple sleep latency test, starting from 200 mg in the morning. Modafinil opposes the impairment of cognitive (in particular, memory), psychomotor and neurosensory performance induced by sleep deprivation. This activity is observed in the absence of any modifications of appetite or behaviour.

Morning administration of 200 mg does not appear to affect nocturnal sleep. Administration of 100 mg morning and noon may prolong the subjective time taken to fall asleep. Evening administration may disturb sleep. This pharmacodynamic activity does not appear to affect the autonomic nervous system.

CLINICAL TRIALS

Studies reported here were multicenter, randomized, double-blind, placebo-controlled parallel-group clinical trials. The efficacy criteria reported for the trials included:

- Maintenance of Wakefulness Test (MWT), which quantitatively measures the patient’s ability to resist sleep and maintain wakefulness. The patients were asked to attempt to remain awake without using extraordinary measures. The test was terminated after 20 minutes if no sleep occurred or 10 minutes after sleep onset.
- Clinical Global Impression of Change (CGI-C), which is a 7-point scale ranging from “Very Much Worse” to “Very Much Improved” from baseline; it was assessed by an independent clinician who had no access to any data about the patients other than a measure of their baseline severity.
- Epworth Sleepiness Scale (ESS), which is a recall-based questionnaire devised to provide a measurement of the subject’s general level of day-to-day sleepiness, or preferably, sleep propensity.

Narcolepsy:

The effectiveness of modafinil in reducing the excessive sleepiness (ES) associated with narcolepsy was established in two US 9-week, multicentre, placebo-controlled, two-dose (200mg per day and 400mg per day) parallel-group, double-blind studies of outpatients who met the ICD-9 and American Sleep Disorders Association criteria for narcolepsy (which are also consistent with the American Psychiatric Association DSM-IV criteria). These criteria included either:

- Recurrent daytime naps or lapses into sleep that occur almost daily for at least three months, plus sudden bilateral loss of postural muscle tone in association with intense emotion (cataplexy), or
NEW ZEALAND DATA SHEET

- A complaint of excessive sleepiness or sudden muscle weakness with associated features: sleep paralysis, hypnagogic hallucinations, automatic behaviours, disrupted major sleep episode; and polysomnography demonstrating one of the following: sleep latency less than 10 minutes or rapid eye movement (REM) sleep latency less than 20 minutes.

In addition, for entry into these studies, all patients were required to have objectively documented excessive daytime sleepiness, a Multiple Sleep Latency Test with two or more sleep onset REM periods, and the absence of any clinically significant active medical or psychiatric disorder.

In both studies, the primary measures of effectiveness were:
1. sleep latency, as assessed by the MWT, and
2. the change in the patient’s overall disease status, as measured by the CGI-C.

For a successful trial, both measures had to show significant improvement.

Patients in both modafinil treatment groups were able to stay awake longer than those receiving placebo and were rated by an independent clinician as having a significant improvement in illness. A statistically significantly enhanced ability to remain awake was shown on the MWT and the CGI-C scale (see Tables 2 and 3).

Obstructive Sleep Apnoea/Hypopnoea Syndrome (OSAHS):

The results from two major phase 3 clinical trials of modafinil in patients with OSAHS are presented in Tables 2 and 3. The effectiveness of MODAVIGIL® in reducing the excessive sleepiness associated with OSAHS was established in two clinical trials. In both studies, patients were enrolled who met the International Classification of Sleep Disorders (ICSD) criteria for OSAHS (which are also consistent with the American Psychiatric Association DSM-IV criteria). These criteria include either, 1) excessive sleepiness or insomnia, plus frequent episodes of impaired breathing during sleep, and associated features such as loud snoring, morning headaches and dry mouth upon awakening; or 2) excessive sleepiness or insomnia and polysomnography demonstrating one of the following: more than five obstructive apnoeas, each greater than 10 seconds in duration, per hour of sleep and one or more of the following: frequent arousals from sleep associated with the apnoeas, bradytachycardia, and arterial oxygen desaturation in association with the apnoeas. In addition, for entry into these studies, all patients were required to have excessive sleepiness as demonstrated by a score ≥10 on the Epworth Sleepiness Scale, despite treatment with continuous positive airway pressure (CPAP). Evidence that CPAP was effective in reducing episodes of apnoea/hypopnea was required along with documentation of CPAP use.

Study 303 (n = 327) assessed the efficacy and safety of two doses of modafinil (200 mg and 400 mg per day) in the treatment of excessive sleepiness in patients with established OSAHS, despite partial or satisfactory use of continuous positive airway pressure (CPAP) therapy. The primary efficacy variables for study 303 were MWT and CGI-C. Study 402 (n = 157) provides supportive data for modafinil 400mg per day in the treatment of excessive sleepiness in patients with established OSAHS, despite the use of effective CPAP therapy. The primary efficacy variable for study 402 was ESS.

Clinically significant improvements were reported for each parameter and for both doses of modafinil compared to placebo in Study 303 out to 12 weeks’ double-blind treatment, and Study 402 out to 4 weeks’ double-blind treatment (see Tables 2 and 3).

In a 12 month open-label extension period for Study 303 in which patients titrated their daily dose of modafinil according to clinical response, ESS scores remained consistently improved compared to baseline values in both those previously on modafinil and those previously on placebo.
For OSAHS, modafinil has been shown to produce clinically meaningful reductions in excessive sleepiness and its adverse effects on quality of life, in both the short and long term.

**Shift Work Sleep Disorder (SWSD):**

Two clinical trials conducted in patients with shift work sleep disorder provide information on the efficacy of modafinil in this indication. The moderate to severe subgroup of patients with SWSD for whom modafinil is indicated is defined by the inclusion criteria in the pivotal clinical trials. These criteria included a CGI-S (Clinical Global Impression of Severity) rating of at least “moderately ill” (relating to ES on shift nights) at baseline, a mean sleep latency of no more than 6 minutes on the Multiple Sleep Latency Test (MSLT) and no more than 87.5% sleep efficiency (time sleeping/time in bed). All patients met the International Classification of Sleep Disorders (ICSD) criteria for chronic SWSD (which are consistent with the American Psychiatric Association DSM-IV criteria for Circadian Rhythm Sleep Disorder: Shift Work Type).

Patients were enrolled if they worked at least 5 night shifts per month (of which at least 3 nights were consecutive) and planned to maintain this schedule for the duration of the double-blind portion of the study. Each night shift was no longer than 12 hours in duration and included at least 6 hours between the hours of 2200 and 0800. Patients with any other disorder that might account for their excessive sleepiness were excluded.

Placebo or modafinil was taken 30 to 60 minutes before each night shift. Having worked three consecutive night shifts, patients were admitted to the sleep centre for a fourth, simulated night shift (= a study visit), during which the various efficacy parameters were assessed.

Study 305 (n = 209) evaluated the efficacy and safety of 12 weeks’ therapy with modafinil at a dose of 200 mg as treatment for adults with excessive sleepiness associated with chronic shift work sleep disorder. The primary efficacy variables were MSL-MSLT and CGI-C. Statistically significant improvements were seen for patients in the modafinil group when compared to patients in the placebo group for both of the primary endpoint measures (see Tables 2 and 4).

Study 306 (n = 278) evaluated the safety and impact on Quality of Life of 12 weeks of modafinil therapy at dosages of 200mg or 300mg once daily as treatment for adults with excessive sleepiness associated with shift work sleep disorder. The potential impact of modafinil treatment on quality-of-life was assessed by measuring the mean changes from baseline to week 12 using the following measures:

- Functional Outcomes of Sleep Questionnaire (FOSQ)
- 36-Item Short Form Health Survey (SF-36)

In Study 306, modafinil treatment appeared to have a clinically meaningful effect on patient quality of life as assessed by the FOSQ. For the patients in the modafinil 300 mg/day group, improvement from baseline to week 12 was statistically significant for the total score (p = 0.0126), and for the individual scores for vigilance (p = 0.0123), activity level (p = 0.0055) and general productivity (p = 0.0041) when compared with placebo. Although not statistically significant, the p-values for the change from baseline for the modafinil-200mg/day treatment group showed a trend toward significance.

Improvement was observed in the mental component score of SF-36 at all time points for patients in the modafinil-treated groups compared with the placebo-treated group. Statistical significance was observed at endpoint with the modafinil 300 mg/day group for the mental component summary (p = 0.0113), vitality (p < 0.0001) and role emotion (p = 0.0444) when compared with placebo.
In a 12-month open label extension period for Study 306, improvements in FOSQ total score and in the SF-36 mental composite score at endpoint were of the same magnitude as those seen in the double-blind period, and were considered clinically meaningful.

In SWSD, modafinil has been shown to produce clinically meaningful reductions in excessive sleepiness and had positive impact on quality of life, in both the short and long term.

### Table 2. Summary of the pivotal US clinical studies of modafinil as measured by defined CGI-C responder rates

<table>
<thead>
<tr>
<th>CGI-C number (% improved at endpoint)</th>
<th>Study</th>
<th>400 mg</th>
<th>200 mg</th>
<th>Placebo</th>
</tr>
</thead>
<tbody>
<tr>
<td>Narcolepsy Study 301</td>
<td></td>
<td>62 (72)</td>
<td>61 (64)</td>
<td>34 (37)</td>
</tr>
<tr>
<td>p-values compared to placebo</td>
<td></td>
<td><em>p = 0.0001</em></td>
<td><em>p = 0.0001</em></td>
<td>not applicable</td>
</tr>
<tr>
<td>Narcolepsy Study 302</td>
<td></td>
<td>52 (60)</td>
<td>48 (58)</td>
<td>33 (38)</td>
</tr>
<tr>
<td>p-values compared to placebo</td>
<td></td>
<td><em>p = 0.0026</em></td>
<td><em>p = 0.0044</em></td>
<td>not applicable</td>
</tr>
<tr>
<td>OSAHS Study 303</td>
<td></td>
<td>63 (68%)</td>
<td>60 (61%)</td>
<td>37 (37%)</td>
</tr>
<tr>
<td>p-values compared to placebo</td>
<td></td>
<td><em>p = 0.0001</em></td>
<td><em>p = 0.0000</em></td>
<td>not applicable</td>
</tr>
<tr>
<td>SWSD Study 305</td>
<td></td>
<td>--</td>
<td>66 (74%)</td>
<td>37 (36%)</td>
</tr>
<tr>
<td>p-values compared to placebo</td>
<td></td>
<td>--</td>
<td><em>p = 0.0001</em></td>
<td>not applicable</td>
</tr>
</tbody>
</table>

Table 3: Summary of clinical trial data in patients with narcolepsy or OSAHS following treatment with modafinil

Data in changes from baseline to endpoint for modafinil vs placebo, unless otherwise stated:

- **a** Positive value = modafinil better than placebo, negative value = modafinil worse than placebo
- **b** Difference in absolute value over placebo at endpoint

### Table 4. SWSD study 305 sleep latency (mins) from MSLT at endpoint

<table>
<thead>
<tr>
<th></th>
<th>Modafinil 200mg (N=88)</th>
<th>Placebo (N=96)</th>
<th>p-value (ANOVA)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline</td>
<td>mean ± sd: 2.07 ± 1.53</td>
<td>2.04 ± 1.82</td>
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<tr>
<td>Endpoint</td>
<td>mean ± sd: 3.77 ± 4.32</td>
<td>2.37 ± 2.73</td>
<td></td>
</tr>
<tr>
<td>Change at endpoint</td>
<td>mean ± sd: 1.70 ± 3.79</td>
<td>0.34 ± 2.77</td>
<td>0.0022</td>
</tr>
<tr>
<td></td>
<td>range: -4.1 to 14.0</td>
<td>-10.3 to 8.1</td>
<td></td>
</tr>
</tbody>
</table>
5.2 Pharmacokinetic properties

Modafinil is a racemic compound, whose enantiomers have different pharmacokinetics (e.g. the half-life of the \(l\)-isomer is approximately three times that of the \(d\)-isomer in humans). The enantiomers do not interconvert. At steady state, total exposure to the \(l\)-isomer is approximately three times that of the \(d\)-isomer. The trough concentration (\(C_{\text{minss}}\)) of circulating modafinil after once daily dosing consists of 90% of the \(l\)-isomer and 10% of the \(d\)-isomer.

Absorption and Distribution

Modafinil is slowly absorbed with an absorption half-life of approximately 1 hour. Peak plasma concentrations (\(C_{\text{max}}\)) of approximately 3.3mg/L are reached 3 hours (\(t_{\text{max}}\)) after administration of a 200 mg dose. Both the area under the plasma concentration curve (AUC), and the peak plasma concentration show dose-proportionality in the 50 to 400 mg range. The absolute oral bioavailability could not be determined due to the aqueous insolubility (< 1mg/mL) of modafinil, which precluded intravenous administration. Food has no effect on the overall bioavailability of modafinil, however, its absorption (\(t_{\text{max}}\)) may be delayed by approximately one hour if taken with food.

Modafinil is well distributed in body tissue with an apparent volume of distribution (~0.9 L/kg) larger than the volume of total body water (0.6 L/kg). Modafinil is weakly bound to plasma proteins (62%), mainly to albumin. At serum concentrations obtained at steady state after doses of 200 mg/day, modafinil exhibits no displacement of protein binding of warfarin, diazepam, or propranolol.

Metabolism and Excretion

The major route of elimination (~90%) is metabolism, primarily by the liver, with subsequent renal elimination of the metabolites. The elimination half-life of modafinil after multiple doses is about 10-12 hours. Urine alkalinisation has no effect on the elimination of modafinil.

Metabolism occurs through hydrolytic deamination, S-oxidation, aromatic ring hydroxylation, and glucuronide conjugation. Less than 10% of an administered dose is excreted as the parent compound. In a clinical study using radiolabelled modafinil, a total of 81% of the administered radioactivity was recovered in 11 days post-dose, predominantly in the urine (80% vs. 1% in the faeces).

The chief metabolite (40-50% of the dose) is acid modafinil, which has no pharmacological activity. The excretion of modafinil and its metabolites is chiefly renal, with a small proportion being eliminated unchanged (< 10%).

Only two metabolites reach appreciable concentrations in plasma, i.e., acid modafinil and modafinil sulfone. In preclinical models, modafinil acid, modafinil sulfone, 2-[[diphenylmethyl]sulfonyl]acetic acid and 4-hydroxy modafinil, were inactive or did not appear to mediate the arousal effects of modafinil.

In humans, modafinil shows a possible induction effect on its own metabolism after chronic administration of doses ≥ 400 mg/day. In vitro studies with human hepatocytes and liver microsomes have shown induction of metabolising enzymes CYP3A4 and CYP1A1/2, and inhibition of CYP2C19 (see section 4.5 Interaction with other medicines and other forms of interaction).

Special Populations

Children: The pharmacokinetics of modafinil have not been studied in children.
**Age effect:** A slight decrease (~20%) in the oral clearance of modafinil was observed in subjects with a mean age of 63 years (range: 53 to 73 years). The clearance of modafinil may be reduced in the elderly.

**Gender Effect:** The pharmacokinetics of modafinil are not affected by gender.

**Race effect:** The influence of race on the pharmacokinetics of modafinil has not been studied.

**Renal impairment:** The pharmacokinetics of modafinil were not significantly influenced in patients with severe chronic renal failure (creatinine clearance ≤ 20mL/min), but the exposure to modafinil acid (an inactive metabolite) was increased 9-fold.

**Hepatic impairment:** The oral clearance of modafinil was decreased by about 60% and the steady state concentration was doubled in patients with severe chronic hepatic impairment.

### 5.3 Preclinical safety data

Toxicology studies by single and repeated dosing have revealed no particular toxic action in animals.

Modafinil is not considered to be mutagenic or carcinogenic.

Reproductive toxicity studies conducted in rats and rabbits showed an increased incidence in skeletal variations (changes in the numbers of ribs and delayed ossification), embryo-fetal lethality (peri-implantation loss and resorptions) and some evidence of an increase in stillbirths (rats only), in the absence of maternal toxicity, at clinically relevant exposures.

There was no effect on fertility and no evidence of teratogenic potential at systemic exposures equivalent to the maximum recommended human dose.

Reproduction toxicity studies revealed no effect on fertility, nor any teratogenic effect, nor any effect on viability, growth or development of the offspring.

Animal exposure to modafinil, based on actual plasma levels in the general toxicology, reproductive and carcinogenicity studies, was less than or similar to that expected in humans. This circumstance is the result of metabolic auto-induction noted in the pre-clinical studies. However, animal exposure on a mg/kg dose basis to modafinil in the general toxicology, reproductive and carcinogenicity studies was greater than the expected exposure, calculated on a similar basis, in humans.

In the rat peri-post-natal study, modafinil concentration in milk was about 11.5 times higher than in plasma.

### 6 PHARMACEUTICAL PARTICULARS

#### 6.1 List of excipients

Modafinil tablets contain the following excipients:

- Lactose monohydrate
- Crospovidone
- Povidone K30
- Lactose Anhydrous
- Talc
- Silica, Colloidal Anhydrous
- Sodium Stearyl fumarate.
6.2 Incompatibilities
Not applicable.

6.3 Shelf life
24 months

6.4 Special precautions for storage
Store below 25 °C

6.5 Nature and contents of container
Blister packs of 30 tablets (3 blisters strips containing 10 tablets).

6.6 Special precautions for disposal and other handling
Any unused medicine or waste material should be disposed of in accordance with local requirements.

7 MEDICINE SCHEDULE
Prescription Medicine

8 SPONSOR
Max Health Ltd
PO Box 65 231
Mairangi Bay
Auckland 0754
Telephone: (09) 815 2664.

9 DATE OF FIRST APPROVAL
10 October 2019

10 DATE OF REVISION OF THE TEXT
10 October 2019

SUMMARY TABLE OF CHANGES

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<th>Summary of new information</th>
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