

(19) AUSTRALIAN PATENT OFFICE

(54) Title
Detection of candida

(51) 6 International Patent Classification(s)
G01N 033/569 C07K 001/36
G01N 033/571

(21) Application No: 2002248996 (22) Application Date: 2002.04.23

(87) WIPO No: W002/088741

(30) Priority Data

(31) Number	(32) Date	(33) Country
09/841,188	2001.04.25	US 7

(43) Publication Date : 2002.11.11
(43) Publication Journal Date : 2003.04.17

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(56) Related Art
Sinclair et al (1987) Med. Lab. Sciences 44: 137-140

(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property Organization
International Bureau



(43) International Publication Date
7 November 2002 (07.11.2002)

PCT

(10) International Publication Number
WO 02/088741 A1

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- (21) International Application Number: PCT/AU02/00507 (81) Designated States (*national*): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LI, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW.
- (22) International Filing Date: 23 April 2002 (23.04.2002)
- (25) Filing Language: English
- (26) Publication Language: English
- (30) Priority Data: 09/841,188 25 April 2001 (25.04.2001) US (84) Designated States (*regional*): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).
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- Published:**
— with international search report
- For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.*



WO 02/088741 A1

(54) Title: DETECTION OF CANDIDA

(57) Abstract: The present invention relates to a method and a means of diagnosing Candida infection. In particular the present invention relates to a method of diagnosing Candida infection by measuring the levels of antibody to Candida cytoplasmic antigen present in a biological sample taken from a subject at risk of, or suspected to be suffering from a Candida infection.

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DETECTION OF CANDIDAFIELD OF THE INVENTION

5 The present invention relates to a method and a means of diagnosing *Candida* infection. In particular the present invention relates to a method of diagnosing *Candida* infection which is both sensitive and rapid.

BACKGROUND OF THE INVENTION

10 *Candida* is the most commonly identified causative agent of oral or vaginal thrush. However, over the last few decades *Candida* has emerged as a significant cause of life-threatening infections in hospital patients. Ironically the increasing incidence of these "invasive" or
15 "systemic" *Candida* infections has been advances in modern medicine. Patients that are now surviving major injuries, surgery, cancers and organ transplants are vulnerable to life-threatening *Candida* infections. In the United States, *Candida* is now the fourth most common cause of
20 blood infections in hospitals.

The major problem with systemic *Candida* infections is that there are few definitive clinical signs or symptoms. Treatment is largely based on suspicion rather than a definitive diagnosis. Even with the
25 availability of anti-fungal drugs such as fluconazole a high mortality rate (30 to 70%) is associated with systemic *Candida* infections. The high rate of mortality is largely due to the rapid onset of infection and a rapidly fatal outcome. Without an accurate diagnosis the
30 infection often goes unnoticed until it is too late to effectively treat. This has led to a comment by clinicians that *Candida* infections are usually diagnosed at autopsy. Accordingly, there is a need for a rapid diagnostic assay that is capable of early diagnosis of
35 *Candida* infection so that appropriate treatment may be instituted thereby reducing the mortality rate.

The main difficulty in the diagnosis of *Candida*

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infections is that being a commensal, mere isolation of *Candida* from body surfaces, or orifices, is not diagnostic of an infection. Culture of *Candida* from blood or deep tissue is still the main method of diagnosis of systemic *Candida* infections. However, it can take several days for a culture to become positive, by then it may be too late to effectively treat the infection. Also, false positives may occur due to contamination from superficial body sites. Of more importance, is the observation that in up to fifty percent of autopsy proven cases of systemic candidiasis, blood cultures were negative and therefore of no diagnostic value.

Nuclear magnetic resonance (NMR) and radioisotope scanning have been used to detect *Candida* infections in tissues and organs. However, those methods are not useful for early diagnosis.

Recently analysis of the *Candida* metabolite arabinitol was proposed as a diagnostic tool. However, as arabinitol is produced by the human body, further clinical studies have cast doubt on its value.

The polymerase chain reaction (PCR) has also been used in the diagnosis of invasive *Candida* infections. However, PCR has not established itself as a useful diagnostic method for *Candida* for the same reasons as outlined above ie *Candida* is a ubiquitously present microorganism and false positives, due to superficial contamination, are prevalent.

Immunoassays are the established procedures for the diagnosis of many types of infectious diseases. Immunoassays have the advantage that they are rapid and have a standardised assay format. Immunoassays can be designed to either detect *Candida* antigens, or host antibodies reactive against *Candida* antigens. Several immunoassays are commercially available for the detection of *Candida* antigens in sera or other body fluids. However, these assays lack either sensitivity or specificity or both.

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Immunoassays have been developed based on the detection of immunodominant *Candida* antigens. *Candida* mannan is a highly immunogenic cell wall antigen. However, as *Candida* is a commensal, most individuals have antibody to *Candida* mannan, so its usefulness in the diagnosis of systemic infection is limited. The applicant has now surprisingly found that a more discriminatory assay for *Candida* than previously used is the detection of cytoplasmic antigen. The advantage of this diagnostic assay is that antibody to this cytoplasmic antigen is only produced in response to an actual infection. The applicant has further demonstrated that the use of a combination of cytoplasmic antigen with other antigens is very predicative of *Candida* infection.

Accordingly, the present invention overcomes or at least alleviates the problems normally associated with diagnosing *Candida* infection.

SUMMARY OF THE INVENTION

In its most general aspect, the invention disclosed herein provides a simple and rapid method for diagnosis of *Candida* infection. The method of diagnosis of *Candida* infection may be used to screen large numbers of samples for possible infection.

Accordingly, in one aspect, the invention provides a method of diagnosing *Candida* infection, comprising the steps of:

a). obtaining a biological sample from a subject at risk of, or suspected to be suffering from, *Candida* infection, and

b). measuring the levels of antibody to *Candida* cytoplasmic antigen present in the biological sample.

Antibody levels may be measured using known techniques of immunology including enzyme-linked immunoassay (ELISA or EIA), biligand binding (sandwich technique), fluorometric assay, chemiluminescent assay, immunochromatography, radialimmunodiffusion or

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radioimmunoassay (RIA). ELISA, immunochromatography or chemiluminescent assay methods are particularly preferred, since these are quick, sensitive, and specific, and are readily automated for large-scale use. These methods also
5 provide quantitative determinations.

The diagnostic method utilises antigens expressed by *Candida*, especially cytoplasmic antigen. The antigens isolated from *Candida* as disclosed herein may, in certain embodiments of the diagnostic method of the present
10 invention, be immobilised on an inert surface, embedded in a gel, or may be conjugated to a molecule which imparts colour, fluorescence or radioactivity to the antigen.

In a second aspect, the invention provides a method for assessing the prognosis of *Candida* infection,
15 comprising the steps of measuring the levels of antibody to *Candida* cytoplasmic antigen in a biological sample.

Persons skilled in the art will appreciate that the techniques disclosed herein may be used on any type of biological sample. Preferable the biological sample is
20 selected from the group consisting of bone marrow, plasma, spinal fluid, lymph fluid, the external sections of the skin from respiratory, intestinal, and genitourinary tracts, tears, saliva, milk, blood; both whole blood and sera, blood cells, tumours and organs. Most preferably
25 the biological sample is sera.

Biological samples that may be analysed by the method of the present invention can also be obtained via swabs, shunts or the like. The biological samples may be analysed directly, or may be treated prior to testing by,
30 for example, concentration or pH adjustment.

In a third aspect, the present invention further provides a method of detecting the presence or absence of a *Candida* antibody comprising the steps of:

- a). exposing a biological sample, which may
35 include a *Candida* antibody, to an isolated cytoplasmic *Candida* antigen; and
- b). detecting the reaction between antibody and

antigen.

In an especially preferred embodiment of the present invention the diagnostic assay further utilises other *Candida* antigens in combination with the cytoplasmic
5 antigen. In particular the cell wall antigen (including mannose) and/or purified immunodominant antigen (enolase) are utilised.

Accordingly, in a forth aspect of the present invention there is provided a method of diagnosing *Candida*
10 infection, comprising the steps of:

- a). obtaining a biological sample from a subject at risk of, or suspected to be suffering from, *Candida* infection, and
- b). measuring the levels of antibody present in
15 the biological sample to *Candida* cytoplasmic antigen in combination with measuring the levels of antibody to either cell wall antigen or immunodominant antigen (enolase) or both.

The reagents and means of diagnosis of the present invention may also be embodied in a kit for use in
20 a diagnostics laboratory or may be adapted and automated for analysing large numbers of samples at a central receiving centre.

Accordingly, in a fifth aspect the invention provides a kit when used for detecting the presence or
25 absence of a *Candida* antibody in a biological sample, comprising:

- a). a biological sample collection device;
- b). a cytoplasmic *Candida* antigen; and
- 30 c). means for detecting reaction between the antibody and antigen in the sample.

Suitable buffering agents and ionic salts may also be included in the kit.

In a sixth aspect the invention provides a method
35 of preparing a cytoplasmic antigen comprising the step of removing lipoproteins by chloroform extraction.

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BRIEF DESCRIPTION OF THE FIGURES

Figure 1 shows a coomassie blue stained SDS-PAGE with major protein bands of the *Candida* cytoplasmic antigen fraction observed at 55kDa, 35 to 45kDa region, 30kDa and 20kDa.

Figure 2 shows a single coomassie blue band of 48kDa corresponding to the expected size of the enolase antigen.

Figure 3 shows a coomassie blue stained gel of the clarified cell wall antigen preparation. A broad smear of stain can be seen ranging in size from 90kDa to 200kDa

Figure 4 shows a number of sera screened against the *Candida* cytoplasmic antigen preparation.

Figure 5 shows antibody reactivity to the three *Candida* antigens - cytoplasmic, cell wall and immunodominant antigens, using negative control sera.

Figure 6 shows antibody reactivity to the three *Candida* antigens - cytoplasmic, cell wall and immunodominant antigens, using sera from patients with superficial candidiasis.

Figure 7 shows antibody reactivity to the three *Candida* antigens - cytoplasmic, cell wall and immunodominant antigens, using sera from patients with systemic candidiasis.

Figure 8 shows the error bar of the Applicant antigen test values in the different blood culture patients (95%CI).

Figure 9 shows an error plot of the mean *Candida* antibody values measured by the Applicant antigen test in both the blood culture positive and negative groups of patients (95% confidence interval).

Figure 10 shows an error bar graph of the Applicant antigen test data for invasive candidiasis and healthy controls.

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ABBREVIATIONS USED

	EDTA	Ethylenediaminetetraacetic acid
	EIA	Enzyme immunoassay
5	ELISA	Enzyme-linked immunosorbent assay
	RIA	Radioimmunoassay
	BSA	Bovine serum albumin
	DMSO	Dimethyl sulfoxide
	β -Me	β -mercaptoethanol
10	TMB	3,3',5,5'-tetramethyl-benzidine

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

The practice of the present invention employs, unless otherwise indicated, conventional molecular biology, cellular biology, and immunoassay techniques within the skill of the art. Such techniques are well known to the skilled worker, and are explained fully in the literature. See, e.g., Harlow and Lane, "Antibodies: A Laboratory Manual" (1988); Maniatis, Fritsch & Sambrook, "Molecular Cloning: A Laboratory Manual" (1982); "Animal Cell Culture" (R.I. Freshney, ed., 1986); "Immobilised Cells and Enzymes" (IRL Press, 1986); B. Perbal, "A Practical Guide to Molecular Cloning" (1984); Sambrook, et al., "Molecular Cloning: a Laboratory Manual" (1989) and Ausubel, F. et al., 1989-1999, "Current Protocols in Molecular Biology" (Green Publishing, New York).

In describing the present invention, the following terminology is used in accordance with the definitions set out below.

As used herein, a "biological sample" refers to a sample of tissue or fluid isolated from a individual, including but not limited to bone marrow, plasma, serum, spinal fluid, lymph fluid, the external sections of the skin, respiratory, intestinal, and genitourinary tracts, tears, saliva, milk, blood; both whole blood and anti-coagulated whole blood, blood cells, tumours, organs, and also includes samples of *in vivo* cell culture

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constituents, including but not limited to conditioned medium resulting from the growth of cells in cell culture medium, putatively *Candida* infected cells, recombinant cells, and cell components.

5 "Human tissue" is an aggregate of human cells which may constitute a solid mass. This term also encompasses a suspension of human cells, such as blood cells, or a human cell line.

For the purposes of this specification it will be
10 clearly understood that the word "comprising" means "including but not limited to", and that the word "comprises" has a corresponding meaning.

It will be clearly understood that, although a number of prior art publications are referred to herein,
15 this reference does not constitute an admission that any of these documents forms part of the common general knowledge in the art, in Australia or in any other country.

Persons skilled in the art will appreciate that
20 any number of different immunoassays may be used in the present invention. For example, the *Candida* antigens disclosed herein may be used in antibody capture assays, antigen capture assays, wherein the antigen/antibody complex forms a "special" class of antigen or two-antibody
25 sandwich assays.

TECHNIQUES USED FOR ANTIGEN PREPARATION

The term "*Candida* antigen" as used here means any one of the three separate types of *Candida* antigen
30 utilised in the present invention, namely, cell wall antigen (including mannose), total cytoplasmic antigen (mannose depleted) or purified immunodominant antigen (enolase). Use of the term "*Candida* antigens" means that all three antigens were involved or could be utilised. A
35 number of techniques may be used to prepare the *Candida* antigens including biochemical extraction, column chromatography, Gel fractionation, gene cloning,

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differential precipitation, filtration, dialysis or centrifugation; however, the preferred techniques are those disclosed herein. Briefly, these techniques involve either mechanical, chemical or enzymatic lysis of *Candida* 5 cells, followed by separation of insoluble cell walls from soluble cytoplasmic fraction by centrifugation, filtration and dialysis. Chemical treatment of cell wall fraction to release cell wall antigens followed by centrifugation and dialysis. Filtration and organic extraction of soluble 10 cytoplasmic cell extract. Separation of mannoproteins by ConA affinity chromatography. Purification of the immunodominant enolase antigen from the soluble cytoplasmic extract by anion and cation affinity chromatography. It will be appreciated by those skilled 15 in the art that other techniques, or modifications or variations of the above techniques, may be adopted without adversely affecting the spirit of the present invention.

TECHNIQUES USED FOR ANTIBODY PREPARATION AND LABELLING

20 Antiserum to the *Candida* antigens disclosed herein may be produced in a host animal such as rabbit or sheep. The serum fraction containing the antibody may be isolated by standard techniques. This antiserum may be employed in several of the embodiments of the invention 25 hereinafter set forth, or a more sensitive and specific antibody might be obtained by further purification of the serum by electrophoresis, high-speed centrifugation or the like. Ultimately, large quantities of highly specific monoclonal antibody may be produced by means of the 30 hybrid-myeloma techniques by methods known to those skilled in the art.

Certain embodiments of the present invention employ antibody to the *Candida* antigens immobilised on cellulose, agarose, sephadex or glass beads or other 35 similar inert surfaces such as metal, plastic or ceramic which do not interfere with subsequent reaction. Adsorption, Br-CN activation or other techniques known in

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the art may be employed to immobilise the antibody.

Other embodiments of the present invention employ the antibody to the *Candida* antigens conjugated to a chromophoric (highly coloured) molecule, an enzochromic (an enzyme which produces colour upon addition of reagents) molecule, fluorochromic (fluorescent) molecule or a luminogenic (luminescent) molecule.

The conjugate of antibody with enzyme is made using techniques known in the prior art. (For references, see Avrameas, S. and Uriel, J., in *Comptes Rendus Hebdomadaires des Seances de l'Academie des Sciences*, vol. 262, p. 2543, (1966); Nakane, P.K. and Pierce, G.B., in *Journal of Histochemistry and Cytochemistry*, vol. 14, p. 929, (1966); Nakane, P.K., in *Methods in Enzymology*, vol. 37, p. 133, (1975)).

Chromophoric molecules that may be used are 2,3-dinitrobenzene (DNB) salts, dinitrophenol (DNP) and methyl and butyl orange. Other suitable chromophoric agents are well known in the art. Enzochromic molecules that may be conjugated with the antibody are enzymes that give colour with appropriate reagents. Examples are alkaline phosphatase (ALP) which develops colour with nitrophenyl phosphate (NPP), glucose oxidase with glucose, and D-galactopyranoside. These and other examples are well known in the art. Examples of fluorogenic agents are 2,4-dinitrofluorobenzene and "pipsyl" derivatives. Luminogenic molecules may be conjugated to antibodies by the method of Branchini, et al. (*Biochem. Biophys. Res. Commun.* 97, 334 [1980]). The term "chromophoric" hereinafter is intended to include "enzochromic", "fluorochromic" and "luminogenic" molecules as well.

Certain embodiments of the invention also utilise *Candida* antibody tagged with a radioactive element. I^{125} conjugated by means of the chloramine-T procedure is a common example, but other methods known in the art may also be employed.

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TECHNIQUES USED FOR ANTIGEN IMMOBILISATION AND LABELLING

Antigen molecules may be immobilised on a solid carrier by a variety of methods known in the art, including covalent coupling, direct adsorption, physical entrapment and attachment to a protein-coated surface. For references describing the methodology, see Silman, I. H. and Katchalski, E. in Annual Review of Biochemistry, Vol. 35, p. 873 (1966); Melrose, G. J. H., in Review of Pure and Applied Chemistry, Vol. 21, p. 83, (1971); and Cuatrecasas, P. and Anfinsen, C. B., in Methods in Enzymology, Vol. 22, (1971).

Lai et al. (German OS No. 2,539,657, U.S. Pat. No. 4,066,512) discloses a method of attachment to a protein-coated surface. In this method, the internal and external surfaces of a microporous membrane are first coated with a water-insoluble protein such as zein, collagen, fibrinogen, keratin, glutelin, polyisoleucine, polytryptophan, polyphenylalanine, polytyrosine, or copolymers of leucine with p-amino phenylalanine. Such a coating renders the membrane capable of immobilising a wide variety of biologically active proteins including enzymes, antigens, and antibodies. A microporous structure is defined as one having more than 50% of its total volume in the form of pores ranging in size from 25 nanometres to 25 micrometers, preferably from 25 nanometres to 14 micrometers. A pore size range from 25 nanometres to 5 micrometers is employed in most applications herein. Uncoated microporous membranes have as much as 70 to 75% of their volume as pore space. The pores permit liquid flow through the membrane. After being coated by zein, for example, the pore space is reduced 5 to 10% with the result that the structure retains its essential properties of having a high proportion of its volume as pore space and permitting liquid flow through the pores. The structure has a large surface area in contact with any solution contained within the pores.

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Such a coated membrane, having immobilised antigen or antibody, provides a compact, easy to manipulate carrier for the immobilised antigen or antibody. Its integral structure permits removal of bound
5 from unbound components by simple mechanical means.

Non-specific binding may be minimised by interposing a second stage immobilisation step, in which an immunochemically neutral protein is immobilised to the filter. Immobilisation therefore occurs in two stages
10 according to a preferred embodiment of the invention: a first stage in which the desired immunochemical component is immobilised, and a second stage, following the completion of the first, in which an immunochemically
15 neutral protein such as fetal calf serum or bovine gamma globulin is next immobilised. The term immunochemically neutral is defined in terms of the specific components of the assay. Any protein, which does not combine immunochemically with a component of the assay or with one
20 of the reagents, is considered immunochemically neutral, even though such protein might be immunochemically reactive in another system.

Where the substance to be detected is an antibody, the immunochemically reactive moiety of the conjugate must be an antibody capable of binding
25 immunochemically with the antibody to be tested. Such antibodies may be obtained by immunising an animal with the antibody or immunoglobulin fraction of serum from the animal in which the antibody to be tested originated. For
30 example, where the antibody to be tested is a human antibody, a goat antibody against human antibody is obtained from the serum of a goat immunised against human immunoglobulin (antibody). The enzyme moiety may be any
35 enzyme capable of catalysing a reaction which can be detected by any method known to those skilled in the art, and which retains its activity after conjugation with antibody. Horseradish peroxidase is preferred because of its convenience and suitability to a wide range of

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applications. It is well known that the enzyme catalyses the oxidations of a variety of organic compounds in the presence of hydrogen peroxide. Many such organic substrates are chromogenic, ie. undergo a colour change upon oxidation.

It has been found in the present invention that the purity of the enzyme preparation used in the formation of conjugate has an effect on the degree of non-specific binding. The greater the purity of the enzyme preparation, the less the non-specific binding. In part, the reduction is made possible because, the total amount of conjugate protein required is reduced as the specific activity of the enzyme is increased. The opportunity for non-specific binding is therefore reduced as well. In the preferred embodiment, the use of a highly purified peroxidase preparation has been found to significantly reduce the amount of colour reaction observed in control samples as compared with known positives.

20 TECHNIQUES USED FOR CANDIDA ANTIBODY DETECTION

Antibody Capture Technique

A *Candida* antigen prepared by the techniques disclosed herein is immobilised, preferably on an inert surface such as PVC, paper or a similar fibulous mat. The immobilised *Candida* antigen is then put into contact with a sample suspected of containing *Candida* antibody. In the case of aqueous samples such as blood or urine, the solution is buffered and ionic salts may be present at optimum concentration for *Candida* antibody-*Candida* antigen interaction. TRIS or borate buffered phosphate at pH 7.5 to 9.0 and ionic strength about 0.010 to 0.5, for example, are suitable buffering agents and ionic salts. The inert surface with *Candida* antigen or *Candida* antigen-*Candida* antibody complex thereon is next put into contact with antibody to *Candida* antigen conjugated to a chromophormic molecule. Preferably the *Candida* antigen is in solution

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buffered at pH from about 7.5 to 9.0 and ionic concentration equivalent to about 0.01M to about 0.1M NaCl. After careful rinsing under water or with suitable surfactants such as Tween 20 to remove excess coloured antibody, the inert surface is inspected for colour, fluorescence or luminescence directly or after addition of colour-developing agents. Colour on the inert surface indicates interaction between immobilised *Candida* antigen-*Candida* antibody complex in solution. A control may be run for colour comparison.

This technique may be adapted to clinical use by employing *Candida* antigens tagged with radioactive elements and observing either depletion of activity in solution or uptake on solid support of radioactivity. This embodiment is highly sensitive and rapid and suitable for large numbers of samples.

Enzyme-Linked Immunoassay-ELISA

A solution comprising *Candida* antibody conjugated to enzyme which forms colour with developing reagents and buffer and ionic salts suitable for reaction between *Candida* antigen and the *Candida* antibody is put into contact and allowed to react with *Candida* antigen immobilised, preferably, on an inert surface such as PVC, paper strip or glass bead. The amount of enzochromic conjugated *Candida* antibody is sufficient to saturate about 50% of the reactive sites on the immobilized antibody. The inert surface with antibody-*Candida* antigen enzyme complex is put into contact with buffered sample suspected of containing *Candida*, said sample having an unknown amount of *Candida* antibody. The colour of the resultant immobilised antibody-*Candida* antigen-enzyme complex on the strip after colour developing reagents are added is observed in comparison to a control strip which has not been treated with sample containing *Candida* antibody. Dilution in colour on inert surface treated with sample means presence of *Candida* antibody in the

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unknown sample.

This method may be adapted for clinical use by contacting samples and immobilised enzyme, preferably in tubes which may be centrifuged and watching developing colour spectrophotometrically. This embodiment is very sensitive and rapid.

Radialimmunodiffusion-Precipitin Reaction

One of the *Candida* antigens is suspended in a softened gelatinous medium such as agar or agarose along with buffers and salts to maintain pH between about 6.0 to 9.0 and ionic strength between about 0.01M to 0.5M for optimal antigen-antibody interaction. The suspending medium of U.S. Pat. No. 4,259,207 is a suitable example. The mixture is spread out to harden on a test plate or, preferably, poured into a disc-shaped container such as an Octolony plate. A small amount of sample is placed on the solidified gel, preferably in a centre well and the plate or disc is allowed to stand preferably covered for a period of hours. Diffusion of sample into the surrounding area occurs during this period. If the *Candida* antibody is present, it reacts with the embedded *Candida* antigen and causes an opaque area in a radial pattern about the point of application of sample. A control can be run for comparison. Calibration of an amount of *Candida* antibody in the sample, if desired, can be obtained by controlling temperature, time and size of sample and comparing the resultant size of radial area with one of known concentration.

Radioimmunoassay

A *Candida* antigen of the present invention is immobilised on an inert surface such as glass beads in a separation column. A portion of *Candida* antigen is conjugated to a radioactive element, preferably I^{125} and allowed to react with the immobilised *Candida* antigen in an amount sufficient to, saturate 50% of the binding

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sites. The immobilised *Candida* antigen-enzyme complex is put into contact with a sample suspected of containing *Candida* antibody, the sample being buffered between pH 6-9 and containing total ionic salts about 0.05 to 0.5M for optimal reaction conditions for formation of *Candida* antigen-antibody complex. The *Candida* antibody is eluted from the antigen and the eluant is measured for radioactivity. Loss of activity compared to a control indicates *Candida* antibody in the sample.

10

Haemagglutination

Candida antibody may be assayed through standard haemagglutination techniques with *Candida* antigen to antibody used as sensitising agent.

15

It is to be understood that methods described hereinabove for assay of *Candida* antibody employing coloured reagents have been presented most specifically for application where neither trained personnel nor sophisticated instruments are available. These methods, however, may be adapted for use in a clinical setting where large numbers of samples are to be assayed by substituting radioactive elements for chromogenic conjugated molecules.

20

It is also to be understood that the term "colour" is not to be interpreted as being limited to the narrow visible range of the electromagnetic spectrum, but is meant to include wavelengths which may be measured by standard spectrophotographic instruments such as spectrophotometers and absorption and emission colourimeters in both the uv and the ir range.

25

Although it is contemplated that the methods of the present invention are to be applied to biological fluids themselves, the sensitivity and specificity of the method can be improved by culture of the fluids preferably on medium selective for *Candida* prior to testing.

30

Sensitivity may also be improved by preliminary treatment of biological samples with lysing agents such as

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isotonic solution, sound, or lysozyme to release *Candida* antibody into the extracellular environment. U.S. Pat. No. 4,166,765, for example, discloses suitable lysing procedures for biological samples containing bacteria.

5 Any lysing agent may be employed which does not interfere with subsequent enzyme activity.

ASSAYS EMBODIED IN KIT FORM

The diagnostic method and means of the present invention may be embodied in the form of a kit for use by individuals for self-diagnosis of *Candida* in the privacy of their homes.

The kit comprises a means for sample collection, the *Candida* antigen to *Candida* antibody and a means for detecting reaction between sample and *Candida* antigen.

In embodiments adapted for clinical use, electrophoretic separation techniques such as isoelectric focusing or zone electrophoresis which are based on differences of both size and charge distribution between products and reactants may likewise be used to separate products from reactants. Products separated electrophoretically may be detected by characteristic locations compared to standards or may be identified by colour or immunochemically. Resinous beads of charged surfaces may also be used to separate products and reactants.

The means for detecting reaction in the case of immunoassay in a preferred embodiment of the invention is a gelatinous medium in which the *Candida* antigen to antibody is suspended. The gelatinous medium is in a transparent glass or plastic container and comprises buffer and ionic salts for optimal conditions for formation of the *Candida* antigen-antibody complex. Reaction is noted as a transparent area radiating from the central point at which the sample is applied.

The means for detecting reaction in another preferred embodiment comprising immunoassay is the *Candida*

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antigen to *Candida* antibody conjugated to a chromophore in a sealed, sterile packet along with buffer and ionic salts. For assay, the contents of the packet are diluted with water in a marked tube supplied in the kit. Included
5 also in this embodiment is the antigen to *Candida* antibody immobilised on an inert surface. For assay, the inert surface with immobilised *Candida* antigen is put into contact with sample and then with the solution of chromophore-conjugated *Candida* anti-IgA antibody, protein
10 A or protein G. The inert surface is inspected for colour, which indicates *Candida*.

In a particularly preferred embodiment, the kit of the present invention is provided in the form of an immunochromatographic test strip device. There are many
15 patents that cover a number of technologies, formats, reagents and materials that may be of great value in the development and production of immunochromatographic test strip devices. For example, US Patent No. 5,075,078, International Patent Application No. WO95/16207, US Patent
20 No. 5,654,162 and European Patent No. 0810436A1. The assay methods used with the devices disclosed in these patents are essentially the same. A ligand specific for the analyte (normally, but not necessarily an antibody [Ab]) is immobilised to a membrane such as nitrocellulose.
25 The detector reagent, typically an antibody coupled to latex or colloidal metal, is deposited (but remains unbound) into the conjugate pad. When sample (urine, plasma, whole blood, etc.) is added to the sample pad, it rapidly wets through to the conjugate pad and the detector
30 reagent is solubilised. The detector reagent begins to move with the sample flow front up the membrane strip. Analyte that is present in the sample will be bound by the antibody that is coupled to the detector reagent. As the sample passes over the zone to which the capture reagent
35 has been immobilised, the analyte detector reagent complex is trapped. Colour develops in proportion to the amount of analyte present in the sample.

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In the present case, while the above principles are the same, rather than detecting analyte *per se*, the immunochromatographic test strip device would detect antibody. In such situations, it would be the antigen(s) disclosed herein which would be immobilised onto membranes, sample pads, reagent pads and other porous media rather than antibody. There is a wealth of information regarding the development of such devices including methods of binding antigen/antibodies to nitrocellulose and the like and detecting such bound material. See for example, Towbin et al. 1979, Proc. Natl. Acad. Sci. USA 76:4350, the entirety of which is included herein by reference.

Although the invention has been described with reference to presently preferred embodiments, it should be understood that various modifications can be made without departing from the spirit of the invention. Moreover, the following examples are offered by way of illustration only and are not intended to limit the invention in any manner. All patent and literature references cited herein are expressly incorporated.

EXAMPLE 1 PREPARATION OF CANDIDA ANTIGEN

The following three types of *Candida* antigen were prepared:

- 1). Cell wall antigen (including mannose);
- 2). Total cytoplasmic antigen (mannose depleted);

and

- 3). Purified immunodominant antigen (enolase).

A clinical isolate of the *Candida albicans*, was obtained from a patient with vaginal thrush. The identity of the *Candida* species was confirmed with the use of an API 20C Auxonogram strip (API System S.A., France). The *C. albicans* isolate was designated KEMH5.

200ml YEPD culture medium (1% yeast extract, 2% peptone, 2% D-glucose) was inoculated with the isolate as a starter culture and incubated for 24h at 30°C with

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aeration. The starter culture was then used to inoculate a 10L YEPD culture incubated under similar conditions in a 23L Bio-Flo Fermenter IV System (New Brunswick Scientific, Edison, NJ).

5 The *Candida* culture was harvested from the Bio-Flo fermenter system and separated from culture medium with the use of a Pellicon filtration cassette (Millipore, USA). Concentrated cells were separated from residual medium by centrifugation in 500ml centrifuge flasks for
10 15min at 1,660 x g and 4°C. The supernatant was discarded and the pelleted cells were resuspended in protein extraction buffer (20mM bis-Tris, pH 6.5). The yeast cells were then centrifuged as described previously, resuspended and pooled for further processing.

15 *Candida* cells were ruptured mechanically with the use of a Dynomill® (WAB, Switzerland). Milling was continued until 99% cell disruption was obtained. The soluble *Candida* cell extracts were collected and dispensed into 50ml centrifuge tubes. The extracts were centrifuged
20 for 12h at 8,517 x g and 4°C to precipitate insoluble cell walls. The supernatants containing the soluble cytoplasmic antigen fraction were recovered and passed through a 0.45µm filter membrane.

The filtrates were then extracted with an equal
25 volume of chilled chloroform. Following centrifugation at 4°C for 15min at 1,036 x g the upper aqueous phase was aspirated and transferred to a dialysis tube. The soluble cytoplasmic protein fractions were dialysed in column binding buffer (20mM Tris/HCl, pH 7.4, 0.5M NaCl, 1mM
30 MnCl₂.4H₂O, 1mM CaCl₂) for 12h in preparation for chromatography.

The soluble cytoplasmic antigen fraction was depleted of contaminating soluble cell wall mannoprotein by Con A-Sepharose chromatography. The dialysed
35 cytoplasmic antigen fraction was filtered through a 0.45µm filter. 50ml of the dialysed extract was applied onto a Con A-Sepharose column (2.6 x 12.5 cm) equilibrated in

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binding buffer at a flow rate of 4ml/min. The unbound flow-through fraction (non-glycosylated proteins) was collected. Bound mannoproteins were eluted with 0.5M α -methyl mannoside in binding buffer. This step was performed before the next run and to clean the column before storage.

The soluble cytoplasmic antigen fraction was dialysed overnight against 20mM Tris.Cl, pH7.4. An estimate of the quantity of protein in solution was performed using the Bio-Rad[®] (Bradford) microassay procedure in accordance with the manufacturers instructions. A portion of the cytoplasmic antigen extract was analysed by SDS-PAGE.

As shown in Figure 1 there was a number of major protein bands observed which varied in size from approx 20kDa up to approx 60kDa in size. The major staining bands being at 55kDa, four bands in the 35 to 45kDa region, 30kDa and 20kDa. This was in stark contrast to the large number of Coomassie blue staining bands in the original crude lysate prior to organic extraction and Con A-Sepharose chromatography.

Purification of the enolase antigen was conducted in the same fashion as the soluble *Candida* cytoplasmic antigen except that it was not subjected to Con A-Sepharose chromatography. Instead, following dialysis and filtering through a 0.20 μ m syringe filter (cellulose acetate), the filtered extracts were applied to a Pharmacia Biotech XK 50/20 chromatography column packed with Pharmacia Biotech Source 15Q quaternary ammonium anion exchanger (Pharmacia LKB, Uppsala, Sweden). The column was equilibrated prior to chromatography with column binding buffer 'A' (20mM bis-Tris, pH 6.5). Anion exchange chromatography of the crude extracts was controlled and recorded using the Bio-Rad[®] Econo[®] system (Bio-Rad Laboratories, USA). Bound protein was eluted from the column with a salt gradient of buffer 'B' (1M NaCl in buffer 'A', pH 6.5). The recovered fractionated

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proteins were analysed by an enzyme activity assay.

The active enzyme enolase hydrolyses D(+)-2-phosphoglyceric acid (PGA) to phosphoenolpyruvate (PEP). The production of PEP can be monitored by

5 spectrophotometry at 240nm. 20µl of protein solution was combined with 1ml of enolase substrate solution (50mM Tris-HCl pH 7.4, 2.7mM magnesium acetate, 1.0mM EDTA, 1.2mM D(+)-2-phosphoglyceric acid) in a quartz cuvette and the change of absorbance recorded at 1min intervals. The
10 specific activity was defined as the conversion of 1µmol of PGA to PEP per min per mg protein. An estimate of the quantity of protein in solution was performed using the Bio-Rad® (Bradford) microassay procedure.

Eluate fractions containing enolase activity were
15 selected and dialysed for 12h at 25°C in hpH_2O . The dialysed fractions were recovered and filtered through a 0.20µm syringe filter. The filtrate was concentrated ten-fold by evaporation under vacuum for 5h. The concentrated samples were dialysed with binding buffer 'A' (10mM sodium
20 acetate, pH 4.7) immediately prior to application to a Pharmacia Biotech Mono S HR10/10 chromatography column packed with methyl sulphamate cation exchanger (Pharmacia LKB, Uppsala, Sweden). Cation exchange chromatography was performed using the Bio-Rad® Biologic system. Bound
25 protein fractions were eluted from the column with a salt gradient of buffer 'B' (1M NaCl in buffer 'A', pH 4.7). Fractions containing enolase activity were identified by the enzyme activity assay described above.

Figure 2 shows a single Coomassie blue band of
30 48kDa corresponding to the expected size of the enolase antigen. The identification of the 48kDa antigen as the glycolytic enzyme enolase was confirmed by an enolase activity assay.

Purification of the cell wall antigen was
35 conducted as follows: the precipitated insoluble cell walls were collected following centrifugation as described above. The cell walls were washed with hpH_2O then

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collected by centrifugation at 6,000 rpm. This step was repeated three times or until the supernatant was no longer cloudy. This ensured any residual soluble cytoplasmic antigen was removed from the cell wall preparation. The washed cell wall pellet was then resuspend in 10mM Phosphate buffer pH7.4 containing 1% v/v β -Me and incubated for 30min at 37°C in a shaker to solubilise the cell wall antigens. The sample was then centrifuged for 5min at 8,000 rpm and the pellet was then discarded. The supernatant was transferred into a fresh tube and recentrifuged (5min at 8,000 rpm). The supernatant containing the solubilised cell wall antigen was then dialysed in hpH_2O for 48h at 4°C (four changes of water), or until no odour was detected. Following dialysis the sample was centrifuged three times 5min at 8,000 rpm to remove any residual particular matter.

Following clarification the cell wall antigen preparation was analysed by SDS-PAGE. The resulting Coomassie blue stained gel is presented in Figure 3. A broad smear of stain is seen ranging in size from 90 kDa to 200 kDa. The lack of discrete protein bands is typical of mannoproteins, where differences in the number of mannose groups added to the protein base results in a variety of molecular weights.

25

EXAMPLE 2 ENZYME LINKED IMMUNOSORBENT ASSAYS (ELISAS)

A serum panel was collected from 1998 to 2000 from various patients with *Candida* infections. Negative control (Control) sera (n=20) were obtained from the Red Cross Blood Bank, Perth, Australia and was obtained from healthy males in the 19 to 25 year age group. Sera (n=13) from patients with recurrent vulvo vaginal candidiasis (VVC) were obtained from King Edward Memorial Hospital, Perth, Australia. Sera (n=108) from patients with oral candidiasis were obtained from Clinipath Ltd and the UWA Dental School, Perth, Australia. Sera (n=39) from patients (n=28) with systemic candidiasis were obtained

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from Princess Margaret Hospital, Perth, Australia and Prince of Wales Hospital, Sydney, Australia.

In the case of patients with oral and vaginal *Candida* infection, confirmation of infection was made by
5 physical examination and by culture of *Candida* organisms from the relevant body site. In the case of patients with systemic infection, confirmation of infection was through positive blood culture or biopsy. In all cases the immune status of the patient was unknown.

10 Sera from patients with either superficial or systemic candidiasis were screened by ELISA using trays coated with the *Candida* cytoplasmic antigen. The protein content of each antigen preparation was determined using a commercial assay (BioRad) with BSA as a standard. A
15 series of ELISAs were performed to determine the optimal coating concentration for each antigen (data not shown). The optimal coating concentration being that which gave the greatest discrimination between a positive and a negative control serum. For each antigen the optimum
20 coating concentration was determined to be 2µg/ml.

A 96 well C8 strip microtitre plate (Greiner GmbH, Germany), was coated with either *Candida* cell wall antigen, cytoplasmic antigen, or purified enolase antigen as prepared in Example 1. 50µl of a 2.0µg/ml solution of
25 the antigen was diluted in coating buffer (0.1M NaHCO₃, pH 9.3) and added to individual wells. The plates were incubated for 12h at 4°C then equilibrated to ambient temperature. After equilibrating the plates to ambient temperature, coating solution was decanted and the plate
30 tapped dried. Plates were inverted on paper towel to drain. Alternatively excess coating solution was aspirated by the automated plate washer (Dynatech Laboratories, Chantilly VA, USA). It was important not to wash the plate at this stage.

35 A volume of 300µl of blocking solution (PBS pH 7.3, 2% (w/v) BSA (ICN, Australia), 0.01% (w/v) Tween 20), was applied to each well and incubated at 25°C for 90min.

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Blocking solution was decanted and the plate tapped dried. Plates were inverted on paper towel to drain and tapped dried for a second time. At this stage plates were either used immediately, or dried for storage. Plates to be
5 dried were placed inverted in a sealable container such as a plastic food container with a number of silica gel desiccant sachets for 48h. The inclusion of approximately 20 small desiccant sachets was adequate for the drying of 6 coated ELISA micro-well trays. Dried plates were sealed
10 into heat-sealed packets with a single desiccant sachet and labelled. Plates were stored at 4°C until required. Packets containing plates were equilibrated to ambient temperature before opening.

Human test sera diluted 1/100 in blocking
15 solution was dispensed into wells in 50µl aliquot's and incubated at 37°C for 30min. The primary antibody solution was aspirated and wells were washed six times in PBS-Tween 20. The plates were inverted on paper towels and allowed to drain for 10min. The plates were then tapped dried.

20 A volume of 100µl of a horseradish peroxidase anti-human IgG conjugate diluted 1/10,000 in blocking solution was dispensed to each well. Secondary antibody solution was incubated at 37°C for 30min. The secondary antibody solution was aspirated and wells were washed six
25 times in PBS-Tween 20. Plates were inverted on paper towel to drain for 10min and then tapped dried. Plates were inverted on paper towel for a second time and allowed to drain for 5min. Plates were then tapped dried. Particular care was employed to ensure that all traces of
30 secondary conjugate solution was removed as residual conjugate was established as the major factor responsible for disparity of results (Dynatech Laboratories Inc, USA).

A volume of 100µl of TMB liquid substrate solution was dispensed into each well and developed at 25°C
35 for 10min. The reaction was terminated with the addition of 100µl of 1M phosphoric acid or 1M H₂SO₄. The absorbance values for each well were measured at 450nm, reference

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620nm with a MRX automated plate reader.

Each immunoassay was performed in triplicate and the mean value of absorbance was used. The absorbances are shown as a Scatter diagram in Figure 4. Three groups of patients with *Candida* infections were analysed. The first group were patients with systemic candidiasis (Systemics), the second group had oral candidiasis (Oral) and the third group had vulvovaginal candidiasis (VVC). Blood bank sera (Control) from males in the 19 to 25 year age group, who were at low risk of having an undetected or subclinical *Candida* infection were used as a control. The cut-off absorbance ($OD_{450} = 0.22$) was the mean value of the negative control sera. From these data the cytoplasmic antigen ELISA had a sensitivity of 89% and a specificity of 95%. This is higher than that reported for other *Candida* serological tests (Zoller et al., 1991. J. Clin. Micro. 29:1860-1867).

To further increase the sensitivity of the *Candida* ELISA multiple antigens were used. These were the cell wall, cytoplasmic and native enolase (described above).

The use of multiple antigens increased the sensitivity of the *Candida* ELISA. It also provided greater discrimination between superficial and systemic infection. Six negative control sera (serum obtained from healthy males in the 19 to 25 year age group) were used in ELISAs with microtitre tray wells individually coated with the three *Candida* antigens. For each serum the antibody titre to each of the three antigens was below that of the cut-off line (Figure 5). This line is the cut off value assigned based on a comparison of the average antibody titres of sera from control patients versus those of candidiasis patients. The value plotted on the y-axis of the graph is the ratio of the cut-off absorbance divided into the absorbance of the test serum.

Serum obtained from 6 patients with superficial candidiasis was then reacted in the ELISA. Again the

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absorbance value of each serum was divided by the absorbance of the cut-off (Figure 6). The characteristic antibody response of the sera from patients with superficial candidiasis was a high titre against the cell wall antigen preparation (1.5 to 2 times the cut-off value). The antibody reactivity to the complete cytoplasmic antigen preparation was positive in most cases (1 to 1.5 times the cut-off). In contrast the antibody titre to the enolase antigen was below or equal to that of the cut-off. There is a correlation between the antibody titre to the internal *Candida* antigens (cytoplasmic and enolase) and the severity of the superficial infection (data not shown). However, the severity of the infection in the six patients analysed was not known.

Six sera taken from patients with systemic candidiasis (confirmed by positive blood culture) were analysed by ELISA. The results are presented in Figure 7. In the case of the patients with systemic candidiasis the antibody response to the cell wall antigen preparation was positive (1.5 to 2 times the cut-off value). Also, the antibody titres to the internal *Candida* antigens (cytoplasmic and enolase) were also positive (1.5 to 2.5 times cut-off value).

25 CONCLUSIONS

The *Candida* mannan depleted cytoplasmic antigen preparation disclosed herein can be used to identify patients with *Candida* infections. The sensitivity and specificity using an ELISA with microtitre trays coated with this antigen is greater than that obtained by other *Candida* diagnostic tests. Further, the ELISA assay format disclosed herein is easier to perform, more robust and more rapid than formats used in other available *Candida* diagnostic assays. The ELISA format also has the advantage that it is quantifiable. This enables the patient to be monitored over a period of time and changes in the titre of the antibody response to the *Candida*

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antigens recorded. The ability of the test to monitor overtime the antibody titre to *Candida* antigens has a prognostic value in terms of measuring the patient's response to antifungal drugs and in the overall survival prospects of the patient. Another advantage of the cytoplasmic antigen preparation is that the method developed to produce the antigen is simpler and more rapid than other available procedures (eg. compare with that of Zoller et al., 1991, *supra*).

10

EXAMPLE 3 CLINICAL EVALUATION IN FRANCE

Clinical evaluation of the triple antigen test kit as described in Examples 1 and 2 was undertaken in the Department of Parasitology and Medical Mycology at the University of Grenoble Faculty of Medicine, Grenoble, France using stored sera.

Sera from two groups of patients were analysed: those that were blood culture positive and those that were blood culture negative. When possible, sera were taken before, at the time of and after the first day of positive blood culture to be tested. The blood culture negative group was divided into 3 subgroups: Patients that were colonised with *Candida* and were serology positive, patients that were colonised with *Candida* and were serology negative, and patients that were not colonised with *Candida* and serology negative. The sera were obtained from patients hospitalised between 1998 and 2000.

The triple antigen ELIZA test ("the Applicant antigen test") was performed according to Example 2. The cut-off calibrator sera was obtained by pooling sera taken from males in 19 to 25 age group who had no history of *Candida* infections.

Table 1 shows that the Applicant antigen test was positive in 15 out of 19 patients who had a positive blood culture.

35

TABLE 1

APPLICANT TRIPLE ANTIGEN TEST AS USED IN THE FRENCH STUDY

Patient Group	Patient ID	Serum ID	Candida species	Date of serum relative to first +ve culture	Applicant Triple Ag Abs	Applicant Triple Ag Abs/ Cut off (0.46) ratio	Applicant Triple Ag Score (0.46 cut-off)	Serology-immunofluorescence IFI	IEP Pasteur	IEP FSK	Ag emie	
Candidemia	AMI	C1	C.g	-4	1.597	3.5	+++	+++				
	AMI	C2		+3	1.519	3.3	+++	++	+++	+++		
	BRIG	C3	C.a	-2	0.385	0.8	-	-				
	BRIG	C4		+10	0.325	0.7	-	-				
	COE	C5	C.t	-13	0.405	0.9	-	-				
	COE	C6		+1	0.734	1.6	+	+++				
	COE	C7		+29	0.632	1.4	+	-				
	COH	C8	C.g	-14	0.597	1.3	+	-				
	COH	C9		+9	0.661	1.4	+	-				
	COH	C10		+65	0.391	0.9	-	-				
	COM	C11	C.g	+1	1.805	3.9	+++	+	+++	++		
	COM	C12		+19	1.862	4.0	+++	++	++	+++		
	CON	C13	C.g	-27	0.732	1.6	+	-			+	
	CON	C14		+1	0.5	1.1	(+)	-			++	
	CON	C15		+8	0.367	0.8	-	-			++	
	DA SI	C17	C.a	+2	1.805	3.9	+++	+	++++	++		
	DA SI	C18		+70	1.277	2.8	++	++	++	(+)		
	FER	C19	C.a	-35	0.693	1.5	+	-				
	FER	C20		+2	0.368	0.8	-	-			-	
	FER	C21		+16	0.229	0.5	-	-				
	FON	C22	C.a	-46	0.51	1.1	(+)	-				
	FON	C23		+3	1.899	4.1	+++	+++				
	FON	C24		+27	1.854	4.0	+++	+++				
	HAM	C25	C.a	+1	1.083	2.4	++	+	+	+	+	
	HAM	C26		+31	1.168	2.5	++	+	++	(+)		
	HEN	C27	C.t	+2	0.324	0.7	-	-				
	HEN	C28		+7	0.646	1.4	+	-				
	HEN	C29		+40	0.432	0.9	-	-				
	KHA	C30	C.a	-13	0.332	0.7	-	-				
	KHA	C31		+2	1.553	3.4	+++	+				
	KHA	C32		+27	1.393	3.0	+++	+				
	LON	C33	C.a	-2	0.341	0.7	-	-				
	LON	C34		+6	0.447	1.0	(+)	-				
	LON	C35		+61	0.35	0.8	-	-				
	MAN	C36	C.a	-28	0.505	1.1	(+)	-				
	MAN	C37		+5	0.288	0.6	-	-				
	MAN	C38		+72	0.199	0.4	-	-				
	NI	C39		C.t & C.k	+3	0.223	0.5	-	-			
	NI	C40			+9	0.368	0.8	-	++	+	+	

5

Patient Group	Patient ID	Serum ID	Candida species	Date of serum relative to first +ve culture	Applicant Triple Ag Abs	Applicant Triple Ag Abs/ cut off (0.46) ratio	Applicant Triple Ag Score (0.46 cut-off)	Serology-immunofluorescence IPI	IEP Pasteur	IEP FSK	Ag-emie
	PAS	C41	C.a	+5	0.865	1.9	+	+	++	+	
	PASa	C42		+32	1.279	2.8	++	+	++	++	+
	PIL	C43	C.p	-2	0.495	1.1	(+)	-			
	PIL	C44		+51	0.831	1.8	+	+			
	RAM	C45	C.t	+5	1.414	3.1	+++	++			
	RAM	C46		+23	1.114	2.4	++	+			
	NOI	C47	?	0	0.611	1.3	+	+++			
Hospital patients that are colonised but have negative Candida serology	ABE	D21	No info.	No info.	0.748	1.6	+	+			
	FRE	D22	No info.	No info.	0.454	1.0	-	-			
	BEN	D23	C.a	urine	0.331	0.7	-	-			
	BER	D24	C.t	mouth/fae	0.463	1.0	-	-			
	BOM	D25	C.a	broncal/fae	1.046	2.3	++	++			
	CAP	D25	No info.	No info.	0.658	1.4	+	-			
	CAR	D27	C.a	trachea	0.933	2.0	++	+			
	CHE	D28	C.t	urine	1.376	3.0	+++	-			
	FER	D29	C.a	urine/faeces	0.363	0.8	-	-			
	GIN	D30	C.g	urine/faeces	0.663	1.4	+	-			
	PER	D31	C.a	thorax drain	0.378	0.8	-	-			
Hospital patients that are non-colonised and have negative Candida serology	BEN	D32	C. spp	urine	0.469	1.0	-	-			
	BON	D33	-		0.44	1.0	-	-			
	CIA	D34	C.a	urine	0.92	2.0	++	++			
	DAVID										
	CAR	D35	-		0.651	1.4	+	-			
	PEL	D36	-		0.752	1.6	+	-			
	DI M	D37	-		0.489	1.1	(+)	-			
	FEU	D38	-		0.633	1.4	+	-			
	FOG	D39	C.a	thorax drain	1.095	2.4	++	+			
	MOR	D40			0.38	0.8	-	-			
GO	D41			0.677	1.5	+	-				

Patient Group	Patient ID	Serum ID	Candida species	Date of serum relative to first +ve culture	Applicant Triple Ag Abs	Applicant Triple Ag Abs/ Cut off (0.46) ratio	Applicant Triple Ag Score (0.46 cut-off)	Serology - immunofluorescence IPI	IEP Pasteur	IEP FSK	Ag-emie
Hospital patients that are colonised and have positive Candida serology	ALL	D42	C. spp	urine	1.177	2.6	++	+			
	BAR	D43	C.g, C.a, C.t	septic shock	1.375	3.0	+++	+++			
	BOE	D44	C.g	urine/mouth	1.096	2.4	++	++			
	BUI	D45	No info.	No info.	1.125	2.4	++	++			
	COL	D46	No info.	No info.	1.062	2.3	++	+++			
	DAG	D47	No info.	No info.	0.705	1.5	+	-			
	BE	D48	C.a & C.t	mouth/trachea	1.123	2.4	++	+++			
	GEN	D49	No info.	No info.	1.426	3.1	+++	++++			
	GEN	D50	No info.	No info.	1.489	3.2	+++	++++			
	LEC	D51	No info.	No info.	1.668	3.6	+++	++++			
LECr	D52	No info.	No info.	1.62	3.5	+++	++++				

Legend:

5

Candida Culture Sp.	Applicant Ab	IFI	IEP Pasteur	IEP FSK	Ag-emie
C.a = C. albicans	<10 = -	<20 = -	1 arc = +	1 arc = +	1/2 dil = + 1/4 dil =
C.g = C. glabrata	10-20 = +	20 = +	2 arc = ++	2 arc = ++	++
C.k = C. kefir	20-30 = ++	40 = ++	3 arc = +++	3 arc = +++	+++
C.p = C. parapsilosis	30-40 = +++	80 = +++	4 arc = ++++	4 arc = ++++	++++
C.t = C. tropicalis		160 = ++++			
		320 = ++++			

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Of the 12 patients who had sera taken before or on the day of the first positive blood culture, 8 gave a positive (or low positive) result. When compared with other serology tests used by the French group 12 out of 19 patients were
5 positive by the immunofluorescence (IFI) serology test. All but one of these was positive using the Applicant antigen test. One patient was also positive by the Applicant antigen test, but negative by IFI. All of the 5
10 patients that tested positive by IEP Pasteur, IEP FSK or Ag-emie serology tests were also positive by the Applicant antigen test.

It is possible that some of the patients that were negative by both the Applicant and the IFI test may have had a transient candidemia due to central line
15 contaminations.

Six of 11 patients that were know to be colonised, but had negative serology were positive by the Applicant antigen test. Two of the positive patients were also positive by IFI. Of the 10 non-colonised hospital
20 patients with negative serology six were positive by the Applicant antigen test, two of these positive patients were also positive by IFI. All nine patients that were colonised patients with positive serology were positive by the Applicant antigen test. These data compared to 8 out
25 of 9 patients that were positive by IFI. The only IFI negative sample was a low positive by the Applicant antigen test.

The statistical analysis of these data is presented in Figure 8 and Table 2.

TABLE 2

5

Category	Mean (Units)	95% Confidence Interval
Candidemia Patients	21.79 ^{a,b}	16.25 - 27.33
Colonised + negative Serology	14.55 ^{a,c}	9.67 - 19.42
Non-colonised + negative serology	14.2 ^{b,d}	10.68 - 17.72
Colonised + positive serology	27.27 ^{c,d}	23.12 - 31.43

a. p = 0.71

b. p = 0.58

c & d. p <0.01

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In the candidemia patients with positive blood culture to *Candida*, the mean of their *Candida* antibody levels detected by Applicant antigen test was 21.79 (16.25 - 27.33 95% CI). Using the Independent Samples T-test, the *p* value was 0.71 between the candidemia group and the colonised group with negative serology to *Candida*. The *p* value was 0.58 between the means of the candidemia group and the non-colonised group that was negative for *Candida* serology.

10 For the negative blood culture patients, the patients in the groups that were negative for *Candida* serology had generally lower *Candida* antibody levels detected by the Applicant antigen test. The mean antibody levels were 14.55 units in the colonised group (9.67 - 15 19.42, 95% CI) and 14.2 units in the non-colonised group (10.68-17.72, 95% CI). These levels were significantly lowered ($p < 0.01$) from the mean antibody levels in the group of patients that had positive *Candida* colonisation culture results and positive *Candida* serology, mean 27.27 20 (23.12-31.43, 95% CI). These are clearly seen in the error bars in Figure 8.

Overall, there was a good correlation with the Applicant antigen test and other tests used. There was also a good correlation with the titre of antibody 25 detected by the Applicant antigen test and the level of positiveness of the other tests ie., a patient that had a high positive result with the Applicant antigen test also had a similar result with the other tests used (eg, patients AMI, COM, DA SI, FON, PAS and RAM). Similarly 30 patients that were negative or low positive with the Applicant antigen test were also negative or weak positive by the other tests (eg, patients BRIG, FER, HEN, LON and MAN). It was noted that some of the blood culture negative patients were positive by the Applicant antigen 35 test, which demonstrated the great sensitivity of the Applicant antigen test.

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EXAMPLE 4 CLINICAL EVALUATION IN SPAIN

A similar clinical evaluation to that undertaken in Example 3 was conducted by Professors Guillermo Quindós, MD, PhD, María Dolores Moragues, PhD, and José Pontón, PhD, Department of Immunology, Microbiology and Parasitology at the Faculty of Medicine, University of País Vasco, Bilbao, Spain.

The retrospective study sera were obtained from 11 patients (Table 3 - Patients 1.1 to 1.32) with invasive candidiasis as defined by positive blood culture or by histology and positive tissue biopsy. The "blood culture negative" group consisted of sera from 12 patients (Table 4 - Patients 2.2 to 2.53) selected on the basis of the patients have a risk of invasive candidiasis but having negative blood cultures. Between 3 and 5 sera were tested per patient. For patients with microbiologically proven candidiasis the sera were taken before, at the time of and after the positive blood culture. For the blood culture negative group the sera were taken at various times during hospitalisation. As well as sera from hospital patients, sera from three healthy blood donors were also tested (Table 5). Also a group of fresh sera were prospectively tested from 5 patients, two with positive *Candida* blood cultures and 3 without (Table 5).

Table 3

Patients With Positive Blood Culture

Patient	Day extraction	Triple Ag	Platelia Ag	Platelia Ab	Spanish Anti-B	Spanish Anti-GT	Candida species & Outcome
1.01	-21	+	+	+	++++	+	C. albicans
1.01	-11	+	+	+	++++	++	
1.01	0	+	+	+	++++	++	
1.01	7	+	++	++	+++	+++	
1.01	13	+++	+	++	+++	+++	
1.11	-1	+++	++	+++	++++	++	C. glabrata
1.11	8	+++	++	+++	++++	+++	
1.11	21	+++	+	+	+++	+++	Exitus d45
1.11	29	+++	+	+++	++++	++++	
1.17	-11	+++	++	+++	++++	-	C. albicans
1.17	-4	+++	+	+++	+++	+	
1.17	0	+++	+	++	+++	+	Exitus d15
1.17	3	+	+	+	+++	+	
1.17	5	-	+	+	+++	+	
1.18	-6	++	(+)	+	+++	-	
1.18	2	++	+	+	++	-	
1.18	6	+	+++	+++	+++	-	
1.18	13	+	+++	-	+	+	
1.19	-3	-	-	-	+	-	C. albicans
1.19	0	-	+	-	+	-	
1.19	4	-	+	-	+	-	Exitus d7
1.19	7	+	+	-	++	-	
1.22	-11	+	+	+	+	+	C. albicans
1.22	-7	+	+	-	+	+	
1.22	0	+	+	+	++	+	
1.22	2	+	+	+	+	+	
1.22	7	+	+	+	+	+	
1.25	-2	-	+	-	++	-	C. albicans
1.25	1	+	-	+	+++	-	
1.25	4	+	-	-	++	-	
1.25	14	+	(+)	-	++	-	
1.25	26	+	+	-	++	-	
1.26	-9	-	+	(+)	++	-	C. parapsilosis
1.26	-5	+	+	+	+++	-	
1.26	0	+	+	+	+++	-	
1.26	9	+	++	+	+++	-	
1.26	16	+	+	+	++	-	

Patient	Day extraction	Triple Ag	Platelia Ag	Platelia Ab	Spanish Anti-B	Spanish Anti-GT	Candida species & Outcome
1.30	-25	+	(+)	?		+?	<i>C. albicans</i>
1.30	-4	+	+	+++		++++	
1.30	0	+	-	+++		++++	
1.30	3	+	-	?		+?	
1.30	56	+	+	?		-?	Exitus d70
1.31	-4	++	-	++		+?	<i>C. albicans</i>
1.31	-1	+	+	++		++++	
1.31	6	++	(+)	++		++++	
1.31	13	++	-	++		++++	Exitus d32
1.32	-19	-	-	+		-	<i>C. parapsilosis</i>
1.32	-17	-	+	+		-	
1.32	0	-	-	+		-	
1.32	7	-	+	+		-	
1.32	15	-	+	+		-	Discharge d33

Scoring:

Applicant Ab	Platelia Ag	Platelia Ab	Spanish B-Ab	Spanish GT Ab
0-10 = -	<0.5 = -	<1=-	<20=-	<20=-
10-20 = +	0.5-5=+	1-10=+	20-80=+	20-200=+
20-30 = ++	5-10=++	10-20=++	80-600=++	200-600=++
>30 = +++	>10=+++	>20=+++	600-5000=+++	600-1200=+++
			>5000=++++	>1200=++++

Table 4

Patients With Negative Blood Culture

5

	Patient Day extraction	Applicant Triple Ag	Platelia Ag	Platelia Ab	Spanish Anti-B	Spanish Anti-GT
2.2	1	++	+			-
2.2	5	++	+			-
2.2	6	++	+			(+)
2.4	1	+	(+)			ND
2.4	15	++	+			ND
2.4	17	+	+			ND
2.7	1	+	-	-	+	-
2.7	4	++	+	+	+++	-
2.7	8	+	+	+	+++	-
2.7	11	+	+	+	+++	-
2.7	15	+	+	+	+++	-
2.10	1	+	-		+++	-
2.10	3	++	-		+++	-
2.10	7	++	-		+++	-
2.10	11	++	-		++++	-
2.10	15	++	-		+++	-
2.14	1	++	(+)	+	+++	-
2.14	3	++	-	+	+++	-
2.14	6	++	-	+	+++	-
2.14	9	++	(+)	+	+++	-
2.14	12	++	+	+	+++	-
2.18	1	-	-			-
2.18	4	+	+			-
2.18	8	+	-			-
2.18	12	+	+			-
2.18	22	+	+			+
2.26	1	+	-			-
2.26	9	+	+			+
2.26	16	+	-			+
2.26	23	+	+			+
2.26	30	+	+			+
2.49	1	-	-			-
2.49	11	+	-			-
2.49	15	-	-			-
2.49	18	-	-			(+)
2.49	27	+	-			+

	Patient Day	extraction	Applicant Triple Ag	Platelia Ag	Platelia Ab	Spanish Anti-B	Spanish Anti-GT
2.50	1		+	(+)			+
2.50	9		+	+			(+)
2.50	15		+	-			+
2.50	22		+	-			+
2.50	26		+	-			+
2.51	1		-	-			-
2.51	11		-	-			(+)
2.51	18		-	+			-
2.51	22		-	+			-
2.51	29		+	-			-
2.52	1		+	-			(+)
2.52	8		+	-			-
2.52	11		+	-			(+)
2.52	15		+	-			(+)
2.52	18		+	-			(+)
2.53	1		+	-			-
2.53	8		+	-			-
2.53	11		+	+			(+)
2.53	18		+	-			-
2.53	22		+	-			-

Scoring:

Rockeby Ab	Platelia Ag	Platelia Ab	Spanish B-Ab	Spanish GT Ab
0-10 = -	<0.5 = -	<1=-	<20=-	<20=-
10-20 = +	0.5-5=+	1-10=+	20-80=+	20-200=+
20-30 = ++	5-10=++	10-20=++	80-600=++	200-600=++
>30 = +++	>10=+++	>20=+++	600-5000=+++	600-1200=+++
			>5000=++++	>1200=++++

Table 5

New Patients and Blood Donor Controls

	Patient Day extraction	Blood Culture	Applicant RESULT	Platelia Ag	Platelia Ab	Spanish Anti-B	Spanish Anti-GT
B. Donor 1	N/A	-	-	ND	ND	ND	ND
B. Donor 2	N/A	-	-	ND	ND	ND	ND
B. Donor 3	N/A	-	-	ND	ND	ND	ND
New Patients							
1	N/A	-	-	ND	ND	ND	ND
2	N/A	-	(+)	ND	ND	ND	ND
3	N/A	Asperg	+	ND	ND	ND	ND
4	?	C.g	+	ND	ND	ND	ND
4	?	C.g	+++	ND	ND	ND	ND
5	?	C.a	++	ND	ND	ND	ND
5	?	C.a	+++	ND	ND	ND	ND

5

Legend:

Culture Species Applicant Ab
 C.a=C. albicans 0-10 = -
 C.g=C. glabrata 10-20 = +
 20-30 = ++
 >30 = +++

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Table 6 summarises the original Spanish data split into the two groups of patients, one blood culture positive and the other blood culture negative. Of the blood culture positive group, the Applicant antigen test
5 identified 8/11 patients as positive before they became blood culture positive. Ultimately 10/11 patients were positive with the Applicant antigen test. Only one patient (1.32) remained negative. This patient was also
10 negative by the Spanish germ tube antibody test and was only transiently positive by the Platelia (BioRad) mannan antigen test. It may be possible that this patient had a transient candidemia.

Table 6

Summary of Spanish Data

	Applicant Triple Ag	Platelia Ag	Platelia Ab	Spanish Anti-B	Spanish Anti-GI
Culture Positive Patients					
Negative result	1	0	1	0	4
Positive before culture	8	11	9	8	6
Positive after culture	2		1	0	1
Total Patients tested	11	11	11	8	11
Culture Negative Patients					
Negative result	0	3	0	0	6
Positive result	12	9	2	3	5
Total Patients tested	12	12	2	3	11
Patient ID					
1.01	pos	pos	pos	pos	pos
1.11	pos	pos	pos	pos	pos
1.17	pos	pos	pos	pos	pos
1.18	pos	pos	pos	pos	(pos)
1.19	(pos)	pos	neg	pos	neg
1.22	pos	pos	pos	pos	pos
1.25	pos	pos	(pos)	pos	neg
1.26	pos	pos	pos	pos	neg
1.30	pos	pos	pos	ND	pos
1.31	pos	pos	pos	ND	pos
1.32	neg	pos	pos	ND	neg
Patient ID					
2.02	pos	pos			neg
2.04	pos	pos			
2.07	pos	pos	pos	pos	neg
2.10	pos	neg		pos	neg
2.14	pos	(pos)	pos	pos	neg
2.18	pos	pos			(pos)
2.26	pos	pos			pos
2.49	pos	neg			(pos)
2.50	pos	(pos)			pos
2.51	(pos)	pos			neg
2.52	pos	neg			neg
2.53	pos	(pos)			neg

5

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The Applicant antigen test identified all 12 of the blood culture negative patient group as being positive for *Candida* antibody. In comparison, the Platelia Mannan antigen test identified 9/12 patients as being positive compared to 5/11 by the Spanish Germ tube antibody test. The main problem with the blood culture negative patient group was that there was no other confirmation of diagnosis.

Overall, there was good correlation with the results of the Applicant antigen test and that of the other serology tests used by the Spanish group. Where a patient was strongly positive by the Applicant antigen test (ie.: patient's 1.11, 1.17, 1.18 and 1.31), they were also strongly positive by the other tests. Also, where sera were negative or low positive by the Applicant antigen test, they are also usually negative or weakly positive by the other tests. For example, patients 1.19, 1.22, 1.25, 1.26, 1.32, 2.18, 2.26, 2.49, 2.50, 2.51, 2.52 and 2.53.

Where fresh sera was analysed (Table 5), there was a perfect correlation with the Applicant antigen test and whether the sera was blood culture positive or negative.

The statistical analysis of these data is presented in Figure 9 and Table 7. From the error plot diagram in Figure 9, it is evident that the group of patients with positive blood culture have a higher *Candida* antibody levels detected by Syscan3 (mean 25.86, 95% CI: 16.28 - 35.44) as compared to the patients with negative blood culture as a group (mean 17.30, 95% CI: 13.42 - 21.19). Comparing the means using the Independent Samples T-Test, the difference between the two groups is statistically significant at $p = 0.087$.

Table 7

Mean and 95% Confidence Interval of Mean of Applicant
Antigen Test Scores

5

Category	Mean (Units)	95% Confidence Interval
Positive Blood Culture	25.86*	16.28 - 35.44
Negative Blood Culture	17.30*	13.42 - 21.19

* p = 0.087

EXAMPLE 5 CLINICAL EVALUATION IN AUSTRALIA

Sera collected from patients with invasive candidiasis was obtained from an Australian hospital (1997 to 1998), the patients had haematological malignancies (n=24). Control sera were collected from males 18 to 25 years of age (n=20) with no history of *Candida* infection. The patient sera were tested with the Applicant antigen test as described in Example 2. Each sera was tested in triplicate and the average reading used. The average absorbance reading for each serum was divided by that of the "cut-off" calibrator serum supplied with the Applicant antigen test. This value was then multiplied by 10 to give a value in arbitrary units.

The results of the Applicant antigen test using a value of 20 units (two times the cut-off calibrator serum value) or above as defining a positive sample is presented in Table 8.

Table 8

5 Results of the Applicant antigen test using 20 units as a cut-off

	Invasive Candidiasis	Healthy Controls	Total
<i>Test Positive</i>	20	0	20
<i>Test Negative</i>	4	20	24
Total	24	20	44

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With the Applicant antigen test using 20 units as cut-off, the specificity of the test was 100% and sensitivity was 83.3%. -Positive predictive value was 83.3% and negative predictive value was 100%. When the
5 value of the test considered positive was set at 10 units or 1 times the value of the cut-off sera absorbance, the specificity of the test decreased, but the sensitivity increased (Table 9). The specificity was 90%, sensitivity
87.5%. Positive predictive factor increased to 91.3%,
10 while negative predictive factor decreased to 85.7%.

Table 9

5 Results of the Applicant antigen test using 10 units as a cut-off

	Invasive Candidiasis	Healthy Controls	Total
Test Positive	21	2	23
Test Negative	3	18	21
Total	24	20	44

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The results of the Applicant antigen test using sera from patients with invasive candidiasis are presented in Table 10. Only one of the negative results came from a patient who was blood culture positive. Two of the four
5 negative samples were from patients with central line contaminations. Three of the four negative test results came from patients with *Candida parapsilosis* infections, the other being *C. albicans*.

Table 10

Sera from Patients with Invasive Candidiasis Test with
Applicant Antigen Test

5

Patient	Abs (420 nm)	Units	Result	Site of isolation	Candida spp.
A	1.69	61	Positive	blood culture	parapsilosis
B	0.15	5	Negative	blood culture	parapsilosis
C	1.16	42	Positive	peritoneal cavity	guilliermondii
D	1.85	67	Positive	blood culture	albicans
E	1.16	42	Positive	blood culture	albicans
F	1.49	54	Positive	peritoneal cavity	glabrata
G	0.56	20	Positive	peritoneal cavity	parapsilosis
I	1.24	45	Positive	cathater	albicans
K	0.98	35	Positive	oesophagus	albicans
L	3.46	124	Positive	peritoneal cavity	albicans
M	0.19	7	Negative	central line	albicans
N	1.44	52	Positive	wound	albicans
P	0.55	20	Positive	sputum	glabrata
Q	1.12	40	Positive	sputum	tropicalis
R	1.02	37	Positive	central line	albicans
S	0.88	32	Positive	blood culture	albicans
T	1.6	58	Positive	blood culture	glabrata
U	0.22	8	Negative	central line	parapsilosis
V	0.55	20	Positive	urine	parapsilosis
W	0.31	11	Negative	peritoneal cavity	parapsilosis
X	0.59	21	Positive	central line	albicans
Y	0.85	31	Positive	blood culture	albicans
Z	1.06	38	Positive	bronch	albicans
ZA	1.05	38	Positive	urine	tropicalis

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The Applicant antigen test data for the invasive candidiasis group and healthy controls are presented in the error bar diagram as seen in Figure 10 and Table 11. In Figure 10, the group with invasive candidiasis has a higher mean (31.45 units) as compared to the healthy blood donor group (7.52 units). This difference was statistically significant ($p < 0.01$). The 95% confidence interval range of the means was higher as well in the invasive candidiasis group (23.57 - 39.33 units), as compared to the healthy donor group (6.92 - 8.12 units).

Table 11

Mean and 95% Confidence Interval of the Mean of Patients
with Invasive Candidiasis and healthy blood donors

5

Group	Mean (Units)	95% CI of mean(Units)
Invasive Candidiasis	31.45	23.57 - 39.33
Healthy blood donors	7.52	6.92 - 8.12

- 53 -

In this study the Applicant antigen test was used to test sera from patients with invasive candidiasis, superficial candidiasis (oral or vaginal thrush) and healthy male controls. As a commensal organism, healthy individuals can have a measurable antibody titre to *Candida* antigens. In order to differentiate between normal and infection associated antibody levels a cut-off calibrator serum was supplied. The absorbance of the serum being tested was divided by the cut-off calibrator serum absorbance and multiplied by 10 to give an arbitrary unit value. Using a value of 20 units or above as an indicator of a positive test gave the greatest discrimination between the patient group with invasive candidiasis and the healthy controls (positive predictive value of 83%, negative predictive value 100%). If the value at which a sample was considered positive was lowered to 10 units (ie. the cut-off calibrator value), the positive predictive value increased slightly to 87.5% but the negative predictive value decreased to 90%.

Only one patient with a positive blood culture returned a negative test result with the Applicant antigen test. Two out of the four negative sera were from patients with a central line contamination. This could therefore reflect a transient infection in these patients, which may not provoke an antibody response. It is of interest that 3 of the 4 negative tests were due to *C. parapsilosis* infections. This organism is frequently associated with biofilms, which may shield it from the host immune response.

In conclusion, the Applicant antigen test is a rapid, reliable and easy test to perform. It showed good sensitivity and specificity in the diagnosis of invasive and severe superficial *Candida* infections.

THE CLAIMS DEFINING THE INVENTION ARE AS FOLLOWS:

1. A method of diagnosing *Candida* infection, comprising the steps of:
 - 5 a). obtaining a biological sample from a subject at risk of, or suspected to be suffering from, *Candida* infection;
 - b). preparing a composition comprising antigen consisting of a soluble cytoplasmic antigen preparation
10 which is mannose depleted and which consists essentially of *Candida* antigens of molecular weights 55 kDa, 30 kDa and 20 kDa;
 - c). contacting said antigen composition with said biological sample; and
 - 15 d). using a detection system to determine if antibodies from the biological sample are bound to said antigen composition by means of an antigen/antibody reaction.
- 20 2. A method according to claim 1, wherein the antigen composition further comprises one or more antigens selected from the group consisting of cell wall and enolase antigen.
- 25 3. A method according to claim 1 or claim 2, wherein step d) is a detection system selected from the group consisting of enzyme-linked immunoassay (ELISA or EIA), biligand binding (sandwich technique), fluorometric assay, chemiluminescent assay, radialimmunodiffusion and
30 radioimmunoassay (RIA).
4. A method according to claim 1 or claim 2, wherein step d) is by ELISA or chemiluminescent assay.
- 35 5. A method according to any of claims 1-4, further comprising the step of binding the antigen composition to a solid phase either by adsorptive binding, covalent binding, or via a ligand already bound to the solid phase.
- 40 6. A method according to any one of claims 1-5, further comprising the step of using secondary labelled antibodies

to detect the antibodies to *Candida* present in the biological samples.

5 7. A method according to claim 6, further comprising the step of labelling the secondary antibodies with a label selected from the group consisting of fluorescent dye, radioisotope and enzyme, or combinations thereof.

10 8. A method according to claim 7, wherein the secondary antibody is labelled via bound ligands.

15 9. A method according to claim 1, wherein detection in the detection system is selected from the group consisting of colour development, chemiluminescence, fluorescence and radioactivity, or combinations thereof.

20 10. A method according to any one of claims 1-9, further comprising the step of performing the detection of antibodies by a method selected from the group consisting of qualitative detection and quantitative detection or combination thereof.

25 11. A method according to claim 7, further comprising the step of directly labelling the secondary antibody.

12. A method according to claim 7, further comprising the step of indirectly labelling the secondary antibody.

30 13. A method according to any one of claims 1-12, wherein the antigen composition is either immobilised on an inert surface, embedded in a gel, or conjugated to a molecule.

35 14. A method according to claim 13, wherein the molecule imparts colour, fluorescence or radioactivity to the antigen.

40 15. A method according to any one of claims 1-14, wherein the biological sample is selected from the group consisting of bone marrow, plasma, spinal fluid, lymph fluid, skin, tears, saliva, milk, blood, serum, blood cells, tumours and organs.

16. A method according to claim 15, wherein the skin consists of external sections selected from the group consisting of respiratory, intestinal, and genitourinary tracts.

5

17. A method according to claim 15, wherein the biological sample is serum.

18. A kit when used for detecting the presence or absence of a *Candida* antibody in a biological sample, comprising:

- 10 a). a biological sample collection device;
b). composition comprising antigen consisting of a soluble cytoplasmic antigen preparation which is mannose depleted and which consists essentially of antigens for
15 detecting antibodies to *Candida* of molecular weights 55 kDa, 30 kDa and 20 kDa;
c). means for detecting reaction between the antibody in the sample and antigen composition.

19. A kit according to claim 18, further comprising buffering agents and ionic salts.

20. A composition comprising antigen consisting of a soluble cytoplasmic antigen preparation which is mannose
25 depleted and which consists essentially of antigens for detecting antibodies to *Candida* of molecular weights 55 kDa, 30 kDa and 20 kDa.

30 Dated this 17th day of August 2004

ROCKEY BIOMED LIMITED

By their Patent Attorneys

GRIFFITH HACK

Fellows Institute of Patent and

35 Trade Mark Attorneys of Australia

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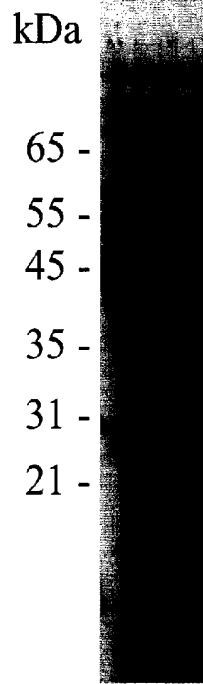


FIGURE 1

2/10

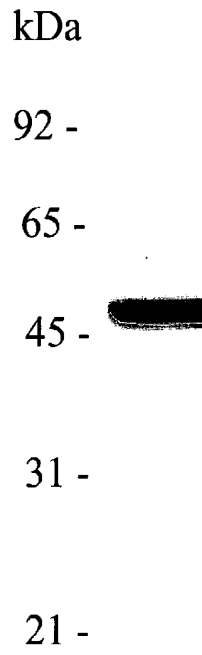


FIGURE 2

3/10

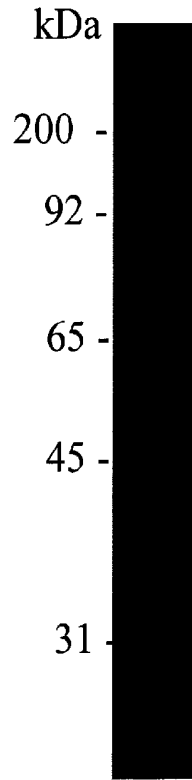


FIGURE 3

4/10

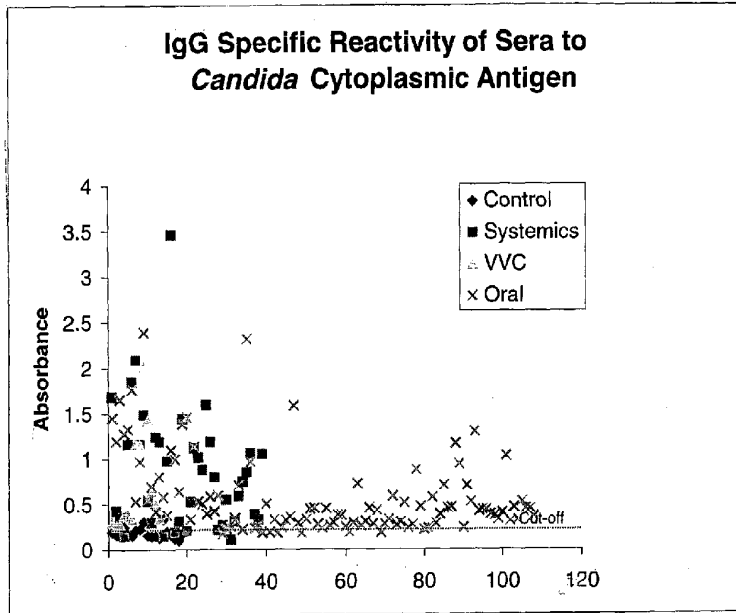


FIGURE 4

5/10

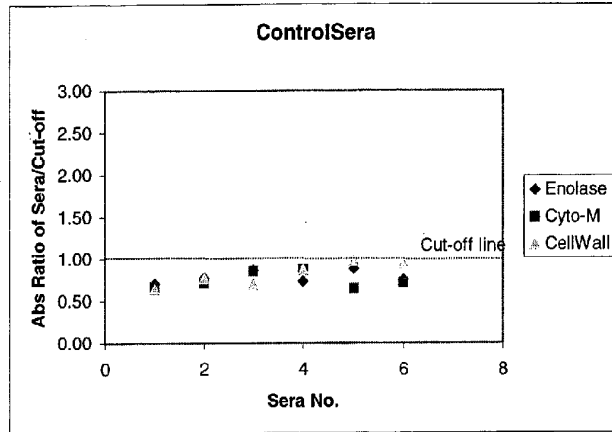


FIGURE 5

6/10

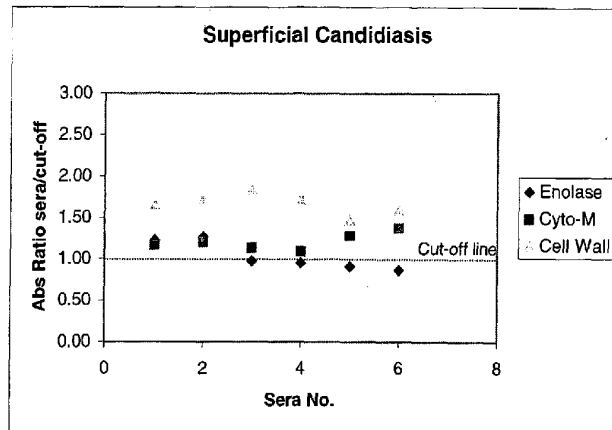


FIGURE 6

7/10

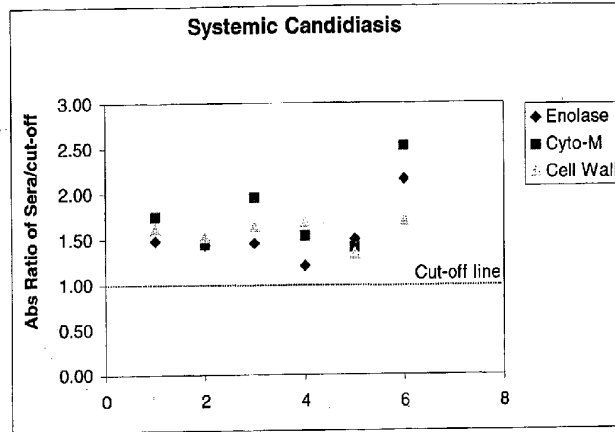


FIGURE 7

8/10

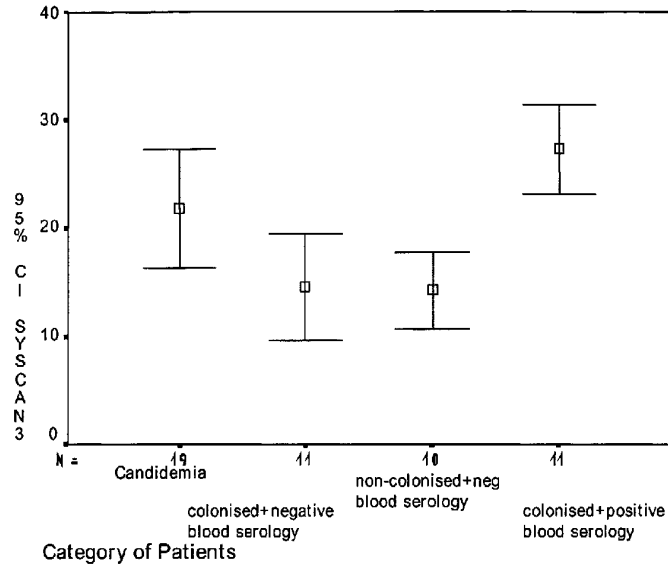


FIGURE 8

9/10

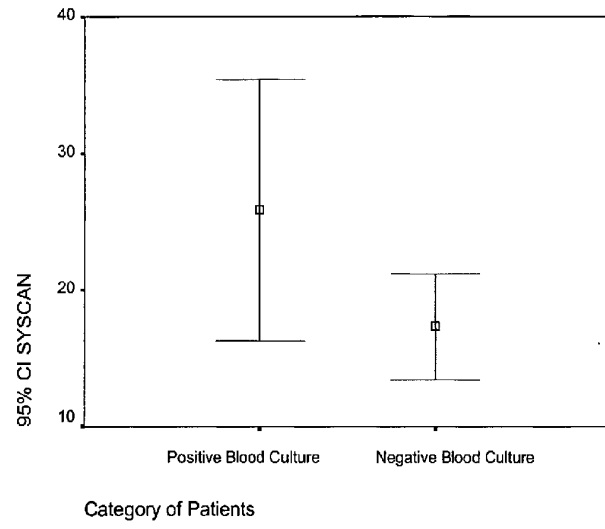


FIGURE 9

10/10

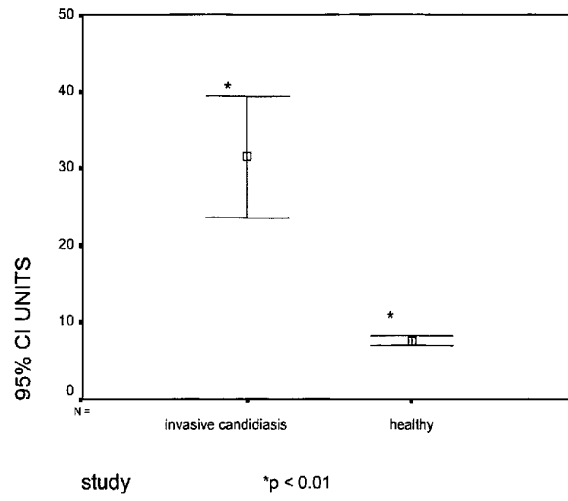


FIGURE 10