radiopaque contrast medium for oral or rectal administration only. Each mL contains 660 mg diatrizoate meglumine and 100 mg diatrizoate sodium; pH has been adjusted to 6.0 to 7.6 with sodium hydroxide. Each mL contains approximately 4.8 mg (0.21 mEq) sodium and 367 mg organically bound iodine. Inactive ingredients: edetate disodium, flavor, polysorbate 80, purified water, saccharin sodium, simethicone, and sodium citrate.

Diatrizoate meglumine is designated chemically as 1-deoxy-1-(methylamino)-D-glucitol 3,5-diacetamido-2,4,6-triiodo-benzoate (salt); diatrizoate sodium is monosodium 3,5-diacetamido-2,4,6-triiodobenzoate. Structural formulas:

\[
\text{dariatzoate meglumine} \quad C_{11}H_{8}I_{3}N_{2}NaO_{4} \\
\text{dariatzoate sodium} \quad C_{11}H_{9}N_{2}O_{4}C_{2}H_{4}N_{2}O_{5} \quad \text{MW 809.13}
\]

Organically Bound Iodine: 47.1% CAS-131-49-7

Organically Bound Iodine: 59.9% CAS-737-31-5

**CLINICAL PHARMACOLOGY**

The most important characteristic of contrast media is the iodine content. The relatively high atomic weight of iodine contributes sufficient radiodensity for radiographic contrast with surrounding tissues. Diagnostic enteral radiopaque agents have few known pharmacological effects. Diatrizoate meglumine and diatrizoate sodium exert a mild laxative effect attributable to their high osmolarity. Diatrizoate meglumine and diatrizoate sodium are sparingly absorbed from the intact gastrointestinal tract, and therefore permit gastrointestinal examination and delineation after oral or rectal administration. Oral administration is used for radiographic examination of the esophagus, stomach and proximal small intestine. Rectal administration is used for examination of the colon; however, visualization of the distal small bowel is generally unsatisfactory, since the hyperosmoticity of the medium causes intraluminal diffusion of fluid and subsequent dilution of the medium. Enough absorption from the gastrointestinal tract to permit incidental irradiation of the urinary tract may be expected (see Precautions); this should also be considered when thyroid testing is being contemplated, since iodine-mediated thyrotoxic effects may occur (see Precautions).

**INDICATIONS AND USAGE**

Gastrografin (Diatrizoate Meglumine and Diatrizoate Sodium Solution) is indicated for radiographic examination of segments of the gastrointestinal tract (esophagus, stomach, proximal small intestine, and colon). The preparation is particularly indicated when a more viscous agent such as barium sulfate, which is not water-soluble, is not feasible or is potentially dangerous. Gastrografin may also be used as an adjunct to contrast enhancement in computed tomography of the torso (body imaging); the preparation is indicated, in conjunction with intravenous administration of a radiopaque contrast agent, when unenhanced imaging may not provide sufficient definition to permit visualization of normal loops of bowel from adjacent organs or areas of suspected pathology.

**CONTRAINDICATIONS**

There are no absolute contraindications to the use of Gastrografin (Diatrizoate Meglumine and Diatrizoate Sodium Solution) (see Precautions, General).

**WARNINGS**

A 1 in 4.6 (1:4.6) dilution of Gastrografin yields an approximately isotonic 16.5 percent diatrizoate salts solution; less dilute solutions are hypertonic and may lead to intraluminal movement of fluid with resulting hypovolemia. In young or debilitated children and in elderly cachectic persons, the loss of plasma fluid may be sufficient to cause shock. See DOSAGE AND ADMINISTRATION for recommended dilutions that must be used for infants and young children (under 10 kg) and for dehydrated or debilitated patients. Electrolyte disturbances must be corrected prior to using hypertonic solutions. In debilitated patients and in patients with electrolyte imbalances, postprocedural monitoring of hydration, serum osmolality, electrolytes and dehydration status is essential. In pediatrics or severely debilitated patients, the maintenance of an open intravenous fluid line should be considered. In patients with severe circulatory insufficiency, the possibility of accidental aspiration into the atelectatic lung or into a tracheo-esophageal fistula should be avoided.

The possibility of accidental aspiration into the trachea or into a tracheo-esophageal fistula following ingestion or instillation, could result in serious pulmonary complications (e.g., pulmonary edema or pneumonitis) even though the patient's condition may be promptly expected. Bronchial entry of any orally administered contrast medium causes a pulmonary effusion. Therefore, pulmonary entry by aspiration and use in patients with esophagotracheal fistula should be avoided.

**PRECAUTIONS**

**General**

Diagnostic procedures which involve the use of radiopaque contrast agents should be carried out under the direction of personnel with the prerequisite training and with a thorough knowledge of the particular procedure to be performed. Appropriate facilities should be available for coping with any complication of administration, as well as for treatment of reaction to the contrast medium.

The possibility of a reaction should always be considered. Patients at increased risk include those with a history of a previous reaction to a contrast medium, patients with a known sensitivity to iodine per se, and patients with a known clinical hypersensitivity (bronchial asthma, hay fever, and food allergies). A positive history of allergies or hypersensitivity does not arbitrarily contraindicate the use of a contrast agent when a diagnostic procedure is thought essential, but caution should be exercised (see Adverse Reactions, and Precautions, Information for the Patient).

**Rectal administration of undiluted Gastrografin (Diatrizoate Meglumine and Diatrizoate Sodium Solution) in any patient, particularly with large doses and/or in those with overstentention, has been reported to be associated with mucosal irritation.

Cases of hyperthyroidism have been reported with the use of oral contrast media. Some of these patients reportedly had multinodular goiters which may have been responsible for the increases in hormone synthesis in response to excess iodine. Administration of an intravascular iodinated radiopaque diagnostic agent to a hyperthyroid patient precipitated thyroid storm; a similar situation could follow administration of oral preparations of ioddides. Therefore, caution should be exercised when administering enteral gastrointestinal radiopaque agents to hyperthyroid and euthyroid goiterous patients.

Consideration should be given to the potential for precipitation of water-soluble iodine as crystals under condition that may promote hyperacidity (i.e., fasting, emotional upset, or stress). Harmful effects directly attributable to precipitate formation have not been reported. However, the possibility of interpreting the precipitate radiologically as an anatomical abnormality (e.g., dilated loop of the small intestine) or injury, should be kept in mind.

**Information for the Patient**

Patients should receive the following information and instructions:

1. This drug has been prescribed to perform an x-ray of the gastrointestinal tract.
2. Inform the physician if pregnant or if allergic to iodine, any foods, or x-ray materials.
3. The iodine in diatrizoate salts may interfere with some thyroid tests if these are needed in the future. Inform the attending physician at that time about this gastrointestinal study.
4. This drug may cause abdominal cramping, nausea, vomiting, diarrhea, skin rashes, itching, heartburn, dizziness, or headache in some patients, but most reactions are mild and pass quickly.

**Drug/Laboratory Test Interactions**

**Thyroid Function Tests**

The results of protein bound iodine (PBI) and radioactive iodine uptake studies, which depend on iodine estimations, will not accurately reflect thyroid function for six months, and possibly as long as one year, following the administration of diagnostic enteral radiopaque media. Thyroid function tests, if indicated, generally should be performed prior to the administration of any iodinated agent. However, thyroid function can be evaluated after use of these agents by using T₃ uptake and total or free thyroxine (T₄) assays which are not dependent on iodine estimations.

**Pancreatic Tests**

Small quantities of contrast medium in the intestinal tract may cause false low trypsin values when determined spectrophotometrically. Therefore, duodenal instillation should not precede pancreatic function tests involving spectrophotometric trypsin assays.

Any test which might be affected by contrast media should be performed prior to administration of the contrast medium.

**Carcinogenesis, Mutagenesis, Impairment of Fertility**

Long-term studies in animals have not been performed to evaluate carcinogenic or mutagenic potential or possible impairment of fertility in males or females.
Pregnancy Category B
When administered intravenously, diatrizoate salts cross the placenta and are evenly distributed in fetal tissues. No teratogenic effects attributable to diatrizoate meglumine or diatrizoate sodium have been observed in teratology studies performed in animals. There are, however, no adequate and well-controlled studies in pregnant women. Because small amounts of these agents may be absorbed, and animal teratology studies are not always predictive of human response, these agents should be used during pregnancy only when clearly needed.

Procedures including radiation involve a certain risk related to the exposure of the fetus.

Nursing Mothers
Diatrizoate meglumine is excreted in breast milk following intravascular administration. Because small amounts of enteral gastrointestinal radiopaque agents may be absorbed following oral or rectal administration, caution should be exercised when they are administered to a nursing woman.

Pediatric Use
See WARNINGS, and PRECAUTIONS, General.

Local injury to the colonic mucosa, particularly in the presence of underlying disease which interferes with intestinal viability, has been reported in cases where recommended doses and dilutions (see DOSAGE AND ADMINISTRATION) were not used; when extemporaneous dosage is elected, the polysorbate 80 level in the dose may be a contributing factor to injury.

ADVERSE REACTIONS
Most adverse reactions to enteral diagnostic radiopaque agents are mild and transitory. Nausea, vomiting and/or diarrhea, urticaria with erythema, hypoxia, acute dyspnea, tachyarrhythmia, and anaphylaxis have occurred following ingestion of the contrast medium, particularly when high concentrations of large volumes of solution are administered. Severe changes in serum osmolality and electrolyte concentrations may produce shock-like states (see WARNINGS). It should be kept in mind that serious or anaphylactoid reactions that may occur with intravascular administration of radiopaque contrast agents are theoretically possible following administration by other routes.

OVERDOSAGE
See WARNINGS regarding potential hypovolemia, hypotension, or shock. The maintenance of an open intravenous fluid line for rehydration may be advisable. See DOSAGE AND ADMINISTRATION for appropriate doses and dilutions. Treatment of an overdose should be directed toward the support of all vital functions, and prompt institution of symptomatic therapy.

DOSE AND ADMINISTRATION
General
This medium is not to be used for the preparation of solutions for parenteral administration. Oral or rectal administration only.

The routine preparatory measures employed for barium studies are also appropriate for this agent. For pediatric and severely cachectic patients the maintenance of an intravenous fluid line may be advisable.

Radiographic Examination of Segments of the Gastrointestinal Tract
Oral Administration: Adult oral dosage may range from 30 to 90 mL (11 to 33 g iodine), depending on the nature of the examination and the size of the patient. For infants and children less than 5 years of age, 30 mL (11 g iodine) are usually adequate; for children 5 to 10 years of age, the suggested dose is 60 mL (22 g iodine). These pediatric doses may be diluted 1:1, if desired, with water, carbonated beverage, milk, or mineral oil. When used in infants, the solution may be given in a nursing bottle. Pediatric doses may also be used in dehydrated and/or debilitated adult patients. A 1:1 dilution is also recommended when the contrast medium is used in elderly cachectic individuals.

For very young (under 10 kg) and debilitated children the dose should be diluted: 1 part Gastrografin (Diatrizoate Meglumine and Diatrizoate Sodium Solution) in 3 parts water is recommended. For Enemas or Enterostomy Instillations: Gastrografin should be diluted when it is used for enemas and enterostomy instillations. When used as an enema, the suggested dilution for adults is 240 mL (88 g iodine) in 1,000 mL of tap water. For children under 5 years of age, a 1:5 dilution in tap water is suggested; for children over 5 years of age, 90 mL (33 g iodine) in 500 mL of tap water is a suitable dilution.

Tomography (Body Imaging)
A usual adult dose is 240 mL of a dilute Gastrografin solution prepared by diluting 25 mL (9.17 g iodine) to one liter with tap water. Less dilute solutions (up to 77 mL (28.26 g iodine) diluted to one liter with tap water] may be used when indicated. The dose is administered orally about 15 to 30 minutes prior to imaging in order to permit the contrast medium to reach the pelvic loops. HOW SUPPLIED
Gastrografin (Diatrizoate Meglumine and Diatrizoate Sodium Solution USP) is available in packages of: Twenty-four 30 mL single dose bottles (NDC 0270-0445-35). Twelve 120 mL bottles (NDC 0270-0445-40).

Storage
Protect from light. Store at 20-25°C (68-77°F) [See USP]; avoid excessive heat.
| NOM DU PRODUIT / PRODUCT NAME: **GASTROGRAFIN** |
| NUMÉRO PRODUIT / PRODUCT NUMBER: **CW1336-2** |
| NUMÉRO PHARMACODE / PHARMACODE NUMBER: **1149** |

| SPECIFICATIONS |
| COULEURS / COLORS: **BLACK** |
| TYPE DE PAPIER / PAPER TYPE: **OFFSET** |
| POIDS DU PAPIER / PAPER WEIGHT: **80 M** |
| DIMENSIONS (MM): À PLAT / FLAT: **381 x 254** |
| PLIÉ / FOLDED: **76,2 x 31,8** |
| POINTS DE COLLE / GLUE SPOTS: **N/A** |
| CONFORMITÉ DU PLIAGE / CONFORMITY OF FOLDING: |

| APPROBATION / APPROVAL |
| E-Z-E-M |
| APP.: DATE: |
| APP.: DATE: |
| APP.: DATE: |
| APP.: DATE: |