

INSTRUCTIONS FOR USE (ENGLISH).....2



# **ENGLISH**

### DESCRIPTION

EmboGod<sup>ath</sup> Microspheres are biocompatible, hydrophilic, non-absorbable, precisely calibrated acrylic polymer microspheres impregnated with portone gelatin and are available in a large range of sizes and concentrations. EmboGod Microspheres are coloured to facilitate visualization during handling. These spheres are designed to offer controlled, targeted embolization.

#### IMPLANTABLE DEVICE MATERIALS TABLE

Material	Duration of exposure	Level of patient exposure (maximum solid content by syringe)
Trisacryl Copolymer	Long-Term (> 30days)	159 ± 6 mg
Gelatin	Long-Term (> 30days)	23 ± 1 mg
Colloidal gold	Long-Term (> 30days)	1.5 mg

#### HOW SUPPLIED

20-mL prefilled syringe with a standard Luer-lock tip, individually packaged in blister tray sealed by a Tyvek® peel- away lid. Plastic screw cap and plunger. Elastomer three-skirt plunger joint.

Contents: 1 mL or 2 mL of microspheres in sterile, pyrogen-free, 0.9% NaCl solution.

# INTENDED USE / INDICATIONS

EmboGold Microspheres are designed to occlude blood vessels, for therapeutic or preoperative purposes, in the following procedures:

- Embolisation of hypervascular tumours and processes, including uterine fibroids, meningiomas, liver tumours.
- Embolisation of the prostate arteries for relief of symptoms related to Benign Prostatic Hyperplasia.
   Embolisation of arteriovenous malformations.
- Haemostatic embolisation.

 $40\text{-}120\,\mu\text{m}$  microspheres are more specifically designed for embolisation of meningiomas and liver tumours.

#### CLINICAL BENEFITS

Embolisation with EmboGold Microspheres is a minimally invasive treatment that is effective:

 For women with uterine fibroids for relief of related symptoms including heavy menstrual bleeding, pelvic pain or pressure, and/or urinary dysfunction, and for improvement of quality of life.

- For patients with hypervascular tumours, including liver tumours, for relief of related symptoms and for delay of the disease progression.
- For patients with meningioma, for reduction of intraoperative blood loss during resection procedure.
- For men with benign prostatic hyperplasia (BPH) for relief of related lower urinary tract symptoms (LUTS), such as urinary frequency, inability to urinate, incomplete emptying of bladder, difficulty starting urination, and straining to urinate or weak urine stream, and for improvement of quality of life.
- For patients with arteriovenous malformations for relief of related symptoms.
   For patients with haemorrhage for immediate and long-
- term bleeding control.

  For a copy of this device's current European Summary of Safety and Clinical Performance (SSCP), please on the European da-

For a copy of this device's current European Summary of Safety and Clinical Performance (SSCP), please go the European database on medical devices (Eudamed), where it is linked to the basic UDI-DI. https://ec.europa.eu/tools/eudamed. Basic UDI-DI: 088445048793EH.

Alternatively, download a copy of the SSCP from: https://www.merit.com/sscp

# MAGNETIC RESONANCE IMAGING

EmboGold Microspheres are made of tris-acryl polymer impregnated with porcine gelatin and colored with gold and magnetic resonance (MR) are compatible.

# CONTRAINDICATIONS

All indications

- Patients unable to tolerate vascular occlusion procedures.
- Vascular anatomy or blood flow precluding correct catheter
- Presence of arteries supplying the lesion not large enough to accept EmboGold Microspheres
- Presence of collateral vessel pathways potentially endangering normal territories during embolization
   Presence or likely onset of vasospasm.

placement or embolic agent injection.

- Vascular resistance peripheral to the feeding arteries precluding passage of EmboGold Microspheres into the lesion
   In large diameter arteriovenous shunts (i.e., where the blood
- does not pass through an arterial/capillary/venous transition but directly from an artery to a vein)
- High-flow arteriovenous shunts or with a diameter greater than the selected microspheres.
- Presence of severe atheromatous disease.
   Patients with known allergy to gelatin and/or gold.

 $40\text{--}120\,\mu m$  and  $100\text{--}300\,\mu m$  microspheres are not recommended for use in the bronchial circulation.

### **UFE Specific Contraindications**

- Pregnant women
  - Suspected pelvic inflammatory disease or any other active pelvic infection
- Any malignancy of the pelvic region
   Endometrial neoplasia or hyperplasia
- Presence of one or more submucosal fibroid(s) with more than
- 50% growth into the uterine cavity
   Presence of pedunculated serosal fibroid as the dominant fibroid(s)
- Fibroids with significant collateral feeding by vessels other than the uterine arteries

#### PAF SPECIFIC CONTRAINDICATIONS

- Active urinary tract infection or prostatitis
- Prostate cancer
   Bladder cancer
- Chronic renal failure
- Chronic renal failure
- Bladder atonia, neurogenic bladder disorder, or other neurological disorder impacting bladder function
- Bladder stones
- Urinary obstruction due to causes other than BPH, including urethral stricture

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Excessive vessel tortuosity or severe atherosclerosis

## **Neurological Specific Contraindications**

- Presence of patent extra-to-intracranial anastomoses or shunts Presence of end arteries leading directly to cranial nerves
- In any vasculature where EmboGold Microspheres could pass directly into the internal carotid artery, vertebral artery, intracranial vasculature or the above listed vessels

#### POTENTIAL COMPLICATIONS All indications

Vascular embolisation is a high-risk procedure. Complications may occur at any time during or after the procedure, and may include, but are not limited to, the following:

- Complications related to catheterization (e.g. haematoma at the site of entry, clot formation at the tip of the catheter and subsequent dislodgement, nerve and/or circulatory injuries which may result in leg injury, infection)
- Vessel or lesion rupture and haemorrhage
- Occlusion of vessels in healthy territories
- Paralysis resulting from untargeted embolization or ischemic
- iniury from adiacent tissue edema
- Stroke or cerebral infarction
- Ischaemia at an undesirable location, including ischaemic stroke, ischaemic infarction (including myocardial infarction), and tissue necrosis

- Blindness, hearing loss, loss of smell, and/or paralysis Capillary bed occlusion and tissue damage
- Death
- Undesirable reflux or passage of EmboGold Microspheres into normal arteries adjacent to the targeted lesion or through the lesion into other arteries or arterial beds, such as the internal
- carotid artery, pulmonary, or coronary circulations Pulmonary embolism due to arterial venous shunting
- Vasospasm
- Recanalization Foreign body reaction necessitating medical intervention.
- Infection necessitating medical intervention. Allergic reaction to medications (e.g. analgesics)
- Allergic reaction due to contrast media or embolic material Allergic response to gold
- Cutaneous irritations (e.g. rash), possibly delayed from the time of embolization
- Post-embolization syndrome, such as transient pain. nausea, vomiting, fever, possibly delayed from the time of embolization
- Transient hypertensive episode
- Additional information is found in the Warnings section.

# **UFE Specific Potential Complications**

The most frequently anticipated post procedure complications are abdominal pain, discomfort, fever and/or nausea,

- collectively known as "Post-embolization Syndrome." Some patients may also experience constipation. This is generally managed with prescription or over-the-counter medications.
- Premature ovarian failure (i.e., menopause)
- Amenorrhea
- Infection of the pelvic region Uterine/ovarian necrosis
- Phlehitis
- Deep vein thrombosis with or without pulmonary embolism Vaginal discharge
- Tissue passage, fibroid sloughing, or fibroid expulsion post UFE Post-LIFF intervention to remove perrotic fibroid tissue
- Vagal reaction
- Hysterectomy

# **PAE Specific Potential Complications**

- Non-targeted embolization of the rectum, bladder, scrotum, penis, or other areas
- The most frequent post-procedure complication includes "Post-PAE Syndrome", which includes nausea, vomiting, fever, pelvic pain, burning sensation, dysuria, and frequent or urgent urination
- Skin burn (radiation exposure) from prolonged fluoroscopy time
- Blood in urine, semen, or stool
- Bladder spasm

- Urinary tract infection Urinary retention
- Constipation
- Urethral obstruction

# **Neurological Specific Potential Complications**

- Ischemic stroke or ischemic infarction Neurological deficits, including cranial nerve palsies
- PRECAUTION

# All indications

- DO NOT LISE THIS PREFILLED SYRINGE TO DIRECTLY INJECT EMBOGOLD MICROSPHERES, THIS IS A "RESERVOIR" SYRINGE PLEASE REFER TO INSTRUCTIONS PARAGRAPH.
- EmboGold Microspheres must only be used by specialist physicians trained in vascular embolisation procedures. The size and quantity of microspheres must be carefully selected according to the lesion to be treated, entirely under the physician's responsibility. Only the physician can decide the most appropriate time to stop the injection of microspheres.
- Patients with known allergy to contrast medium may require corticosteroids prior to embolization.
- Additional evaluations or precautions may be necessary in managing periprocedural care for patients with the following conditions:
  - Bleeding diathesis or hypercoagulative state

- Immunocompromise.
- Do not use if blister tray, peel-away film, screw cap or syringe appear damaged.
- This is a disposable product. Discard opened syringes after use.
- For single patient use only Contents supplied sterile Never reuse, reprocess, or resterilize. Reuse, reprocessing or
- resterilization may compromise the structural integrity of the device and/ or lead to device failure, which in turn may result in patient injury, illness, or death. Reuse, reprocessing or resterilization may also create a risk of contamination of the device and/ or cause patient infection or cross infection including, but not limited to, the transmission of infectious disease(s) from one patient to another. Contamination of the device may lead to injury, illness or death of the patient. All procedures must be performed according to accepted asentic technique.
- any other application.

# **UFE Specific Precautions**

There is an increased chance of retro-migration of EmboGold Microspheres into unintended blood vessels as uterine artery flow diminishes. Embolization should be stopped when the vasculature surrounding the fibroid can no longer be visualized but before complete stasis in the uterine artery.

The syringe is intended for embolic use only. Do not use for

 UFE should only be performed by specialist physicians who have received appropriate training for treatment of uterine leiomyomata (fibroids).

#### Liver tumour Specific Precautions

There is no known incompatibility between EmboGold
Microspheres and chemotherapeutics used for the treatment
of liver tumours.

# PAE Specific Precautions

- The PAE procedure should only be performed by specialist physicians who have received appropriate training.
- Collateral circulation may be present and can dilate and supply adjacent arteries as resistance within the prostatic bed increases. Therefore, there is potential for severe complications with nontargeted embolization.
- There is an increased chance of retro-migration of EmboGold Microspheres into unintended blood vessels as prostatic artery flow diminishes. Embolization should be stopped when the vasculature surrounding the prostate can no longer be visualized but before complete stasis in the prostatic artery.

### **Haemostatic indication Specific Precautions**

 Embolization of the splenic artery may be associated with inferior yena cava thrombus.

#### WARNINGS

#### All indications

- EmboGold Microspheres contain gelatin of porcine origin, and, therefore, could cause an immune reaction in patients who are hyperesnikive to collagen or gelatin. Careful consideration should be given prior to using this product in patients who are suspected to be allergic to injections containing gelatin stabilizers.
- Studies have shown that EmboGold Microspheres do not form aggregates, and as a result, penetrate deeper into the vasculature as compared to similarly sized PVA particles. Care must be taken to choose larger sized EmboGold Microspheres when embolizing arteriovenous malformations with large shunts to avoid passage of the spheres into the pulmonary or coronary circulation.
- Some of the EmboGold Microspheres may be slightly outside
  of the range, so the physician should be sure to carefully
  select the size of EmboGold Microspheres according to the
  size of the target vessels at the desired level of occlusion in
  the vasculature and after consideration of the arteriovenous
  angiographic appearance. EmboGold Microspheres size should
  be selected to prevent passage from artery to vein.
   The roll or file FemboGold Microspheres (with the wishle)
- The color of the EmboGold Microspheres could be visible through the skin if injected into arteries feeding superficial tissues.

- Because of the significant complications of misembolisation, extreme caution should be used for any procedures involving the extracranial circulation encompassing the head and neck, and the physician should carefully weigh the potential benefits of using embolisation against the risks and potential complications of the procedure. These complications can include blindness, hearing loss, loss of smell, paralysis and death.
- Serious radiation induced skin injury may occur to the patient
  due to long periods of fluoroscopic exposure, large patient
  diameter, angled x-ray projections, and multiple image
  recording runs or radiographs. Refer to your facility's clinical
  protocol to ensure the proper radiation dose is applied for
  each specific type of procedure performed. Physicians should
  monitor patients that may be at risk.
- Onset of radiation-induced injury to the patient may be delayed. Patients should be counseled on potential radiation side effects and whom they should contact if they show symptoms.
- Pay careful attention for signs of mistargeted embolisation.
   During injection carefully monitor patient vital signs to include SaO2 (e.g. hypoxia, CNS changes). Consider terminating the procedure, investigating for possible shunting, or increasing microsphere size if any signs of mistargeting occur or patient symbotms develon.

Consider upsizing the microspheres if angiographic evidence of embolisation does not quickly appear evident during injection of the microspheres.

#### **UFE Specific Warnings**

#### Warnings About UFE and Pregnancy

- The effects of UFE on the ability to become pregnant and carry a fetus to term, and on the development of the fetus, have not been determined. Therefore, this procedure should only be performed on women who do not intend future pregnancy.
- Women who become pregnant following UFE may be at increased risk for postpartum haemorrhage, preterm delivery, cesarean delivery, and malpresentation.
   Devascularization of the uterine myometrium resulting from
- Devascularization of the uterine myometrium resulting fro UFE may theoretically put women who become pregnant following UFE at increased risk of uterine rupture.

# Other UFE Warnings

- When using EmboGold Microspheres for uterine fibroid embolization, do not use microspheres smaller than 500 microns.
- An appropriate gynecologic work-up should be performed on all patients presenting for embolization of uterine fibroids (e.g., gynecologic history, fibroid imaging, endometrial sampling to rule out carcinoma in patients with abnormal menstrual bleeding).

The diagnosis of uterine sarroma could be delayed by taking a nonsurgical approach (such as UFE) to treating fibroids. It is important to pay dose attention to warning signs for sarroma (e.g., rapid tumor growth, postmenopausal with new uterine enlargement, MRI findings) and to conduct a more thorough work-up of such patients pior to recommending UFE. Recurrent or continued tumor growth following UFE should be considered a potential warning sign for sarroma and surgery should be considered.

#### PAE Specific Warnings

- A thorough clinical evaluation should be performed on all patients presenting for embolization for BPH (e.g., urinalysis, digital rectal exam, symptom scores, prostate imaging, prostate-specific antigen test, transrectal ultrasound) to rule out prostate cancer.
- Because of the tortuous vessels and duplicative feeding arteries in the pelvic area, extreme caution should be used when performing prostatic artery embolization (PAE). Complications of mistargeted embolization include ischemia of the rectum, bladder, scrotum, penis or other areas.
- When using EmboGold Microspheres for prostatic artery embolization, do not use microspheres smaller than 100 microns. It is recommended to use 300-500 microns.

# Warnings About PAE and Fertility

The effects of PAE on fertility have not been determined.
Therefore, this procedure should not be performed on men wanting to father a child

# Haemostatic Specific Warnings

 Since EmboGold Microspheres have not been evaluated to control bleeding or haemorrhaging for neurovascular indications, they should not be used for this purpose in the neurovasculature

## Warnings about use of small microspheres

- Careful consideration should be given whenever use is contemplated of embolic agents that are smaller in diameter than the resolution capability of your imaging equipment. The presence of arteriovenous anastomoses, branch vessels leading away from the target area or emergent vessels not evident prior to embolization can lead to mistargeted embolization and severe commiscations.
- Microspheres smaller than 100 microns will generally migrate distal to anastomotic feeders and therefore are more likely to terminate circulation to distal tissue. Greater potential ischaemic injury results from use of smaller sized microspheres and consideration must be given to the consequence of this injury prior to embolization. The potential consequences

- include swelling, necrosis, paralysis, abscess and/or stronger post embolization syndrome.
- Post-embolization swelling may result in ischaemia to tissue adjacent to target area. Care must be given to avoid ischaemiaintolerant, nontargeted tissue such as nervous tissue.

- Carefully evaluate the vascular network associated with the lesion using high resolution imaging prior to beginning the
- embolization procedure.

  EmboGold Microspheres are available in a range of sizes.

  Because of the potential for misembolisation and the inherent
  variability in sphere sizes, the physician should be sure to
  carefully select the size of Embodold Microspheres according
  to the size of the target vessels at the desired level of occlusion
  in the vacculature.
- of the vasculature.

  Carefully settler the size of microspheres according to the size of the vessels identified and the catheter used. EmboGold Microspheres are flexible microspheres that support temporary compression by 20 to 30% to facilitate passage through microcatheters. Studies have shown a direct correlation between the size of microspheres and the size of occluded vessels.

- Choose a delivery catheter based on the size of the target vessel and the microsphere size being used. Refer to below table for catheters and EmboGold Microspheres sizes compatibility.
- EmboGold Microspheres are not radiopaque. It is recommended that the embolization be monitored using fluoroscopic visualization by adding the appropriate amount of non-ionic contrast medium to the physiologic suspension fluid.

# To Deliver EmboGold Microspheres

- Inspect packaging and syringe before use to ensure that they are intact. The external surface of the syringe is sterile.
- According to aseptic technique, open the peel-away film beginning at the tip, and peel back the film completely to the base. Gently tip the sterile vial on the sterile field, avoiding contact with any parts previously sealed.
- Unscrew the cap of the EmboGold Microspheres prefilled
- This highly recommended to add non-ionic contrast agent to monitor the injection adiologically. Gently draw non-ionic contrast agent directly into the reservoir syringe. The ideal suspension is usually obtained with a mixture of 50% non-ionic contrast agent and 50% saline solution. To obtain a homogeneous suspension of EmboGold Microspheres, gently inveret the 20 m Syringe several times. Non-ionic contrast

- agent and 0.9% NaCl solution can be added in the same proportions to obtain a more diluted suspension.
- Do not use the 20mL prefilled syringe to inject EmboGold
  Microspheres through the catheter, as a catheter occlusion
  may result
- may result.

  Remove all air from the syringe and connect it to one hub of
- the three-way stopcock.

  Wait several minutes to allow the EmboGold Microspheres to suspend in the solution.

  Draw up the suspension using a 1mL or 3mL injection syringe connected to another hub of the three-way stopcock. Avoid
- back and forth movements to reduce the risk of introducing air into the system. Check that the desired quantity and concentration of microspheres are used.

  Remove all air from the syringe.

  Introduce the delivery catheter into the target vessel according to standard techniques. Position the catheter tip as close as
- possible to the treatment site to avoid inadvertent occlusion of normal vessels.

  Screw the syringe onto the hub of the catheter, using the male Luer-lock connector of the stopcock.
- Open stopcock to connect the injection syringe with the catheter.
- Under continuous fluoroscopic control, slowly infuse microspheres into the blood stream while observing the contrast flow rate. If there is no effect on the flow rate, repeat

- the delivery process with additional injections of the EmboGold Microspheres/contrast solution. Consider using larger sized EmboGold Microspheres if the initial injections do not alter the contrast flow rate. If the EmboGold Microspheres/contrast solution requires re-suspension, gently invert the 20 mL syringe several times.
- Always inject under free flow conditions. Reflux of microspheres can induce immediate ischaemia of healthy tissues or vessels.
   Continue influsion until the desired devaccularization is.
- obtained. Studies have shown that EmboGold Microspheres penetrate more distally into the lesion than PVA particles of similar size. Reduction of the arterial blood supply to the lesion is therefore more progressive. Exercise conservative judgment in determining the embolization endpoint.
- Femoral puncture can result in arterial spasm. This may predispose to femoral thrombosis (e.g., leg injury). Femoral patency should be re-assessed prior to final catheter removal.
- At the end of the infusion, remove the catheter while maintaining gentle suction to avoid dislodging any residual microspheres still within the catheter lumen, then close the three-way stoocock.
- Apply pressure to the puncture site until hemostasis is complete.
- Discard any remaining EmboGold Microspheres and the used syringes.

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- Additional UFE Specific Instructions
   When embolizing uterine fibroids, choose an EmboGold
   Microsohere of 500 microns or greater.
- At the discretion of the physician, pneumatic compression devices may be used for patients currently taking hormone therapy, uterine volume >1000cc, and patients that are overweight to lower the risk of deep vein thrombosis.
- Embolization should be stopped when the vasculature surrounding the fibroid can no longer be visualized but before complete stasis in the uterine artery. There is an increased chance of retro-migration of EmboGold Microsphere into unintended blood vessels as uterine artery flow diminishes.

### Additional PAE Specific Instructions

- For prostatic artery embolization, it is recommended to use EmboGold Microspheres 300-500 microns.
- A Foley catheter, with its balloon inflated with a mixture of non-ionic contrast and saline, may be placed prior to PAE for use as a landmark during the embolization procedure.
- · PAE can be performed by either radial or femoral access.

### Additional AVM Specific Instructions

 When embolizing arteriovenous malformations (AVMs), choose an EmboGold Microsphere size that will occlude the nidus without passing through the AVM.

#### CONSERVATION / STORAGE/DISPOSAL

- EmboGold Microspheres must be stored in a cool, dry and dark place in their original syringe and packaging.
- . Use by the date indicated on the syringe label.
- Do not freeze.
   Do not resterilize.
- After use, EmboGold Microspheres must be disposed as per hospitals contaminated waste circuit.

	Size Range (µm)	Minimum Catheter ID	Color Code	1 mL	2 mL
	40-120	0.016" (0.41 mm)	Orange	S110EG	S120EG
ĺ	100-300	0.017" (0.43 mm)	Yellow	S210EG	S220EG
	300-500	0.018" (0.46 mm)	Blue	S410EG	S420EG
	500-700	0.020" (0.51 mm)	Red	S610EG	S620EG
ĺ	700-900	0.027" (0.69 mm)	Green	S810EG	S820EG
	900-1200	0.038" (0.97 mm)	Purple	S1010EG	S1020EG

### PATIENT COUNSELING INFORMATION

- Patients should have a clear understanding prior to embolization of who will provide their post procedure care and whom to contact in case of an emergency after embolization.
- Embolization patients should have an understanding of the potential benefits, risks, and adverse events associated with embolization. In particular, patients should understand that there is a chance their symptoms will not improve following embolization.

# Information on packaging



	8	Do not resterilize
<b>®</b>		Do not use if package is damaged and consult instruction for use
	类	Keep away from sunlight
	<del>*</del>	Keep dry
	8	Single use
	<u> </u>	Caution
	Ж	Non-pyrogenic
	STERILE	Sterilized using steam
	o°c ∦	Lower limit of temperature

MD	Medical Device
UDI	Unique Device Identifier
	Single sterile barrier system with protective packaging inside
[]i	Consult Instructions for Use
<b>†</b> ?	Patient identification
₩,	Healthcare centre or doctor
[31]	Date
<u></u>	Patient information website
C €2797	CE mark - Notified body identification: 2797

	Cautio
R <sub>Z</sub> ONLY	device
	nhysic

on: Federal (USA) law restricts this e to use by or on the order of a licensed physician.

In the European Union, any serious incident that has occurred in relation to the device should be reported to the manufacturer and the competent authority of the applicable Member State.





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For Patent Coverage, See www.merit.com/patents

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