#### Diprofol® 2% Propofol 2%

For IV Administration

### Diprofol 2% (propofol 20 mg/ml) 2. Qualitative and quantitative composition

1. Name of the medicinal product

### Diprofol 20 mg/ml, emulsion for injection or infusion, contains 20 mg/ml propofol.

The emulsion contains as excipients soybean oil

and egg lecithin, which are also used in intravenous feeding. It contains no preservatives. For the full list of excipients, see Section 6.1.

3. Pharmaceutical form Emulsion for injection or infusion.

### Diprofol is a white, aqueous isotonic oil-in-water

emulsion for intravenous administration.

4. Clinical particulars 4.1 Therapeutic indications

### Diprofol 2% is a short-acting intravenous general

<u>Posology</u>

## anaesthetic for

• induction and maintenance of general anaesthesia in adults and children > 3 years.

- sedation of ventilated patients > 16 years of age in the intensive care unit.
- sedation for diagnostic and surgical procedures. alone or in combination with local or regional
- anaesthesia in adults and children > 3 years. 4.2 Posology and method of administration

#### 4.2.1 Induction of General Anaesthesia Propofol must be used only in well equipped hospitals or medical centers by doctors trained in anaesthesia or the treatment of intensive care

patients. Continual monitoring of the circulation and the respiration (for example ECG pulse oxymeter) is necessary. Provisions for prevention of airway obstruction, artificial respiration and other resuscitation provisions must be immediately available at all times. As regards sedation during surgical or diagnostic operations propofol must not be administered by the same person who performs the surgical or diagnostic operation. Additional analgesics are generally necessary in combination with propofol.

by infusion. Administration of Diprofol 2% by bolus

Diprofol 2% may be used to induce anaesthesia by infusion but only in those patients who will receive Diprofol 2% for maintenance of anaesthesia. In unpremedicated and premedicated patients, it is

recommended that Diprofol 2% should be titrated

(approximately 2ml [40mg] every 10 seconds in an average healthy adult by infusion) against the response of the patient until the clinical signs show the onset of anaesthesia. Most adult patients aged less than 55 years are likely to require1.5-2.5mg/ kg of Diprofol 2%. The total dose required can be reduced by lower rates of administration (1-2.5ml/ min [20-50mg/min]).

cardiac function, the dosage requirements will be less and the total dose of Diprofol 2% may be reduced to a minimum of 1 mg/kg body weight. In these patients lower rates of administration should be applied (approximately 1 ml, corresponding to In older people the dose requirement for induction

should be given at a slower rate and titrated against the response. Paediatric population Diprofol 2% is not indicated for induction of anaesthesia in children less than 3 years of age. For induction of anaesthesia in children over 3

### years of age, Diprofol 2% should be titrated slowly

until clinical signs show the onset of anaesthesia. The dose should be adjusted according to age and/or bodyweight. Most patients over 8 years of age require approximately 2.5mg/kg bodyweight of Diprofol 2% for induction of anaesthesia. In younger children, dose requirements may be higher (2.5-For ASA 3 and 4 patients lower doses are Anaesthesia can be maintained by administering

clinical signs of light anaesthesia. Administration of Diprofol 2% by bolus injection is not recommended.

it is therefore important to maintain Diprofol 2% administration until the end of the procedure. Adults The required rate of administration varies considerably between patients, but rates in the region of 4-12 mg/kg/h usually maintain satisfactory anaesthesia.

### When Diprofol 2% is used for maintenance of anaesthesia the rate of infusion or 'target

ASA grades 3 and 4 will require further reductions in dose and dose rate. Rapid bolus administration (single or repeated) should not be used in older people as this may lead to cardiorespiratory depression. Paediatric population

3 years of age by administering Diprofol 2% by infusion to maintain the depth of anaesthesia required. The required rate of administration varies considerably between patients but rates in the region of 9-15 mg/kg/h usually achieve satisfactory anaesthesia. In younger children, dose

of 0.3-4 mg/kg/h of Diprofol 2% (See Section 4.4 Special warnings and precautions for use).

#### For sedation during intensive care it is advised that Diprofol 2% should be administered by continuous infusion. The infusion rate should be determined

by the desired depth of sedation. In most patients sufficient sedation can be obtained with a dosage Diprofol 2% is not indicated for sedation in intensive care of patients of 16 years of age or younger (see Section 4.3 Contraindications). It is recommended that blood lipid levels be monitored should Diprofol 2% be administered to patients thought to be at particular risk of fat overload.

1.0 ml of Diprofol 2% contains approximately 0.1g If the duration of sedation is in excess of 3 days, lipids should be monitored in all patients.

When Diprofol 2% is used for sedation of

# anaesthesia the rate of infusion should also be

to cardiorespiratory depression. Paediatric population Diprofol 2% is contraindicated for the sedation of ventilated children aged 16 years or younger receiving intensive care. 4.2.4 Sedation for Surgical and Diagnostic **Procedures Adults** 

To provide sedation for surgical and diagnostic

Most patients will require 0.5-1 mg/kg over 1-5 minutes for onset of sedation. Maintenance of sedation may be accomplished by titrating Diprofol 2% infusion to the desired level of sedation - most patients will require 1.5-4.5 mg/ kg/h. In patients of ASA grades 3 and 4 the rate of administration and dosage may need to be reduced. According to required dose, alternatively Diprofol 1 %

reduced. Patients of ASA grades 3 and 4 will require further reductions in dose and dose rate. Rapid bolus administration (single or repeated) should not be used in older people as this may lead to cardiorespiratory depression.

In children over 3 years of age, doses and administration rates should be adjusted according to the required depth of sedation and the clinical

response. Most paediatric patients require1-2 mg/ kg body weight of Diprofol 2% for onset of sedation. Maintenance of sedation may be accomplished by titrating Diprofol 2% infusion to the desired level of sedation. Most patients require1.5-9 mg/kg/h Diprofol 2%. In ASA 3 and 4 patients lower doses may be required. 4.2.5 Method of Administration

## Method of administration Propofol 20 mg/ml should be administered

undiluted intravenously. Propofol 20 mg/ml must not be mixed with injection or infusion fluids. However, simultaneous administration of propofol

or sodium chloride 0.9% via a Y - connector close to the injection site is possible. Ampoules and vials should be shaken before use. Before use the neck of the ampoule and the rubber stopper of the infusion vial must be disinfected with

aseptically into a sterile syringe or infusion system and then administered directly. During the infusion period the sterility of both propofol and the infusion system should be maintained.

Medicines or fluids that are added to a running propofol infusion must be added close to the

Diprofol 2% may be used to induce anaesthesia injection is not recommended.

In patients over this age and in patients of ASA grades III and IV, especially those with impaired

20 mg every 10 seconds). **Elderly** of anaesthesia with Diprofol 2% is reduced. The reduction should take into account of the physical status and age of the patient. The reduced dose

4mg/kg bodyweight). recommended (see also Section 4.4). 4.2.2 Maintenance of General Anaesthesia Diprofol 2% by continuous infusion to prevent the

Recovery from anaesthesia is typically rapid and

concentration' should also be reduced. Patients of

#### Diprofol 2% is not indicated for maintenance of anaesthesia in children less than 3 years of age. Anaesthesia can be maintained in children over

requirements may be higher. For ASA 3 and 4 patients lower doses are recommended (see also Section 4.2.3 Sedation During Intensive Care

appropriately if the monitoring indicates that fat is being inadequately cleared from the body. If the patient is receiving other intravenous lipid concurrently, a reduction in quantity should be

Administration of Diprofol 2% should be adjusted

made in order to take account of the amount of lipid infused as part of the Diprofol 2% formulation:

reduced. Patients of ASA grades 3 and 4 will require further reductions in dose and dose rate. Rapid bolus administration (single or repeated) should not be used in older people as this may lead

### procedures, rates of administration should be individualised and titrated to clinical response.

**Elderly** 

may be used. When Diprofol 2% is used for sedation the rate of

infusion or 'target concentration' should also be

Paediatric population Diprofol 2% is not indicated for surgical and diagnostic procedures in children aged less than 3 years.

20 mg/ml together with an infusion of glucose 5%

medicinal alcohol (spray or tissues). After use, any remaining medicine must be destroyed.

Propofol does not contain any preservatives and promotes the growth of micro-organisms. After opening of an ampoule or piercing of a vial, the contents must therefore immediately be put filters. The contents of an ampoule or a vial of propofol and any syringe of propofol are intended for single administration to one patient. Any remaining medicine must be destroyed after use. Infusion of undiluted propofol 20 mg/ml When propofol is administered by means of a continuous infusion, control of the infusion rate by means of a burette, drop counter, syringe pump or

cannula. Propofol must not be administered via

infusion systems that are provided with microbial

the case for parenteral administration of all kinds of fat emulsions, the duration of use of one infusion system for a continuous infusion with propofol must remain limited to 12 hours. The infusion system and the container must be removed and replaced after a maximum of 12 hours. Residues of propofol left over at the end of the infusion period or after changing of the system must be destroyed. In order to diminish pain at the beginning of the injection of propofol 20 mg/ml for induction of

volumetric infusion pump is recommended. As is

general anaesthesia, lidocaine can be injected directly before injection of propofol 20 mg/ml. Before administering the muscle relaxant atracurium, after administration of propofol, through the same infusion system, it is recommended to flush out the

infusion system. **Duration of administration** Propofol can be administered for a maximum of 7 days.

## 4.3 Contraindications

Hypersensitivity to the active substance or to any of the excipients listed in section 6.1.

## Diprofol 2% contains soybean oil and should not be

used in patients who are hypersensitive to peanut or soya. Diprofol 2% must not be used in patients of 16

years of age or younger for sedation in intensive care (see section 4.4). 4.4 Special warnings and precautions for use Diprofol 2% should be given by those trained in

anaesthesia (or, where appropriate, doctors trained

Patients should be constantly monitored and

facilities for maintenance of a patient airway, artificial

in the care of patients in Intensive Care).

ventilation and oxygen enrichment and other resuscitative facilities should be readily available at all times. Diprofol 2% should not be administered by the person conducting the diagnostic or surgical procedure. Abuse of, and dependence on Diprofol 2%,

been reported. As with other general anaesthetics, the administration of Diprofol 2% without airway care may result in fatal respiratory complications. When Diprofol 2% is administered for conscious sedation, for surgical and diagnostic procedures, patients should be continually monitored for early signs of hypotension, airway obstruction and oxygen

predominantly by health care professionals, have

and use of premedicants and other agents. Occasionally, hypotension may require use of intravenous fluids and reduction of the rate of administration of Diprofol 2% during the period of anaesthetic maintenance.

During induction of anaesthesia, hypotension and

transient apnoea may occur depending on the dose

with other sedative agents, when Diprofol 2% is used for sedation during operative procedures, involuntary patient movements may occur. During procedures requiring immobility these movements maybe hazardous to the operative site.

An adequate period is needed prior to discharge

of the patient to ensure full recovery after use of

Diprofol 2%. Very rarely the use of Diprofol 2% may

be associated with the development of a period of post-operative unconsciousness, which may be accompanied by an increase in muscle tone. This may or may not be preceded by a period of wakefulness. Although recovery is spontaneous, appropriate care of an unconscious patient should be administered. Diprofol 2% induced impairment is not generally detectable beyond 12 hours. The effects of Diprofol 2%, the procedure, concomitant medications, the age and the condition of the patient should be

• The advisability of being accompanied on leaving the place of administration The timing of recommencement of skilled or hazardous tasks such as driving The use of other agents that may sedate (e.g., benzodiazepines, opiates, alcohol)

considered when advising patients on:

As with other intravenous anaesthetic agents, caution should be applied in patients, with cardiac, respiratory, renal or hepatic impairment or in

hypovolaemic or debilitated patients. Diprofol 2%

clearance is blood flow dependent; therefore,

concomitant medication that reduces cardiac output will also reduce Diprofol 2% clearance. Diprofol 2% lacks vagolytic activity and has been associated with reports of bradycardia (occasionally profound) and also asystole. The intravenous administration of an anticholinergic agent before induction, or during maintenance of anaesthesia should be considered, especially in situations where vagal tone is likely to predominate or when Diprofol

2% is used in conjunction with other agents likely to

When Diprofol 2% is administered to an epileptic

Appropriate care should be applied in patients with

disorders of fat metabolism and in other conditions

where lipid emulsions must be used cautiously (see

patient, there may be a risk of convulsion.

cause a bradycardia.

section 4.2).

Use is not recommended with electroconvulsive As with other anaesthetics sexual disinhibition may

occur during recovery. The benefits and risks of the proposed procedure should be considered prior to proceeding with repeated or prolonged use (>3 hours) of propofol in

young children (< 3 years) and in pregnant women

as there have been reports of neurotoxicity in

The use of Diprofol 2% is not indicated in newborn

preclinical studies, see Section 5.3.

Paediatric population

infants as this patient population has not been fully investigated. Pharmacokinetic data (see Section 5.2) indicate that clearance is considerably reduced in neonates and has a very high inter-individual variability. Relative overdose could occur on administering doses recommended for older children and result in severe cardio vascular depression. Diprofol 2% is not recommended for use in children <3

years of age due to difficulty in titrating small volumes.

Advisory statements concerning Intensive Care

#### Propofol must not be used in patients of 16 years of age or younger for sedation for intensive care as the safety and efficacy of propofol for sedation in this age group have not been demonstrated (see Section 4.3).

**Unit management** 

Use of propofol emulsion infusions for ICU sedation has been associated with a constellation of metabolic derangements and organ system failures that may result in death. Reports have been received of combinations of the following: Metabolic acidosis, Rhabdomyolysis, Hyperkalaemia, Hepatomegaly, Renal failure, Hyperlipidaemia, Cardiac arrhythmia, Brugada-type ECG (elevated ST-segment and coved T-wave) and rapidly progressive Cardiac failure usually unresponsive to inotropic supportive reatment. Combinations of these events ha referred to as the Propofol Infusion Syndrome. These events were mostly seen in patients with

delivery to tissues; serious neurological injury and/or sepsis; high dosages of one or more of the following pharmacological agents - vasoconstrictors, steroids, inotropes and/or Diprofol 2% (usually at dose rates greater than 4mg/kg/h for more than 48 hours). Prescribers should be alert to these events in patients with the above risk factors and immediately discontinue propofol when the above signs develop. All sedative and therapeutic agents used in the intensive care unit (ICU), should be titrated to maintain optimal oxygen delivery and haemodynamic parameters. Patients with raised intra-cranial pressure (ICP) should be given appropriate treatment to support the cerebral perfusion pressure during

these treatment modifications.

exceed the dosage of 4mg/kg/h.

sodium diet.

disorders of fat metabolism and in other conditions where lipid emulsions must be used cautiously. It is recommended that blood lipid levels should be monitored if propofol is administered to patients thought to be at particular risk of fat overload. Administration of propofol should be adjusted

Treating physicians are reminded if possible not to

Appropriate care should be applied in patients with

**Additional Precautions** Caution should be taken when treating patients be susceptible to exacerbations of their disorder care. Maintenance of normothermia, provision of carbohydrates and good hydration are recommended for such patients. The early presentations of mitochondrial disease exacerbation and of the 'Propofol Infusion Syndrome' maybe similar. Diprofol 2% contains no antimicrobial preservatives and supports growth of micro-organisms.

the vial seal. Administration must commence

serious head injuries and children with respiratory tract infections who received dosages in excess of those advised in adults for sedation in the intensive The following appear to be the major risk factors for the development of these events: decreased oxygen

appropriately if the monitoring indicates that fat is being inadequately cleared from the body. If the patient is receiving other intravenous lipid concurrently, a reduction in quantity should be made in order to take account of the amount of lipid infused as part of the propofol formulation; 1.0mL of Diprofol 2% contains approximately 0.1g of fat.

with mitochondrial disease. These patients may when undergoing anaesthesia, surgery and ICU

Diprofol 2% contains 0.06 mg sodium per ml. To be

taken into consideration by patients on a controlled

who are predisposed to zinc deficiency, such as those with burns, diarrhoea and/or major sepsis. When Diprofol 2% is to be aspirated, it must be drawn aseptically into a sterile syringe or giving set immediately after opening the ampoule or breaking

EDTA chelates metal ions, including zinc, and reduces

microbial growth rates. The need for supplemental

zinc should be considered during prolonged

administration of Diprofol 2%, particularly in patients

Diprofol 2% and infusion equipment throughout the infusion period. Any infusion fluids added to the Diprofol 2% line must be administered close to the cannula site. Diprofol 2% must not be administered via a microbiological filter. Diprofol 2% and any syringe containing Diprofol

without delay. Asepsis must be maintained for both

2% are for single use in an individual patient. In accordance with established guidelines for other lipid emulsions, a single infusion of Diprofol 2% must not exceed 12 hours. At the end of the procedure or at 12 hours, whichever is the sooner, both the reservoir of Diprofol 2% and the infusion line must be discarded and replaced as appropriate. 4.5 Interaction with other medicinal products

### and other forms of interaction Diprofol 2% has been used in association with

spinal and epidural anaesthesia and with commonly premedicants, neuromuscular blocking drugs, inhalational agents and analgesic agents; no pharmacological incompatibility has been encountered. Lower doses of Diprofol 2% may be required where general anaesthesia is used as an adjunct to regional anaesthetic techniques. Profound hypotension has been reported following anaesthetic with propofol in patients treated with rifampicin. The concurrent administration of other CNS depressants such as pre-medication drugs,

inhalation agents, analgesic agents may add to the sedative, anaesthetic and cardiorespiratory depressant effects of Diprofol 2% (see Section 4.4). A need for lower propofol doses has been observed in patients taking valproate. When used

concomitantly, a dose reduction of propofol may be considered. 4.6 Fertility, pregnancy and lactation

The safety of Diprofol 2% during pregnancy has not been established. Studies in animals have shown

should not be given to pregnant women except when absolutely necessary. Diprofol 2% crosses the placenta and can cause neonatal depression. Diprofol 2% can, however, be used during an induced abortion. Diprofol 2% crosses the placenta and can cause neonatal depression. It should not be used for

obstetric anaesthesia. **Breast-feeding** Studies of breast-feeding mothers showed that small

#### quantities of Diprofol 2% are excreted in human milk. Women should therefore not breast-feed for

24 hours after administration of Diprofol 2%. Milk produced during this period should be discarded. 4.7 Effects on ability to drive and use machines Diprofol 2% has moderate influence on the ability to drive and use machines. Patients should be advised that performance at skilled tasks, such as

driving and operating machinery, may be impaired

Diprofol 2% induced impairment is not generally

detectable beyond 12 hours (see Section 4.4).

for some time after general anaesthesia.

4.8 Undesirable effects General

## Induction and maintenance of anaesthesia or sedation is generally smooth with minimal evidence

of excitation. The most commonly reported ADRs are pharmacologically predictable side effects of an anaesthetic/sedative agent, such as hypotension. The nature, severity and incidence of adverse events observed in patients receiving Diprofol 2% may be related to the condition of the recipients and the operative or therapeutic procedures being undertaken. The following definitions of frequencies are used: Very common (≥1/10), common (≥1/100 to <1/10), uncommon ( $\geq 1/1,000$  to <1/100), rare ( $\geq 1/10,000$ 

to <1/1,000), very rare (<1/10,000) and not known (cannot be estimated from the available data). Table of Adverse Drug Reactions

System Organ Class	Frequency	Undesirable Effects
Immune system disorders	Very rare	Anaphylaxis – may include angioedema, bronchospasm, erythema and hypotension
Metabolism and nutrition disorders	Frequency not known (9)	Metabolic acidosis <sup>(5)</sup> , Hyperkalaemia <sup>(5)</sup> , Hyperlipidaemia <sup>(5)</sup>
Psychiatric disorders	Not known (9)	Euphoric mood. Drug abuse and drug dependence <sup>(8)</sup>
Nervous system disorders	Common	Headache during recovery phase
	Rare	Epileptiform movements, including convulsions and opisthotonus during induction, maintenance and recovery
	Very rare	Postoperative unconsciousness
	Not known (9)	Involuntary movements
Cardiac disorders	Common	Bradycardia (1)
	Very rare	Pulmonary oedema
	Not known (9)	Cardiac arrhythmia <sup>(5)</sup> , cardiac failure <sup>(5),(7)</sup>
Vascular disorders	Common	Hypotension (2)
uisoruers	Uncommon	Thrombosis and phlebitis
Respiratory, thoracic and mediastinal disorders  Gastrointestinal disorders	Common	Transient apnoea during induction
	Not known (9)	Respiratory depression (dose dependent)
	Common	Nausea and vomiting during recovery phase
	Very rare	Pancreatitis
Hepatobiliary disorders	Not known (9)	Hepatomegaly <sup>(5)</sup>
Musculoskeletal and connective tissue disorders	Not known (9)	Rhabdomyolysis (3),(5)
Renal and urinary disorders	Very rare	Discolouration of urine following prolonged administration
	Not known (9)	Renal failure (5)
Reproductive system and breast disorders	Very rare	Sexual disinhibition
	Not known	Priapism
General disorders and administration site conditions	Very common	Local pain on induction (4)
	Very rare	Tissue necrosis (10) following accidental extravascular administration
	Not known (9)	Local pain, swelling, following accidental extravascular administration
Investigations	Not known (9)	Brugada type ECG (5),(6)
Injury, poisoning and procedural complications	Very rare	Postoperative fever
1. Serious bradyor reports of pro 2. Occasionally, h fluids and red. 3. Very rare report Diprofol has be ICU sedation. 4. May be minimized by ti 5. Combinations o Syndrome", ma	gression to as iypotension may action of the ac s of rhabdomyoly en given at dose zed by using the fossa. With Dipr he co-administra if these events, iy be seen in se	. There have been isolated ystole.  require use of intravenous lministration rate of Diprofol. sis have been received where es greater than 4 mg/kg/hr for elarger veins of the forearm of ol 1% local pain can also be atton of lidocaine. reported as "Propofol Infusion priously ill patients who often elevents,

outcome) in adults. The cardiac failure in such cases was usually unresponsive to inotropic supportive treatment. Abuse of and drug dependence on propofol, predominantly by healthcare professionals.

Not known as it cannot be estimated from the available clinical

been reported where tissue viability has

6. Brugada-type ECG- elevated ST- segment and coved T-wave Rapidly progressive cardiac failure (in some cases with fatal

see section 4.4

trial data. 10 Necrosis has

been impaired.

Dystonia/dyskinesia have been reported.

Local The local pain which may occur during the induction phase can be minimized by the use of the larger veins of the forearm and antecubital fossa.

Thrombosis and phlebitis are rare. Accidental clinical extravasation and animal studies showed

minimal tissue reaction. Intra-arterial injection in animals did not induce local tissue effects. Reporting of suspected adverse reactions

Reporting suspected adverse reactions after authorization of the medicinal product is important. It allows continued monitoring of the benefit/risk

balance of the medicinal product.

#### Any suspected adverse events should be reported to the Ministry of Health according to the National Regulation by using an online form: https://sideeffects.health.gov.il

4.9 Overdose Accidental overdosage is likely to cause cardiorespiratory depression. Respiratory depression should be treated by artificial ventilation with oxygen. Cardiovascular depression would require lowering of the patient's head and, if severe, use of plasma expanders and pressor agents. 5. Pharmacological properties

general

#### Pharmacotherapeutic group: Other anaesthetics ATC code: N01AX10

5.1 Pharmacodynamic properties

Mechanism of action Propofol (2,6-diisopropylphenol) is a short-acting general anaesthetic agent with a rapid onset of action of approximately 30 seconds. Recovery

from anaesthesia is usually rapid. The mechanism of action, like all general anaesthetics, is poorly

understood. However, propofol is thought to produce its sedative/anaesthetic effects by the positive modulation of the inhibitory function of the

neurotransmitter GABA through the ligand-gated GABA<sub>A</sub> receptors. Pharmacodynamic effects In general, falls in mean arterial blood pressure and slight changes in heart rate are observed when Diprofol 2% is administered for induction and maintenance of anaesthesia. However, the

haemodynamic parameters normally remain relatively stable during maintenance and the incidence of untoward haemodynamic changes is Although ventilatory depression can occur following administration of Diprofol 2%, any effects are qualitatively similar to those of other intravenous

Diprofol 2% reduces cerebral blood flow, intracranial pressure and cerebral metabolism. The reduction in intracranial pressure is greater in patients with an elevated baseline intracranial pressure. Clinical efficacy and safety

anaesthetic agents and are readily manageable in

clear headed with a low incidence of headache and post-operative nausea and vomiting.

adrenocortical hormones.

clinical practice.

In general, there is less post-operative nausea and vomiting following anaesthesia with Diprofol 2% than following anaesthesia with inhalational agents. There is evidence that this may be related

to a reduced emetic potential of propofol. Diprofol 2%, at the concentrations likely to occur clinically, does not inhibit the synthesis of

Recovery from anaesthesia is usually rapid and

Teratology studies in rats and rabbits showed no teratogenic effects. reproductive toxicity (see section 5.3). Diprofol 2%

Paediatric population

## When Diprofol 2% is used to maintain anaesthesia,

efficacy.

blood concentrations asymptotically approach the steady-state value for the given administration rate. Distribution Propofol is extensively distributed and rapidly

Limited studies on the duration of propofol based

anaesthesia in children indicate safety and efficacy

is unchanged up to duration of 4 hours. Literature

evidence of use in children documents use for

prolonged procedures without changes in safety or

5.2 Pharmacokinetic properties

cleared from the body (total body clearance 1.5– 2 litres/minute).

#### <u>Elimination</u> The decline in propofol concentrations following

a bolus dose or following the termination of an infusion can be described by a three compartment open model with very rapid distribution (half-life 2-4 minutes), rapid elimination (half-life 30-60 minutes), and a slower final phase, representative of redistribution of propofol from poorly perfused tissue. Clearance occurs by metabolic processes, mainly in the liver where it is blood flow dependent, to form inactive conjugates of propofol and its corresponding quinol, which are excreted in urine.

After a single dose of 3 mg/kg intravenously, propofol clearance/kg body weight increased with age as follows: Median clearance was considerably lower in neonates <1 month old

(n=25) (20 ml/kg/min) compared to older children (n=36, age range 4 months-7 years). Additionally, inter-individual variability was considerable in neonates (range 3.7-78 ml/kg/min). Due to these limited trial data that indicates a large variability, no dose recommendations can be given for this Median propofol clearance in older aged children after a single 3 mg/kg bolus was 37.5 ml/min/kg (4–24 months) (n=8), 38.7 ml/min/kg (11–43 months)

(n=6), 48 ml/min/kg (1-3 years) (n=12), 28.2 ml/min/ kg (4-7 years) (n=10) as compared with 23.6 ml/min/ kg in adults (n=6). pharmacokinetics are linear over recommended range of infusion rates of Diprofol 2%.

5.3 Preclinical safety data Published studies in animals (including primates) at doses resulting in light to moderate anaesthesia

### demonstrate that the use of anaesthetic agents during the period of rapid brain growth or

synaptogenesis results in cell loss in the developing brain that can be associated with prolonged cognitive deficiencies. Based on comparisons across species, the window of vulnerability to these changes is believed to correlate with exposures in the third trimester through the first several months of life, but may extend out to approximately 3 years of age in humans. In neonatal primates, exposure to 3 hours of an anaesthetic regimen that produced a light surgical plane of anaesthesia did not increase neuronal cell loss, however, treatment regimens of 5 hours

or longer increased neuronal cell loss. The clinical significance of these nonclinical findings is not known, and healthcare providers should balance the benefits of appropriate anaesthesia in young children less than 3 years of age and pregnant women who require procedures against the potential risks suggested by the preclinical data. Propofol is a drug on which extensive clinical experience has been obtained. All relevant information for the prescriber is provided elsewhere in this document.

6. Pharmaceutical particulars 6.1 List of excipients Soybean oil, glycerol, egg lecithin, oleic acid, sodium hydroxide and water for injection.

## 6.2 Incompatibilities

## Diprofol 2% should not be mixed prior to

administration with injections or infusion fluids. However, Diprofol 2% may be administered via a Y-piece connector close to the injection site with the

products mentioned in section 4.2. The neuromuscular blocking agent, atracurium,

should not be given through the same intravenous line as Diprofol 2% without prior flushing. 6.3 Shelf life Shelf life of the product as packaged for sale The expiry date of the product is indicated on the

6.4 Special precautions for storage Protect from light.

In use precautions:

packaging materials. Shelf life after dilution

Store below 25°C. Do not freeze. Vials that their contents have been frozen can no

### longer be used. 6.5 Nature and contents of container Diprofol 20 mg/ml: Glass vials of 50 ml

Diprofol 2% should not be diluted.

6.6 Special precautions for disposal and other handling

Vials must be shaken before use. Any portion of the contents remaining after use should be discarded.

#### If two layers remain visible after shaking, the product should not be used. Additional precautions:

Diprofol 2% contains no antimicrobial preservatives and supports growth of micro-organisms. Asepsis must be maintained for both Diprofol 2% and infusion equipment throughout the infusion period. Any drugs

or fluids added to the Diprofol 2% infusion line must be administered close to the cannula site. Diprofol 2% must not be administered via a microbiological filter. Diprofol 2% and any syringe containing Diprofol 2%

are for single use in an individual patient. For use in long-term maintenance of anaesthesia or sedation in intensive care it is recommended that the infusion line and reservoir of Diprofol 2% be discarded and replaced at regular intervals. 7. Manufacturer

Synthon Hispania S.L., Barcelona, Spain

14 Hakitor St., Haifa Bay 2624761 9. Registration Number

## 125.15.30376.00 Revised in June 2020.

8. License Holder Taro International Ltd.,