



**Case Study** 

NAME • NICOLE

AGE • 32

**DIAGNOSIS** • UNRESECTABLE HCC, TERMINAL

TREATMENT INTENT • THERASPHERE Y-90 NEOADJUVANT TO RESECTION

LOCATION • U OF CHICAGO

A 6-month prognosis. And one option that provided a second chance.

When Nicole was diagnosed with liver cancer, her life turned upside down. The first medical opinion she received was that her tumor was inoperable. The second was the same. But the third was from the multi-disciplinary team at University of Chicago Medicine, who recommended TheraSphere.

The treatment worked and shrunk the tumor, paving the way for resection. Thanks to Dr. Baker, the entire team and the success of the TheraSphere Y-90 radiation, Nicole went from hopeless to healthy, from shock to survival. "The idea of the hybrid approach where we are able to give high dosage of radiation to a tumor to affect kill and also give lower doses to the liver surrounding it to affect hypertrophy ... was the only good option to get her to surgery."

Dr. Talia Baker, Liver Tumor Clinic Director at University of Chicago Medicine



Scan the QR code to see Nicole's full story.



## **THERASPHERE**<sup>™</sup> Y-90 Glass Microspheres

TheraSphere's powerful glass microspheres are engineered to shrink and destroy tumors with pinpoint precision - sparing healthy tissue while preserving future treatment options, such as surgical resection, transplant or systemic agents.

"Nicole's story tells us you should never say never working in these important multi-disciplinary pathways. It is essential to get the patient the best outcome."

Dr. Talia Baker, Liver Tumor Clinic Director at University of Chicago Medi cine Visit TheraSphere.com to learn more about TheraSphere Y-90 Glass Microspheres

## TheraSphere<sup>™</sup> Yttrium-90 Glass Microspheres

INDICATION FOR USE: TheraSphere is indicated for use as selective internal radiation therapy (SIRT) for local tumor control of solitary tumors (1-8 cm in diameter), in patients with unresectable hepatocellular carcinoma (HCC). Child- Pugh Score A cirrhosis, well-compensated liver function, no macrovascular invasion, and good performance status. CONTRAINDICATIONS: TheraSphere is contraindicated in patients: whose Tc-99m macroaggregated albumin (MAA) hepatic arterial perfusion scintigraphy shows any depos to the gastrointestinal tract that may not be corrected by angiographic techniques • who show shunting of blood to the lungs that could result in delivery of greater than 16.5 mCi ((0,6) GGA) ((0,1)) of the lungs, addition pneumonitis has been seen rarely in patients receiving dataset to the lungs greater than 30 cm a single treatment is in whom hepatic artery catheterization is contraindicated, such as patients with vascular abnormalities or bleeding diathesis • who have pulmonary insufficiency (conventionally defined by an arterial oxygen pressure (Pa, O2) of < 60 mmHg, or oxygen saturation (Sa, O2) of < 90%) or severe liver dysfunction, including hepatic encephalopathy, clinically evident ascites or treatment with diuretics for ascites • with portal vein thrombosis (PVT) Type 4 involvement and lack of Tc-99m MAA deposition on the PVT seen on the Tc-99m MAA imaging with >70% tumor replacement in the liver • with comorbidities or poor overall health (= g, ECG performance status rating > 2) which may make the patient a poor candidate for locoregional radiation treatment • who are pregnant. **WARNINGS**: The following pre-treatment, high-risk factors (disease characteristics) have been associated with serious adverse events demend possibly related to use of the device: infiltrative tumor type • tumor modules too numerous to count • AST or AIT > 5 times ULN • bilirubin > 2 mg/dl • tumor volume > 50% combined with albumin < 3 g/dL. Keep the TheraSphere dose vial upright and stored in its lead pot before and during patient treatment, except as required for radiation measurement. Do not open the dose vial acrylic shield prior to patient treatment. Post-treatment, waste materials require caution to prevent contamination and beta shielding due to residual glass microspheres. PRECAUTIONS: GENERAL PRECAUTIONS: As in any intra-arterial procedure, aseptic technique should be practiced, and care should be taken to ensure to residual glass microspheres. minimum patient anesthesia exposure extraneous to therapeutic objective. • Consideration of patient comorbidities should be used when determining the type and volume of minimum patient an estimate by Double extrained to the product of extrained to the entry of the PRECAUTION IN VULNERABLE PATIENTS: No effectiveness or safety data are available to support the use of the device in children or breast-feeding women. ENDOCRINE DISRUPTION CARCINOGENICITY, MUTAGENICITY, TOXICITY TO REPRODUCTION: Ideally the use of this radioactive device in women of childbearing capability should be performed during the first few (approximately 10) days following the onset of menses. RADIATION SAFETY: Radioactive products should be used only by healthcare professionals who are qualified by specific training in the safe use and handling of radionuclides and whose experience and training have been approved by the appropriate government agency authorized to license the use of radionuclides. • As in the use of any radioactive material, ensure minimum radiation exposure to the patient extraneous to the therapeutic objective, and to minimize radiation exposure to workers and others in contact with the patient. RELEASE AND POST-TREATMENT PRECAUTIONS: Post treatment patient care: use universal precautions for body fluid contact. Trace Y-90 may be detectible in blood and urine; handle with gloves and dispose as normal body fluids. The radiation field is expected to be less than 1 mem/h (10 µSV/h) at 3 ft (1 m) from the patient's abdomen. Supplemental shielding and segregation of the patient are not required to maintain exposure to others below regulated limits. • Release ad a fright min the patient's about the patient as an ending and segregation on the patient are not required to maintain exposure to one sector regulated to measure are not required to maintain exposure to one sector regulated to maintain exposure to one sector regulated to they can minimize their radiation exposure by avoiding prolonged time (>12 hours per day) within 1 ft (0.3 m) of the patient's abdomen for the first week post therapy. Patients should be advised that radiation emitted from the patient may be detectible at security screening (e.g., international travel). • Special precautions post-administration: If the patient requires aboves that to about the treatment with the part of th tissue. As a result, patients with compromised liver function may be at greater risk of liver function impairment and hence could experience complications. Clinical side effects usually occur within the first 4 to 6 weeks after treatment. Based on clinical trial data, literature reviews and post market surveillance, adverse events potentially associated with treatment using Y-90 microspheres, including TheraSphere, may include the following: Allergic reaction + Altered liver function, acute or chronic + Anorexia + Anxiety + Ascites + Bile Duct injury Bleeding/Hemorrhage + Chills / rigors + Cholexytis (inflammentory or infectious) + Colitis + Death + Dehydration + Diarrhea + Dizziness + Dyspnea + Edema (any location) Electrolyte above common log e concervators (minimum above or miceuous) e concervators are concervators (minimum above or miceuous) e concervators (m decompensation • Pulmonary edema • Pulmonary fibrosis • Radiation hepatitis • Radiation induced disease, acute • Radio Embolization Induced Liver Disease (REILD) • Sepsis Supraventicular armythmia • Thrombosis (arterial or venous) • Tumor inflammation (including nucleor used = haudo + nucleor issis syndrome • Vomiting • Weight loss. Complications related to the administration procedure itself may include: Allergic reaction: Arterial injury including vessel dissection • Aspiration premumonia • Bruising/bleeding/hematoma at site • Constipation/abdominal distension • Fatigue • Flushing • Infection • Nausea • Nerve damage. CAUTION: Federal (USA) law restricts this device to sale by or on order of a physician. PI-992004-AA Note: Dose to liver does not exceed 150 Gv.

TheraSphere is a registered trademark of Theragenics Corporation, used under license by Boston Scientific Medical Device Limited, a wholly owned indirect subsidiary of Boston Scientific Corporation.

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