The mast cell in early rat adjuvant arthritis

A. GRYFE,* P. M. SANDERS,† AND D. L. GARDNER‡
Division of Experimental Pathology, Kennedy Institute of Rheumatology, London

Adjuvant arthritis is a prominent feature of the disease which follows 9 to 14 days after normal rats are given a single intradermal injection of a mycobacterial adjuvant mixture. Most accounts of the microscopic changes seen in the limb joints have described the late changes, after the disease is clinically established (Pearson and Wood, 1959; Silverstein and Sokoloff, 1960; Pearson, 1963; Glenn and Gray, 1965; Jones and Ward, 1966).

Studies by Jones and Ward (1963) and by Burstein and Waksman (1964a) analysed the initial lesions but disagreed on the characteristics of the cellular changes. The following study was therefore undertaken to clarify the nature of the earliest cellular responses detectable by light microscopy.

Materials and methods

96 male albino Wistar rats weighing from 136 to 277 g. were given a single intradermal injection of 0.05 ml. of an adjuvant mixture into the left hind footpad. Rats were caged randomly in 24 groups of four and maintained on a standard laboratory pellet diet and water ad lib. The animals, which included untreated, normal controls, were examined daily and weighed, and the carpal and tarsal diameters were measured with calipers. Joint inflammation in each limb was assessed daily and recorded on a 4-point scale (Quagliata, Sanders, and Gardner, 1969).

From the 5th to the 12th days, twelve rats, selected at random, were killed each day. The right hind limb was immediately removed, fixed in formal saline solution, and decalcified. 8 to 10 μm. paraffin sections were stained with haematoxylin and eosin (H and E) and examined microscopically for synovitis, myositis, periostitis, periarticular cellulitis, periosteal hyperplasia, and new bone formation. The numbers of mast cells, lymphocytes, histiocytes, and polymorphs infiltrating the tissues, the degree of soft tissue oedema, the amount of inflammatory exudate in joint spaces, and the numbers of cells and quantity of fibrin in the joint space exudate were assessed. The number of intact and of degranulated mast cells in representative areas of the synovial and periarticular tissues was also determined.

Results

Six of the twelve rats killed on the 10th day, ten of twelve on the 11th day, and each of twelve on the 12th day displayed clinical evidence of arthritis in the non-injected right hind limb at the time of death (Fig. 1).

Histological abnormalities were detected in or around all affected joints. Inflammation appeared to arise in the loose connective tissue of the joint and tendon sheath synovium, and in the periarticular loose connective tissues (Figs 2 and 3). The reaction spread directly to adjacent skeletal muscle and periosteum (Figs 4 and 5). The earliest microscopic change recognized was oedema (Fig. 6) observed in two rats at Day 5 and in three, four, and six rats at Days 6, 7, and 8 respectively.

Mast Cells

These were the first to appear in abnormal numbers (Figs 7 and 8); they were numerous in perisynovial and periarticular tissues of normal rats but many more than normal were evident in the non-injected

* Pathologist, Hamilton Health Association, Hamilton, Canada
† Junior Research Fellow, Medical Research Council
‡ Director, Sir Edward Lewis Lecturer, and Head of Division of Experimental Pathology, Kennedy Institute

Requests for reprints should be addressed to D. L. G., Kennedy Institute, Bute Gardens, London, W.6.
feet of rats killed 5 days after the injection of adjuvant. The numbers continued to increase until the 6th day and then declined, at first rapidly and then more gradually. No mast cells were recognized among the synovial cell layers of normal rat limbs, but in a small proportion of treated rats on each day after the 4th, occasional mast cells appeared in this site. No mast cells were seen lying free in joint spaces, whether or not an inflammatory exudate was present. Mast cells devoid of cytoplasmic granules were first identified on Day 11; by Day 12 very few cells with granules were seen.

LYMPHOCYTES
Increased numbers (Fig. 6) appeared constantly on Day 10. A few perivascular lymphocytes were noted on Day 8, but by the 10th day, eight rats exhibited these cells in significantly increased numbers. However, only six of these animals had arthritis. By the 11th day, eleven rats displayed lymphocytes in moderate-to-large numbers in synovial and periar-
ticular tissues; ten of the twelve rats in this group showed clinical disease. All rats killed on Day 12 demonstrated lymphocytic infiltration, first as clusters around blood vessels and then diffusely throughout the tissues.

HISTIOCYTES
These appeared in increased numbers later (Fig. 6). They were observed in only two animals on Day 10 but in eight on Day 11, when their numbers were generally lower than those of the lymphocytes. On Day 12 the number of lymphocytes often exceeded that of histiocytes.

POLYMORPHS
With one exception none was seen until the 10th day when they could be detected in six animals (Figs 6 and 9). At first the number was low, and almost all were confined to vascular lumina. Thereafter, the cells rapidly increased in number but still remained largely within the blood vessels. Not until they were numerous did these cells appear in the surrounding tissue; at this time they could often be seen traversing blood vessel walls. On Day 11, when eleven of the twelve rats showed synovial polymorph infiltration, many of these cells were still intra-
vascular or perivascular. Although the number of polymorphs greatly increased on the 11th and 12th days, they were usually fewer than the lymphocytes and always scarcer than the combined lymphocyte and histiocyte population. When degranulated mast cells appeared, polymorphs were already present in
the tissues. However, when polymorphs were still confined to blood vessels, mast cells invariably possessed intact granules.

The joint exudate consisted mainly of polymorphs enmeshed in fibrin (Fig. 4). Mononuclear cells were either not seen or were sparse and consisted mainly of histiocytes. Lymphocytes were scarce. The few lymphocytes and relatively numerous polymorphs did not reflect the preponderance of lymphocytes in the tissues. The earliest joint exudate appeared in one rat, at Day 10, which had developed clinical arthritis at Day 9. Of the eleven rats with histological evidence of synovitis on Day 11, four had inflammatory exudate in the joint spaces and in three the exudate was present in at least moderate amount. Three of these four rats had arthritis in non-injected limbs on Day 9 and the 4th developed arthritis on Day 10.

No exudate was seen in the joint space in the absence of established histological synovitis, although in one rat killed on Day 12 a scantly exudate appeared when the degree of synovitis was only mild and the number of polymorphs in the synovium small with the majority still intravascular. When fibrin was seen it was always accompanied by polymorphs.
Identification of adjuvant arthritis. It is apparent and basophilic cytoplasm recognized are granules and metachromatic granules in the synovial cell layers lining joint spaces, and degranulation. The first two changes begin at approximately the same time. More than the normal proportion of mast cells is seen by the 5th day, but this number markedly and suddenly increases on Day 6 and then decreases, at first sharply and then more gradually. The literature on rat adjuvant arthritis contains no previous reference to this cell population.

Mast cells are believed to be the major source of histamine in rats (Keller, 1966). Keller has discussed the release of histamine by degranulation and sometimes by cytolysis of mast cells. In the present study, both mechanisms could explain the fall in the numbers of mast cells after the 5th day, the former resulting in mast cells which are more difficult than normal to identify, the latter decreasing the actual numbers. Why mast cells should contribute to the inflammatory reaction in adjuvant disease is more difficult to resolve. According to Sheldon and Bauer (1960), tissue mast cells initiate acute inflammation at many sites of injury. These cells, the histamine released from them, and possibly other products such as heparin and 5-hydroxytryptamine, are considered to take part in anaphylactic immune reactions (Keller, 1966). However, adjuvant disease is believed by most authorities to be due to hypersensitivity of the delayed type (Waksman, Pearson, and Sharp, 1960; Sharp, Waksman, Pearson, and Madoff, 1961; Pearson, 1963; Waksman and Wennersten, 1963; Pearson and Wood, 1964). It remains possible that mast cells contribute to the first part of a biphasic inflammatory reaction in adjuvant disease.

Discussion

It is shown that mast cells are the first cells to appear in abnormal numbers in the inflamed tissues in rat adjuvant arthritis.

Mast cells are normally large, numerous, and conspicuous in many rat tissues, including the synovium and the periarticular structures (Rosate, 1959; Selye, 1965). They can be recognized with ease in sections stained with haematoxylin and eosin and have prominent metachromatic cytoplasmic granules. Identification of mast cells is more difficult after the granules have been lost. Under these conditions they are recognized as swollen cells with a central nucleus and basophilic cytoplasm in which a few peripheral granules are occasionally seen (Parratt and West, 1957).

In adjuvant arthritis, a sequence of changes is noted in the mast cell population. There are increased numbers, apparent infiltration by mast cells of the synovial cell layers lining joint spaces, and degranulation. The first two changes begin at approximately the same time. More than the normal proportion of mast cells is seen by the 5th day, but this number markedly and suddenly increases on Day 6 and then decreases, at first sharply and then more gradually. The literature on rat adjuvant arthritis contains no previous reference to this cell population.

Mast cells are believed to be the major source of histamine in rats (Keller, 1966). Keller has discussed the release of histamine by degranulation and sometimes by cytolysis of mast cells. In the present study, both mechanisms could explain the fall in the numbers of mast cells after the 5th day, the former resulting in mast cells which are more difficult than normal to identify, the latter decreasing the actual numbers. Why mast cells should contribute to the inflammatory reaction in adjuvant disease is more difficult to resolve. According to Sheldon and Bauer (1960), tissue mast cells initiate acute inflammation at many sites of injury. These cells, the histamine released from them, and possibly other products such as heparin and 5-hydroxytryptamine, are considered to take part in anaphylactic immune reactions (Keller, 1966). However, adjuvant disease is believed by most authorities to be due to hypersensitivity of the delayed type (Waksman, Pearson, and Sharp, 1960; Sharp, Waksman, Pearson, and Madoff, 1961; Pearson, 1963; Waksman and Wennersten, 1963; Pearson and Wood, 1964). It remains possible that mast cells contribute to the first part of a biphasic inflammatory reaction in adjuvant disease.

FIG. 7 Distribution of mast cells in right hind foot connective tissue 6 days after injection of mycobacterial adjuvant into the opposite foot. Mast cells stand out as small black bodies. Toluidine blue. ×150.
Degranulation of mast cells in adjuvant arthritis is not observed until polymorphs have crossed vessel walls and gained access to the tissues. This confirms the findings of Janoff and Schaefer (1967) who demonstrated that polypeptides from the lysosomes of polymorphs effectively degranulated mast cells. It also suggests that the increase in vascular permeability, with oedema, found as the earliest histological feature of adjuvant arthritis, is not necessarily dependent upon the release of mast cell histamine.

In normal human synovium, mast cells do not occur in the layers of synoviocytes (Barnett, Davies, and MacConaill, 1961); nor were they seen in our control animals. In the injected rats, very occasional mast cells appear to have gained access to the layers of synovial cells lining joint cavities, but none was observed within the joint spaces. This is not the case with other cells, particularly polymorphs, which constitute the predominant cell population in joint exudates. Intra-articular histiocytes, morphologically similar to and possibly arising from the synovial cell lining, are much fewer in number and lymphocytes are uncommon in joint exudates. A preponderance of mononuclear cells in the synovium and of polymorphs in the synovial fluids, of course, a common finding in other forms of animal and of human arthritis (Gardner, 1965; 1969). The histological features of the joint effusion of adjuvant arthritis are those of a nonspecific acute inflammatory exude. Polymorphs appear later than

**FIG. 8** Venule in inflamed connective tissue in early rat adjuvant arthritis. Mast cells are seen as large cells with ovoid or round nuclei (top right and lower right). H and E. × 540.

**FIG. 9** Polymorphs in early adjuvant arthritis, accumulating locally in venule (centre) and migrating into adjacent connective tissue. H and E. × 320.
lymphocytes in this inflammatory reaction and they accumulate within and engorge blood vessels before gaining access to the tissues. Lymphocytes do not accumulate within vessels in the inflammation of adjuvant arthritis; they are first seen in a perivascular location from which they migrate to the tissue spaces. When polymorphs migrate, they do not appear to delay in a perivascular location but quickly enter the tissue spaces.

It is confirmed that the inflammation of adjuvant disease begins in synovial and periarticular connective tissue and that it spreads directly to involve skeletal muscle and periosteum, the sequence observed by Jones and Ward (1963). Oedema, an early sign of inflammation, can be identified at the site of the developing inflammatory reaction before cellular infiltration (Movat, 1966), a sequence noted in the present study.

The precise mechanism of the inflammation of adjuvant arthritis has not yet been defined. Acid-fast bacilli have been shown within adjuvant droplets in primary and secondary lesions (Akamatsu, Nishizawa, Watanabe, and Kumagai, 1966) and radioactivity has been demonstrated around joints after the intradermal injection of labelled adjuvant (Jones and Ward, 1964), suggesting that components of adjuvant are conveyed to, and may localize in, synovial tissue. The dissemination of adjuvant or of the products of adjuvant alone could incite an inflammatory reaction, a view supported by Lack (1968). Thus, Newbould (1964) showed that removal of lymph nodes draining an injection site within 5 days of the administration of adjuvant prevented the development of secondary lesions, while lymphadenectomy on or after the 7th day did not. This suggested that, if adjuvant dissemination were necessary for initiation of the inflammatory process, this dissemination would probably occur by the 5th or 6th day. That this time is critical was also confirmed by Quaglialo and others (1968; 1969) who demonstrated that either Rubidomycin or an antilymphocytic serum would prevent adjuvant disease only if given before the 5th day.

The theory that adjuvant disease is due to delayed hypersensitivity is persuasive (Waksman and others, 1960; Sharp and others, 1961; Pearson, 1963; Waksman and Wennersten, 1963; Pearson and Wood, 1964). The appearance of lymphocytes before polymorphs supports this view and that of Burstein and Waksman (1964a). These authors point out that the discrepancies between their findings and those of Jones and Ward (1963) may arise first from the latter’s failure to examine joint sections 1 to 2 days before the onset of clinical arthritis, and secondly from the differences in strains of rat, in adjuvant mixtures, and in routes of inoculation employed. With autoradiographic methods, Burstein and Waksman (1964b) have shown that the lymphocytes which appear in the tissues are haematogenous in origin and that a significant proportion of histiocytes are derived from these lymphocytes. This appears to explain why histiocytes appear later than lymphocytes. The combined lymphocytes and histiocytes usually outnumber the polymorphs, even in well-established lesions.

Summary

A survey has been made of the cell populations of the joints and periarticular tissues of rats during the phase of onset of experimental adjuvant arthritis. The investigation began before clinical evidence of arthritis was identifiable and extended from the 5th to the 12th day after the injection of adjuvant.

The phenomena of inflammation were first recognized in synovial and perisynovial zones; they then spread to adjacent tissue. Mast cells were the first to appear in excess, although oedema was the earliest disturbance suspected microscopically. Aggregates of lymphocytes were seen before polymorphs accumulated in extravascular planes, and were first recognized as perivascular clusters. With the spread of polymorphs into perisynovial tissues, histiocytes infiltrated these zones and mast cells became degranulated. An inflammatory joint exudate appeared only after synovitis was established; polymorphs predominated in these effusions and mononuclear cells in the tissues.

This work would not have been possible without the generous support of the Arthritis and Rheumatism Council for Research.

We are grateful to Mr. George Munroe for expert technical assistance and to Mrs. Mei Ling Foo for help with the preparation of the manuscript.

References


——— (1964b) Ibid., 37, 195 (The pathogenesis of adjuvant disease in the rat. II. A radioautographic study of early lesions with the use of H\(^2\)-thymidine).


JONES, R. S., and WARD, J. R. (1963) Arthr. and Rheum., 6, 23 (Studies on adjuvant-induced polyarthritis in rats. II. Histogenesis of joint and visceral lesions).

— — (1964) 'Proceedings of the IVth International Symposium on Reticulo-endothelial systems, Otsukyoto, Japan', p. 298 (Tissue distribution of radioisotopically labelled components of adjuvant).


— — (1964) J. exp. Med., 120, 547 (Passive transfer of adjuvant arthritis by lymph node or spleen cells).


ROSATE, A. (1959) Biol. lat. (Milano), 12, 451 (Mastocytes (mast cells) of the synovial stratum in normal and experimentally immobilized diarthroses) (Italian).


WAKSMAN, B. H., PEARSON, C. M., and SHARP, J. T. (1960) J. Immunol., 85, 403 (Studies of polyarthritis and other lesions induced in rats by injection of mycobacterial adjuvant. II. Evidence that the disease is a disseminated immunologic response to exogenous antigen).

The mast cell in early rat adjuvant arthritis.

A Gryfe, P M Sanders and D L Gardner

*Ann Rheum Dis* 1971 30: 24-30
doi: 10.1136/ard.30.1.24

Updated information and services can be found at:
http://ard.bmj.com/content/30/1/24.citation

**Email alerting service**

Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

**Notes**

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/