Effect of yttrium 90 on experimental allergic arthritis in rabbits

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Meier-Ruge, W., Müller, W., and Pavelka, K. (1976). Annals of the Rheumatic Diseases, 35, 60–66. Effect of yttrium 90 on experimental allergic arthritis in rabbits. Seventeen rabbits were immunized with complete Freund’s adjuvant and bovine serum albumin by the method of Dumonde and Glynne (1962), as modified by Cooke and Jasin (1972). Fifteen weeks after allergic arthritis developed in the knee joint, 8 animals were given an injection of 200 µCi yttrium 90 (90Y) into the left joint cavity; 7 were injected with 400 µCi. The animals were sacrificed at 2, 4, 8, 12, and 16 weeks, and at 6 and 12 months after the injection. The right knee joint served as control for assessment of untreated allergic arthritis. Morphological control of the severity of the arthritis was provided by sacrificing 2 uninjected animals 13 weeks after immunization.

After 13 weeks the allergic arthritis progressed to severe inflammation of the knee joint marked by massive round-cell infiltration, oedema, and proliferation of synovial mesothelium in the synovial villi and joint capsule.

Treatment with 90Y was effective 2 weeks after injection with the disappearance of inflammatory oedema and marked regression of round-cell infiltration. This was accompanied by degeneration of the synovial mesothelium and fibrosis of the subsynovial tissue and synovial vessels as a secondary effect of the radiation.

In the animals with severe allergic arthritis, the healing effects of 90Y were more marked than the secondary effects of the radiation. The secondary effects were dose-dependent and consisted of patchy necrosis of the chondrocytes in the fibrocartilage, in the meniscus, in the cruciate ligaments, and in the tangential cartilage of the joint. There was also localized bone-marrow necrosis in the tibia adjacent to the joint.

Treatment with 90Y of arthritic knee joints with the lowest effective dose of the isotope—if necessary with repeated application—seems justified. A single large dose does not have a greater therapeutic effect and causes more radiation damage to the joint.

In view of the possible secondary effects in the joint, the indication for 90Y therapy should be restricted, particularly in young patients, to cases of chronic relapsing arthritis unresponsive to other treatment.

Recently the local treatment of arthritis by intra-articular injections of radioactive isotopes has become more common (Prichard, Bridgman, and Bleehen, 1970; Oka and others, 1971; Jalava, 1973; Müller, Fridrich, and Pavelka, 1974a). Because of its optimal radiation properties and low absorption from the joint capsule (Webb, Lowe, and Bluestone, 1969; Gumpel, Williams, and Glass, 1973; Stevenson and others, 1973), the preferred isotope for treating a large joint is yttrium 90 (90Y). For smaller joints, other isotopes with low beta-energy are used, such as gold 198, erbium 169, and rhenium 186 (Delbarre and others, 1968, 1973; Müller, Fridrich, and Pavelka, 1974b).

As our earlier studies on the normal rabbit knee joint have shown (Pavelka and others, 1975) 90Y induces fibrosis of the subsynovial connective tissue and synovial villi. The occurrence of these changes, similar to those of joint aging, suggested that treatment of the inflammation with 90Y might cause radiation-induced secondary effects which would exacerbate rather than reverse the pathological joint changes, thereby limiting the effectiveness of isotope treatment in man. We therefore induced adjuvant...
arthritis in rabbits and compared the morphological changes occurring in untreated animals with those in animals given local injections of $^{90}\text{Y}$.  

Methods

Seventeen male rabbits, weighing 2.0–2.5 kg, were immunized by the method of Dumonde and Glynn (1962), as modified by Cooke and Jasin (1972). Under mild anaesthesia (Numal), 1 ml of complete Freund's adjuvant (Difco Ltd., corresponding to 2 mg of lyophilized mycobacteria) and 1 ml of bovine serum albumin (Behring AG, type HD 05, corresponding to 5 mg) were injected intracutaneously under aseptic conditions into each of 4 shaved skin areas of the back. All the animals tolerated the immunization well.

Three weeks after inoculation, the animals were given an injection into each knee joint cavity of 2.5 mg albumin in 0.5 ml physiological saline. Fifteen weeks later 8 animals were injected with 200 µCi $^{90}\text{Y}$ into the left knee joint cavity and 7 animals were injected with 400 µCi $^{90}\text{Y}$. The right knee joint served as untreated arthritis control in all the animals. Two animals were not injected with $^{90}\text{Y}$ and observations were made of the severity of the arthritis 13 weeks after immunization. The complete plan of the trial is shown in Fig. 1.

Histological Technique

The knee joint was opened by removing the patella and the joint was fixed for 7 days in Bouin's solution, which was changed twice during the week. After 4 days of fixation the joint was sawn through to give a transverse midsection disc giving a complete picture of the meniscus. The sections were then fixed for the remaining 3 days in Bouin's solution, followed by dehydration of the sections for 4 days in methanol. Without decalcification, they were then embedded in methylmethacrylate by Burkhardt's method (1966a,b) (100 ml destabilized methylmethacrylate, 3.5 g benzol peroxide, and 25 mg Plastoid N). After soaking the sections for 3 days with repeated changing of the methacrylate, the mixture was allowed to polymerize at 37°C in vacuo by the methods of Burkhardt (1966a,b), Gardner (1972), and Vitali (1970).

The polycrystalline resin blocks were cut with a cut-all Zeiss microtome to yield sections 4 µm thick which were placed on gelatin-coated slides, covered with cigarette paper, pressed with a small linoleum roller, and dried at about 40°C.

Before staining, methacrylate was removed from the sections with benzene and they were rehydrated with a series of aqueous ethanol dilutions of decreasing strength. Staining was carried out as for paraffin sections using the following stains: (1) jaune solide (haematoxylin-acid fuchsin-fast yellow); (2) Giemsa; (3) toluidine blue (metachromatic staining); (4) azure eosin (Nocht-Maximov).

Results

Examination of the 2 untreated animals showed that severe arthritis of the knee joint developed 13 weeks after immunization. This was marked by massive round-cell, and to some extent plasma-cell, infiltration of the synovial villi with proliferation of the synovial mesothelium (Fig. 2). The latter also showed

FIG. 1 Experimental course of allergic arthritis in the rabbit with and without $^{90}\text{Y}$ treatment. 

A = Immunization with Freund's adjuvant and albumin; B = 2.5 mg albumin injection into the knee joint; C = 200 µCi or 400 µCi $^{90}\text{Y}$ injection into the left knee joint

FIG. 2 Allergic arthritis 12 weeks after immunization. Synovial villi of the knee joint showing massive round-cell infiltration and proliferation of synovial mesothelium. Paraffin; haematoxylin and eosin. ×120
uneven round-cell infiltration, with oedema of the
neighbouring connective tissue. The thin-walled
vessels of the subsynovial tissue were hyperaemic and
markedly dilated. In the joint cavity there was a
moderately protein-rich exudate with a low cell
content.

The arthritis was still highly active 17 weeks
after immunization. The synovial villi continued
to show round-cell infiltration, oedema, and to
some extent fibrinoid swellings. Massive proliferation
of the synovial mesothelium could be seen in the
recess of the joint capsule. A prominent feature was
the focal inflammatory synovial changes. In the
region of the meniscus the synovial villi showed
proliferation of fibrocytes and fibroblasts, a sign
of a chronic disease process, and focal proliferation
of the synovial mesothelium.

In the animals which 2 weeks previously, i.e. 17
weeks after immunization, had been given injections
of 200 or 400 μCi 90Y into the joint cavity, the inflam-
mation had regressed. The round-cell infiltration
and oedema in the joint capsule and synovial villi
could no longer be seen, and such infiltrations as were
present were small and mostly insignificant. To some
extent the synovial cells showed pyknosis or con-
siderable structural breakdown. Signs of synovial
regeneration could be seen, particularly in the region
of the plicae of the joint capsule, though this was
accompanied by severe fibrosis of the subsynovial
tissue of the joint capsule (Fig. 3). The vessels showed
connective-tissue 'sheathing' or thickening of the
walls.

In the treated joints, especially those injected with
400 μCi 90Y, pyknosis was observed in the fibrous
tissue of the cruciate ligaments and the cartilage
of the meniscus together with karyolysis of a few chon-
drocytes (Fig. 4). A notable additional finding due
to the 90Y treatment was a loss of erythropoietic
bone marrow in the region of the tibial attachment
of the cruciate ligaments.

Nineteen weeks after immunization the treated
joints showed the characteristic signs of irradiation
injury with fibrosis of the synovial membrane and
vessels. The synovial mesothelium was partially
regenerated. In the region of the cruciate ligaments
there were small defects in the tangential layer of the

![FIG. 3 Femoral parts of the joint capsule in allergic arthritis (31 weeks after immunization)](image1)

![FIG. 4 Meniscus 2 weeks after exposure to 400 μCi 90Y. Degeneration of chondrocytes with pyknosis and karyolysis. Polyacrylic resin; toluidine blue. ×300)](image2)
femoral joint cartilage. A few proliferating cartilage cells could be seen in the fibrocartilage. This was a much more prominent finding in the animals treated with 400 μCi 90Y 8 weeks before being sacrificed, i.e. 23 weeks after immunization. At this stage there was severe degeneration of cruciate ligament with localized necrosis. In addition there was incipient regeneration of cartilage cells marked by the appearance of giant cartilage cells and polynuclear chondrocytic giant cells.

Sixteen weeks after 90Y injection of the knee joint, i.e. 31 weeks after immunization, slight signs of arthritis with localized round-cell infiltration in the synovial villi were still to be seen (Fig. 5). The surrounding area of the joint capsule showed fibrous changes. In line with other observations on joints treated with 90Y there were fibrotic changes in the vessels, some of which showed collagenous degeneration and ectasia of the walls.

This relatively normal appearance of the joint contrasted sharply with that of the untreated joint on the other side. There was massive proliferation of the synovial villi in the plicae of the joint capsule, which was enlarged. Both the synovial villi and the subsynovial layers of the joint capsule exhibited diffuse round-cell infiltration (Fig. 6, left) consisting of plasma cell and lymphoid cell elements as well as protein-storing phagocytes. In addition there was proliferation of synovial mesothelium of inflammatory origin. Some of the synovial villi showed deposits of fibrin. Localized fibrinoid degeneration of the tangential cartilage on the tibial side of the knee joint was also observed.

The most serious damage was in the region of the cruciate ligaments (Fig. 6), where fibrinoid necrosis was seen alongside massive round-cell infiltration of synovial villi. Degenerated and polynuclear chondrocytes were present in the fibrous tissue of the cruciate ligaments, together with fibrinoid degeneration and foci of histiocytic and fibroblastic cells (Fig. 6, right).

These arthritic lesions were still in evidence at 9 and 15 months after immunization, though now mainly confined to the plicae of the capsule, to synovial villi at the base of the meniscus, and less so to the region of the cruciate ligaments. They were marked by considerable collections of phagocytes among the round-cell infiltrations. The plicae of the capsule were enlarged and marked by the presence of synovial villi showing various degrees of inflammatory injury. As in the earlier stages of the arthritis, the border of the synovial mesothelium was seen to be widened.

FIG. 5 (Left) 16 weeks after treatment with 90Y. Slight residual arthritis in the plicae of the joint capsule. Collections of round-cells in synovial villi. Regenerating synovial mesothelium. Polyacrylic resin; toluidine blue. ×72. (Right) Contralateral side with allergic arthritis. Massive proliferation of synovial villi showing round-cell infiltration. Spreading of synovial mesothelium. Polyacrylic resin; toluidine blue. ×28
A few foci of fibrinoid connective-tissue degeneration were present in parts of the synovial mesothelium that had been cut tangentially.

Six and 12 months after injection, the capsules of the treated joints displayed in most cases a normal or only slightly inflammatory degree of infiltration. The synovial villi showed fibrosis of varying severity. In one animal with severe allergic arthritis the inflammation, though still active, had markedly regressed 12 months after treatment with $^{90}$Y. Destruction of joint cartilage at the periphery and in the region of the cruciate ligaments was considerably less in the treated joints than in the untreated ones (Fig. 7).

As far as the therapeutic effect was concerned, there was no fundamental difference between a dosage of 200 $\mu$Ci and 400 $\mu$Ci $^{90}$Y. In the animals treated with 400 $\mu$Ci $^{90}$Y there was evidence of radiation damage in the tibial bone marrow near the joint as well as lesions in the tangential layer of the peripheral joint cartilage and in the region of the cruciate ligaments. The most serious radiation damage was in the cruciate ligaments themselves.

Attention must finally be drawn to a general observation made on the synovial membrane of the rabbit knee joint. In healthy rabbits 1–2 years old the synovial mesothelium is found principally in the region of the synovial villi in the femoral and tibial plicae of the joint and at the base of the meniscus. In animals with adjuvant arthritis the synovial mesothelium gradually extends over the whole of the joint capsule.

**Discussion**

The results show that the local injection of $^{90}$Y can be regarded as an effective therapeutic method in allergic arthritis in the rabbit. The inflammatory oedema disappears as early as 2 weeks after injection of $^{90}$Y. Radiation from the isotope causes fibrosis of the subsynovial connective tissue of the joint capsule and of the synovial villi. It also affects the complex vascular system of the synovial membrane, with closure of some of the vessels occurring.

Signs of degeneration in the synovial mesothelium are seen only during the 2 weeks after $^{90}$Y injection. Then there is rapid and extensive regeneration of the synovial membrane, though the extent to which the synovial mesothelium has spread over the joint capsule decreases with the increasing age of the rabbit. As noted above, in principle the synovial mesothelium of the nonarthritic rabbit joint is confined mainly to the region of the synovial villi in
the plicae of the knee joint capsule, to the base of the meniscus, and to the synovial villi in the region of the cruciate ligaments. As allergic arthritis develops the synovial mesothelium proliferates, spreads, and gradually extends over the whole of the joint capsule. This proliferation is directly related to the extent of subsynovial round-cell infiltration and of fibrinoid swelling of the connective tissue of the joint capsule.

Small lesions in the tangential layer of the joint cartilage were seen in the joints treated with $^{90}$Y particularly when the larger dose of 400 $\mu$Ci had been injected. This was often conspicuous in the region of the cruciate ligaments and at the periphery of the joint surfaces. In addition, minor radiation damage occurred in the region of the cruciate ligaments with loss of erythropoietic bone marrow in the tibia. It is near these ligaments that the joint cavity is least protected by cartilage and bone (Jalava, 1974). When the untreated joints are compared with treated joints one year after injection of $^{90}$Y, the arthritic destruction is seen to be much greater in the untreated joints. However, using a dose of 400 $\mu$Ci $^{90}$Y results in necrosis in the region of the meniscus and cruciate ligaments, with involvement of chondrocytes and reactive chondrocytic giant cells as well as of giant cells in the fibrocartilage of these ligaments. On the other hand, though these changes are important, they are of little significance in comparison with the severe destruction brought about in the region of the cruciate ligaments by allergic arthritis. In less severe arthritis the radiation damage to the cruciate ligaments is the most prominent lesion.

There is no fundamental difference between doses of 200 $\mu$Ci and 400 $\mu$Ci $^{90}$Y. In severe arthritis neither dose suppresses the disease process entirely, for some degree of inflammation persists even if slight. Kalliomäki, Jalava, and Möttönen (1974) carried out similar studies on rats (Jalava, 1974) using a different experimental design.

The following conclusions can be drawn with regard to the therapeutic effects obtained with the Dumonde-Glynn (1962) experimental model of rabbit allergic arthritis.

1. Allergic arthritis of the rabbit can be effectively treated with $^{90}$Y.
2. In severe arthritis the secondary radiation effects are less severe than the primary effects with this model.
(3) In order to minimize the secondary effects of radiation the smallest effective dose should be used, if necessary with repeated application (Jalava, 1974).

(4) In view of the appearance of the known secondary effects of therapy with $^{90}$Y on the joint cartilage and on the fibrocartilage of the cruciate ligaments, the indications for $^{90}$Y therapy should be restricted. $^{90}$Y should be reserved for cases of chronic arthritis unresponsive to other treatment and should be restricted in young patients because of the long life expectancy (de la Chapelle and others, 1972; Dolphin, 1973; Stevenson and others, 1973; Ingrand, 1973; Jalava and Salonius, 1974).

References


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