MOVICOL[®] Ready To Take Oral Solution (macrogol 3350 and electrolytes)

NAME OF THE MEDICINE:

MOVICOL Ready To Take, Oral Solution.

DESCRIPTION:

A clear colourless to light yellow, free flowing liquid.

Each 25 mL of MOVICOL Ready To Take oral solution contains:

Macrogol 3350	13.125 g
Sodium chloride	350.8 mg
Sodium bicarbonate	178.6 mg
Potassium chloride	50.2 mg

The concentration of electrolyte ions in each 25 mL sachet is:

Sodium	325 mmol/L
Chloride	267 mmol/L
Potassium	27 mmol/L
Bicarbonate	85 mmol/L

This corresponds to the following amount of each electrolyte in each 25 mL dose:

8.125 mmol
6.675 mmol
0.675 mmol
2.125 mmol

MOVICOL Ready To Take oral solution al so contains sucralos e (E955), purified water, and strawberry-banana flavour, which contains natural flavouring substances, flavouring preparations (including celery oil) and propylene glycol.

PHARMACOLOGY:

Macrogol 3350 exerts an osmotic action in t he gut, which induces a laxative effect. Macrogol 3350 increases the stool volume , which triggers colon motility via neuromuscular pathways. The physiological consequence is an improved propulsive colonic transportation of the softened stool s and a facilitatio n of the defaecation. Electrolytes combined with macrogol 3350 are exc hanged across the intestinal barrier (mucosa) with serum electrolytes and excreted in faec al water without net gain or loss of sodium, potassium and water.

Macrogol 3350 is unchanged along the gut. It is virtually unabsorbed from the gastrointestinal tract. Any macrogol 3350 that is absorbed is excreted via the urine.

The laxative action of macrogol has a time course which will vary according to the severity of the constipation being treated.

INDICATIONS:

For use in adults and children over 12 years of age for effective relief from constipation and treatment of chronic constipation.

CONTRAINDICATIONS:

Intestinal perforation or obstruction due to structural or functional disorder of the gut wall, ileus and severe inflammatory conditions of the intestinal tract, such as Crohn's disease, ulcerative colitis and toxic megacolon.

Known hypersensitivity to macrogol or any of the ingredients.

PRECAUTIONS:

Adverse reactions are possible as described under *Adverse Reactions*. If patients develop any symptoms indicating shifts of fluid/electrolytes (eg. oedema, shortness of breath, increasing fatigue, dehydration, cardiac failure) MOVICOL Ready To Take should be stopped immediately and electrolytes measured, and any abnormality should be treated appropriately.

The absorption of other medicines could transiently be reduced due to a decrease in gastro-intestinal transit time induced by MOVICOL Ready To Take (see interactions with other drugs).

As with all laxatives, pr olonged use is not usually recommended and may lead to dependence. If prolonged use is neces sary, it should only be under medical supervision. Extended use may be necess ary in the care of patients with severe chronic or resistant constipat ion, secondary to multiple sclerosis or Parkinson's Disease, or induced by regular constipation ng medication, in particular opioids and antimuscarinics. Patients should be advised to drink plenty of water. They should also increase fibre in the diet, except in the case of medication-induced constipation.

Use in pregnancy:

Category B1. Drugs which hav e been taken by only a limited num ber of pregnant women and women of child bearing age, without an increase in the frequency of malformation or other direct or indirect harmful effects on the h uman fetus having been observed. Studies in animals have not shown evidence of an increased occurrence of fetal damage.

There were no direct embryotoxic or terat ogenic effects in rats at maternally toxic doses up to 40 g/kg/day, 51x the maxim um recommended dose in humans for chronic constipation.

Indirect effects, including reduc tion in fetal and pla cental weights, reduced fetal viability and abortions, were noted in the rabbit at doses below the maximum recommended human dose. Rabbits are particularly sensitive to the effects of GI acting substances, and the findings are considered most likely a reflection of poor maternal condition as a result of an exaggerated pharmacodynamic response rather than direct embryofetal toxicity. There was no indication of a teratogenic effect.

Use in Lactation:

No effects on the breastfed newborn/inf ant are anticipated since the systemic exposure of the breast-feeding woman to macrogol 3350 is negligible. MOVICOL Ready to Take can be used during breast-feeding. <u>Use in children:</u> Not recommended. Alternative MOVICOL products are available for children.

Mutagenicity and carcinogenicity:

Preclinical studies provide evidence that macrogol 3350 has no significant systemic toxicity potential, based on conventional studies of pharmacology, repeated dos e toxicity, genotoxicity and toxicity to reproduction.

There are long-term animal toxic ity and carcinogenicity studies involving m acrogol 3350. Results from these and ot her toxicity studies using high levels of orally administered high m olecular weight macrogol s provide evidence of safety at the recommended therapeutic dose.

INTERACTIONS WITH OTHER MEDICINES:

There is a possibility that the absorption of other medicines could be transiently reduced during use with MOVICOL Ready To Take (see Precautions). The rehave been isolated reports of decreased efficacy with some concomitantly administered medicinal products, e.g. anti-epileptics. A theoretical potential also e xists for decreased absorption (rate and extent) of drugs which are generally poorly absorbed or are contained in sustained or modified release dosage forms. This is more likely to occur if MOVICOL Ready T o Take is overdosed to induce watery diarrhoea.

ADVERSE REACTIONS:

Reactions related to the gastroint estinal tract occur most commonly. These reactions may occur as a consequenc e of expansion of the contents of the gastrointestinal tract, and an increase in motility due to the pharmacologic effects of MOVICOL Ready To Take. Diarrhoea usually responds to dose reduction.

System Order Class	Adverse Event
Immune system disorders	Allergic reactions, including anaphylactic reactions, dyspnoea, and skin reactions (see below).
Metabolism and nutrition disorders	Electrolyte disturbances, particularly hyperkalaemia and hypokalaemia.
Nervous system disorders	Headache.
Gastrointestinal disorders	Abdominal pain, diarrhoea, vomiting, nausea, dyspepsia, abdominal distension, borborygmi, flatulence, anorectal discomfort.
General disorders and administration site conditions	Peripheral oedema.
Skin and subcutaneous tissue disorders	Allergic skin reactions including angioedema, urticaria, pruritus, rash, erythema.

DOSAGE AND ADMINISTRATION:

Movicol Ready To Take should be used dire ctly from the sachet. This product does not need to be diluted with water.

The fluid content of MOVICOL Ready to Take does not replace regular fluid intake and adequate fluid intake must be maintained. It is recommended that patients drink a glass of water or other fluid after taking MOVICOL Ready To Take.

Adults and children 12 years and older:

<u>Constipation</u>: One sachet of MOVICOL Ready T o Take once daily. This may be increased to 2 – 3 sachets daily, if required according to individual response.

Children under 12 years of age:

MOVICOL Ready To Take is not recommended for use in children below the age of 12 years (see PRECAUTIONS). Alternative MOVICOL products are available for children.

Patients with impaired cardiovascular function: No more than two sachet s should be taken in any one hour.

Patients with renal insufficiency: No dosage change is necessary for treatment of constipation.

OVERDOSAGE:

Severe pain or distension can be treated by nasogastric aspiration. Extensive fluid loss by diarrhoea or vomiting may require co rrection of electrolyte disturbances. For information on the management of overdose, contact the Poisons Information Centre on 13 11 26 (Australia)

PRESENTATION AND STORAGE CONDITIONS:

MOVICOL Ready To Take is a liquid for oral solution that does not require dilution before use. Each 25 mL contains 13.125 g of macrogol 3350 and electrolytes. It is supplied in 25 mL sachets, in boxes of 10, 20, 30 and 50 sachets. Not all pack sizes may be marketed.

Store below 30°C. Do not refrigerate or freeze.

NAME AND ADDRESS OF SPONSOR:

Norgine Pty Ltd. 3/14 Rodborough Road, Frenchs Forest NSW 2086.

POISON SCHEDULE

MOVICOL Ready To Take is a PHARMACY MEDICINE (S2).

DATE OF FIRST INCLUSION IN THE ARTG:

3 May 2016

DATE OF MOST RECENT AMENDMENT:

Not applicable.