Title: PROCESSES AND APPARATUS FOR CARRYING OUT SPECIFIC BINDING ASSAYS

Abstract

A process for carrying out a specific binding assay (for example an immunoassay) in which (a) a sample under assay, possibly containing a substance being tested for, is reacted with (b) a specific binding partner for the substance being tested for, immobilised on a solid support, and (c) a specific binding partner for the substance being tested for which is conjugated to a detectable marker, thereby to form a complex by reaction between whatever quantities are present of the substance being tested for with reagents (b) and (c), in which the marker is immobilised to the support via the substance being tested for, and is detected or assayed as an index of the quantity present in the sample (a) of any of the substance being tested for; characterised in that reaction ingredients (a), (b) and (c) are all mixed in a single step for reaction in a single reaction liquid, and competitive interference between the binding reactions of the substance being tested for and reagents (b) and (c) is avoided either by use of an antibody of narrow and different, non-interfering specificity, such as a monoclonal antibody, to avoid the interference, or by use of a slow-release form of reagent (c). The apparatus for carrying out the assays has a handling piece (13) with several displacers (11) on which specific binding agents are immobilised.
FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

<table>
<thead>
<tr>
<th>Code</th>
<th>Country</th>
<th>Code</th>
<th>Country</th>
</tr>
</thead>
<tbody>
<tr>
<td>AT</td>
<td>Austria</td>
<td>KP</td>
<td>Democratic People’s Republic of Korea</td>
</tr>
<tr>
<td>AL</td>
<td>Australia</td>
<td>LI</td>
<td>Liechtenstein</td>
</tr>
<tr>
<td>BR</td>
<td>Brazil</td>
<td>LU</td>
<td>Luxembourg</td>
</tr>
<tr>
<td>CF</td>
<td>Central African Republic</td>
<td>MC</td>
<td>Monaco</td>
</tr>
<tr>
<td>CG</td>
<td>Congo</td>
<td>MG</td>
<td>Madagascar</td>
</tr>
<tr>
<td>CH</td>
<td>Switzerland</td>
<td>MW</td>
<td>Malawi</td>
</tr>
<tr>
<td>CM</td>
<td>Cameroon</td>
<td>NL</td>
<td>Netherlands</td>
</tr>
<tr>
<td>DE</td>
<td>Germany, Federal Republic of</td>
<td>NO</td>
<td>Norway</td>
</tr>
<tr>
<td>DK</td>
<td>Denmark</td>
<td>RO</td>
<td>Romania</td>
</tr>
<tr>
<td>FI</td>
<td>Finland</td>
<td>SE</td>
<td>Sweden</td>
</tr>
<tr>
<td>FR</td>
<td>France</td>
<td>SN</td>
<td>Senegal</td>
</tr>
<tr>
<td>GA</td>
<td>Gabon</td>
<td>SL</td>
<td>Soviet Union</td>
</tr>
<tr>
<td>GB</td>
<td>United Kingdom</td>
<td>TD</td>
<td>Chad</td>
</tr>
<tr>
<td>HU</td>
<td>Hungary</td>
<td>TG</td>
<td>Togo</td>
</tr>
<tr>
<td>JP</td>
<td>Japan</td>
<td>US</td>
<td>United States of America</td>
</tr>
</tbody>
</table>
PROCESSES AND APPARATUS FOR CARRYING OUT SPECIFIC BINDING ASSAYS

This invention relates to processes for carrying out specific binding assays, e.g. immunoassays, and to apparatus, e.g. test kits, for carrying out these processes.

In particular embodiments the invention is applicable to enzyme-linked and fluorescent-marker-linked specific binding assays, including immunoassays.

A wide variety of immunoassays and other specific binding assays is already known.

An important group of such assays comprises those in which a ternary complex is formed between a specific adsorbent, the material under assay, and a marker-conjugated material with specific binding capacity for the material under assay ("conjugate"). Certain assays of this kind have been called "sandwich" or "antiglobulin" assays. They have the property that the quantity of marker becoming fixed to the specific adsorbent is directly rather than inversely related to the quantity of the material under assay that participates in the ternary complex, and this can be simply measured after separation of the immobilised material from the remaining free marked conjugate, in whatever manner is appropriate to the marker in use.

However, as appears for example from the above-mentioned Habermann (1970) publication and G.B. specification No. 1,363,565, the performance of these assays is not without difficulty: they demand either a number of successive manipulation steps to carry out the assay reactions, or else suffer from low sensitivity, which careful choice of reagents has not so far been able to overcome.

Commercial test kits are available comprising essentially:
(a) a plate consisting of an array of tubes or pre-formed wells which are coated with an antibody or antigen as the case may be;
(b) an enzyme linked to an appropriate antibody (a so-called conjugate) against a substance to be detected if present in a test sample and
(c) a substrate for the determination of the activity of the enzyme.

One standard procedure for conducting an assay for antigen or antibody involves:
(1) determining the working dilution for the test sample;
(2) removing any excess of antibody or antigen used to sensitise the wells;
(3) washing the wells;
(4) introducing a proportion of the suitably diluted test sample;
(5) incubating for about two hours to allow the substance to be detected in the test sample to bind to the sensitising substance;
(6) washing the wells to remove unreacted material;
(7) introducing the suitably diluted conjugate (incubate for about 2 hours);
(8) washing the wells to remove unreacted material;
(9) adding a solution of the substrate;
(10) incubating until a suitable intensity of colour develops as a result of the reaction of the substrate with the enzyme;
(11) stopping the reaction, e.g. with a strong alkali and
(12) measuring the optical density of the reacted substrate solution.

This procedure is also time-consuming since each of the several antibody/antigen reactions requires several hours to reach equilibrium. In practice, shorter incubation times are used but only at the expense of sensitivity and/or economy.

According to the results of the present work, it is believed that an obstacle to the use of fewer assay steps is an unwanted interference with the formation of the desired immobilised complex which can originate in interfering reactions between two of the components. By using a specific binding agent of selected narrow specificity, or in slow-release form, such interference can be avoided, and high-sensitivity assays carried out using fewer manipulation steps.

According to this invention there is provided a process for carrying out a specific binding assay (for example an immunoassay) in which (a) a sample under assay, possibly containing a substance being tested for, is reacted with
(b) a specific binding partner for the substance being tested for, immobilised on a solid support, and (c) a specific binding partner for the substance being tested for which is conjugated to a detectable marker, thereby to form a complex by reaction between whatever quantities are present of the substance being tested for with reagents (b) and (c), in which the marker is immobilised to the support via the substance being tested for, and is detected or assayed as an index of the quantity present in the sample (a) of any of the substance being tested for; characterised in that reaction ingredients (a), (b) and (c) are all mixed in a single step for reaction in a single reaction liquid, and competitive interference between the binding reactions of the substance being tested for and reagents (b) and (c) is avoided either by use of an antibody of narrow specificity, such as a monoclonal antibody, to avoid the interference, or by use of a slow-release form of reagent (c).

The narrow specificity required of the antibody is a capacity to bind specifically with the substance under test but without preventing the binding reaction between the substance under test and its other specific binding partner. Such an antibody can be selected out of a number of antibodies with an affinity for the substance under test, by using normal methods to verify the progress of a binding reaction between the other specific binding partner and a complex previously formed between the substance to be tested in the assay and the narrow-specificity antibody to be selected.

According to a preferred embodiment of the present invention, the conjugate between antibody and the enzyme or other marker, and/or the antibody (if any) which is coupled to the solid surface, comprises a monoclonal antibody or other antibody of sufficiently narrow specificity to ensure that the desired assay reaction or reactions are not impeded
by competition between the conjugate and the immunosorbent in their reactions with whatever quantities are present of the substance being tested for in the sample under assay. Monoclonal antibody of sufficiently narrow specificity can, for example, be produced as antibody derived from a line of antibody-producing cells, derived from a single antibody-producing progenitor cell or cells. Such a line can, for example, be produced by known cell fusion, culture and isolation techniques using very pure antigens as comparative material.

Alternatively, in many cases antibody of sufficiently narrow specificity can be obtained in the (polyclonal) immunoglobulins of antisera raised against discrete chemical or physical molecular fragments of the material under test, for example, antibody against Fc fragments (or against smaller peptide fragments) of immunoglobulins to be tested for, or against sub-units or peptides of protein antigens to be tested for. The object in each case is to ensure substantial freedom from interference which can arise particularly, for example, in carrying out immunoassays of the "sandwich" or "antiglobulin" test configurations.

In a "sandwich" test configuration, antigen under test can be specifically adsorbed to a first antibody bound to a solid surface, and a second antibody carrying an enzymic or other (e.g. fluorescent or radioactive) marker is specifically bound to the adsorbed antigen under test. Marker specifically so bound is used for measurement and determination of the antigen under test, e.g. by direct measurement, such as radiometry or fluorimetry, or exposure of enzyme marker to substrate followed by product measurement. Thus, in preferred sandwich tests, the two antibodies used can have different, non-interfering specificity with respect to the same antigen under test.

In an "antiglobulin" test configuration, sometimes also referred to as a "sandwich" test configuration, the position
is analogous: the material under test is itself an immunoglobulin; the material bound to a solid surface is its corresponding antigen or hapten; and the material carrying the marker is an antiglobulin corresponding to the species and immunoglobulin type of the antibody under test. In preferred antiglobulin tests, the antiglobulin can have sufficiently narrow specificity as not to interfere with the subsequent adsorption of its corresponding globulin to the insolubilised antigen.

If antibodies from ordinary antisera raised against unmodified antigen (polyclonal antibodies) are used in sandwich or antiglobulin tests, there is a very likely risk that if all ingredients are mixed in a single step there will be interference between the two specific adsorption reactions. When such tests are carried out according to the present invention, using apparatus as described herein, such interference can be avoided either by using antibodies of narrow specificity as described, or else by ensuring that the binding of test material to the solid surface takes place before exposure of test material to the other (marker-conjugated) binding agent if there is a risk that binding by that other agent would prevent subsequent adsorption to the solid surface. Such a sequence can be ensured by arranging for slow release of the other (marker-conjugated) binding agent.

Particular instances of suitable assay specificities, antibody specificities, and slow-release forms of conjugated reagent (c) are described for example below.

It has also been found that in carrying out such specific binding assays, a worthwhile improvement in reaction kinetics can be obtained if the reaction liquid containing ingredients (a), (b) and (c) is contained in a well or cup of which the majority of the volume is occupied by a displacer body. (The use of inserts of various rod or ball shaped forms is known in connection with other kinds of
immunoassay, as described in G.B. Specification Nos. 1,414,479 and 1,485,729.

The displacer body can, for example, be of a shape substantially complementary to and slightly smaller than that of the cup or well, so that the liquid phase containing one of the specific binding reagents is approximately in the form of a shell occupying the space between the displacer and the cup or well. The displacer can be loose-fitting and not fixedly mounted, i.e. movable relatively to the cup or well, so that by relative motion between displacer and well the liquid between them can be given a stirring or agitation motion.

For example, a round well can have a round displacer therein with an external diameter slightly smaller than the diameter of the well. The presence of the displacer can reduce the space available for liquid in the well by a factor of for example 2-10, e.g. 3-8, comparing volumes based on similar liquid levels in the well, e.g. when filled to its normal operating level, or its maximum capacity. For example, a microtitre well designed to have 300 microlitre of liquid filled into it during a normal assay, can be used with a displacer leaving 30-150 microlitre liquid space, e.g. 50-100 microlitre.

The use of wells or cups together with displacers as described herein can improve the efficiency of the assay reaction steps because, in the first place, it allows more concentrated reagents to be used with no increase in the weight of reagent or decrease in the size of the microtitre wells, compared with the normal conditions encountered in microtitre wells of given size; and in the second place, it increases the sensitised surface area available to react with a given liquid reagent volume, so that comparatively faster adsorption kinetics can be achieved without having to increase specific reagent density on the sensitised surface or encountering problems of crowding.
A set of displacer bodies can be preferably present in certain embodiments of the invention, e.g. as an integral part of a lid which can be fitted onto a microtitre plate, e.g. a standard plate of 8 x 12 wells. The set can be large enough to fit all the wells of the plate or a sub-set thereof, e.g. a row. The dimensions of the displacers and the volume of liquid to be dispensed into the well can be chosen relative to the well in the manner described above, and preferably so that the liquid to be tested is in contact with substantially the major part and preferably the whole inner surface of the well.

The immobilised specific binding partner (reagent (b)) for the substance to be assayed can be immobilised on the wall of the well or cup in which the assay reaction takes place. Alternatively, according to a feature of the invention independently capable of providing advantage and convenience in use, a liquid displacer, for example in the form of a stick, peg or stud, for dipping into a liquid assay reagent, can have an immunosorbent surface. This allows the portion of the assay materials needing to be carried over from one reagent to the next, and the associated manipulations, to be handled more easily than when the sensitised surface is part of a hollow well. An alternative form for such a liquid displacer body is a tuft of bristles or leaves of suitable material, or equivalent body with large surface area. A further alternative form is a stud or peg with relatively hollowed-out and projecting portions of its surface, e.g. with grooves and associated ribs, e.g. annular grooves. Such an arrangement can give robustness, increased sensitised surface area, and better reactivity.

Test apparatus according to related embodiments of the invention can thus comprise a set of sensitised liquid-displacer bodies of one or more of such forms, joined to a common handling-bar, link or lid, and for use in combination
with a complementary set or sets of wells containing any
of the remaining materials used in the assay. The several
displacer bodies of the set can have the same or different
sensitisation so that one or a plurality of different assay
types can be carried through simultaneously. If desired,
the displacer bodies can be removably and exchangeably
mounted on the handling bar, link, or lid, so that sets of
desired specificity can be built up at will from a common
stock for carrying out large numbers of tests according to
a desired pattern.

One advantage of such arrangements is that a number of
displacer bodies can be sensitised in the same body of liquid
reagent, avoiding fluctuating conditions of concentration,
etc., resulting from dosing aliquots into wells.

The displacer bodies can, in this embodiment, be of any
material suitable for the preparation of an immunosorbent by
covalent bonding or adsorption: e.g. polystyrene, nylon, or
cellulose acetate. (The nature of the displacer surface
does not matter provided it is inert, when the sensitisation
is to be on the well surface rather than the displacer
surface.) Linkage of antibodies, antigens, etc., to the
displacer bodies can be carried out by linking methods known
in themselves, e.g. partial acid hydrolysis of nylon surface,
substitution of exposed amine surface with glutaraldehyde,
and coupling of material to be bound, e.g. antibody or antigen,
to immobilised aldehyde groups. Suitable methods among a
wide variety are given for example by Inman & Hornby (1972)
Biochem.J. 129, 255; Campbell, Hornby & Morris (1975)
P.E.B.S. Letters 78, 251; and G.B. patent specifications
Nos. 1,470,955 and 1,485,122.

It can be seen that the invention also provides a kit of
test materials for carrying out a specific protein-binding
assay, comprising (i) an immobilised specific binding
partner for a substance to be tested for, carried on a solid
support, and (ii) a marker-conjugated specific binding partner for the substance to be tested for, which can be added to a reaction liquid contacting immobilised reagent (i) either as a slow-release form, or in any form provided that the specific binding partners in reagents (i) and (ii) include an antibody of narrow specificity so that reagents (i) and (ii) do not interfere with each other's binding reactions with the substance to be tested. Optionally the kit can also comprise materials for later estimation of the amount of marker immobilised during the assay.

Reagent (i) can be immobilised on either a displacer body for a reaction well, or on a reaction well wall, as described above. A slow-release form of reagent (ii) can be for example a sucrose or equivalent glaze on a complementary surface of either the displacer or the well wall, also as described above. The narrow-specificity antibody can be selected for example from monoclonal antibodies in the manner already described.

The invention is further illustrated by reference to the following details and accompanying drawings, of which:

Figure 1 is a vertical cross-section of an arrangement of container and displacer for carrying out specific binding assays according to the invention.

Figure 2 diagrammatically shows an array of several of the arrangements shown in Figure 1.

Figures 3 and 4 show in diagrammatic side view and part-sectional end view, an assembly of displacers for carrying out certain embodiments of the invention.

Figure 1 of the drawing shows in vertical cross-section a round-bottom plastics microtitre well-shape vessel 1 containing a round-section round-bottom displacer body 2 of slightly smaller and complementary shape. The displacer 2 can be of glass or plastics material. The normal capacity of well 1 is for a 300 microlitre sample. The presence of the displacer 2 reduces the effective capacity to 50
microlitre. This quantity of liquid 3 forms a shell in contact with the whole of the surface area of the normal capacity of the well and with the displacer body 2. Thus the surface-to-volume ratio of liquid 3 is several times greater than that of the body of liquid which would fill the normal capacity of the well in the absence of displacer 2.

Displacer 2 can be left loose so that during use of the well for assay any agitation given to the well 1 causes stirring of liquid 3. Alternatively, displacer 2 can be directly driven. In one arrangement, shown in Figure 2, a tray 4 containing a 2-dimensional array of wells similar to well 1 has a lid 5 to which a complementary set of displacer bodies similar to cisplacer 2 is mounted. The displacers may be integrally formed with lid 5, e.g. in the form of hollow moulded protrusions. In an alternative arrangement the fixed set of displacers may form a row or rows only, less than the full array.

In use, for example, for immunoassay of the ELISA type, the bottoms and lower parts of the walls of wells 1 are coated with immunological reagent to form immunosorbent, and 50 microlitre quantities of the materials under test, in suitable dilutions and media, are contacted with the well walls and bottoms, by the use of the displacer bodies 2. Immunological reagent conjugated to enzyme marker, if not already dispersed throughout the reaction liquid, is added to the reaction liquid in an alternative way: The displacers can be coated with a conjugate of a detectable enzyme or other marker with an immunological reagent, e.g. antigen or antibody. Suitable quantities of the conjugate can be dried onto the tips of the displacer bodies, preferably in a glaze, e.g. of sucrose, to prevent aggregation of conjugate molecules and ensure a gradual and slow solubilisation of the conjugate thereby avoiding saturation of the immunosorbent with the conjugate before the former gets the chance to react with the substance to be
detected. The test is then performed by dispensing an appropriate
volume of test sample consisting, for example, of patient's serum
into a well, followed by inserting a displacer body into the well.
Thus, the sample and conjugate can be added almost simultaneously,
thereby reducing the time-consuming sequence of operations.
After the slow release of the conjugated material from the dried
glaze on the tips of the displacers 2, formed for example, by
evaporation of a solution of the conjugate and sucrose, (or after
formation of the assay complex from predispersed reagents) the
amount of specifically bound enzyme marker is measured in the usual
way. The dosimetry and quantitation of the reagents for the test
are carried out according to well-established methods and in
themselves form no part of the invention.

Figures 3 and 4 of the accompanying drawings show, in
diagrammatic side view and end (part-sectional) view, an assembly
of displacer bodies for carrying out specific binding assays
according to a further and advantageous arrangement given by way of
example. The assembly of such displacer bodies can form part of
composite apparatus according to the invention comprising such an
assembly (or a single such displacer according to the invention)
together with a complementary well or wells for carrying the liquids
used in the assays. Either (preferably) the displacer, or the well,
carries immobilised specific binding agent.

Referring to Figures 3 and 4, the characteristic component of
the apparatus for carrying out the assay in each well is a liquid
displacer 11 in the form of a stud, stick or peg as shown for dipping
into liquid in an assay well as described herein. The displacer 11
is formed, e.g. by moulding or turning in a suitable material such as
nylon, cellulose acetate or polystyrene. One preferred form is a
nylon peg, of for example, about 5 mm greatest diameter, or slightly
narrower than the internal diameter of an associated microtitre
well, and about 15-20 mm height to project out of such a well. About,
for example, 8 millimetre of a lower extremity 12, or other lower
portion of displacer 11, can conveniently be provided with grooves,
a screw thread, chamfering, or other irregularity to increase its
surface area, not shown in detail in the drawings, which in Figures
3 and 4 indicate only a slight overall narrowing of grooved extremity 12 of displacer 11 due to the formation of such grooves.

Each displacer 11 is mounted on a handling piece 13 which can have either one displacer 11 or as many as desired mounted thereon, e.g. in a one-dimensional array, as shown, or a two-dimensional array, corresponding to the arrangement of Figure 2. The displacers 11 can be formed either as one piece with the handling piece or as separate mountable parts which can, if desired, be exchangeable. Illustrated is a socket-and-screw arrangement for exchangeably mounting displacers 11 of the type described at, for example, about 9 millimetre intervals, corresponding to well spacings in the complementary microtitre plate. The handling piece can be of any suitable material, e.g. plastics such as nylon, etc., or metal, e.g. brass, copper or stainless steel.

Biological/chemical conditioning of the displacer 11, e.g. the lower grooved extremity 12 thereof, can be carried out for example as follows to suit the material for carrying out specific binding assays such as immunoassay. The displacer can, if desired, be used as described in Example 1, but alternatively and preferably, it can be made to carry an immunosorbent or other specific binding adsorbent. Antibody or antigen, for example, of type suitable for the desired assay configuration which in itself forms no part of this invention, can be attached by physical adsorption (especially when using polystyrene) or (preferably) by covalent linkage to grooved extremity 12, e.g. by methods already referred to herein.

The assay is then carried out, for example, as follows. Reagent and/or test solution is placed in the well, to a volume sufficient to fill the well only when the displacer is inserted (50-200 microlitre for example). With the displacer present the binding reactions take place, and washing can be achieved simply by removing the displacers
and rinsing them, e.g. under running tap water.

Even though two binding events may be required (e.g. test antibody binds to immobilised antigen, then enzyme-conjugated antiglobulin binds to the test antibody) it is possible to complete the binding sequence in a single step. This requires the careful choice of reagents, as described above, so that the conjugate antibodies do not compete for all the binding sites on the test material and prevent it from binding to the immunosorbent.

The presence of the displacers in the wells can provide for the important operation of stirring the fluid samples. Not only does this allow a more rapid binding reaction, it also allows the conjugate to be pre-dried into the wells in a firm and stable state, such that it can only be reconstituted by means of such stirring. The stirring might be continued for the duration of the whole binding step, or might be started only after the first binding reaction has been completed. A suitable form of coating is for the well bottoms to be given a dried sucrose glaze containing the conjugate. This can release slowly enough to achieve the desired sequence of binding reactions, as discussed herein above.

After the binding reactions have been completed the displacers can be washed and transferred to another set of wells containing a solution of enzyme substrate, where they remain until the positive controls have developed an adequate colour (and negative controls are still adequately blank). In a standard ELISA, the enzyme/substrate reaction has to be stopped by adding a reagent such as 2M sodium hydroxide solution, but even this step can be dispensed with when the arrangement of this invention is used. The reaction can be terminated simply by removing the displacers from the substrate solutions; the substrate-containing wells can then be simply placed in a standard micro-ELISA reader for the measurement and recording of results.
In existing commercial ELISA systems the immunosorbent component (usually the microtitre wells or equivalent) is necessarily a disposable item, but in a system according to this invention, possible covalent attachment or antigens can allow the immunosorbent to be re-usable (if it is desired to be so). The binding reactions can be reversed by soaking the displacers in a releasing agent such as 4M magnesium chloride, or sodium dodecyl sulphate, whilst the covalently attached reagent remains intact. This arrangement can bring benefits in terms of precision, reliability, speed and/or sensitivity. For the manufacturer and user alike there can be important economies in the use of immunoreagents which are often scarce and expensive.

A further optional feature which can be present in the above-described format of this invention is that of the precision heating of individual reaction mixtures. The rates of the binding and of the enzyme/substrate reactions are dependent on temperature; temperature differences across a microtitre plate can cause related response variations, so creating serious defects in calibration and reproducibility. Normally, ELISA reactions are speeded up by placing the tray in a standard air incubator, but this can be less than satisfactory, because the rate at which wells are heated often depends on their position in the tray. The rate of heat exchange in an air incubator is rather slow: the whole plate warms up relatively slowly. Individual heaters can be made (for example, from precision resistance thermometers used in a heating mode rather than a measurement mode), and inserted into the centre of the displacer bodies, for example those illustrated in Figures 1-4, and, in that position, can ensure a rapid and efficient delivery of heat through the displacer body surface. This system (which incorporates suitable thermostatic control of a kind well known in itself) can therefore provide faster reactions and reduce well-to-well variation at the same time as dispensing with the need for an incubator.
Example 1

An illustrative scheme of preparing materials for immunoassay and carrying out the assay according to the invention is described below, for use with the displacers shown in Figures 3–4. The particular assay configuration illustrated happens to be an enzyme-linked antiglobulin test directed to assaying antibodies against soya protein. It will be clear that other configurations and specificities can be used and the arrangements of this invention applied to them, as for example many of those mentioned in US Patents Nos. 3,654,090, 3,971,932, 3,839,153, 3,850,752, 3,879,262 and 3,996,345.

Preparation of Soya Protein-linked Displacer Pegs

Nylon pegs as described above in relation to Figures 3–4 are immersed in 3.6M HCl and held at 48°C for 35 minutes after which they are washed under running tap water and then soaked in distilled water. Next, they are immersed in chilled 12.5% aqueous glutaraldehyde and kept at 4–8°C for 20 minutes. This reaction is stopped by washing the pegs under running tap water and then soaking them in a large volume of distilled water for at least 2 hours.

After this activation sequence, the pegs are placed in the wells of a microtitre tray, together with 200 µl of a solution of soya protein (300 µg per ml, in phosphate buffered saline with Tween detergent (PBST) + 0.1% thimerosal). The trays and pegs are placed in a sealed container and the reaction continued at 37°C for 54 hours.

Finally, unused active sites are blocked by transferring the pegs into another set of wells containing 200 µl of 1.5M ethanolamine, where they are maintained at room temperature for 24 hours. Before use, they are soaked in PBS + 0.15% Tween for a further 3 days at room temperature.

In an alternative preparative method, the hydrolysis and glutaraldehyde treatment are identical but, instead of 300 µg/ml of protein being linked to the surface, the pegs are treated with poly-L-lysine (PLL) at 100 µg/ml for 18
hours at 37°C. After a thorough wash in tap water and then PBS, the linked PLL is activated by treatment with 12.5% glutaraldehyde (18 hours at 37°C). Finally the soya protein is linked to the glutaraldehyde-substituted PLL by incubating the treated pegs in 200 μl of soya protein at 100 μg/ml, for 18 hours at 37°C. Excess active groups are blocked by treatment in 1.5M ethanolamine at pH 8.5 for seven hours.

In further alternative procedures, the nylon can be treated with dimethyl-1,3-propane diamine to cleave nylon polymer chains, the free amine groups being then substituted with glutaraldehyde, and optionally after further insertion of amino and aldehyde "spacer" groups, the protein desired to be coupled is linked to the terminal aldehyde.

In a further alternative method, not involving polymer chain cleavage of the nylon, displacers are soaked in PBS + 2% sodium dodecyl sulphate and then allowed to dry in the air at room temperature overnight. They are 0-alkylated by being immersed in 100% dimethyl sulphate for 3 minutes at room temperature. The reaction is stopped by washing the pegs in ice cold ethanol and then cold water. Immediately thereafter the pegs are immersed in a solution of soya protein (100 μg/ml) and kept at room temperature for 3 days. Finally they are transferred to a 10mM solution of ethanolamine where they are maintained at room temperature for a further 2 days, before a final wash in Tris/HCl buffer, pH 8.0. Alternatively, the alkylating agent used can be triethylxonium tetrafluoroborate, as described by Inman and Hornby (1972) cited above. Suitable coupling methods from which choice can be made are mentioned in, for example, GB Specifications 1,316,990, 1,470,955, 1,485,122-3, and USP 3,817,837 (cols. 31-34, for example). Preparation of Enzyme Labelled Sheep-anti Rabbit IgG

Rabbit globulin (prepared from rabbit serum by sodium sulphate precipitation) is immobilised onto cyanogen bromide-activated sepharose 4B (Pharmacia) by the normal procedure
described by the manufacturers. A high-titre sheep anti-
serum to rabbit globulin (Seward Laboratory, Immunostics 
(Trade Mark) Product BS 01) is passed down the column and 
the non-bound fall through fraction rejected. The bound 
antibodies are released by passing 0.5M acetic acid down the 
column. The protein-containing fractions are pooled and 
dialysed against PBS to give an antibody preparation 
which is conjugated with enzyme by the following 
procedure.

The following method is based on that described by 
Engvall and Perlmann (1971) (Immmunochemistry, 8, 871). 
The IgG fraction of sheep antiserum to rabbit IgG (Seward 
Laboratory, Immunostics (Trade Mark), Product BS 01) is 
adjusted to a concentration of 5 mg/ml. A 0.1 ml aliquot 
of this is used to dissolve 1.5 mg of Sigma type VII 
alkaline phosphatase (supplied as a suspension in 3.2M 
ammonium sulphate). The resultant solution is dialysed 
against phosphate buffered saline (PBS) and then mixed with 
5 µl of a 25% glutaraldehyde solution (aqueous). The 
reaction with glutaraldehyde is continued for 2½ hours at 
room temperature and then stopped by dilution to 1 ml with 
PBS and extensive dialysis. Finally, the product is diluted 
to 10 ml in 0.05M tris/HCl buffer (pH 8.0) containing 50 
mg/ml ovalbumin, 0.2 mg/ml magnesium chloride, 0.2% sodium 
azide and 0.2% merthiolate. The conjugate can be 
applied to a microtitre well bottom by making it concentrated 
in sucrose solution and a 5-50 microlitre aliquot is then 
evaporated to a hard sucrose glaze at 37°C for slow release 
on stirring.

Preparation of Soya Antigen
The protein used as an antigen is prepared from soya 
bean flour by a method based on the procedure described by 
bean flour (10 g) is dispersed in PBS (100 ml) and after 

thorough mixing the suspension is centrifuged to yield a clear 
supernatant solution.
An aliquot of this solution is mixed with 30 g of urea to give a final volume of 50 ml. This solution is heated in a steam bath for 1 hour before 1 ml of 2-mercapto-ethanol was added. Steaming is continued for a further 45 minutes. After being cooled to room temperature the solution is dialysed, first against running tap-water for 2 hours and then against 2 x 5 litres 0.15M sodium chloride. The final dialysed solution swells to 80 ml and has a very slightly opalescent appearance.

Preparation of Rabbit Antibodies to Renatured Soya Protein

Rabbits are given an intra-muscular injection of renatured soya antigen (2 mg) in Freund's complete adjuvant emulsified as a multiple water-in-oil in water system. The volume given to each animal is 2.0 ml. Subcutaneous booster injections of antigen in saline (0.25-0.5 ml) are given 31, 100, 108 and 115 days after the initial inoculation. Serum samples are taken at various times through this schedule, and used as sources of antibody to be tested for by the following assay schedule.

Immunoassay using the Sensitised (Soya-protein-linked) Displacer

Rabbit antisera and the sheep-anti rabbit/enzyme conjugate are diluted appropriately in PBST and then the following sequence of operations was followed.

1. Diluted antisera placed in wells, followed immediately by sensitised displacer pegs.
2. Displacer pegs + wells (containing a sucrose glaze of conjugate) + sample placed in sealed container and incubated at 37°C for about 90 minutes. Agitation is carried out but only after about half the incubation period has elapsed.
3. Displacer pegs removed and rinsed under running tap water, then PBST.
4. Washed displacer pegs placed in wells containing substrate for enzyme (Sigma 104 (Trade Mark)).
5. Displacer pegs + wells + substrate placed in a sealed container and incubated at 37°C for 45 minutes.
6. Displacer pegs removed and optical densities of substrate samples measured.

The standardisation and calibration of this test is carried out by methods which are in themselves well established and form no part of this invention.

Example 2

An alternative assay and set of assay materials for the detection or estimation of anti-(soya) antibodies uses a monoclonal antibody. These materials and the assay are carried out as Example 1 except that: A mouse monoclonal anti-(rabbit IgG) with good affinity is used in place of the (polyclonal) sheep antiserum to rabbit IgG previously used. Instead of the IgG fraction of the antiserum prefiously to be mixed with the enzyme and conjugated, an ascitic fluid fraction containing the monoclonal antibody is used: this fraction is prepared by affinity chromatography on the same kind of rabbit globulin-conjugated Sepharose gel as in Example 1, but the elution is carried out afterwards with 4M MgCl₂ solution instead of 0.5 M acetic acid. The final conjugate preparation is added to the well directly in step 1 instead of being applied as a sucrose glaze, and the incubation time in step 2 can be reduced to 1 hour. It is found that this assay provides not only a gain in time and simplicity but can also yield a more accurate and reliable relationship between final colour development in steps 5-6 and quantity of anti-(soya) antisera added in step 1.

Example 3

An assay specific for K-type immunoglobulin G and the preparation of materials for it can be carried out as follows.

Pegs moulded of nylon 66 in the form illustrated in Figure 3 are washed in alcohol and distilled water. Mouse monoclonal anti-(human K-chain) antibody is immobilised on to them by exposure of the pegs at room temperature for a few hours of overnight to a 20 microgram/ml solution of the
antibody in 0.01 M aqueous phosphate buffer at pH 8, kept
free of other protein and of any detergent. The preparation
of monoclonal antibody to K chain (and also of the antibody
used below directed to \( \gamma \)-chain) is derived from a corresponding
ascitic fluid by affinity chromatography on a column for
example of Affigel (Biorad, Trade Mark) conjugated to human
IgG, in otherwise the same way as in Example 1 or 2. (About
10 mg IgG is used for conjugation of 1 ml of gel, and at least
2 ml of gel is used for chromatography of 1 ml of ascitic
fluid.)

A mouse monoclonal (human \( \gamma \)-chain) antibody is conjugated
with alkaline phosphatase as in Example 1, to form the other
specific binding reagent in the assay.

The general procedure of Example 1 or 2, was then used
together with these materials, except that the non-covalent
nature of the absorption of antibody to form immunosorbent
made it advisable to use as reaction buffers at pH 7.1
phosphate/saline/Tween/ovalbumin buffer (0.15% phosphate,
0.85% NaCl, 1.5 ml/ml Tween, 1 mg/ml ovalbumin). This
provided a specific and sensitive assay for K-type human
IgG. An alternative arrangement for the assay of this
Example involves omitting the conjugate from step 2,
incubating the pegs and test sample about 60 minutes,
washing, then incubating the pegs about 60 minutes with
conjugate dissolved in a similar buffer before going to
step 3. Both assay methods give advantage in allowing
highly specific determination of the \( \gamma \)-K subpopulation of
IgG molecules, because the narrow specificity avoids the
interfering cross-reactions which would have rendered such
specific assays impossible using less specific reagents.

It was found in one series of trials that the assay
result could be made linear in the range 10-100 ng \( K \)-IgG
per 0.2 ml assay reaction volume, (using about 50 ng
conjugate based on starting antibody content).

A corresponding assay for \( \lambda \)-type IgG can be easily
carried out if in the preparation of the immunosorbent, i.e.

SUBSTITUTE SHEET
the nylon 66 pegs with antibody immobilised thereon, a mouse monoclonal antibody directed against -chains is used instead.

It can be seen that the particular methods and materials described above can be varied in specificity to provide a wide range of assays sharing the convenience of fewer manipulation steps and freedom from interfering side reactions in use.

Among the monoclonal antibodies that can be obtained according to published procedures and selected for use as assay materials in the present invention are for example anti-rotavirus, anti-(human) IgE, anti-(bovine) IgG, anti-\(\beta_2\)-fetoprotein, anti-(human and other) IgA, anti-(thyroid-binding globulin), anti-\(\beta_2\) microglobulin, anti-pregnanediol, anti-oestradiol and anti-(human) IgM antibodies, and others. According to the invention these reagents can be used to provide the basis of highly-specific assays for the corresponding specific binding partner which may correspond to the whole population or a specific subpopulation of the type of material under assay: any of the above-described assay procedures are usable for this purpose.

Not all of the corresponding assays need be immuno-assays in the strictest sense. For example, using monoclonal anti-(TBG), a corresponding specific binding assay involves immobilised \(T_3\) or \(T_4\) as the specific binding partner of the TBG being tested for, and the use of a selected marker-conjugated monoclonal anti-TBG enables a sensitive "sandwich" type assay to be carried out in few steps without interference between immune complex formation and the binding of TBG to \(T_3\) or \(T_4\).

An assay for specific immune complexes can be carried out in an analogous way by the use of immobilised conglutinin and a marker-conjugated monoclonal antibody specific to the component of the immune complexes desired to be assayed.
it will be well understood that the other arrangements of this invention are also susceptible of wide variation and application. For example, a set of displacers as shown in Figures 3-4 can comprise a handling piece or holder fitted with, e.g., eight pegs, the displacers being appropriately sensitised for use in the assay of human antibodies to any one or more of, for example, toxoplasma, rubella virus, cytomegalovirus and herpesvirus. The displacers can have corresponding antigen fixed thereto, and a marker-conjugated anti-(human globulin) is used in the test which is chemically of the known antiglobulin format, with reagents provided and calibrated accordingly. It can be specially convenient if all four antigen sensitisations are represented in duplicate, each on two displacers of the assembly, shown in Figures 3-4. Then such an assembly can, for example, be carried through assay steps designed to assay eight aliquots of a single patient sample, in duplicate for each of the four above-mentioned antibodies. Such an arrangement can be especially convenient in that the remainder of the test tools and reagents, other than the displacers carrying specific sensitisation, can be qualitatively similar for all the assays, for example, the conjugate can be an anti-(human globulin) antibody coupled to a suitable marker enzyme, and common means can be used for detecting the enzyme. These components can be suitably provided in microtitre wells prepared in conjunction with the sensitised displacer pegs, if desired.

Other immunoassay arrangements according to the invention can be directed in specificity towards, for example, immunoglobulin E, (otherwise described as reagin-immunoglobulin), for which an antiglobulin assay can in format resemble the radioimmunoassay using antiglobulin described in USP 3,720,760, with optionally the substitution of another kind of marker conjugating the anti-IgE or other labelled antibodies used, e.g. a
fluorescent or especially an enzymic marker, used as described above. Where markers other than enzyme markers are used, the appropriate respective known coupling techniques are used. Analogous assay arrangements can be made for clinical diagnostic use, e.g. plasma C-reactive protein, urinary β₂-microglobulin or Factor VIII. For these antigenic test materials it can be convenient to use a sandwich-type assay format.

Other modifications and variations will be apparent to the skilled reader. In particular, the vesicle-bound marker systems described in specification EP-0014530 can be used to provide marking for the assays of the present invention. Reference is also made to this specification and the literature cited therein for further examples of sandwich and antiglobulin type assay configurations.
1. A process for carrying out a specific binding assay (for example an immunoassay) in which (a) a sample under assay, possibly containing a substance being tested for, is reacted with (b) a specific binding partner for the substance being tested for, immobilised on a solid support, and (c) a specific binding partner for the substance being tested for which is conjugated to a detectable marker, thereby to form a complex by reaction between whatever quantities are present of the substance being tested for with reagents (b) and (c), in which the marker is immobilised to the support via the substance being tested for, and is detected or assayed as an index of the quantity present in the sample (a) of any of the substance being tested for; characterised in that reaction ingredients (a), (b) and (c) are all mixed in a single step for reaction in a single reaction liquid, and competitive interference between the binding reactions of the substance being tested for and reagents (b) and (c) is avoided either by use of an antibody of narrow and different, non-interfering specificity, such as a monoclonal antibody, to avoid the interference, or by use of a slow-release form of reagent (c).

2. A process according to claim 1, characterised in that a monoclonal antibody immobilised on a solid surface to form an immunosorbent is comprised in reagent (b).

3. A process according to claim 1 or 2, characterised in that a monoclonal antibody in marker-conjugated form is comprised in reagent (c).

4. A process according to claim 1, 2 or 2, characterised in that reagent (c) is used in the reaction in slow-release form such as a glaze coated on a surface in contact with the reaction liquid.
5. A process according to any of claims 1-4, characterised in that the specific binding reagent (b) is immobilised on the surface of a displacer body (2) which occupies a majority of the volume of a well or cup (1) containing the aqueous liquid in which the specific binding reaction takes place.

6. A process according to claim 5, in which the displacer body (2) has a shape substantially complementary to and slightly smaller than that of the well or cup (1), preferably reducing the volume available in the well or cup (1) for occupation by the aqueous reaction liquid by a factor of 2-10.

7. A process according to any of claims 1-6, characterised in that the immobilised specific binding reagent (b) is either an immobilised antibody or an immobilised substance to which an immunoglobulin under assay can specifically bind, and marker conjugated-reagent (c) comprises either another antibody specific to the binding partner of antibody reagent (b), or an antiglobulin specific for an immunoglobulin under assay.

8. A process for carrying out a specific binding assay in which (a) a sample under assay, possibly containing a substance being tested for, is reacted with (b) an antibody of corresponding specificity immobilised on a solid support and (c) an antibody (also with binding specificity for the substance being tested for) conjugated to a detectable marker, thereby to form a complex between whatever quantities are present of the substance being tested for with reagents (b) and (c), and in which after the binding reaction the amount of detectable marker bound is directly or indirectly assayed, characterised in that the antibodies comprised in reagents (b) and (c) are monoclonal antibodies.
9. Materials for carrying out specific binding assay according to any of claims 1-8, comprising (i) an immobilised specific binding partner for a substance to be tested for, carried on a solid support, and (ii) a marker-conjugated specific binding partner for the substance to be tested for, which can be added to a reaction liquid contacting immobilised reagent (i) either as a slow-release form, or in any form provided that the specific binding partners in reagents (i) and (ii) include an antibody of narrow specificity so that reagents (i) and (ii) do not interfere with each other's binding reactions with the substance to be tested.

10. Processes and materials for carrying out specific binding assay, substantially as described above with reference to the joint and several features of the accompanying specification and drawings.

11. A process for carrying out immunoassay in which a sample under assay is reacted with two specific binding reagents of which one or each is a monoclonal antibody of narrow and different, non-interfering specificity to ensure that the desired assay reactions do not impede each other.
**INTERNATIONAL SEARCH REPORT**

**I. CLASSIFICATION OF SUBJECT MATTER**

According to International Patent Classification (IPC) or to both National Classification and IPC

Int.Cl. 3: G 01 N 33/54

**II. FIELDS SEARCHED**

Minimum Documentation Searched

<table>
<thead>
<tr>
<th>Classification System</th>
<th>Classification Symbols</th>
</tr>
</thead>
<tbody>
<tr>
<td>Int.Cl. 3</td>
<td>G 01 N 33/54; G 01 N 33/56; A 61 K 39/395</td>
</tr>
</tbody>
</table>

Documentation Searched other than Minimum Documentation to the Extent that such Documents are Included in the Fields Searched

**III. DOCUMENTS CONSIDERED TO BE RELEVANT**

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of Document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to Claim No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>GB, A, 2029571, published March 19, 1980, see page 1, lines 45-53, claims 1,4,5 and 8, Bio-Rad Laboratories Inc.</td>
<td>1,7,9,10</td>
<td></td>
</tr>
<tr>
<td>GB, A, 2016687, published September 26, 1979, see page 1, line 64 – page 2, line 4, claims 1,4,7 and 8, Abbott Laboratories</td>
<td>1,4,7,9,10</td>
<td></td>
</tr>
<tr>
<td>GB, A, 2034466, published June 4, 1980, see page 1, lines 15-36, claims 1,2,4, 8,9,11,15,17,23,24 and 26, Ames-Yissum Ltd.</td>
<td>1,5,7,10</td>
<td></td>
</tr>
<tr>
<td>GB, A, 1562957, published March 19, 1980, see page 3, lines 29-59, figures 1 and 2, Immuno Aktiengesellschaft für Chemischmedizinische Produkte</td>
<td>5,6</td>
<td></td>
</tr>
<tr>
<td>GB, A, 1584129, published February 4, 1981, see the whole document, R.N. Piasio et al.</td>
<td>1,5,6,10</td>
<td></td>
</tr>
</tbody>
</table>

* Special categories of cited documents:
  
  "A" document defining the general state of the art
  
  "E" earlier document but published on or after the international filing date
  
  "L" document cited for special reason other than those referred to in the other categories
  
  "O" document referring to an oral disclosure, use, exhibition or other means
  
  "P" document published prior to the international filing date but on or after the priority date claimed
  
  "T" later document published on or after the priority date and not in conflict with the application, but cited to understand the principle or theory underlying the invention
  
  "X" document of particular relevance

**IV. CERTIFICATION**

Date of the Actual Completion of the International Search: October 2, 1981

Date of Mailing of this International Search Report: October 15, 1981

International Searching Authority: EUROPEAN PATENT OFFICE, Branch at The Hague

P.O.Box 5818 Patentlaan 2
2280 HV RIJSWIJK (ZH), The Netherlands

Signature of Authorized Officer: G.L.M. Kruydenberg
### FURTHER INFORMATION CONTINUED FROM THE SECOND SHEET

| E | EP, A, 0027008, published April 15, 1981, see the whole document, Ventrex Laboratories Inc. | 1,5,6,7,10 |
| A | US, A, 3645090, published April 4, 1972, A.H.W.M. Schuurs et al. cited in the application | |

### V. OBSERVATIONS WHERE CERTAIN CLAIMS WERE FOUND UNSEARCHABLE 10

This international search report has not been established in respect of certain claims under Article 17(2) (a) for the following reasons:

1. Claim numbers, because they relate to subject matter not required to be searched by this Authority, namely:

2. Claim numbers, because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:

### VI. OBSERVATIONS WHERE UNITY OF INVENTION IS LACKING 11

This International Searching Authority found multiple inventions in this international application as follows:

1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims of the international application.

2. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims of the international application for which fees were paid, specifically claims:

3. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claim numbers:

### Remark on Protest

- The additional search fees were accompanied by applicant's protest.
- No protest accompanied the payment of additional search fees.