in dermatology site of predilection is an unreliable criterion.

All three diseases may attack the mucous membranes of the mouth and eyes, but in *E. bulbosum* the chief distribution of the eruption falls on the backs and palms of the hands and similar situations on the feet and the extensor surfaces of the knees and elbows. In *Dermatitis herpetiformis* the eruption may come anywhere, with perhaps a special liability to attack the axillae and groins, and *Pemphigus vulgaris* has, as far as my experience goes, no favourite sites. Subjectively, *E. bulbosum* is usually associated with burning and tingling, and *D. herpetiformis* nearly always with maddening pruritus. Pemphigus may either give rise to a sense of soreness or of itching, though it is fair to admit that some observers classify all pruriginous pemphigoid eruptions with *D. herpetiformis*.

Finally, I should like to refer to two other forms of generalised acute eruption. Dermatologists have generally taken the suffix "ide" as a convenient term to denote that the eruption is of haematogenous spread—e.g., syphilide, tuberculide. Of recent years it has been observed on the continent, where the occurrence is fairly common, and also in this country, where it is, I think, rare, that in certain cases of ringworm of the scalp a generalised, non-progressive eruption of minute scaling papules, occurring either singly or in ringed groups, may appear on the trunk. Without going into controversial detail, I may say that the doctrine brought forward is that this eruption is due to the entry of particles of the fungus into the blood stream, and the subsequent deposition therefrom into the skin. Hence it has been named "Trichophytide." It somewhat resembles the eruption commonly called dry seborrhoeic eczema of the trunk.

The second form is analogous to this, and was described by me. In certain cases of impetigo of the scalp in children one finds also a similar eruption of minute follicular scaly papules occurring singly or in groups and coming out acutely on the trunk. It is not bullous or even vesicular, and just as the "Trichophytide" does not resemble the original, local inoculation of ringworm of the scalp, in like manner this eruption, secondary to the impetigo, does not resemble the true bullous lesion of inoculated impetigo. By analogy with the "Trichophytide" I have called this the "Streptococcide." It is far from uncommon in England, and the only thing that surprises me is that it had not been described before I called attention to it. The diagnosis in both cases is, of course, simple, provided that due attention is paid to the general examination of the patient's skin.

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**STREPTOCOCCI: THEIR TOXINS AND ANTITOXINS.**

**BY**

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Our time will be spent mainly in considering the streptococci, and chiefly the pathogenic ones, under the headings of classification and pathology—i.e., what the streptococci are and what they do. It is probably fair to say that the attention of bacteriologists has recently, and more fruitfully, become more centred on what these organisms do than on what they are.

**CLASSIFICATION.**

In all scientific work accurate classification is of fundamental importance, and many attempts have been made in the past to classify these streptococci by the careful investigations of eminent workers in bacteriology; light has gradually dawned, and we are to-day almost in sight of a rational view of the group.

We may for the sake of clearness take some historical liberties and consider each of these attempts in sequence.

The first attempts were naturally based on morphology from the time when Pasteur drew his historical chain of cocci on the blackboard at a meeting in Paris, and soon such names as longus and brevis arose. Throughout all this work attempts were constantly being made to link these various groupings with the various diseases produced by the cocci. Thus, *Streptococcus longus* was thought by von Lingelsheim, 1899, to be pathogenic, and the short-chained variety, *brevis*, much less so.

Next, cultural characters were carefully explored and groupings were arranged on the basis of fermentation of carbohydrates, &c.; now arose the names *pyogenes, fecalis*, &c. This work is largely due to English bacteriologists—Gordon, Andrews, and Horder (1902–1906).

Blood plates were next (Schottmüller, 1903, Th. Smith and Brown, 1915) extensively used and a division was made into: (1) the non-haemolytic group; (2) those giving a green ring, the *viridans* group; and finally (3) the haemolytic group, which produce a clear ring around the colony, a so-called "hemolysis," though the appearance suggests that there has been also decolorisation. This latter group gives hemolysis of red blood cells in suspension in a test-tube. It was clearly pointed out by these workers that there was a transition from cocci giving rapid and complete hemolysis to those with feeble haemolytic power, but the

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* Lecture delivered at St. Mary's Hospital, May, 1927.
grouping “hæmolytic” and “non-hæmolytic” is convenient.

If one requires a bird’s-eye view of the results up to this period, that given by Park and Williams is perhaps one of the most convenient, with its division into hæmolytic and non-hæmolytic groups. The beta group of Smith and Brown includes *S. pyogenes, anginosus,* and *equi*; the alpha group is divided into two groups: those producing methæmoglobin—e.g., *faecalis* and *salivarius*—and those not producing methæmoglobin—e.g., the gamma type, *anaëmolyticus,* &c.

The next attack during the immediate past has been by agglutination and absorption methods (Dochez, 1919, Gordon, &c., and recently by Smith, Griffith, and James, &c.). The technical difficulties were great, and only within the past few years has it proved practicable by various methods of agglutination in buffered broth, rapid subculturing, or rapid microscopic testing, to work with sufficiently stable emulsions to give reasonably consistent results. It was at first thought that by agglutination we might be able to separate clearly the pathologically defined groups—e.g., scarlet fever, erysipelas, puerperal fever, &c., but later work has shown that by careful agglutination and absorption work the streptococci can be subdivided into a number of main groups and probably a considerable number of subgroups.

**Overlapping of Groups.**

The groups definitely overlap. Thus, the scarlet fever strains can be fairly clearly divided into three main groups and an unknown number of smaller groups, but some cultures derived from puerperal fever and erysipelas are indistinguishable from those in the main scarlet fever groups.

Finally, since the use by the Dicks (1924) of the intradermic method in human beings, the attempts have been mainly directed to seeing what the pathogenic actions of the various streptococci are, and whether any classification can be based thereon. But here again we are faced with great confusion. At first it was thought that the scarlet fever streptococcus would produce a scarlet fever toxin and would on injection into animals cause the production of a scarlet fever antitoxin; similarly it was thought that the puerperal streptococcus would give a characteristic toxin and antitoxin, that “cellulitis” and “septicæmic” strains would similarly give clearly distinguishable toxins and antitoxins.

Amoss and Birkhaug concluded as the result of their work that the erysipelas streptococcus had clearly specific toxin-antitoxin relationships quite independent of the scarlet fever streptococcus. It is not certain that further experience will confirm this clear specificity.

It is already clear that there is some overlap amongst the main groups. How much is not yet determined. Some workers go so far as to conjecture that there is one antigen only—i.e., that the toxins of the streptococci causing scarlet fever “septic sore-throat,” erysipelas, puerperal fever, cellulitis, &c., are all one and the same toxin, and that whatever toxin used be the same antitoxin is obtained.

**Methods of Investigation.**

There are several methods of investigating this problem, and they are all being actively pursued at the present time. If we take the *S. scarlatinae* of Dochez and the Dicks and make a toxin from it, we find that when the toxin is injected in suitable doses intradermally into human beings, some give a positive red reaction, and others give no response—the so-called negative reaction. A group of people who have had scarlet fever will have a much higher percentage of negative reactions than those who have never had scarlet fever or been in contact with it.

(1) If we can obtain large groups of people who have had, e.g., “septic sore-throat” or puerperal fever, we can test them with the scarlet fever toxin and with a culture filtrate or “toxin” made from cultures of streptococcus obtained from puerperal fever or tonsillitis. If we end by finding that all the people who give negative reactions to the scarlet fever toxin also give negative reactions to the tonsillitis and puerperal fever toxins, and those giving a positive reaction to one toxin give a positive to the other two, we shall be justified in assuming that there is a close antigenic relationship between the three toxins and presumably between the three diseases. The results from this line of research are at present too meagre to justify any confident conclusion, but apparently there is a considerable degree of agreement in the groups—i.e., the majority of people who give a positive response to the Dick scarlet fever test will also probably give a positive response to the injection of the other toxins, and “Dick negative reactors” will usually be “negative” to the other toxins.

(2) We may test the hypothesis in another way—make an antitoxin to one of the streptococci—e.g., scarlet fever antitoxin, and test its effect on streptococci obtained from the other diseases and on their toxins. Parish and Okell have shown that, in the rabbit test, scarlet fever antitoxin has a significant protective action against a number of hemolytic streptococci obtained from other diseases.

(3) We may mix the scarlet fever antitoxin with the toxin of, e.g., puerperal fever, and inject the mixture into the skin of subjects who give a positive response to the injection of “puerperal toxin.” It is found in practice (Eagles, McLaughlin) that a considerable degree of overlap occurs; thus scarlet fever antitoxin will neutralise tonsillitis toxin or puerperal toxin in a considerable number of instances.

(4) We may use one antitoxin for the treatment of the other diseases—e.g., scarlet fever antitoxin for tonsillitis, puerperal fever, &c. Here again the

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1 Pathogenic Micro-organisms, 1925, p. 303.
results are too few to justify any definite conclusion, but evidence giving some support to the "unitarian" hypothesis is beginning to accumulate.

(5) We may find a dose of the culture of — e.g., scarlet fever streptococcus that will kill laboratory animals — e.g., rabbits, and see if the other antitoxins give any protection. The table which my colleagues Parish and Okell have kindly allowed me to use shows that these other antitoxins have considerable protective effect against the scarlet fever streptococcus and other heterologous haemolytic streptococci.

(6) We may inject intradermally into a scarlet fever rash, antitoxins made from cultures of — e.g., puerperal or cellulitis strains, to see if the Schultz-Charlton blanching results. We have reports showing that in a few recent observations definite blanching was so produced.

(7) We may inquire whether people who are immunised with scarlet fever toxin until they give a negative reaction to the Dick test will be immune against tonsillitis or puerperal fevers, &c.

THE "UNITARIAN" VIEW.

Where the story will lead us shall not know for a year or two. But it is of extraordinary interest that at present an amount (though not all — cf. Park, Blake, and co-workers, &c.) of recent immunological evidence seems to be almost leaning towards the hypothesis that all pathogenic streptococci are identical in their pathogenic and "toxic" action. But if the implications of this "unitarian" hypothesis conflict with the very large body of evidence enshrined in clinical medicine, we must be very hesitant to adopt the hypothesis. If immunology hints that follicular tonsillitis and puerperal fever, for example, are caused by the same organism as scarlet fever, we must inquire what clinical evidence there is in support of this view. With regard to tonsillitis, various clinicians, after a lifelong experience of the exanthemata, have inclined to the view that one may have scarlet-fever-without-rash — i.e., "septic sore-throat" or follicular tonsillitis; and others have maintained that there is a close relationship between puerperal fever and scarlet fever. Goodall and Washburn \(^2\) wrote thus: "Nurses suffering from erysipelas have conveyed puerperal fever to lying-in women; and medical men and nurses have contracted erysipelas when in attendance upon cases of puerperal fever."

So far we may account the clinicians on the side of the hypothesis. But if these diseases are essentially identical, one disease should give rise to the other. Can epidemiology help us here? Can we on the one hand find communities in which one of these diseases was common and yet never gave rise to the other diseases?

In the old military epidemics of "septic sore-throat" in former times, did the introduction of this disease into a new batch of troops lead to the outbreak of sore-throat or also of scarlet fever and erysipelas, &c.? When puerperal fever raged a century ago, so that midwifery hospitals had to be closed, did erysipelas and scarlet fever rage simultaneously in those hospitals? I have been unable to discover the evidence dealing with these points which almost certainly exists in medical literature. Can we find clear instances in which an isolated community free of these diseases became infected with, say, scarlet fever through the introduction of one case of tonsillitis, puerperal fever, erysipelas, or cellulitis into its midst?

I lived for some years in a rather isolated small town overseas, in which I saw puerperal fever and follicular tonsillitis from time to time, but never a case of scarlet fever, nor had one been recorded in the history of the town so far as I could find.

It may be that a streptococcal disease of any kind requires the streptococcus plus another factor. It will be remembered that for years the \(B.\) \(suispestifer\) was thought to be the cause of swine fever in pigs, for by the injection of this organism a disease apparently identical with swine fever can be produced. Dorsett in his masterly work showed that only one feature was lacking — i.e., infectivity. The pigs injected with the culture developed what appeared to be the typical disease, but his great discovery was that these pigs did not infect others, whereas the natural disease was highly infectious. Further investigation showed that a filterable virus was the cause of the naturally occurring highly infectious swine fever, and that the other bacillus was so common in pigs that, when the real swine fever disease occurred, the bacillus infected the animal’s body and produced the characteristic intestinal lesions. It is possible to have a herd of pigs infected with the virus — i.e., true highly infectious swine fever, and another herd infected only with the \(B.\) \(suispestifer\), showing fever and intestinal lesions. So close is the alliance and so universal is the bacillus that it is to-day safe in England for administrative purposes to diagnose the presence of swine fever because intestinal lesions produced by another infecting agent — the \(B.\) \(suispestifer\) are present.

The Dicks have described in human volunteers the production of true clinical scarlet fever following the application of their culture of streptococcus. The crucial experiment is lacking. Would the patient have infected other people? In other words, had he the natural infectious disease? It is doubtful if this experiment will ever be done. In its absence we must depend on the collateral lines of evidence above referred to.

We may sum up by saying that some of the lines of investigation at present being pursued point to the suggestion that all the ordinary diseases caused by haemolytic streptococci are different manifestations of the same disease. It is by no means certain that clinical and epidemiological evidence will support this hypothesis. For the moment, however, we must keep an open mind,

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\(^2\) Infectious Disease, 1896, p. 330.
and, in the treatment of the diseases caused by the haemolytic streptococci, be prepared to use whatever serum is on reasonable clinical evidence found to cure the given disease.

**Present Knowledge.**

It would perhaps be useful at this stage to take a general view of our knowledge in this field. Where the ground is shifting under our feet from day to day, it is of some service to consider which landmarks are so firmly established as to be probably permanent, which have been set up or are in process of being erected, and which may probably not stand the stress of time.

Streptococci mainly of the haemolytic variety, and with various fermentation reactions, are constantly present in various "septic" conditions. It is practically certain that they are the real cause, for we know from experiment on human volunteers that with cultures of *S. erysipelas* characteristic erysipelas has been produced. The production of experimental erysipelas with cultures of streptococci obtained from lesions of erysipelas, or from septic ones (presumably *S. pyogenes*), was for some time a method of "treatment" of inoperable cancer. (In this connexion an old experiment by Koch and Petruschky records that in a volunteer erysipelas was produced by the injection of streptococci isolated from erysipelas. After recovery, experimental erysipelas was produced again on the same area. The experiment was repeated ten times.)

The many unfortunate infections of pathologists, after accidental finger-pricks while doing autopsies or while handling cultures of streptococci, leave little doubt that the streptococci are the real cause of these various manifestations of lymphaengitis, cellulitis, and even septicemia. Similarly, we are entitled to believe that streptococci are the cause of severe puerperal fever with septicaemia.

With regard to the *viridans* group of haemolytic streptococci, they are so frequently found in certain pathological conditions, notably subacute endocarditis with bacteriæmia, that it is reasonable to believe that they are the cause thereof.

With regard to scarlet fever, the Dicks and Nicolle believe that they produced typical scarlet fever in human subjects by the inoculation of cultures of the *S. scarlatinae*. They have not shown, what from a coldly scientific point of view would be interesting and valuable, that the disease so produced is infectious to others and therefore in every respect resembles natural scarlet fever.

It is further reasonable to believe that most of the symptoms at least of uncomplicated scarlet fever are due to the "toxin" of the *S. scarlatinae*, for, by the injection of sterile filtrate made from a culture of the streptococcus, subcultured perhaps hundreds of times since its isolation from the human subject, one can produce pyrexia, headache, "strawberry tongue," the characteristic throat condition, albuminuria and rash followed by peeling—in other words, a pathological condition indistinguishable from naturally occurring scarlet fever. It is practically certain that this so-called "scarlatinoid syndrome" is non-infectious.

It appears to be established beyond doubt that the response to the Dick test is related to immunity against scarlet fever, and that scarlet fever antitoxin is effective in the treatment of uncomplicated scarlet fever; it is probably without effect on the late septic or pyogenic conditions occurring in scarlet fever. Further, by immunising with Dick toxin one can make positive reactors negative to the Dick test, and this negative reaction indicates a reasonably high immunity against scarlet fever.

When we consider the more recent work, we find a distinct suggestion that the toxins and antitoxins of all the haemolytic streptococci are so closely related as almost to amount to identity. But whether this view will be entitled to a "land mark," time and further experience alone can tell. Current clinical views may be summarised under some such headings as the following:—

(a) *Experimental evidence.*—(1) *S. pyogenes* when injected has caused at least local septic conditions. (2) *S. erysipelas* when injected has caused erysipelas. (3) *S. scarlatinae* when rubbed on the tonsils has caused scarlet fever.

(b) *Conclusions reasonably based on much clinical observation.*—Streptococci, usually haemolytic, are the cause of puerperal fever, mastoiditis, septic pneumonia, &c. *S. viridans* is the usual cause of subacute infectious endocarditis.

(c) *Current hypotheses.*—Truth dependent on experience in the future. Because streptococci occur in dental septic conditions, they cause constitutional symptoms. Because streptococci (enterococci, &c.), occur in the bowel in large numbers they cause diarrhoea, &c.

Although the title of our prescribed subject deals with the toxins and antitoxins of the streptococci, I may be permitted to refer to the other aspect of streptococcal attack, the so-called "septic" manifestations. A hypothesis which we might use for the moment for the sake of clearness is that the *S. scarlatinae* has two modes of attack—the "toxic" and the "septic." In violent epidemics, it is the early acute toxic attack which kills most patients; in mild epidemics it is the later "pyrogenic" or "septic" manifestations, in which the living organisms settle down locally in the glands, joints, mastoid, &c., that cause the greatest harm.

It is a curious thing that in the Parish-Okell rabbit method, scarlet fever antitoxin gives complete protection against the first "toxic" attack of the streptococcus, which is rapidly fatal to unprotected rabbits, but that this antitoxin apparently gives but small or no protection against the development of the later septic lesions in joints, &c. Similarly, it is probable that no scarlet fever antiserum now available has any direct effect on the late septic complications of scarlet fever. It is probable that the solutions of this difficult part of the problem will be found in the investigation of cellular
immunity with which the Medical School of St. Mary's Hospital has been for so long identified, and that future work may lead to a convergence of the "cellular" and "humoral" aspects of immunological research of the activities of the streptococci.

**Summary.**

The examination of agglutination, toxin and antitoxin relationships of the various "pathogenic" streptococci suggests that there is a very close relationship between the members of the group. The further analysis of this close relationship is the problem before us.

**Rheumatism.**

*By Charles Sundell, M.D. Lond., Physician in Charge of the Children's Department, Prince of Wales's General Hospital; Senior Physician to the Seamen's Hospital, Greenwich.*

For the purpose of this lecture it is proposed to confine the conception of rheumatism within definite limits excluding both osteo-arthritis with its atrophic and hypertrophic bone changes and rheumatoid arthritis with its rarefaction of bone, muscular atrophy, and its unexplained relation to absorption from septic foci obvious or occult. When these diseases are excluded there remains a large group of rheumatic conditions with which alone it is proposed to deal to-day.

Rheumatism is met with in two forms, acute and chronic. There is evidence that these are variations of one underlying state differing only in the mode and intensity of the response of the tissues to the irritation of a single morbid cause.

At first sight it is, perhaps, surprising that the dramatic signs and symptoms of rheumatic fever in a child should be regarded as identical in origin with the lumbago of his grandparent. But experience has much support to lead to this view. Careful inquiry into past history will again and again reveal the fact that an adult sufferer from fibrositis or "neuritis" has been marked as the prey of rheumatism during the years of his early childhood by growing pains, tonsillitis, or acute fibrule arthritis. It is not uncommon for the progress of a definite acute rheumatic fever into a slow crippling chronic rheumatic process to occur, or for alternations of acute joint affections and attacks of chronic fibrosic and muscular pains to be seen in the same individual. Such observations emphasise the latency of the rheumatic state, the chronicity of the disease, and its similarity in this particular to gout.

In connexion with the latency of this disease mention may be made of its frequent appearance in an occult form, sapping energy and destroying the sense of well-being. Children are particularly prone to this manifestation; they may suffer for weeks or months from fatigue before definite limb pains or joint swelling make their appearance. Delayed or restless sleep at night may be followed by intense sleepiness in the morning and lack of energy during the day. Such symptoms of ill-health in a child should always arouse the suspicion of rheumatism and the call for extra rest be generously answered. In the adult a very similar state of affairs is often seen. For weeks or months before the appearance of undoubted rheumatic signs there may exist a lowering of mental and physical vitality that makes work and play burdensome and clouds all joy in life. Under appropriate antirheumatic treatment these unpleasant symptoms are among the earliest to disappear.

Rheumatism may be described as a chronic state of the body characterised by certain chemical and physiological stigmata, essentially chronic in its course, liable to periods of great activity, marked by long periods of latency and by slowly progressing phases of slight activity, influencing particularly fibrous and synovial and muscle tissues, and producing a definite lowering of vitality both physical and nervous.

**Causes of Rheumatism.**

Infection with germs from mucous surface is for many a satisfactory explanation of the origin of this disease. The supporters of this view must, however, explain why: (1) case-to-case infection is unknown; (2) why members of certain families are peculiarly prone to the disease in successive generations; (3) why the great antirheumatic specific, salicylate, so powerful in rheumatism is so powerless in other infections with the same or closely allied organisms.

Careful considerations of problems such as these force one to the conclusion that the upholders of the infection theory see only part of the truth. If germs play a part in rheumatism, as they undoubtedly do in rheumatoid arthritis, may it not be that their rôle is a subsidiary one and that the real fundamental cause of the disease is to be sought in some aberration of body chemistry and physiology that renders the tissues vulnerable to germ attack?

Three features of the rheumatic state are of great interest and importance: (1) During periods of latency, quiescence, or merely fibrositic pain the skin is abnormally dry. (2) During such periods the temperature is always abnormally low. (3) During spontaneous active rheumatism and under the influence of artificially induced fever sweating is profuse and strongly acid in reaction. These points merit a little elaboration. The dry skin of the rheumatic is sometimes obvious; sometimes questioning is necessary to elicit the fact that sweating is always very slight or that
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