1) NAME OF THE MEDICINAL PRODUCT: VERDYE

2) QUALITATIVE AND QUANTITATIVE COMPOSITION: Each vial contains 25 mg powder of Indocyanine Green (as Monosodium Salt), to be reconstituted with 5 ml of water for injection.

1 ml of the reconstituted solution for injection contains 5 mg Indocyanine Green.

For full list of excipients see section 6.1.

3) PHARMACEUTICAL FORM: Powder for solution for injection. Dark-green powder.

4) CLINICAL PARTICULARS

4.1) Therapeutic indications: This medicinal product is for diagnostic use only.

In cardiac, circulatory and micro-circulatory diagnostics:
- measurement of circulation times
- measurement of cardiac output and stroke volume
- measurement of end-systolic ventricular volume
- measurement of intrathoracic blood volume
- measurement of circulating blood volume
- measurement of partial volumes
- measurement of organ perfusion - measurement of intra-and extracardiac shunts
- measurement of intracardiac valvular insufficiency
- measurement of peripheral perfusion (e.g. perfusion of extremities, eyes)

In liver function diagnostics:
- measurement of excretory function of the liver
- determination of partial functions of the liver

4.2) Posology and method of administration: Before administration the powder must be reconstituted with water for injection. The reconstituted solution is clear and free from visible particles, see section 6.6.

Diagnostic procedures with VERDYE should be performed under the supervision of a physician. VERDYE is intended for intravenous injection via an injection needle, a central or peripheral catheter or cardiac catheter.

The administration and site of VERDYE are of critical importance for the quality of the measurements. In principle, for obtaining optimal quality first pass indicator dilution curves, the injection should be as close as possible to the vascular bed, organ or tissue of interest.

On peripheral injection the injection should be made immediately after application of tourniquet and the arm should be raised after release of tourniquet. This ensures rapid transport of the dye from the site of injection and peripheral injection is then practically equivalent to central venous injection.

Dosage:

Single dose per measurement in adults, elderly, children:

Cardiac, circulatory, micro-circulatory and tissue perfusion diagnostics as well as cerebral blood flow: 0.1 to 0.3 mg/kg body weight as bolus injection.

Liver function diagnostics: 0.25 – 0.5 mg/kg body weight as bolus injection

Ophthalmic angiography: 0.1 to 0.3 mg/kg body weight as bolus injection

Total daily dose: Adults, elderly, adolescents 11-18 years:
The total daily dose of VERDYE should be kept below 5 mg/kg body weight.
Children 2 – 11 years: The total daily dose should be kept below 2.5 mg/kg body weight.
Children 0 - 2 years: The total daily dose should be kept below 1.25 mg/kg body weight.

Methods of measurement: The absorption and emission maximum of indocyanine green are both in the near infrared range, the absorption maximum at 800 nm and the emission maximum for fluorescence measurement at 830 nm.

In vitro-tests indocyanine green remains stable in human serum for several days.

Dissolved in water, indocyanine green shows no detectable decomposition at least for a few hours.

Measurement of cardiac, circulatory, and cerebral blood flow and liver function: Areas under the first pass curve, transit time, half-life, plasma disappearance rate and retention rate of VERDYE can be determined

a. non-invasively by pulse dye densitometry or near infrared spectroscopy
b. invasively by fiberoptic probes/catheters in suitable vessels
c. conventionally by determination of the concentration either by continuous withdrawal of heparinised blood through a cuvette densitometer or by collection of blood samples and measurement of the plasma concentration in a photometer.

Evaluation of fundus perfusion in ophthalmic angiography: The perfusion of the fundus of the eye can be determined and quantified by ophthalmic fluorescence angiography.

Measurement of tissue perfusion: Tissue perfusion of the superficial tissue layers can be made visible and quantified by near infrared fluorescence video angiography.

4.3) Contraindications:

VERDYE is contraindicated for safety reasons in:

• patients with hypersensitivity to indocyanine green or to sodium iodide unless special precautions are taken,
• patients with hypersensitivity to iodine,
• patients with hyper-thyroidism, patients with autonomic thyroid adenomas
• as in-vitro experiments have shown that indocyanine green displaces bilirubin from its protein binding, VERDYE should not be used in premature infants or neonates in whom an exchange transfusion is indicated due to of hyperbilirubinemia,
• if injection of VERDYE was poorly tolerated in the past it must not be used again, since severe anaphylactic reactions might occur.
4.4) Special warnings and precautions for use:

- Since severe anaphylactic reactions might occur after application of VERDYE, it must only be applied under supervision of a physician.
- Due to an increased incidence of adverse reactions in patients with severe renal insufficiency, VERDYE must only be applied after a careful benefit/risk assessment.
- Heparin preparations containing sodium bisulphite reduce the absorption peak of indocyanine green in plasma and blood and, therefore, should not be used as an anticoagulant for the collection of samples for analysis.
- Indocyanine green is stable in plasma and whole blood so that samples obtained in discontinuous sampling techniques may be read hours later.
- Sterile techniques have to be used in handling the dye solution.

- The iodine content of VERDYE can interfere with thyroid tests performed before or after administration of the dye. Therefore, radio-active iodine uptake studies should not be performed for at least a week following the use of VERDYE.

4.5) Interaction with other medicinal products and other forms of interaction:

Regarding incompatibilities with solvents for dilution see section 6.6.

The clearance of indocyanine green may be altered by medicinal products that interfere with liver function. Probencid and some of its metabolites may be secreted into the bile, and may depress the biliary secretion of indocyanine green which may result in an impaired indocyanine green liver function test.

Concomitant use of certain medicinal products and injectables can alter the absorption. The absorption is reduced by injectables containing sodium bisulphite (particularly in combination with heparin). The following gives an overview of interaction with other medicinal products:

Medicinal products and substances that can reduce absorption:

- anticonvulsants
- bisulphite compounds
- haloperidol
- heroin
- meperidine
- metamizol
- methadone
- morphium
- nitrofurantoin
- opium alkaloids
- phenobarbital
- phenylbutazone

Medicinal products and substances that can increase absorption:

- cyclopropane
- probenecid
- rifamycin

4.6) Fertility, pregnancy and lactation:

**Pregnancy:** Data on a limited number (242) of exposed pregnancies indicate no adverse effects of Indocyanine green on pregnancy or on the health of the foetus / newborn child. To date, no other epidemiological data are available. No studies for reproduction, teratogenicity, or carcinogenic properties in animals are available. The potential risk for humans is unknown.

Caution should be exercised when prescribing to pregnant women. Repeated applications on one day have to be avoided.

**Lactation:** It is not known whether this medicinal product is excreted in human milk. Because many medicinal products are excreted in human milk, caution should be exercised when indocyanine green is administered to a nursing woman.

4.7) Effects on ability to drive and use machines:

- No studies on the effects on the ability to drive and use machines have been performed.

4.8) Undesirable effects:

- Anaphylactic or urticarial reactions have been reported in patients with or without history of allergy to iodides.
- Also in very rare cases coronary artery spasm has been described.
- It is known that injection of indocyanine green preparations can in very rare cases cause nausea and anaphylactoid or anaphylactic reactions (<1/10000).
- In patients with terminal renal insufficiency the possibility that an anaphylactic reaction occurs seems to be increased. Symptoms which should be mentioned are: unrest, feeling of warmth, pruritus, urticaria, acceleration of heart rate, fall in blood pressure and shortness of breath, bronchospasm, flush, cardiac arrest, laryngospasm, facial oedema, nausea. Together with the anaphylactoid reaction, hyperoxygenophilia may occur.
- If, contrary to expectations, symptoms of anaphylaxis do occur, the following immediate measures should be taken:
  - stop further administration of VERDYE leave injection catheter or cannula in the vein
  - keep airways free
  - inject 100-300mg hydrocortisone or a similar preparation by rapid intravenous injection
  - substitute volume with isotonic electrolyte solution
  - give oxygen, monitor circulation
  - slowly administer antihistamines intravenously

The following additional measures are indicated in cases of anaphylactic shock:

- place patient in recumbent position with legs raised
- rapidly substitute volume with e.g. isotonic electrolyte solution (pressure infusion), plasma expanders.
- immediately administer 0.1–0.5mg adrenaline diluted to 10ml with 0.9% saline intravenously (repeat after 10 minutes if necessary).

Urticarial reactions of the skin occurred very rarely (<1/10000).

Two anaphylactic deaths have been reported following indocyanine green administration during cardiac catheterization. One of these was in a patient with a history of penicillin and sulfa allergy. Deaths due to anaphylaxis occurred in less than 1/330000 (estimate) including single reports. 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 effects should be reported to the Ministry of Health according to the National Regulation by using an online form http://forms.gov.il/globaldata/getsequence/getsequence.aspx?formType=AdversEffectMedic@moh.gov.il

4.9) Overdose:

- Up to now no case of medicinal product overdose or laboratory findings accompanying overdose of VERDYE has been reported.

5) PHARMACOLOGICAL PROPERTIES

5.1) Pharmacodynamic properties:

Pharmacotherapeutic group: Other diagnostic agents ; ATC code: V04CX

The active substance in VERDYE is 2-{7-[1,1-dimethyl-3-(4-sulfobutyl)-benz[e]indolin-2-ylidene]-1,3,5-heptatrienyl}-1,1-dimethyl-3-(sulfobutyl)-1H-benz[e]-indolium hydroxide, inner salt, sodium salt.

The molecular formula is C43H47N2NaO6S2. The molecular weight is 774.96 daltons.

Indocyanine green has a sharply defined spectral peak absorption of nearinfrared light at 800 nm in blood plasma or blood. This is the same wavelength with which the optical density of oxygenated haemoglobin in blood approximately equals that of reduced haemoglobin. Therefore, this coincident light absorption makes it possible to measure indocyanine green concentrations in blood, plasma and serum in terms of its optical density at 800 nm, independent of variations in oxygen saturation level.

Indocyanine green permits recording of the indicator-dilution curves for both diagnostic and research purposes.

Indocyanine green does not cross the placental barrier. The volume of distribution corresponds to the blood volume.
After oral or rectal administration indocyanine green is not absorbed from the gut.

**Protein-binding:** Following intravenous injection, indocyanine green is rapidly bound to plasma proteins, of which beta-apolipoprotein B is the principle carrier (95 %).

**Metabolism:** Indocyanine green is not metabolised.

**Elimination:** Plasma disappearance is biphasic, showing an initial elimination half-life t1/2 of 3-4 min and a secondary phase with a dose-dependent t1/2 of approximately 60-80 min.

Indocyanine green is taken up from the plasma almost exclusively by the hepatic parenchymal cells with a maximum rate of uptake (transport maximum: Tm of about 0.1 mg/minute/kg) and is secreted unmetabolized and unconjugated entirely into the bile. The concentration maximum in bile is reached after about 1/2–2 hours depending on the amount injected.

After biliary obstruction, the dye appears in the hepatic lymph, independently of the bile, suggesting that the biliary mucosa is sufficiently intact to prevent diffusion of the dye, though allowing diffusion of bilirubin.

As indocyanine green is not reabsorbed in the intestine there is no enterohepatic circulation.

**5.3) Preclinical safety data:** Acute toxicity: the LD50 after single IV dose was 87 mg/kg in rats, 60 mg/kg in mice, and between 50 mg/kg and 80 mg/kg in rabbits. After dissolution in water for injection and administration by intraperitoneal injection in mice the LD50 was found to be 650 mg/kg body weight. No macroscopic or histopathological changes were observed.

Genetic toxicity: indocyanine green was not found to be mutagenic in the tests performed (Ames test, gene mutation assay - thymidin kinase locus/TK+-/- in mouse lymphoma L5178Y cells, chromosome aberration test in Chinese hamster V79 cells).

No studies for reproduction, teratogenicity, or carcinogenic properties in animals are available but decades of experience in humans have not revealed any incidence of these properties.

6) PHARMACEUTICAL PARTICULARS

6.1. List of excipients: The medicinal product does not contain excipients.

6.2) Incompatibilities: This medicinal product must not be diluted with solutions containing salts (saline, Ringer's solution etc.) as this can lead to precipitation of the dye. This medicinal product must not be mixed with other medicinal products except those mentioned in section 6.6.

6.3) Shelf life: The expiry date of the product is indicated on the label and packaging. Do not use after the expiry date. After reconstitution, the solution should be used immediately, protected from light.

6.4) Special precautions for storage: Do not store above 25°C. Keep vials in the outer carton in order to protect from light.


Carton packs with 1, 2, 3, 4 or 5 vials, each with a content of 25 mg powder for solution for injection. Not all pack sizes may be marketed.

6.6) Special precautions for disposal: This medicinal product should be reconstituted immediately prior to use. This medicinal product is reconstituted by addition of 5 ml water for injection to the vial containing 25 mg of active substance, giving a dark-green solution for injection with a concentration of 5 mg/ml (0.5% w/v).

If an incompatibility is noted in the form of unclear solution then the reconstituted solution should be discarded. Visually inspect the reconstituted solution. Only use clear solutions free from visible particles. This medicinal product is for single use only.

7) Israeli Drug Registration Number(s): 129.12.29280.00 and 129.12.29280.01

8) Manufacturer: Diagnostic Green GmbH, Aschheim-Dornach, Germany [in cooperation with Patheon Italia S.p.A., Monza (MI), Italy and/or Marien-Apotheke, Prien / Chiemsee, Germany].

9) Israeli Marketing Authorization Holder: Concept for Pharmacy Ltd., VAT # 511089997, P.O.B. 2105, Kfar Saba 4464316, Israel.

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