Protein sulphhydril depression during adjuvant arthritis

A. LORBER, J. LEEB, P. CARROLL, JR., W. BAUMGARTNER, AND V. HILL
From the Clinical Immunology and Rheumatic Disease Section, Department of Medicine, University of Southern California, Los Angeles, and the Veterans Administration Hospital, Long Beach, California

Lorber, A., Leeb, J., Carroll, P., Jr., Baumgartner, W., and Hill, V. (1975). Annals of the Rheumatic Diseases, 34, 346-349. Protein sulphhydril depression during adjuvant arthritis. The changes in plasma protein sulphhydril level were measured during the course of adjuvant-induced arthritis in rats. Major depressions in the plasma sulphhydril level occurred at the onset of adjuvant disease, and the extent of the depression was related to the severity of the disease. Plasma sulphhydril levels remain unchanged when the systemic arthritis is suppressed by inclusion of a competing antigen in the adjuvant. Changes in sulphhydril content of the plasma were shown not to be due to weight loss or decrease in plasma protein level.

Deficiency of sulphhydril (SH) levels of plasma proteins has been reported in patients with connective tissue disorders (rheumatoid arthritis, systemic lupus erythematosus, rheumatoid vasculitis) (Lorber, Boyv, and Chang, 1971). A study was conducted using adjuvant arthritis (AA) as a model to gain insight into the significance of this SH depression with respect to the aetiology and/or pathogenesis of such disorders. The systemic nature of AA is indicated by the widespread biochemical disturbances which accompany the chronic joint inflammation. In this sense, AA may be comparable to the above human collagen vascular disorders. The actual development of AA can be readily quantified and distinguished from an initial phase of acute inflammation at the site of injection of the adjuvant, thus providing an excellent experimental vehicle for studying the SH depression during the various phases of the disease.

Specifically, this study attempts to establish the effect on plasma SH values of the initial, acute inflammatory phase characterized by inflammation of the injected joint and of the various phases of AA. Changes in serum SH levels at these various phases of the disease were compared to the arthritis score, changes in plasma protein levels, and body weight. SH values were also determined in animals in which the AA was suppressed by a competing antigen without, however, influencing the initial inflammatory response.

Materials and methods

For all experimental and control groups, 5 male Lewis strain rats (Microbiological Associates) weighing 150–250 g initially, were used. Adjuvant arthritis was induced by intradermal injection of 0·1 ml Freund’s complete adjuvant into a foot pad of the left hind paw. The adjuvant was prepared by suspending Mycobacterium butyricum (Difco Laboratories) in paraffin oil (6·0 mg/ml).

The arthritis score was used as an index of the severity of the AA and was determined by observation of the extent of swelling, lesions, and redness of each joint, including the tail, with scores of 0 to 4 being assigned to each joint. The maximum score for one animal was 20.

Blood samples were collected into heparinized tubes by amputation of the tip of the tail. Plasma protein SH was determined radiochemically by addition of 0·1 ml plasma to 0·2 ml 203Hg p-chloromercuribenzenesulphonic acid (PCMBs, 1·18 mmol/l in 0·09 % NaCl), to a final volume of 1·5 ml with water followed by precipitation with 1·5 ml of 7% trichloroacetic acid (TCA) and centrifugation. Precipitates were washed twice with 3·75 % TCA, and protein bound 203Hg in the precipitates was counted in a well scintillation spectrometer. When larger sample volumes were available, serum sulphhydril levels were also measured by argentometric titration, and no significant differences were found. In those occasional samples which showed signs of haemolysis, corrections were made for heme protein SH in the plasma samples by determining (before TCA precipitation) the heme peak at 415 nm in both the plasma sample and an appropriately diluted haemolysate of cells from the same animal. The haemolysate was analysed for SH in the same manner as the

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Correspondence to: Dr. A. Lorber, Memorial Hospital Medical Center, 2801 Atlantic Avenue, Long Beach, California 90801, U.S.A.
plasma. From the calculated SH per heme OD in the haemolysate and the OD of the heme in the plasma sample, the heme protein SH in the plasma sample was calculated and subtracted from the total protein SH.

Suppression of AA with competing antigen (Pearson and Wood, 1964; Gery and Waksman, 1967) was achieved by homogenizing human IgG (7S fraction) with Freund's complete adjuvant before injection. Booster shots of the IgG alone (in saline) were administered on days 8, 26, and 44. The production of antibody against human IgG was monitored by means of a latex agglutination test (Hyland RA test). Albumin was measured fluorometrically (Rees, Filides, and Laurence, 1954). Total plasma protein was determined by the biuret method (Reinhold, Seligson, Schreiner, Riddle, Sharon, and Vanderan, 1953).

**Results**

The serial changes which occur in plasma SH levels during the course of adjuvant arthritis disease are shown in Fig. 1. It can be seen that the disease can be divided into 5 different phases on the basis of disease activity (Baumgartner, Beck, Lorber, Pearson, and Whitehouse, 1974). During the nonimmune inflammatory period (phase I), a rapid swelling of the injected foot takes place, accompanied by a slight decline in plasma SH levels. No significant swelling is seen in the un.injected feet during this period. The onset of AA is signified by the appearance of marked swelling in all extremities, occurring between 8 and 18 days after injection (phase II). Plasma SH levels show a marked decline during this period, and numerous other signs of disseminated disease (vascular lesions, balanitis, splenomegaly) become evident. Plasma SH levels continue to be markedly depressed (P > 0.05 by the 't' test) during the sustained phase of the disease (phase III), which includes the period from about day 18 to day 48. During the recovery phase of the disease (phase IV, 50–65 days after injection), many of the clinical manifestations of the disease disappear as SH levels return to normal.

The temporary decrease in SH values which accompanied a slight rise in the arthritis score between 70 and 80 days may have been due to a relapse,
which is known to occur in AA. The arthritic score does not return to zero due to permanent joint deformity and osteogenesis.

When the sustained phase of AA (phase III) was blocked by the addition of human IgG to Freund's adjuvant, the decline in plasma SH did not take place nor did the development of systemic lesions or generalized arthritis (Fig. 2). Acute inflammation was limited entirely to the injected joint (phase I). Evidence of humoral immune response to the IgG was shown by the development of antihuman IgG titre, ranging from 160–1280 reciprocal dilutions. This response, however, was not correlated with depletion of plasma protein SH.

Although adjuvant rats lose substantial amounts of body weight during the course of the disease, total plasma protein content remains relatively stable (Fig. 3). Some changes in the albumin/globulin ratio are seen, but similar alterations can be observed in normal rats when restricted to 50% of the normal food intake (which is equal to the most depressed food intake observed during AA). These starved animals suffered weight losses similar to the AA rats and similar alteration in A/G ratio (Zahiri, Gagnon, Ayotte, and Laurin, 1969), but there is no decline in SH content with food deprivation (Table).

Discussion

It appears that the sulphhydryl depression of plasma proteins in adjuvant arthritis is associated with the protracted inflammatory phase of the disease. The major depression of SH level occurs during this phase, and those changes which occur during the acute phase are more moderate, and appear to be transitory. Other biochemical changes have been reported to occur in AA, including depressed serum albumin levels, raised fibrinogen and a-2-macroglobulins, and impaired liver microsomal oxidase activity (Lowe, 1964; Weimer, Wood, and Pearson, 1968; Beck and Whitehouse, 1973). All of these changes, however, are expressed during the initial acute inflammatory phase and can be induced by agents such as croton oil and carrageenan without leading to AA (Goldstein, Shemano, Demeo, and Beiler, 1967; Varsa-Handler, Handler, and Gordon, 1967; Glenn, Gray, and Kooyers, 1965).

The observed SH depression was not due to the acute inflammation, since those rats in which AA was suppressed by the competing antigen did not show any depression. Neither was the depression due to decreased food consumption or loss of body weight, since starvation of normal animals did not lead to SH declines. Although the albumin/globulin ratio did vary during AA, the declines noted in albumin level could not account for more than one half of the decrease in SH, assuming 1 mol SH/mol albumin. Similarly, reports have shown wide fluctuations in plasma SH of rheumatoid arthritic patients, unrelated to major changes in the composition of plasma proteins (Lorber, and others, 1971). This suggests that oxidation of protein SH groups, or formation of new proteins deficient in SH, is occurring. Increased heterogeneity in electrophoretic patterns has previously been reported (Lorber, and others, 1971). Biochemical abnormalities, such as albumin changes and SH depression, may both result from a more basic liver malfunction, the existence of which is indicated by the impaired drug hydroxylation in the liver in AA (Beck and Whitehouse, 1973).

Changes in SH levels similar to those observed in the AA model could be postulated as contributory or aetiological factors for human rheumatoid disease on the following basis:

1. Sulphhydryl depression can lead to abnormal protein configuration (e.g. aggregation of IgG) which in turn serves as an antigenic stimulus for autoimmune process.

2. Depletion of SH groups in the local environment of immunocompetent cells (e.g. in the spleen, thymus, or lymph nodes) could alter or disturb the normal function of lymphocytes residing there. Sulphhydryl groups, and their availability to the many enzyme and membrane systems, may be essential for maintenance of the normal immune state.

3. The activation of complement C3 and neutral collagenase, and loss of normal lysosomal permeability barriers, may also be associated with decrease in protein SH levels.

Although on theoretical grounds the sulphhydryl depletion appears to be a plausible factor in the induction and/or pathogenesis of adjuvant arthritis,
the cause of the depression has not been established. One possible mechanism, which will be discussed in a subsequent publication, is that the SH depression is caused by an increase in oxidative stress which, in turn, leads to damage of the membranes of various cell organelles (lysosomes, mitochondria, etc.). The increase in oxidative stress may be due to the production of hydrogen peroxide as a result of enhanced phagocytic activity of polymorphonuclear leucocytes and macrophages.

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