The Utility of the Microcrystalline Cellulose Sphere as a Particulate Embolic Agent: An Experimental Study

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AJNR Am J Neuroradiol 2000, 21 (6) 1160-1163
http://www.ajnr.org/content/21/6/1160
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BACKGROUND AND PURPOSE: Although various particulate materials have been developed as embolization agents, their biocompatibility remains unclear. We used an animal model to examine the possibility of using FDA-approved microcrystalline cellulose spheres (CELPHERE) as solid embolic material for the permanent occlusion of blood vessels.

METHODS: Angiographic and histologic studies in 12 canine renal arterial systems were conducted to evaluate the performance of CELPHERE beads at 1 hour, and at 4 and 12 weeks after embolization.

RESULTS: The CELPHERE beads traveled to vessels with diameters approximating their own. Larger vessels were occluded by aggregations of beads. There was no disruption of vessel walls and no evidence of perivascular hemorrhage or inflammatory changes.

CONCLUSION: Because CELPHERE beads are easy to handle, highly biocompatible, and have few adverse effects, they are suitable for intravascular applications.

Although we previously developed an ideal particulate embolic material, the cellulose porous bead (CPB) (9–11), the FDA has not yet approved it. Therefore, we continue our quest for a new embolic agent composed of material that could gain FDA approval.

We tested a particulate material, the microcrystalline cellulose sphere (CELPHERE) bead, that meets all of the criteria listed above and, in addition, offers advantages related to both morphologic and biological compatibility with vessels. We carefully examined the properties and characteristics of CELPHERE beads, and studied embolization techniques in a dog model as a preliminary step toward eventual clinical use in arterial embolization.

Methods

Celphere embolization beads (Asahi-Kasei Co. Ltd., Tokyo, Japan) are newly developed microspheres that are manufactured commercially as structural elements of tablets. They are made of 100% Avicel microcrystalline cellulose—a purified, partially depolymerized alpha cellulose derived from purified specialty grades of wood pulp. Microcrystalline cellulose is a primary excipient in pharmaceutical oral dosage formulations because of its very low reactivity to active ingredients. It is white, tasteless, odorless, virtually free of organic and inorganic contaminants, and recognized as safe by experts. It is FDA-approved and is listed in the Drug Master File as “Microcrystalline Cellulose Spheres, CELPHERE”. Microcrystalline cellulose spheres are insoluble in water and most organic solvents; they are resistant to degradation in water and are water-absorbent. The insolubility and water-absorbent properties of the beads result in reduced particle agglomeration. They are fundamentally globoid in shape and come in different di-
ameters (150, 200, 300, 500 μm). They are of uniform and accurately calibrated size; the mean error of their diameter is within 30 μm. Their surface contour is smooth under an electron microscope (Fig 1).

CELPHERE beads of two different sizes (150 and 200 μm in diameter) were suspended in iodinated contrast material in stoppered glass bottles at about 1000 CELPHERE beads per mL and sterilized for 20 minutes in a steam autoclave at 121°C at 11 Pa. The number of beads in the suspension was obtained by counting aliquots with a Coulter multisizer 2 (Coulter Electronics Limited, Beds, England). For each batch, the quality of sterilization was bacteriologically confirmed.

In the in vivo experiments, we used 12 adult mongrel dogs weighing about 15 kg. They were premedicated intramuscularly with 0.01 mg atropine sulfate and 10 mg ketamine chloride per kg. Prior to intratracheal intubation, they were anesthetized with IV injection of 10 mg thiopental sodium per kg. Additional thiopental sodium (5mg/kg) was IV injected as needed. Muscle relaxation was produced with suxamethonium chloride. The experiments were performed under controlled mechanical ventilation. During the experiment, the arterial carbon dioxide pressure was maintained between 35 and 40 mm Hg; the partial pressure of arterial oxygen was kept at 100 mm Hg or higher. The right femoral artery in the groin was punctured, a 4F introducer sheath was placed, and 2000 U heparin was injected. At the end of the procedure, protamine sulfate was injected to reverse the effect of heparin.

Renal preembolization angiograms were obtained using a 4F polyethylene catheter with the tip of the catheter positioned in the orifice of the artery. Under fluoroscopic control, a unilateral embolization of the renal artery was performed slowly until the flow was arrested (usually after 6 mL). Suspended in the embolization volumes were 1000 CELPHERE beads/mL. Selective renal angiograms were obtained immediately and at 1 hour, and again at 4 and 12 weeks after embolization. All dogs, except those sacrificed 1 hour after embolization (n=4), were allowed to recover from anesthesia. The latter were sacrificed 4 weeks (n=4) and 12 weeks (n=4) after the procedure. Two animals in each group underwent embolization with CELPHERE beads measuring 150 and 200 μm in diameter, respectively. The kidneys were removed en bloc, fixed in 10% neutral buffered formalin, cut into 3-micron sections, and stained with hematoxylin & eosin, Giemsa, and Elastica van Gieson.

In all experiments, we followed the Animal Care Guidelines set forth by the Animal Experimentation Ethical Committee of our institution.

Results

None of the dogs manifested any outward signs of abnormality during the pre- and postembolization observation period. Renal angiograms, obtained before embolization, showed typical canine arterial vasculature. CELPHERE produced a smooth reduction in blood flow; severe stasis occurred in the immediate phases after embolization of the main renal artery. Renal angiograms obtained 1 hour and 4 and 12 weeks after embolization revealed complete occlusion of the renal artery (Fig 2).

On examination of the gross kidney specimens, thrombosis of the main renal artery was noted. During the 12 weeks after embolization, a progressive, marked decrease in renal size occurred. Microscopically, CELPHERE beads measuring 200 μm in diameter were distributed throughout the intramedullary arteries; beads measuring 150 μm in diameter were observed in the arteries on the border of the cortex and medulla. The CELPHERE beads traveled to vessels with diameters approximating their own. Larger vessels were occluded by dense aggregates of many beads; there were no interstices (Fig 3A).
There was no damage to vessel walls and no evidence of perivascular hemorrhage. Sections obtained 1 hour after embolization showed fresh thrombus formation. Sections prepared 4 and 12 weeks after embolization showed permanent occlusion of the vessels with infarction. There were no inflammatory changes of the vessel walls. The particles were not resorbed, nor did their appearance change during the 12-week observation period (Fig. 3B).

**Discussion**

We have been testing embolic materials with the goal of detecting some that are superior with respect to ease and safety of use, biocompatibility, and permanence of embolization. We previously developed the solid embolic agent, CPB, which is still undergoing clinical trial. These beads are exceptionally uniform in size, and have a specific gravity similar to blood and net-positive charge. Their long-term safety, however, has not been established. CELPHERE beads are microspheres that are produced commercially; the FDA has approved their use in tablet form for humans.

Embolization with CELPHERE did not result in inflammation to surrounding tissues even 12 weeks after embolization, nor was there any apparent damage to vessel walls, angionecrosis, or focal hemorrhage. Germano et al (12) reported that polyvinyl alcohol (PVA) particles induced a rapid foreign-body response and inflammation in all embolized vessels. This could result in arterial rupture or aneurysm formation. Angionecrosis, focal hemorrhage, and severe vasogenic edema occurred in rare cases. Although foreign-body giant cells are a nonspecific reaction to many foreign materials and do not by themselves indicate toxicity, the presence of localized angionecrosis and focal hemorrhages strongly suggested direct toxicity of the particles (12–15).

The CELPHERE beads used in our study were spherical in shape, their surface was smooth, and it was possible to calibrate their size accurately. The beads migrated distally without clumping, thereby producing a more complete and permanent occlusion. In fact, we encountered no recanalization as late as 12 weeks after embolization.

Because CELPHERE beads come in different sizes (150, 200, 300, 500 μm in diameter with a mean error of 30 μm), the optimal size can be selected on a case-by-case basis. Their tendency to travel to vessels with diameters approximating their own resulted in a high embolic effect. The propensity of some embolic materials (eg, PVA and Sephadex) to become enlarged after introduction into the bloodstream resulted in stretching and, in some cases, rupture of the internal elastic lamina and vessel wall (16). Our experiments showed that the CELPHERE beads retained their size even as late as 12 weeks after injection, and that they produced no injury to the vessel walls.

The apparent density of CELPHERE beads was 0.87–0.97 g/cm³, and the specific gravity was close to that of whole blood. Although CELPHERE beads of 150 and 200 μm diameter traveled easily through the length of the microcatheter (Tracker 18, Guerbet Biomedical, Target Therapeutics), there was slight resistance at the microcatheter connector. To optimize the characteristics of CELPHERE beads for permanent embolization, a desirable improvement would be to reduce their gravity to allow for prolonged suspension.

**Conclusion**

Our animal study showed that CELPHERE beads satisfied most of the requirements of an ideal permanent embolic material. Further slight improvements will make it possible to perform clinical trials with this embolic material.
References


