IMMUNOCHEMICAL MECHANISMS INVOLVED IN PENICILLIN HYPERSENSITIVITY IN EXPERIMENTAL ANIMALS AND IN HUMAN BEINGS

BERNARD B. LEVINE
Assistant Professor of Medicine, New York University School of Medicine, New York, N.Y.

In addition to its importance in clinical medicine, the human penicillin allergy system provides a comparatively well defined haptenic system for the study of fundamental immune mechanisms involved in human allergic diseases. Also, the understanding of the mechanisms involved in clinical allergic reactions to penicillin would in turn allow for more rational approaches to the prediction, diagnosis, prevention and management of these diseases. Penicillin allergies are, of course, immune diseases, i.e., they follow antibody/antigen reactions taking place in relation to tissue. Accordingly, the following kinds of information are important in elucidating their immune mechanisms: (1) the chemical mechanism of antigenicity of penicillin; (2) the antigenic specificities of immune responses to penicillin; (3) the kinds and quantities of antibodies synthesized in response to the administration of penicillin and, (4) the relationship of the nature of the immune responses to the occurrence of clinical allergic reactions. Studies in experimental animals provided a basis for later studies in human beings, and broaden the biological significance of the results obtained. This paper will deal with the chemical mechanisms of antigenicity of penicillin, the antigenic specificities of the immune responses of guinea pigs, rabbits and human beings to administration of penicillin, our early views on the possible immune mechanisms involved in clinical allergic reactions to penicillin, and the clinical implications of these views.

Antigenicity of Benzyl Penicillin

In order for low molecular weight chemicals to induce immune responses, they must first combine irreversibly with tissue macromolecules (most probably, proteins) to form the hapten-protein conjugates which induce the synthesis of specific antibodies. This view is widely accepted among immunologists, and is based on much data which show that of a large number of simple chemicals, only those which combine, through covalent linkages, with proteins or with protein-model compounds are capable of inducing immune responses (Landsteiner, 1945; Gell, Harrington and Pitt-Rivers, 1946; Eisen, 1959). With regard to many drugs which can cause allergic reactions, but would appear from their structural formula to be incapable of reacting irreversibly with proteins, it is reasonable to postulate that either a trace contaminant, a degradation product, or an intermediate metabolite of the drug may be the actual protein-reactive material (Landsteiner, 1945). With regard to the antigenicity of benzyl penicillin (penicillin G, PG), the available evidence indicates that PG first rearranges to form an isomer, D-benzyl-penicillenic acid (BPE) (Fig. 1), which is a highly reactive compound, and which is the protein-reactive compound responsible for the induction of immune responses to PG (Levine, 1960a, 1960b, 1961). This rearrangement appears not to be dependent upon in vivo enzymatic catalyses. It occurs in vitro as well as in vivo, and its rate is increased by low pH and by the presence of cations such as Cu++, Zn++, Fe++ (Florey et al., 1949; Clarke, Johnson and Robinson, 1949). BPE reacts irreversibly mainly with lysine ε-amino groups of proteins to form benzylpenicilloyl-amine haptenic groups (Levine, 1961; Levine and Ovary, 1961). (Fig. 1). Other haptenic groups are formed to a lesser extent, as will be taken up below. Alternatively, it may be considered that PG might react to form benzylpenicilloyl haptenic groups by a direct addition reaction of the β-lactam carbonyl to lysine amine residues of proteins. This view of the antigenicity of PG appears less likely from the following lines of evidence: (1) although this latter reaction does proceed at pH 11.5 with a half life in the order of minutes (Levine, unpublished data), it would proceed very much slower at pH 7.5, probably in the order of 1/10,000th as rapidly as at pH 11.5; (2) PG does rearrange to BPE under physiological conditions in vitro (Levine,
1960b; Clarke et al., 1949) and undoubtedly also in vivo. The rate of rearrangement in vivo is unknown. Based on the data of Eagle (1947) and on arguments given previously (Levine, 1960a), it may be as much as 10% per hour; (3) BPE is an exceedingly chemically reactive compound. In aqueous solution, pH 7.5 at 37°C, it is hydrolyzed at the rate of 11% per minute (Levine, 1961) at least 550 times more rapidly than is PG under identical conditions (Benedict, Schmidt and Coghill, 1946); (4) sensitization of guinea pigs (for allergic contact dermatitis, Levine, 1960a), rabbits (for serum antibodies, Levine, 1961b; 1964a) and human beings (for skin sensitizing antibodies, Levine and Price, 1964) and probably also for serum 7S and 19S antibodies (Levine, unpublished data) induces the formation of antibodies specific for a mixture of diastereoisomers of the benzylpenicilloyl haptenic group rather than for the α-diastereoisomer alone. If the “complete penicillin antigens” which induce antibody formation, were formed by the direct addition of the β-lactam carbonyl of penicillin to amine residues of protein, the resulting benzylpenicilloyl haptenic groups would be formed as a diastereoisomer (Levine, 1962). If the reaction proceeded through the intermediate formation of benzylpenicillenic acid, the benzylpenicilloyl groups would be formed as a diastereomeric mixture (Levine, 1961a). The finding that anti-PG antibodies are specific for a diastereomeric mixture of benzylpenicilloyl groups is thus consistent with the view that PG combines with tissue proteins through the intermediate BPE. An unequivocal proof of this view, however, requires precise measurements of the rate of rearrangement of PG to BPE under conditions which simulate physiological conditions in vitro, e.g. the reaction solution should contain traces of Cu++, etc. These arguments given above, although indicating that PG reacts with tissue proteins primarily through
the reactive intermediate BPE, do not exclude
the possibility that PG may also react directly
with proteins to some extent.

Antigenic Specificities of Immune Responses to
PG in Experimental Animals

Guinea pigs can be made allergic (contact
dermatitis type) to PG by repeated percutaneous
application of solutions of PG in a solvent
containing the anionic detergent Tween 80
(Levine, 1960a).

The haptenic specificities of allergic contact
dermatitis reactions were inferred from the
patterns of allergic cross-reactivity among PG
and its highly purified degradation products, as
well as from the chemical reactivities of PG
and its degradation products with protein-model
compounds (Levine, 1960a, 1960b; Florey,
1949), it is likely that most of the penicillin that
combines irreversibly with tissue proteins form
benzylpenicilloyl haptenic groups, whereas
only a comparatively small fraction of the PG
that reacts forms the penamalactic acid and
penicillamine mixed disulfide haptenic deter-
mindants. On this basis, the benzylpenicilloyl
group has been termed the major antigenic
determinant, and the two mixed disulfide
determinants (including other unidentified trace
haptenic determinants, see below) have been
termed the minor antigenic determinants of
penicillin hypersensitivity (Levine and Price,
1964; Siegel and Levine, 1964).

Antigenic Specificity of Rabbit Anti-benzyl-
penicillin Antibodies

Rabbits can be immunized to PG by injec-
tion of PG emulsified in complete Freund's
adjuvants followed by booster injections of
aqueous penicillin or aqueous suspensions of
procaine \(\text{e}\)-penicillin. Under these condi-
tions, sera containing as much as 400-500 \(\mu\)g/ml.
of antibody protein can be obtained (Levine and
Ovary, 1961). These antibodies are specific
for the benzylpenicilloyl haptenic group as
shown by quantitative precipitin techniques
(Levine and Ovary, 1961), by passive cutaneous
anaphylaxis (PCA) in guinea pigs (Levine and
Ovary, 1961; Levine, 1964a), and by passive
hemaggulination (De Weck, 1962). Benzyl-
penicilloyl haptenic specificity of these
immunological reactions was confirmed by
quantitative hapten inhibition techniques
(Levine and Ovary, 1961; Levine, 1964a; De-
Weck, 1962) using benzylpenicilloyl-\(e\)-amino-
caproate (Levine, 1962) as the univalent hapten.
The anti-benzylpenicilloyl antibodies show
specificity also for the lysine side chain of
proteins through which benzyl-penicilloyl
groups are predominantly bound to protein
Other studies have demonstrated that anti-
benzyl-penicilloyl antibodies produced by
immunization of rabbits with benzylpenicilloyl-
rabbit serum albumin conjugates are specific
also for structural areas of the carrier protein
(carrier specificity) (Levine, 1963). Some
evidence indicates that anti-benzylpenicilloyl
antibodies produced by immunization of rab-
bbits with PG also show carrier specificity to
as yet unidentified carrier proteins (Levine,
1964a). Thus, available evidence indicates that
the bulk of antibodies produced by immuniza-
tion of rabbits with PG show immunological specificity for a rather large antigenic unit comprised of the entire benzylpenicilloyl haptenic group, the lysine side chain, and possibly also for structural configurations of as yet undefined autologous protein carriers to which benzylpenicilloyl groups are bound. No antibodies specific for the benzylpenicillenic acid mixed disulfide or the D-penicillamine mixed disulfide haptenic groups were detected in rabbit anti-PG sera by precipitation and by PCA (Levine and Ovary, 1961; Levine, 1964a), nor could indirect evidence of antibodies specific for the D-benzylpenamaldic acid-mixed disulfide haptenic group be obtained (Levine and Ovary, 1961). DeWeck (1962) stated that he has detected anti-benzylpenicillenic acid disulfide antibodies in rabbit anti-PG sera, but no supporting data was given.

**Antigenic Specificities of Human Anti-Penicillin Antibodies.**

On an operational basis, two different classes of anti-PG antibodies can be detected in human beings, i.e. skin sensitizing antibodies and serum antibodies detectible by passive hemagglutination. From studies in other antigenic systems, skin sensitizing antibodies are believed to be γ1A globulins (Heremans and Vaerman, 1962; Fireman, Vannier and Goodman, 1963). Antigenic specificities of skin sensitizing antibodies can be studied by direct skin test for wheal-and-flare reactions, and by passive transfer of sera from patients with recent allergic reactions to the skin of nonsensitive human recipients. Three kinds of antigenic specificities of skin sensitizing antibodies have been found by these studies:

1. Benzylpenicilloyl-specific antibodies were detected by direct skin tests (Levine and Ovary, 1961; Levine and Price, 1964; Parker, Shapiro, Kern and Eisen, 1962) and by passive transfer techniques (Siegal and Levine, 1964). Skin test reagents which can be used for detection of benzylpenicilloyl (BPO) specific antibodies are multivalent BPO-haptens such as BPO-protein conjugates and BPO-polylysine. Succinylated BPO-polylysine conjugates of poly-L-lysine poly-D-lysine appear, at present, to be the most useful clinical reagents for the detection of BPO-specific skin reactivity. They are effective elicitors of BPO-specific wheal-and-flare reactions (Parker et al., 1962; Levine and Price, 1964), they are virtually non-irritating, and they are incapable of inducing immune responses in experimental animals (Levine, 1964b; Parker and Thiel, 1963), and probably also in man. BPO-specificity of these wheal-and-flare reactions (Levine and Ovary, 1961; Levine and Price, 1964; Parker et al., 1962; Siegal and Levine, 1964) as well as specificity for a diastereoisomeric mixture of BPO groups (Levine and Price, 1964) was confirmed by hapten inhibition experiments, using benzylpenicilloyl aminocaproate (Levine, 1962) as the univalent hapten. Other experiments indicate that the anti-BPO skin-sensitizing antibodies show specificity also for structural configurations of the autologous carrier proteins which induced the synthesis of antibody (carrier specificity) (Levine and Price, 1964). Carrier specificity toward human serum albumin was indicated in one patient (Levine and Price, 1964), towards human gamma globulin in another (Levine and Ovary, 1961), and to as yet unidentified proteins in others (Levine and Price, 1964). Since skin sensitizing antibodies may mediate some kinds of allergic reaction to penicillin, the indication that anti-BPO antibodies are specific also for structural configurations of homologous proteins would suggest that some allergic reactions to penicillin are, in part, auto-immune diseases. The exact clinical significance of carrier specificity is not yet known.

2. Another group of patients show specific wheal-and-flare skin reactivity by direct skin test to crystalline potassium benzylpenicillin (KPG) and to crystalline sodium benzylpenicilloate (free from PG impurity) (Levine and Price, 1964, and unpublished data). Two of these patients (out of eight tested) gave weak reactions also to a multivalent D-penicillamine conjugate (Levine and Price, 1964, and unpublished data). Two other patients were tested and failed to react to D-penicillamine, penilloaldehyde and to benzyl-oxazolone (unpublished data). This pattern of reactivity indicates specificity to the D-benzylpenamaldic-cysteine mixed disulfide and the D-penicillamine-cysteine mixed disulfide groups (Levine and Price, 1964). However, this specificity must be confirmed by more direct experiments. The postulated chemical pathway leading to the formation of these haptenic groups is shown in Fig. 1.

3. The third kind of antigenic specificity was observed by passive transfer technique (Siegal and Levine, 1964). Human skin sites passively sensitized with serum from patients with recent immediate systemic allergic reactions to penicillin, gave wheal-and-flare reactions to crystalline KPG, but not to sodium benzylpenicilloate, nor to
multivalent conjugates of the BPO, the D-benzylpenicillenic acid-disulfide or the D-penicillamine haptenic groups. The antigenic specificities of these skin-sensitizing antibodies have not been determined. Some possibilities are discussed in Siegel and Levine (1964). In addition to these three antigenic specificities, Parker et al. (1962) have reported skin sensitizing antibodies specific for the benzylpenicillenic acid-mixed disulfide group, but this has not as yet been confirmed (Levine, 1964). Accordingly, skin-sensitizing antibodies synthesized by human beings in response to the administration of PG show considerable antigenic heterogeneity. Similar antigenic heterogeneity is seen in allergic contact dermatitis to penicillin (see above).

Hemagglutinating anti-penicillin antibodies are detected by testing serum dilutions with human group O red blood cells (RBC) which had previously reacted with PG (Ley, Harris, Brinkley, Liles and Cahan, 1958; Watson, Jontert and Bennett, 1961; Van Arsdel, O'Rourke, Horan and Kamasaka, 1963, or with benzylpenicillenic acid (Fellner and Levine, unpublished data). Both methods of RBC sensitization result in BPO haptenic groups covalently linked to the RBC membrane, probably through amide linkages with amino residues of membranous proteins. It is not as yet known whether other of the possible penicillins haptenic groups are also coupled to RBC by these methods of sensitization. Hemagglutinating human anti-PG antibodies are both 19S and 7S \( \gamma \)-globulins, with 19S antibodies occurring considerably more frequently (Heremans and Vaerman, 1962; Schwartz and Vaughan, 1963; Fudenberg and German, 1960). Anti-BPO antibodies of the \( \gamma 1A \) class may also give passive hemagglutination, but this has not yet been demonstrated. The haptenic specificity of hemagglutination anti-PG antibodies is predominantly towards the BPO group (Theil and Parker, 1962; Fellner and Levine, unpublished data). Other haptenic specificities, although likely, have not as yet been demonstrated.

Immune Mechanisms of Allergic Reactions to Penicillin

Data pertinent to this problem can be obtained from prospective and retrospective correlations of the immune status of patients with the occurrence of clinical allergic reactions. In these studies, it is important to consider separately the different types of allergic reactions, e.g., immediate systemic, later urticarial, serum-sickness-like, etc., since each kind of reaction may have distinctly different immune mechanisms. Also, it should be considered that non-immune mechanisms (i.e., vascular sensitivity to histamine, or the activity of the reticuloendothelial system in clearing the plasma of antigen-antibody complexes) may play a role in determining whether a patient in whom is present the immunological factors necessary to develop an allergic reaction, does indeed develop an allergic reaction.

It appears probable, at the present time, that immediate systemic allergic reactions to penicillin are mediated by skin-sensitizing (\( \gamma 1A \)) anti-PG antibodies. This statement is based on the following evidence: Siegel has reported that all of 18 patients with recent immediate systemic reactions to penicillin contained in their sera, skin-sensitizing anti-PG antibodies demonstrable by passive transfer (Siegal, 1962). In addition, six other patients with this clinical history all showed direct wheal-and-flare skin reactivity to penicillin skin-test reagents (Fellner and Levine, unpublished data). This close association of anaphylaxis and the presence of skin-sensitizing antibodies has been found in other antigenic systems in human beings (Connell, Sherman and Myers, 1962), and in other mammalian species (Ovary, Benacerraf and Block, 1963; Nussenzweig, Merrymann and Bennaceraff, 1964). Of interest are the findings that the antigenic specificities of skin sensitizing anti-PG antibodies from patients with immediate systemic allergic reactions were mainly to the minor antigenic determinants (i.e. skin reactions were evoked by skin test with KPG alone, or by KPG and sodium benzylpenicilloate, see above) and only comparatively rarely to the benzylpenicilloyl haptenic group (Levine and Price, 1964; Siegal and Levine, 1964; Siegal, 1962). The presence of BPO-specific skin sensitizing antibodies does not necessarily mean that a patient will develop an immediate systemic reaction on administration of penicillin. Rytel, Klion, Arlander and Miller (1963) reported that none of 26 patients with strongly positive wheal-and-flare reactions to penicilloyl-polysynes, and who were immediately thereafter injected with 1.2 million units of benzathine penicillin developed immediate systemic reactions, although two of the 26 developed late urticarial reactions. It is possible that these patients were protected from immediate systemic reactions by the presence of circulating "blocking" antibodies.
Clinical Implications

An important clinical application of these studies is to detect those patients who will have an allergic reaction upon the administration of penicillin. From the considerations discussed above, patients who are liable to develop immediate systemic allergic reactions should show wheal-and-flare reactions on direct skin testing with the proper antigenic reagents, providing that the patients are physiologically capable of manifesting such skin reactions. The available data on the antigenic specificities of skin-sensitizing antibodies found in patients with recent immediate systemic allergic reaction to penicillin indicates that crystalline KPG may be a more effective detector of the potential immediate reactor than is penicilloyl-polyllysine (see above, and Levine and Price, 1964; Siegel and Levine, 1964). However, in order to detect the potential immediate systemic reactor, patients should be tested with both these reagents. A positive wheal-and-flare reaction does not necessarily mean that a patient will have an immediate system reaction on administration of penicillin, but it increases the probability of its occurrence. A negative skin reaction to both test materials appears, at present, to indicate that the patient will be able to receive penicillin without the occurrence of an immediate systemic reaction. However, this view is based on theoretical considerations and on experience with a relatively small number of patients. It is still possible that other skin test materials will have to be added to those now available in order to detect very rare cases of penicillin allergy. Before negative skin tests to the materials now at hand can be accepted as absolutely ruling out the possibility of an immediate systemic allergic reaction, many more patients must be evaluated. At the present time, it is not recommended that prior skin testing of patients about to be treated with penicillin be instituted as a routine clinical procedure. Before this can be done additional information must be obtained on the sensitizing capabilities of the skin test reagents. It is to be noted that negative skin tests to the reagents will not rule out the subsequent occurrence of a late urticarial or serum sickness-like allergic reaction following the administration of penicillin. At present there is no definite way to detect the patient who is more liable to suffer allergic reactions of these types.

Summary

Some current views of the immunological mechanisms of penicillin hypersensitivity are presented. The antigenicity of benzylpenicillin appears to depend in large part upon its ability to rearrange in vivo to its highly reactive isomer D-benzylpenicillinic acid. The postulated chemical pathways leading to the formation of the complete "penicillin antigens," and the identities of the haptenic determinants are presented. The administration of the simple chemical compound, benzylpenicillin, to experimental animals and to man leads to the production of antibodies of at least three different haptenic specificities. Some early views are given on the immune response of human beings to penicillin, on the possible immune mechanisms involved in allergic reactions to penicillin, and on the present clinical applications of these studies.

Fig. 1 is reproduced from J. exp. Med. (1960), 112, 113 by kind permission of the Editor.

REFERENCES

The purely cutaneous reactions to penicillin are nowadays infrequent and relatively unimportant. They consist of varying degrees of acute or chronic non-specific inflammation, with or without abscess formation. They were more commonly encountered with less pure penicillins and depot preparations sometimes injected intradermally or subcutaneously rather than intramuscularly.

All other reactions with manifestations in the skin are part of a general hypersensitivity state, although they may not necessarily be mediated by a specific antibody directed against the drug. In this regard penicillin is similar to sulphonamides in the very diverse types of adverse reaction which it is capable of producing. The factors determining the type of hypersensitivity reaction in an individual patient are mostly obscure, with the possible exception of contact dermatitis. In the latter case, the route of administration—application to the cutaneous or a muco-orificial surface—is of paramount importance.

In the United States, there are well over a hundred preparations of penicillin available to be fed, injected, spread, insufflated and sprayed into every conceivable cavity and surface of the body (Welch, Lewis, Weinstein and Boeckman, 1957). Individuals who have never received penicillin are a fast diminishing group, a result of mass production and the availability of this cheapest of all antibiotics.

**Incidence**

If one accepts any undesired or unexpected sequela as a reaction to the drug, the incidence of penicillin reactions can be very high, being dependent partly on the criteria of the physician. Moore’s (1946) figure is 0.56%, while Cormia, Jacobsen and Smith (1946) registered it as high as 60%. The variation is, of course, partly due to uncritical acceptance of any symptoms and signs, and partly due to the absence of satisfactory methods of proof. Well controlled drug trials have shown that skin eruptions may be seen in patients receiving placebos as frequently as an active agent. Saline injections can induce the rash, arthralgia,
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Bernard B. Levine

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