Transpapillary iridium-192 wire in the treatment of malignant bile duct obstruction

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SUMMARY Twenty four patients with malignant bile duct obstruction were treated with intraluminal radiotherapy using iridium-192 wire inserted through an endoscopically placed nasobiliary catheter. Biliary drainage after treatment was maintained by an endoprosthesis. The median dose of intraluminal radiotherapy was 6000 cGy; two patients with cholangiocarcinoma were given a second course because of disease extension; four patients with pancreatic carcinoma received additional external irradiation (3000 cGy). There was one early death from a cerebrovascular accident (30 day mortality, 4.2%). Cholangitis (30%) was the major early complication and stent blockage (40%) the major late complication; there were no complications directly attributable to radiotherapy. The median survival for patients with pancreatic carcinoma was 250 days and for cholangiocarcinoma, 300 days. This method is technically feasible and may prove safer than the transhepatic technique. The ability of intraluminal irradiation to improve palliation or lengthen survival in patients with malignant bile duct obstruction remains uncertain. Assessment by a prospective, controlled trial is essential.

Intracavitary irradiation is an accepted form of palliation in patients with oesophageal, gastric and rectal malignancy. A high dose of irradiation can be delivered to a limited volume of tissue over a short period - a significant advantage over conventional external radiotherapy. Intraluminal irradiation using iridium or radium placed transhepatically has been used to treat patients with malignant bile duct obstruction. It has been suggested that this form of treatment, together with catheter or endoprosthetic drainage for the relief of jaundice, may double the survival time in patients with hilar cholangiocarcinoma.

Biliary decompression by the transhepatic route has a higher morbidity in malignant bile duct obstruction than the endoscopic transpapillary approach; similar advantages might also be expected using this technique for placement of both the iridium-192 wire and the subsequent endoprosthesis. A method for the transpapillary administration of iridium-192 in patients with malignant bile duct obstruction has been evaluated.

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Methods

PATIENTS Between June 1983 and September 1986, 24 patients were given intraluminal iridium-192 wire irradiation for malignant bile duct obstruction. There were 13 men and 11 women, median age, 69 years (57-87). Obstructive jaundice was the result of carcinoma of the head of pancreas in 11 patients, cholangiocarcinoma in seven (all below the bifurcation), hilar lymph node metastases in three, carcinoma of the gall bladder in two and ampullary carcinoma in one. Histologic confirmation of the diagnosis was obtained in 19 patients (79%). The subsequent clinical course in the remaining five patients was consistent with the radiologic diagnosis of a malignant stricture.

DRAINAGE TECHNIQUE At the initial ERCP a 6 or 7 Fr catheter was passed through the biliary obstruction, rerouted through the nose and drained externally; a barb on the catheter above the stricture prevented downward displacement. After completion of intraluminal irradiation the nasobiliary tube was removed and, at a second ERCP, an endoprosthesis was inserted using a stan-
standard technique\(^2\); since the availability of wide-channel duodenoscopes the stent and nasobiliary catheter have been inserted at the initial procedure (Fig. 1). In one patient the catheter could not be negotiated through the obstruction and was inserted percutaneously; subsequent iridium-192 therapy and endoprosthesis insertion were carried out by this route.

Iridium-192 therapy was administered through the nasobiliary catheter in 23 patients and stents (8–10 Fr) were successfully placed in 20. Surgical bypass (two patients) and percutaneous stent insertion (one patient) were necessary for relief of jaundice in the other three.

**Iridium-192 Wire Loading**

The length of iridium-192 wire used was equivalent to 1.5 times the length of the malignant stricture, allowing for 25% overlap of the stricture at either end. Trimming of the duodenal end of the wire was required for distal strictures to prevent irradiation of the duodenal mucosa.

A movable core, Teflon coated guidewire (0.088 mm diameter, 145 cm long, Medrad TM351451) was used. The stilette was removed and a 1–2 cm nylon thread inserted followed by the length of iridium-192 wire. The stilette was replaced and the iridium-192 wire with the nylon thread were pushed to the sealed distal end of the Teflon sheath (Fig. 2). The loaded guidewire was heat sterilised in an autoclavable transport container.\(^1\)

The upper limit of the malignant stricture was delineated by a nasobiliary cholangiogram and the loaded guidewire was then passed down one limb of a Y-connector (Cook PSFL-PCF-MLL-30) attached to the proximal end of the nasobiliary catheter. Positioning of the iridium-192 segment across the stricture was facilitated by the greater radiodensity of the iridium-192 wire compared with the outer sheath of the guidewire. The guidewire was fixed proximally within its limb of the Y-connector to prevent dislodgement; external bile drainage was established through the other limb. Prophylactic antibiotics were given routinely.

Therapy was designed to administer 6000 cGy with 100% of the dose delivered at a distance of 0.5 cm from the iridium-192 wire. Dislodgement of the nasobiliary catheter occurred in three patients. In one of these, the catheter was repositioned and the course of intraluminal irradiation completed; in the other two no further intraluminal irradiation was attempted (4100 and 4500 cGy delivered). The duration of therapy was determined by the length and radioactivity of the Iridium-192 wire, and the dose rate (estimated from a locally calculated 'escargot' similar to the Paris system\(^3\)). The median duration of intraluminal therapy was 90 hours (70–109).

Four patients, each with carcinoma of the head of pancreas, received a course of external radiotherapy shortly after completion of intraluminal irradiation. Two patients, both with cholangiocarcinoma, underwent a second course of intraluminal irradiation for extension of their disease at 439 and 1065 days.

The field of external radiotherapy was planned while the nasobiliary tube was *in situ* and was administered one to four weeks after completion of the intraluminal irradiation. A total of 3000 cGy was delivered by a 4 MEV linear accelerator in 10 fractions over two weeks using AP/PA fields to a volume not exceeding 10×10 cm even if the tumour exceeded this. The field was planned to ensure that not more than 50% of the right kidney was included in the treatment volume.
There varied intraluminal irradiation developed creas (fistula formation) necessitating surgical treatment. In the 11 patients without early complications the median duration of post-treatment hospitalisation was nine days (1–26); in the presence of early complications (12 patients) this duration was increased to 23.5 days (5–60).

Discussion

The 30-day mortality of 4.2% for intraluminal irradiation and endoscopic stent drainage compares favourably with that for endoscopic biliary drainage alone (8–19%14–17) or palliative surgical bypass (6–19% for pancreatic carcinoma,18–20 11–35% for cholangiocarcinoma21–23). In the only comparable study of iridium-192 therapy, the hospital mortality using the transhepatic approach was 3.3%. The mortality associated with iridium-192 therapy is clearly low and certainly no worse than that for palliative stenting alone.

The incidence of cholangitis (30%) is considerably higher than that after endoscopic stent insertion without intraluminal irradiation (19%18). Karani et al19 reported cholangitis in 12 of 30 patients (40%), it being a contributory factor to death in two cases. Although Molt et al20 did not encounter a single case of cholangitis attributable to transhepatic intraluminal irradiation in 15 patients, cholangitis must still be regarded as a major hazard of iridium-192 therapy by either the transhepatic or transpapillary route.

The increased risk of cholangitis may be because of impaired bile drainage through the nasobiliary tube with the contained guidewire. The simultaneous insertion of stent and nasobiliary tube at the initial ERCP (Fig. 1) might be expected to improve drainage during the period of intraluminal irradiation and reduce the risk of cholangitis; it also eliminates the need for a second ERCP for final stent insertion. At the initial procedure the stent (10–11.5 Fr) is passed over the biliary catheter and positioned across the obstruction; on completion of intraluminal irradiation the pigtail catheter is straightened with a guidewire and withdrawn under fluoroscopic control leaving the stent in situ. This technique has been done successfully in three patients, an insufficient number to determine if the incidence of cholangitis is lowered.

In the absence of a control group treated by biliary stent alone, survival figures after iridium-192 therapy are difficult to interpret. In this and other21 series, however, survival was not significantly greater than after palliative drainage or bypass alone (pancreatic carcinoma, six to seven months14–20 21; cholangiocarcinoma, 8–19 months22–24).

The transpapillary technique of intraluminal irradiation is technically feasible and may prove safer...
than the transhepatic approach. Iridium-192 therapy by either route, however, is associated with a higher incidence of cholangitis than with an endoprosthesis alone. To justify this added risk some benefit in terms of increased longevity or improved quality of life must be shown. A controlled trial of intraluminal irradiation against endoprosthesis alone in patients with malignant bile duct obstruction is required.

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References