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(71) Applicant(s)
Centro De Inmunologia Molecular

(72) Inventor(s)

Jose Enrique Montero Casimiro; Josefa Lombardero Valladares; Rolando Perez Rodriguez; Patricia Sierra Blazquez; Blanca Rosa Tormo Bravo

(74) Agent/Attorney
BALDWIN SHELSTON WATERS, Level 21, 60 Margaret Street, SYDNEY NSW 2000

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31 No. 32009 entre 320 y 322, Reparto Juan de Dios Fraga,
La Lisa, Ciudad Habana 13500 (CU).

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(74) Mandatario: MORENO SAMPER, Olga Lidia; Lex, S.A.,
Avenida 1ra No. 1001, Esquina 10, Miramar, Playa, Ciudad
Habana 11300 (CU).

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(71) Solicitante (para todos los Estados designados salvo US):
CENTRO DE INMUNOLOGIA MOLECULAR (CIM)
[CU/CU]; Calle 216 y 15, Atabey, Playa, Ciudad Habana
12100 (CU).

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(72) Inventores; e

(75) Inventores/solicitantes (sólo US): MONTERO CASIMIRO,
José Enrique [CU/CU]; Calle 314 No. 2916 entre 29 y 31,
Reparto Juan de Dios Fraga, La Lisa, Ciudad Habana 13500
(CU). LOMBARDERO VALLADARES, Josefa [CU/CU];
Calle Agustina No. 70 entre San Miguel y Langueruela, La
Vibora, 10 de Octubre, Ciudad Habana 10700 (CU). PEREZ
RODRIGUEZ, Rolando [CU/CU]; Calle Juan Delgado No.
567, La Vibora, 10 de Octubre, Ciudad Habana 10500 (CU).
SIERRA BLAZQUEZ, Patricia [CU/CU]; Calle 208 No.
1913 entre 19 y 21, Atabey, Playa, Ciudad Habana 12100
(CU). TORMO BRAVO, Blanca Rosa [CU/CU]; Avenida



(54) Title: MONOCLONAL ANTIBODIES ANTI-CD6 AND THEIR USES

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(57) Abstract

Monoclonal antibodies that recognize the CD6 antigen, pharmaceutical compositions that recognizes and that are able to achieve a clinical and histological effectiveness in patients with different clinical types of Psoriasis.

(57) Resumen

Anticuerpos monoclonales que reconocen el antígeno CD6, composiciones farmacéuticas que contengan estos anticuerpos y sean capaces de lograr una efectividad clínica e histológica en pacientes con las diferentes formas clínicas de la Psoriasis.

ABSTRACT

Monoclonal antibodies that recognize the CD6 antigen, pharmaceutical compositions that recognizes and that are able to achieve a clinical and histological effectivity in patients with 5 different clinical types of Psoriasis.

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ANTI-CD6 MONOCLONAL ANTIBODIES AND THEIR USES.

Field of the Invention.

The present invention is related to the field of immunology and particularly with obtaining a pharmaceutical composition that 5 contains a monoclonal antibody that recognizes the CD6 leukocyte differentiation antigen.

Description of the Prior Art.

Monoclonal antibodies (mAbs) have permitted the characterization of molecules of physiological importance expressed on the cell 10 surface. Defined in the cells of the Immune System the "Leukocyte Differentiation Clusters" or antigens (CD) (Schollossman, S.F. et al. (1994) Immunol. Today 15 (3):98). The definition of the role of the CD's in the differentiation and maturation of the lymphoid cells during their ontogenetic development, in the mechanisms of 15 cellular recognition and adhesion and in the mechanisms of activation and proliferation during the immune response have conducted to the use of their respective mAbs in diagnosis and immunotherapy, with promising results (Dantal, J. et al. (1991) Curr. Opin. Immunol. 3:740).

20 The murine mAbs directed against molecules expressed on the cell membrane of human T lymphocytes have contributed to improve the diagnosis of clinical entities in which there is dysfunction of these cells. Moreover they have been used to explore new therapeutic approaches with the purpose of modulating their 25 functional activity as in the cases of Transplant Rejection, Graft Versus Host Disease (GvHD) and Autoimmune Diseases (Waldman, T.A. (1991) Science 252:1657; Waldman, T.A. (1992) Annu. Rev. Immunol. 10: 675)



The CD6 is a not well characterized molecule. It is known that it is a glycoprotein existing in two molecular forms maintained in dynamic equilibrium and differing only in the grade of phosphorylation. In the resting T lymphocytes it is a 5 phosphorylated molecule of 105 kDa, while in activated cells it is a hyperphosphorylated form of 130 kDa (Cárdenas, L. et al. (1990) J. Immunol. 145:1450; Swack, J.A. et al. (1991) J. Biol. Chem. 266: 7137), member of a family of membrane receptors and secretion proteins with a characteristic structure (Kodama, T. et al. (1990) 10 Nature 343: 531; Aruffo, A. et al. (1991) J. Exp. Med. 174: 949). It is expressed on the surface of mature human thymocytes, in T lymphocytes of peripheral blood, constituting the majority of the CD3+ cell population, in a subtype of B lymphocytes and in the 15 neurons of the brain cortex. In the T lymphocytes of peripheral blood it participates in the mechanisms of cellular activation (Reinhartz, E.L. et al. (1982) Cell 30: 735; Kamoun, M. et al (1981) J. Immunol. 127: 987; Mayer, B. et al. (1990) J. Neuroimmunol. 29: 193; Rasmussen, R.A. et al. (1994) J. Immunol. 152: 527). 20 The role of the CD6 molecule in the T cell ontogenesis as well as its possible role in the physiopathology of diseases of different etiology are unknown.

Recently a CD6 ligand was identified and characterized having an extensive cellular distribution in normal tissues such as thymus, 25 spleen, lymph nodes, and skin (Dhavalkumar, D.P. et al. (1995) J. Exp. Med. 181:1563). This molecule, denominated ALCAM (Activated Leukocyte-Cell Adhesion Molecule due to its expression in activated T and B lymphocytes as well as monocytes, is a 100 kDa molecular weight Type I membrane glycoprotein with five extracellular domains 30 similar to those of the immunoglobulins. It can present different



activation levels depending on divalent cations and can mediated heterophylic and homophylic interactions (Bowen, M.A. et al. (1995) J. Exp Med. 181: 2213).

Different anti CD6 mAbs have been used in clinical research for 5 the prevention of the rejection crisis in organ transplantation (Kirkman, R.L. et al. (1983) Transplantation 36: 620) and to deplete bone marrow transplants from lymphocytes for preventing Graft versus Host Disease (GvHD) (Soiffer, R.J. et al. (1992) J. Clin. Oncol. 10:1191). The ior t1 mAb is in a Phase II Clinical 10 Trial for the treatment of Cutaneous T-Cell Lymphomas (Garcia, C.A. et al. (1990) Biotecnologia Aplicada 7(2):176; Faxas, M.E. et al (1993) Biotecnologia Aplicada 10(1): 20).

The ior t1 monoclonal antibody of IgG2a isotype was classified as anti CD6 in the IV International Workshop of Leukocyte 15 Differentiation Antigens, Vienna (1989). This mAb defines an epitope different from the ones recognized by other anti CD6 mAbs. The epitope has a stable conformation and is insensitive to reduction agents, being possibly located in the primary structure of the CD6 molecule (Osorio, L.M. et al. (1994) Cell Immunol. 20 154:123).

This monoclonal antibody has a lower recognition than other CD6 mAbs in peripheral mononuclear cells of healthy donors. The recognition pattern of ior t1 mAb in human cell culture lines of T 25 lymphocytes origin is 47 % in Jurkat cells, 23 % in Molt-4 cells and no recognition of CCRF-CEM cells; of B lymphocytes origin is 9 % in Raji, of erythroblastoid origin is 12 % in K-562 and of myelomonocytic origin is 9 % in U-937. It also recognizes peripheral mononuclear cells of Chronic B Lymphocytic Leukemia (89+/- 4 %) (Garcia C.A. et al. 1992) Biotecnologia Aplicada 30 9(1):70) and lymphocytes of cutaneous lesions in patients with



Cutaneous T-Cell Lymphomas (Rodríguez, T. et al. (1985) Interferón y Biotec. 2(1): 41).

The ior t1 mAb does not inhibit the *in vitro* antigen specific cellular cytotoxicity (Faxas, M.E. et al. (1993) Biotecnología 5 Aplicada 10(1):47). It is capable of activating *in vitro* peripheral blood T lymphocytes of healthy donors. At suboptimum concentrations of OKT3 (anti CD3) the cross-linking with ior t1 induces higher responses than those achieved with other anti CD6 mAbs (Osorio, L.M. et al. (1994) Cell Immunol. 154:123).

10 Psoriasis is a disease whose physiopathology has not been defined (Hunziker, T. et al. (1993) Ther. Umsch. 50(2):110; Elder, J.T. et al. (1994) J. Invest. Dermatol. 102(6):24S). It is characterized by presenting an inflammatory infiltrate in the target organ with predominance of activated T lymphocytes of CD4+ and CD8+ phenotypes 15 (Chang, J.C.C. et al. (1994) Proc. Natl. Acad. Sci. USA 91:9282), as well as strong oligoclonality of the T-Cell receptors, the cells seem to present a marked tendency to migrate to the skin (homing) (Baker, J.N.W.N. et al (1992) Br. J. Dermatol. 127: 205; Menssen, A. et al. (1995) J. Immunol. 155:4078; Valdimarsson, H. et al. 20 (1995) Immunol. Today 16(3):145).

The spontaneous remission of Psoriasis can be predicted if the number of T cells in the skin decrease. Thus, it is suggested they play an important role in the perpetuation of the disease by releasing soluble mediators of the immune response capable of 25 inducing the proliferation of keratinocytes, responsible of clinical manifestations of the disease.

These considerations are supported by facts such as the cure of the disease following allogeneic bone marrow transplantation, the possible HLA association, the improvement, with steroids and 30 specially with immunosuppressors like cyclosporine and the clinical



improvement, although reversible, with anti T cells therapeutic monoclonal antibodies (Griffiths, C.E.M. et al. (1992) Springer Semin Immunopathol. 13:441; Nanney, L.B. et al. (1986) J. Invest. Dermatol. 86(3): 260; Picassia, D.D. et al. (1987) J. Am. Acad. Dermatol. 17(3): 408; Schopf, R.E. et al. (1986) Arch. Dermatol. Res. 279(2): 89; van de Kerkhof, P.C. et al. (1987) Dermatologica 174(5): 224).

The success of immunotherapy with monoclonal antibodies depends on the selection of the target molecule, which should participate in important cellular functions or in the selection of the mAb (Dantal, J. et al. (1991)urr. Opin. Immunol. 3:740). The mAbs evaluated in Psoriasis directed against CD3 (Weinshenker, B.G. et al. (1989) J. Am. Acad. Dermatol. 20:1132) and against CD4 (Poizot-Martin, I. et al. (1991) Lancet 337:1477; Prinz J. et al. (1991) Lancet 338:320; Nicolas, J.F. et al. (1991) Lancet 338: 321) have produced clinical improvement in the patients after multiple high dose endovenous applications, with partial remissions of short duration and early relapse of the symptoms' and signs of the disease in all cases.

Therapeutical application of murine mAbs in multiple doses is associate with some secundary effects and limited clinical utility in patients, these are related with xenogenicity of murine proteins that induce an Human Anti-Mouse Antibodies response (HAMA); humanized antibodies generated by protein engineering have reduced immunogenicity, improved pharmacokinetics and clinical advantage (Winter, G. et al. (1993) Trends-Pharmacol-Sci. 14(5): 139).

Up to now no previous study describing the expression of the CD6 molecule in the T lymphocytes of the inflammatory infiltrate of the skin in Psoriasis nor the possible association of this molecule with the development of the disease has been reported. Additionally



the therapeutic use of an anti CD6 mAb in this disease has not been previously evaluated.

The novelty of the present invention consists in providing topical and systemic pharmaceutical compositions containing an anti CD6 monoclonal antibody and the 5 obtention of an humanized monoclonal antibody anti CD6 for its application in patients with Psoriasis Vulgar, using different administration routes and in different clinical forms of the disease.

According to a first aspect the present invention provides a monoclonal antibody recognizing human CD6 when used for diagnosis and treatment of psoriasis, which is the 10 murine subclone ior t1A produced by the hybridoma of the same name, deposited under accession number ECACC 96112640 with the European Collection of Cell Cultures, as well as any humanized variant obtained thereof.

According to a second aspect the present invention provides use of the 15 monoclonal antibody according to the first aspect for the manufacture of a medicament for the treatment of Psoriasis.

According to a third aspect the present invention provides use of the monoclonal antibody according to the first aspect for the manufacture of a reagent for "in vitro" and "in vivo" diagnosis of Psoriasis.

According to a fourth aspect the present invention provides a pharmaceutical 20 composition for the treatment of Psoriasis which contains the monoclonal antibody according to the first aspect.

According to a fifth aspect the present invention provides a reagent for diagnosis of Psoriasis which contains the monoclonal antibody according to the first aspect.

According to a sixth aspect the present invention provides a method of treating 25 Psoriasis by administering to a subject in need thereof a therapeutically effective amount of a medicament according to the first aspect.

The cell line secreting the monoclonal anti CD6 antibody was deposited with the European Collection of Animal Cell Cultures (ECACC), of Centre for Applied Microbiology and Research, Porton Down, Salisbury, Wiltshire SP4 0JG, United 30 Kingdom, on 26 November, 1996, under accession number 96112640.



DETAILED DESCRIPTION OF THE INVENTION
OBTAINMENT OF THE MONOCLONAL ANTIBODY
Purification of the Murine Monoclonal Antibody

The anti CD6 monoclonal antibody (mAb) can be purified from ascitis fluid by

- 5 Protein A Sepharose, diluted with an equal volume of glycine buffer 1.5 M, NaCl 3 M pH 8.9, equilibrating the matrix with the same buffer. Applying equal volumes of ascitis and buffer elution should be performed at flow speed of 50 mL/h. Wash the column overnight until the zero base line, with the same application buffer.

Afterwards the column is washed with citric acid buffer 0.1 M pH 6 to eliminate

- 10 the IgG1 immunoglobulins, until the base line reaches zero, at a flow speed of 50 mL/h (between 2 and 5 column volumes, approximately 2 hours). Buffer citric acid 0.1 M, pH 5 is applied to elute the IgG2a immunoglobulins (ior t1).

Humanization of the Murine Monoclonal Antibody

Using genetic engineering method variants of the anti CD6 murine mAbs can be

- 15 constructed, such as chimaeric and humanized antibodies, from the variable regions of the light and heavy chains of the murine antibody (Takashi, N. et al. (1982) Cell 29:718; Hieter, P.A. et al. (1980) Cell 29:718).



Description of method of humanization of the Murine Monoclonal Antibody ior t1.

A subclone ior t1A was obtained from the parental murine ior t1 secreting hybridoma cells which recognize the same epitope on the 5 CD6 molecule.

ior t1A was modified in order to decrease its immunogenicity with a procedure (patent # 0699755 E.P. Bul.) which simultaneously reduces immunogenicity of the rodent monoclonal antibody while preserving its ligand binding properties in their entirety. Since the 10 antigenicity of an immunoglobulin is dependent on the presence of T-cell antigenic peptides onto their sequence, the immunogenicity of a xenogenic or allogenic antibody could be reduced by replacing the residues included onto the T-cell antigenic sequences which differ from those usually found in another mammalian species 15 antibodies. Of course, the replacement of residues do not include those involved in to the canonical structures or in the Vernier zone. This judicious replacement of residues have no effect on the structural determinants or on the interdomain contacts, thus, ligand binding properties should be unaffected as a consequence of 20 alterations which are limited to the variable region framework residues.

Analysis of homology of variable regions:

The present procedure makes use of the available sequence data for human antibody variable domains compiled by Kabat et al., 25 "Sequences of proteins of Immunological Interest" Fifth edition., Bethesda, Maryland; National Inst. of Health, 1994.

In the first step the variable domains of the murine ior t1 heavy or light chain are compared with those corresponding variable domains of the human sequences.



The comparison is made by an automated-computarized method (PC-DOS HIBIO PROSIS 06-00, Hitachi.). The most homologous human variable regions are then compared residue for residue to the corresponding murine regions. This will also define the human 5 subgroup to which each mouse sequence most closely resambles.

-Prediction of T-epitopes.

In the second step, the two homologous variable region sequences, mouse and human are analysed for prediction of T-antigenic sequences.

10 The algorithm AMPHI (Bersofsky et al., (1987) J. Immunol 138: 2213) predicts α Helical sequences. The algorithm SOHHA predicts the strip of helix hydrophobicity (Elliott et al., (1987) J.Immunol. 138: 2949). These algorithms predict T-cell presented fragments of antigenic proteins.

15 -Analysis for immunogenicity reduction.

Those residues in the mouse framework which differ from its human counterpart are replaced by the residues present in the human counterpart. This switching occurs only with those residues which are at the T-antigenic sequences.

20 Finally, replacement of those residues responsable for the canonical structures or those involved in the Vernier zone could have a significant effect on the tertiary structure. Hence, they can not be included in the replacement. Additional information about the influence of the proposed replacements on tertiary 25 structure or the binding site, could be obtained from a molecular model of the variable regions.

-The method for constructing and expressing the altered antibody.



The following procedures are used to prepare recombinant DNA sequences which incorporate the CDRs of the murine mAb, both light and heavy chains, into human, frameworks that can be used to transfect mammalian cells for the expression of recombinant antibody less immunogenic and with the antigen specificity of the animal monoclonal antibody:

5 a) Mutagenesis and assembly of variable region domains including CDRs and FRs regions. The PCR-mutagenesis method (Kamman et al., (1989) Nucleic Acids Res. 17: 5404) is preferably used to introduce 10 the changes at different positions.

b) Preparation of an expression vector including one variable region and the corresponding human constant region which upon transfection into cells results in the secretion of protein sufficient for affinity and specificity determinations.

15 c) Co-transfection of heavy and light chain expression vectors in appropriate cell lines.

After about 2 weeks, the cell supernatants are analyzed by ELISA for human IgG production. The samples are then analysed by any method for human IgG capable for binding to specific antigens.

20 Formulation for performing in vitro and in vivo diagnostic studies.

The anti CD6 mAbs can be used for in vitro and in vivo diagnostic purposes in the different clinical forms of Psoriasis, for monitoring patients after topical or systemic treatments, as well 25 as for predicting relapses.

These mAbs purified and dissolved in buffer solution (pH 7.0 +/- 0.5) containing Sodium Azide (0.01-0.2 %) and Bovine Serum Albumin (0.05-0.2 %) can be used to quantify the CD6+ T lymphocytes and/or the expression of this molecule on the surface of the lymphoid 30 cells in biological fluids (i.e. blood, cephaloracidium liquid,



synovial liquid) incubating 50 to 200 mL of the sample with 10 to 30 mL of the mAb at concentrations between 0.1 and 3 mg/mL, for 20 to 30 minutes at 4°C. Followed by washing with buffer solution and incubation with immunoglobulins of another animal species
5 conjugated with fluorescent substances (i.e. Fluorescein, Phycoerythrin). The anti CD6 mAbs can be conjugated directly with fluorescent substances by different methods (Coligan, J.E. et al: (Ed.) Current Protocols in Immunology, National Institutes of Health. Vol. I:5.3.2. Wiley Interscience) and be used with similar
10 purposes previously described, at concentrations between 5 and 30 mg/mL.

Immunohistochemical evaluation of the lesions of patients with Psoriasis.

The immunohistochemical study of cutaneous lesions or other
15 affected tissues (i.e. articulations) of patients with Psoriasis Vulgar can be performed on tissue cryostat sections (i.e. skin) fixed or not in cold acetone, incubating the anti CD6 mAb dissolved between 3 to 10 mg/mL in buffer solution (pH7.0 +/- 0.5) containing Sodium Azide (0.05-0.2 %) and Bovine Serum Albumin (0.05-0.2 %)
20 during 30 minutes. Followed by incubation of the tissue sections with biotinylated anti mouse immunoglobulins (i.e. from sheep, DAKO) and Avidin Biotin Peroxidase Complex (i.e. DAKO) during 30 minutes at room temperature. Finally reaction is developed using 3-amino-9-ethyl-carbazole as chromogen (Sigma) (Hsu, S.M. et al.
25 (1981) J. Histochem. Cytochem. 29:577). Biopsies must be examined by two specialists and the evaluation of the CD6 is adjusted to a scale of points <10 % (+/-), 10-25 % (+), 25-50 % (++)
(+ + +), 50-90 % (+ + + +), 90-100 % (+ + + + +).

Different antibodies to the T lymphocytes CD can be used, anti
30 CD3 (ior t3), anti CD4 (ior t4), anti CD8 (ior t8) and an anti CD45



(ior L3), also an anti Epidermal Growth Factor Receptor mAb (ior egf/r3) as a marker of keratinocytes activation (Mozzanica, N. et al. (1994) Acta Derm. Venereol. Suppl. Stockh. 186:171), which allows the evaluation of details of the characteristics of the 5 inflammatory infiltrate of the lesions during the course of the treatment of the disease.

Immunohistochemical monitoring of the lesions of the treated patients.

Patients treated with anti CD6 monoclonal antibodies can have 10 biopsies of the lesions performed previous to initiating the treatment, during the course of the treatment and after ending the treatment always in the area next to the initial biopsy to evaluate the therapeutical efficacy of the compositions of topical or systemic use. The scale for classifying treatment response can be 15 qualitative in per cent and is established by the index of CD6+ cells over the total CD45+ cells, evaluated in each biopsy.

$$\text{CD6+ cells index} = \frac{\text{Number of CD6+ cells} \times 100}{\text{Number of CD45+ cells}}$$

The index of CD3+ cells, CD4+ cells and CD8+ cells can also be 20 established over the total of CD45+ cells and the index of CD4+ and CD8+ over the total of CD6+ cells.

$$\text{CD3+ cells index} = \frac{\text{Number of CD3+ cell} \times 100}{\text{Number CD45+ cells}}$$

$$\text{CD4+ cells index} = \frac{\text{Number of CD4+ cells} \times 100}{\text{Number of CD45+ cells}}$$

$$\text{CD8+ cells index} = \frac{\text{Number of CD8+ cells} \times 100}{\text{Number of CD45+ cells}}$$

$$\text{CD4+ cells index} = \frac{\text{Number of CD4+ cells} \times 100}{\text{Number of CD6+ cells}}$$



$$\text{CD8+ cells index} = \frac{\text{Number of CD8+ cells} \times 100}{\text{Number of CD6+ cells}}$$

Immunoscintigraphic monitoring of treated patients.

Another form of evaluating the effect of the treatment with the
5 anti CD6 mAb can be the immunoscintigraphic study in the patient of
the expression and distribution of the CD6+ cells during the course
of treatment using between 1 and 5 mg of the same mAb conjugated
with radioactive isotopes such as 99m Technetium, using a
conjugating method like the one described by Mathers, S.J. et al.
10 (1990) J. Nucl. Med. 31(5):692.

Obtainment of therapeutic formulations for topical and systemic use.

The anti CD6 mAbs can be used with therapeutical purposes in
different clinical forms of Psoriasis, both with topical and
15 systemic formulations, in single or multiple doses, with one or
various treatment cycles according to the severity of the disease.

The topical therapeutic formulations with anti CD6 mAb can be
composed by semisolid systems in one or two phases, mainly with
hydrophilic formulations that allow the incorporation of the mAb
20 dissolved in sterile buffer solution ($\text{pH}7.0 \pm 0.5$) in doses
between 0.1 mg and 5 mg per each gram of the product. Formulations
can be elaborated as gels, jelly, ointment, lotions and creams with
a liquid matrix (i.e. water) formulated with gelatin,
carboximethylcellulose or similar substances and bases containing
25 glycerine, calcium ; additionally the compositions may contain
preservatives (i.e. p-hydroxibenzoate) to avoid contamination. The
pH must be physiological so as not to affect the characteristics of
the mAb. These therapeutic compositions should permit the
releasement and penetrability of the mAb in the skin.



Topical treatment should be applied between one and three times a day, over the lesions covered or not and it can be combined with the systemic use (mainly endovenous) of the same mAb with doses between 0.1 and 1 mg/Kg of patient body weight. It should be 5 diluted in physiological solution for endovenous use and administered slowly. The endovenous treatment can be applied independently of the topical administration.

Clinical follow up of treated patients.

10 The clinical evolution of the lesions may be used as the main criteria of the evaluation of therapeutic efficacy.

The main variables of response used for measuring the effects of the treatment may be the improvement of the clinical characteristics of the lesions (infiltration, scales, erythema) and the reduction of the area of the lesions.

15 The degree of severity of the signs of the disease (infiltration, scales and erythema) can be established between the values 0-1-2.

0- no sign.

1- scarce presence.

2- intense presence.

20 The extension of the treated scales should also be considered, measuring 2 of its diameters and calculating the area of the lesion by multiplying the product of the radios (in centimeters) by π (3.14). The dimension of the scale at time 0 represents 100 % and in the subsequent evaluations the per cent of the dimension of the 25 scale is established proportionally.

A PSORIASIS SEVERITY SCORE (PSS) similar to PASI (psoriasis area and severity index) (Fredriksson, T. et al. (1978) Dermatologica 157:238) is obtained with the formula:

infiltration(0-2) + scales(0-2) + erythema(0-2) x % of the
30 6 affected area



Response to treatment is stratified according to the changes in the PSS when completing the times of evaluations, establishing the following categories (Perkins, W. et al. (1993) Br. J. Dermatol. 129:584):

- 5 - Clear (> 90 % improvement in PSS)
- Responders (> 50 % improvement in PSS)
- Non-Responders (< 50 % improvement or deterioration in PSS)
- Worsening (> 50 % increase in PSS)

The evaluation times of the response can be assumed up to 12 weeks from the date of initiating the application of the treatment (pretreatment, weeks 1, 2, 3, 4, 6, 8, 12).

EXAMPLE 1: MURINE VARIABLE REGION OF THE t1A MONOCLONAL ANTIBODY DNA SEQUENCING.

Cytoplasmic RNA was extracted from about 10^6 T1 hybridoma cells 15 as described by Faloro et al (Faloro, J. et al. (1989) Methods in Enzymology 65:718).

The cDNA synthesis reaction consisted of 5 ug RNA, 50 mM Tris-HCl, pH 7.5, 75 mM KCl, 10 mM DTT, 3 mM MgCl₂, 25 pmol CG2AFOR primer

20 (5'GGAAGCTTAGACCGATGGGCCTGTTGTTTG 3') for heavy chain variable region or CK2FOR (5'GGAAGCTTGAAGATGGATACAGTTGGTGCAGC 3') for light chain variable region, 250 uM each of dATP, dTTP, dCTP, dGTP, 15 u ribonuclease inhibitor (RNA guard, Pharmacia) in a total volume of 50ul.

25 Samples were heated at 70°C, for 10 min and slowly cooled to 37°C over a period of 30 min. Then, 100units MMLV reverse transcriptase (BRL) were added and the incubation at 37°C continued for 1 hour.



The VH and VK cDNAs were amplified using the PCR as described by Orlandi et al (Orlandi, R. et al. Proc. Natl. Acad. Sci. USA 86:3833- 3837, (1989)). For PCR amplification of VH, DNA/primer mixtures consisted of 5 ul cDNA, 25 pmoles CG2A FOR
5 (5`GGAAAGCTTAGACCGATGGGGCTGTTGTTTG3`)
and VH1 BACK primers
(5`AGGT (G/C) (A/C) A (A/G) CTGCAG (G/C) AGTC (A/T) GG 3`).

For PCR amplification of VK, DNA/primers mixtures consisted of 5 ul cDNA, 25 pmoles of CK2 FOR
10 (5`GGAAAGCTTGAAGATGGATACAGTTGGTGCAGC 3`)
and
VK10BACK (5`TTGAATTCCAGTGATGTTTGATGACCCA 3`)
primers.

To these mixtures were added 2.5mM each of dATP,dCTP,dTTP, and dGTP, 5 ul constituents of 10X buffer thermolase and 1 unit of Thermolase(ABI) in a final volume of 50 ul. Samples were subjected
15 to 25 thermal cycles at 94⁰C, 30sec; 50⁰C, 30sec; 72⁰C, 1 min; and a last incubation for 5 min at 72⁰C. Amplified VH and VK DNA were purified on Prep. A Gene purification kit (BioRad).

The purified VH and VK DNA were cloned into M13 vector. Clones were sequenced by the dideoxy method using T7 DNA Pol (Pharmacia).
20 See Figure 1 and 2.

EXAMPLE 2: MODIFICATION OF THE VARIABLE DOMAIN SEQUENCES OF ior t1A MURINE MONOCLOINAL ANTIBODY TO HUMANIZE THE PREDICTED T-CELL ANTIGENIC SEQUENCES.

The variable region sequences of heavy and light chains of ior t1A were analyzed for T-cell antigenic sequences. It was made by using the computer algorithm AMPHI, which predict segments of the sequences 11 amino acids in length with an amphipatic helix
25



structure, that is have one side hydrophobic and one side hydrophilic which bind to MHC II molecules.

Onto the variable domain sequence of the heavy chain were predicted 3 segments which are: (It is used Kabat's numbering.).

5 1. FR 1 between amino acids 2-21.
2. CDR1 and FR2 between amino acids 29-43.
3. CDR3 and FR4 between amino acids 95-111.

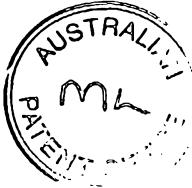
The Figure 1 shows the sequences corresponding to heavy chain.

10 This murine sequence is compared with the immunoglobulin sequences included in the GeneBank and EMBL database. The most homologous human variable region sequence is determined and also the human subgroup to which the murine sequence most closely resembles is defined. In this case the human sequence founded was an IgM belonging to subgroup III of Kabat.

15 Both variable region sequences, human and murine are then compared residue for residue and are selected those residues at FR regions which are not involved in the Vernier zone or with the canonical structures. Therefore they could be changed by those residues at the same position onto the human sequence. The 20 positions 13 and 19 are not modified due to the amino acid Lys is present in the same position in other human immunoglobulins belonging to the same subgroup.

For the heavy chain of murine ior t1A were proposed 4 replacements:

25 1. THR at position 40 by ALA.
2. GLU at position 42 by GLY.
3. THR at position 108 by LEU
4. LEU at position 109 by VAL.



The same procedure applied to the light chain (Figure 2) rendered a set of overlapping segments from aminoacid 2 to aminoacid 69.

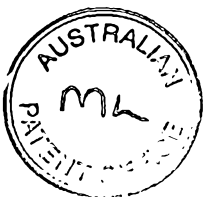
After the analysis we proposed 7 replacements in FRs 1 and 2 at positions: 3, 11, 12, 15, 17, 41 and 43.

- 5 1. LYS at position 3 by GLN
- 2. MET at position 11 by LEU
- 3. TYR at position 12 by SER
- 4. LEU at position 15 by VAL
- 5. GLU at position 17 by ASP
- 10 6. TRP at position 41 by GLY
- 7. SER at position 43 by ALA

EXAMPLE 3: CONSTRUCTION OF MUTANT HEAVY AND LIGHT CHAIN VARIABLE REGION OF ior t1A BY PCR MUTAGENESIS.

15 The changes in the amino acids of mutant heavy and light chain variable region were constructed using PCR mutagenesis (Kammann, M. et al. (1989) Proc.Natl.Acad.Sci. USA, 86: 4220).

Briefly: Two amplification by PCR: the reaction mixture was: 0.5 ul the VH supernatant of single strand DNA cloned in M13, 25 pmoles 20 mutagenic oligo 1 or 2, 25 pmoles mutagenic oligo 3 or 4 primers (See below the primers sequences). To these mixtures were added 2.5mM each of dATP, Dttp, dCTP, and dGTP, 5 μ l constituents of 10X Vent Polymerase buffer (NEB) and 1 unit of Vent DNA Polymerase (NEB) in a final volume of 50 ul. Samples were subjected to 12-15 thermal cycles at 94°C, 30sec; 50°C, 30sec; 75°C, 1 min; and a last 25 incubation for 5 min at 75°C. The products of both PCRs are joined in a second PCR using the outside primers only (3 and 4). Amplified VH DNA were purified on Prep. A Gene purification kit (BioRad).



For the changes in the heavy chain , FR1 in the positions 5, 7, 11, 12 and 13 the primers used, were:

Primer 1:

5' TGG GTT CGC CAG GCT CCG GGG AAG AGG CTG GAG 3'.

5 Primer 3:

5'GTA AAA CGA CGG CCA GT 3'.

These primers are combined in one PCR.

Primer 2:

5'CTC CAG CCT CTT CCC CGG AGC CTG GCG AAC CCA 3'.

10 Primer 4:

5'AGC GGA TAA CAA TTT CAC ACA GGA 3'.

These primers are combine in one PCR. Then, the products of both PCRs are combined in one PCR using 3 and 4 primers.

15 For the changes in the POSITION 108 and 109, the primers designed were:

Primer 1:

5'GGC CAA GGC ACC CTT GTC ACC GTC TCC 3'.

Primer 3:

5'GTA AAA CGA CGG CCA GT 3'.

20 These primers are combined in one PCR.

Primer2:

5'GGA GAC GGT GAC AAG GGT GCC TTG GCC 3'.

Primer 4:

5'AGC GGA TAA CAA TTT CAC ACA GGA 3'.

25 These primers are combined in one PCR. Then, the products of both PCRs are combined in one PCR using 3 and 4 primers.

In the light chain for the changes in the FR1 in the residues 3, 11, 12, 15 and 17 ,the primers were designed as:

PRIMER 1:



5' TGT GAC ATC CAG ATG ACC CAG TCT CCA TCC TCC CTG TCT GCA TCG
CAT CGG TGG GAG ACA GAG TCA CC 3'

PRIMER 3: 5'GTA AAA CGA CGG CCA GT 3'

These primers are combined in one PCR

5 PRIMER 2:

5' GGT GAC TCT GTC TCC CAC CGA TGC AGA CAG GGA GGA TGG AGA CTG
GGT CAT CTG GAT GTC ACA 3'

PRIMER 4: 5'ACT GGC CGT CGT TTT TAC 3'

These primers are combined in one PCR.

10 The products of both PCRs are combined in one PCR using primers
3 and 4.

For the changes in the residues 41 and 43 in the FR2 the primers
were:

PRIMER 1: 5' CAG AAA CCA GGG AAA GCT CCT AAG ACC CTG 3'

15 PRIMER 3: 5' GTA AAA CGA CGG CCA T 3'

These primers are combined in one PCR.

PRIMER 2: 5' CAG GGT CTT AGG AGC TTT CCC TGG TTT CTG 3'

PRIMER 4: 5' ACT GGC CGT CGT TTT TAC 3'

These primers are combined in one PCR.

20 The products of both PCRs are combined in one using primers 3
and 4.

**EXAMPLE 4: PHARMACEUTICAL FORMULATION FOR TOPIC USE IN CUTANEOUS
LESIONS OF PATIENTS WITH PSORIASIS VULGAR.**

The ior t1 mAb was purified and dissolved (50.00 mg) in sterile
25 buffer solution (10 mL pH7.0 +/- 0.5) containing Monobasic Sodium
Phosphate 4.50 mg , Dibasic Sodium Phosphate 18.00 mg , Sodium
Chloride 86.00 mg, Polysorbate 80 1.852 μ L, Water for injection.
The solution containing the mAb was added to a jelly base.



The therapeutic jelly was elaborated with a buffer solution having a composition similar to the one described for the mAb (pH 7.0 +/- 0.5).

**EXAMPLE 5: HISTOLOGICAL STUDY OF CUTANEOUS LESIONS OF PATIENTS
5 WITH PSORIASIS VULGAR.**

The immunohistochemical study of the cutaneous lesions of patients with Psoriasis Vulgar was performed on cryostat sections of skin tissue, using the mAb ior t1 in parallel with different mAbs directed against CD of T lymphocytes. mAbs ior t3 (anti CD3), 10 ior t4 (anti CD4), ior t8 (anti CD8), ior L3 (anti CD45) and DAKO CD6 (anti CD6), as control, as well as ior egf/r3 (anti Epidermal Growth Factor Receptor) were used. Followed by incubation of tissue sections with biotinylated anti mouse immunoglobulins (i.e. from sheep, DAKO) and Avidin Biotin Peroxidase Complex (i.e. DAKO). 15 Finally reaction is developed using 3-amino-9-ethyl-carbazole as chromogen (Sigma). Biopsies were examined by two specialists and the evaluation of the CD6 was adjusted to a scale of points <10 % (+/-), 10-25 % (+), 25-50 % (++) , 50-90 % (+++), 90-100 % (++++) .

20 Patients treated with anti CD6 monoclonal antibodies had biopsies of the lesions performed previous to initiating the treatment and at the end of the third week of treatment in the area next to the initial biopsy . The scale for classifying treatment response was qualitative in per cent and established by the index of CD6+ cells over the total CD45+ cells , evaluated in each biopsy. The per cent of the expression of the anti Epidermal Growth Factor receptor (EGF-R) in the different stratus of the skin was determined as well as the modifications of the histological characteristics typical of the disease.



- In the inflammatory infiltrate characteristic of Psoriasis Vulgar was found to exists an important expression (+++) of the CD6+ T lymphoid phenotype.
- The CD3+ lymphocytes represent approximately 100 % of the CD45+ cells.
- The CD6+ cells represents approximately between 60% and over 90 % of the CD3+ cells.
- The CD6+ cells (ior t1) represents approximately between 30 % and 50 % of the CD6+ cells (DAKO CD6).
- The expression of the EGF-R in keratinocytes was elevated showing a reticulated pattern.

The predominance of the expression of the CD6 molecule in the T lymphocytes of the inflammatory infiltrate results significant since it is a molecule characteristic of activated T lymphocytes.

This makes us believe that this might constitute a leukocitary adhesion molecule of the T lymphocytes, initially activated in the skin by the penetration of exogenous antigens or modified self antigens. Necessary this for the interaction with specific cellular determinants of the skin activated during the response to the said antigens.

EXAMPLE 6: CLINICAL RESPONSE OF PATIENTS WITH PSORIASIS VULGAR TO THE TOPICAL THERAPY WITH ior t1 mAb.

A clinical trial of patients with a diagnosis of Psoriasis Vulgar, in relapse with lesions characteristic of this disease was performed. The study was scheduled in two groups of fourteen patients per group difined according to the jelly received for topical treatment (ior t1 mAb or vehicle). The topical therapeutic formulation was conformed by a jelly base or vehicle (Sodium



Carboximethylcelulose v/v, Propilenglycol, Methylparabeno, Trietanolamine) in which the mAb was incorporated. Treatment was applied two times a day during 21 days without occlusion of the lesions treated.

5 Psoriatic plaque lesions whiten in all the patients treated with ior t1 mAb (Figure 4). This result corresponds with post treatment biopsy performed 21 days after initiating application of the mAb. A decrease of T lymphocyte infiltrate and of the expression of the EGF-R in keratinocytes as well as a regression in the histological 10 signs characteristic of the disease were observed.

EXAMPLE 7: CLINICAL RESPONSE OF PSORIASIS VULGARIS PATIENTS AFTER SCALING-DOWN TOPICAL TREATMENT WITH ior t1 MONOCLONAL ANTIBODY.

A Pilot Clinical Trial in 19 confirmed long-lasting psoriasis vulgaris patients with more than 10% and less than 25 % of their 15 skin affected was performed. Three different groups of 6, 7 and 6 patients received a therapeutic topical formulation containing 0.3, 1 and 3 mg of ior t1A mAb/gram of jelly respectively, in a vehicle jelly consisting of Sodium Carboximethylcelulose A/V, Propylenglicol, Methylparabene, Propylparabene and Triethanolamine.

20 Patients were topically treated 2 times a day during 21 days.

PASI (Psoriatic Area and Severity Index) was scored and analised and human anti mouse antibody (HAMA) response in the sera of patients was also studied. The best results related to clinical response (PASI) and disease free interval were obtained in the 25 group treated with the lower amount of ior t1 mAb (0.3 mg), as well as the HAMA (Human Anti-Mouse Antibody) titres and the amount of patients by group developing it were also higher in that group. Moreover, the presence of anti-idiotype antibodies in patient's sera studied by means of blocking ELISA (Enzyme Linked



Immunosorbent Assay) and FACS (Fluorescent Activating Cell Sorter) were more frequent and much higher in those patients treated with the lower dosis of 0.3mg per gram of jelly.

**EXAMPLE 8:CLINICAL RESPONSE OF ONE PATIENT WITH THE SEVERE FORM
5 OF GENERALIZED PSORIASIS TO THE ENDOVENEOUS TREATMENT WITH THE ior
t1 mAb.**

Female patient, 56 years old with a history of Psoriasis with psoriatic arthropathy diagnosed approximately 17 years ago. In the last 5 years the patient has suffered frequent and intense crisis, 10 causing frequent hospitalizations. Crisis started with erythematous generalized lesions, pain in articulations and muscles, feverish, generalized edemas and malaise. This general status persists and 21 days later new lesions appear in the axillary region, the neck and around the breasts, with ulcerations and 15 infected serous secretion accompanied by fever. Due to the torpid evolution of the disease and the intolerance to all previous treatments including steroid creams, treatment with methotrexate is indicated administering 3 cycles with a total doses of 15 mg. No clinical response was observed.

20 Single endovenous dose of ior t1 mAb at 0.6 mg/Kg of body weight, administered slowly, diluted in 200 mL of Saline Solution 0.9 %.

Simultaneously a therapeutical jelly containing ior t1 mAb at a concentration of 3 mg of mAb/ g of jelly was applied 2 times a day 25 in all the lesions during two days.

The patient starts to improve her general status and dermatological picture around the 6th day of treatment. At day 21 an evaluation was performed and 60 % of the body surface was without lesions and the rest of the skin showed improvement of the



clinical signs of the disease. The patient referred improvement of the symptoms in articulations, having a good general status, normal vital signs and routine laboratory tests.

5 After 30 days the patient maintains complete regression of the symptoms of the disease.

EXAMPLE 10: COMPARED REACTIVITY OF ANTI-CD6 mAbs.

Reactivity of anti-CD6 mAbs on HUT-78 cell line (cutaneous T cell lymphoma) using competitive immunofluorescence assay. Concentration from 0.625 to 20 μ g/mL of murine (ior-t1 and ior-t1A) and humanized (ior-t1A humanized) mAbs competed with 10 biotin-labelled ior-t1 mAb for binding to the CD6 molecule on cell surface. Cells were stained with streptavidin-FITC conjugate (Dako) before flow cytometry analysis using a fluorescence-activated cell sorter (FACScan) equipped with an argon laser (Beckton Dickinson).

As shown in Figure 7, three versions of the antibody showed similar recognition 15 for the specific antigen. Results are expressed as % of stained cells.

BRIEF DESCRIPTION OF THE FIGURES:

FIGURES 1 and 2

Analysis for the modification by way of humanization of the variable regions of heavy and light chains of monoclonal antibody ior t1A.

20 Figure 1: Sequence of the variable region of heavy chain of the murine ior t1A monoclonal antibody.

Figure 2: Sequence of the variable region of light chain of the murine ior t1A monoclonal antibody.

25 A: Sequence of the variable region of heavy chain or light chain of ior t1A murine mAb.

B: Sequence of the variable region of the most homologous human immunoglobulin.

C: Sequence of the modified variable region of ior t1A.

Shading: predicted T-cell antigenic sequences.

30 **Underlined amino acids residues:** amino acids involved in tertiary structure.

Bold font: complementarity determining regions.



Amino acids residues in boxes: replacements which are proposed.

The description is idem for both, heavy and light chains.

FIGURE 3

The results of the expression of the CD6 antigens in lymphocytes of the

5 inflammatory infiltrates characteristic of the cutaneous -----

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2
1



lesions of patients with Psoriasis Vulgar are shown. This evaluation was performed in cryostat sections biopsies of skin affected by plaques lesions localized in the upper and/or lower limbs and/or thorax-abdomen. The histological evaluation was 5 performed previous to the immunohistochemistry study.

FIGURE 4

The evaluation of the therapeutic efficacy of the anti CD6 mAbs used in the treatment of the Psoriasis was performed considering the following variables: infiltration, scales, erythema and the 10 size of the area of the lesion. The great of severity was established between the values zero-1-2. The extension of the treated plaques was established measuring to of its diameters. A Psoriasis Severity Score (PSS) similar to the PASI (Psoriasis Area and Severity Index) was obtained and the response to treatment was 15 stratified according to the changed in the PSS at the end of the designated evaluation time. The following categorizes was established: Clear, Responders, Non-Responders and Worsening.

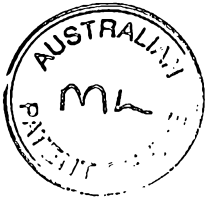
FIGURE 5

Clinical response of patients treated with a topical formulation 20 containing 0.3, 1 or 3 mg of ior t1A mAb/gram of jelly respectively was evaluated by PASI (Psoriatic Area and Severity Index) and the human anti mouse antibody (HAMA) response in the sera of these patients was also studied. The best results related to clinical 25 response (PASI) and disease free interval were obtained in the group treated with the lower amount of ior t1 mAb (0.3 mg), as well as the HAMA titres and the amount of patients by group developing it were also higher in that group. Moreover, the presence of anti-idiotype antibodies in patient's sera were more frequent and much higher in those patients treated with the lower dosis of 0.3mg per 30 gram of jelly.



FIGURE 6

Endovenous treatment was applied with the ior t1 mAb to a 56 years old patient with a history of Psoriasis with psoriatic arthropathy diagnosed approximately 17 years ago. Presenting now a 5 severe form of generalized psoriasis characterized by erithematosquamous generalized lesions, pain in articulations and muscles, feverish, generalized edemas and malaise. This general status did not respond to treatment including methotrexate. Treatment was performed with single endovenous dose of ior t1 mAb 10 at 0.6 mg/Kg of body weight, administered slowly, diluted in 200 mL of Saline Solution 0.9 %. Simultaneously a therapeutical jelly containing ior t1 mAb at a concentration of 3 mg of MAb/ g of jelly was applied 2 times a day in all the lesions during two days, for a total of 224 g. of therapeutic jelly. The clinical response and the 15 immunohistochemistry laboratory results were evaluated weekly. Photographs of the evolution of the cutaneous lesions are shown (the day before and 21 days after the treatment).



THE CLAIMS DEFINING THE INVENTION ARE AS FOLLOWS:-

1. Monoclonal antibody recognizing human CD6 when used for diagnosis and treatment of psoriasis, which is the murine subclone ior t1A produced by the hybridoma of the same name, deposited under accession number ECACC 96112640 with the European 5 Collection of Cell Cultures, as well as any humanized variant obtained thereof.

2. Monoclonal antibody recognizing human CD6 according to claim 1 wherein the said murine subclone ior t1A has a variable region of its heavy chain of sequence:

GLU VAL GLN LEU VAL GLU SER GLY GLY GLY LEU VAL
LYS PRO GLY GLY SER LEU LYS LEU SER CYS ALA ALA
10 SER GLY PHE LYS PHE SER ARG TYR ALA MET SER TRP
VAL ARG GLN THR PRO GLU LYS ARG LEU GLU TRP VAL
ALA THR ILE SER SER GLY GLY SER TYR ILE TYR TYR
PRO ASP SER VAL LYS GLY ARG PHE THR ILE SER ARG
ASP THR SER SER ASN THR ALA TYR MET GLN LEU SER
15 SER LEU ARG SER GLU ASP THR ALA MET TYR TYR CYS
ALA ARG ARG ASP TYR ASP LEU ASP TYR PHE ASP SER
TRP GLY GLN GLY THR THR LEU THR VAL SER SER
and the variable region of its light chain of sequence:

ASP ILE LYS MET THR GLN SER PRO SER SER MET TYR
20 ALA SER LEU GLY GLU ARG VAL THR ILE THR CYS LYS
ALA SER ARG ASP ILE ARG SER TYR LEU THR TRP TYR
GLN GLN LYS PRO TRP LYS SER PRO LYS THR LEU ILE
TYR TYR ALA THR SER LEU ALA ASP GLY VAL PRO SER
ARG PHE SER GLY SER GLY SER GLY GLN ASP TYR SER
25 LEU THR ILE SER SER LEU GLU SER ASP ASP THR ALA
THR TYR TYR CYS LEU GLN HIS GLY GLU SER PRO PHE
THR PHE GLY SER GLY THR LYS LEU GLU ILE LYS ARG
ALA

3. Monoclonal antibody recognizing human CD6 according to claim 1 which is a 30 humanized variant of the murine subclone ior t1A, and the variable region of its heavy chain has the sequence:



GLU VAL GLN LEU VAL GLU SER GLY GLY GLY LEU VAL
LYS PRO GLY GLY SER LEU LYS LEU SER CYS ALA ALA
SER GLY PHE LYS PHE SER ARG TYR ALA MET SER TRP
VAL ARG GLN ALA PRO GLY LYS ARG LEU GLU TRP VAL
5 ALA THR ILE SER SER GLY GLY SER TYR ILE TYR TYR
PRO ASP SER VAL LYS GLY ARG PHE THR ILE SER ARG
ASP ASN VAL LYS ASN THR LEU TYR LEU GLN MET SER
SER LEU ARG SER GLU ASP THR ALA MET TYR TYR CYS
ALA ARG ARG ASP TYR ASP LEU ASP TYR PHE ASP SER
10 TRP GLY GLN GLY THR LEU VAL THR VAL SER SER

and the variable region of its light chain has the sequence:

ASP ILE GLN MET THR GLN SER PRO SER SER LEU SER
ALA SER VAL GLY ASP ARG VAL THR ILE THR CYS LYS
ALA SER ARG ASP ILE ARG SER TYR LEU THR TRP TYR
15 GLN GLN LYS PRO GLY LYS ALA PRO LYS THR LEU ILE
TYR TYR ALA THR SER LEU ALA ASP GLY VAL PRO SER
ARG PHE SER GLY SER GLY SER GLY GLN ASP TYR SER
LEU THR ILE SER SER LEU GLU SER ASP ASP THR ALA
THR TYR TYR CYS LEU GLN HIS GLY GLU SER PRO PHE
20 THR PHE GLY SER GLY THR LYS LEU GLU ILE LYS ARG
ALA

4. Monoclonal antibody according to any one of claims 1 to 3 which recognizes in the human CD6 molecule an epitope of stable conformation and is insensitive to reducing agents.

25 5. Monoclonal antibody according to any one of claims 1 or 2 which is of IgG2a isotype.

6. Humanized monoclonal antibody according to claim 4 which is of IgG1 isotype.

7. Use of the monoclonal antibody according to any one of claims 1 to 6 for the manufacture of a medicament for the treatment of Psoriasis.

30 8. Use of the monoclonal antibody according to any one of claims 1 to 6 for the manufacture of a reagent for "in vitro" and "in vivo" diagnosis of Psoriasis.

9. Pharmaceutical composition for the treatment of Psoriasis which contains the monoclonal antibody of any one of claims 1 to 6.



10. Reagent for diagnosis of Psoriasis which contains the monoclonal antibody of any one of claims 1 to 6.
11. A method of treating Psoriasis by administering to a subject in need thereof a therapeutically effective amount of a monoclonal antibody according to any one of 5 claims 1 to 6.
12. A monoclonal antibody recognising human CD6 when used for diagnosis and treatment of Psoriasis, which is the murine subclone ior t1A produced by the hybridoma of the same name, deposited under accession number ECACC 96112640, substantially as herein described with reference to any one of the Examples but excluding comparative 10 Examples.
13. A humanised monoclonal antibody when used for diagnosis and treatment of Psoriasis, which is the murine subclone ior t1A produced by the hybridoma of the same name, deposited under accession number ECACC 96112640, substantially as herein described with reference to one or more of the Examples but excluding comparative 15 Examples.
14. Use of a monoclonal antibody which is the murine subclone ior t1A produced by the hybridoma of the same name, deposited under accession number ECACC 96112640 for the manufacture of a medicament for the treatment of Psoriasis, substantially as herein described with reference to one or more of the Examples but excluding 20 comparative Examples.
15. Use of a monoclonal antibody which is the murine subclone ior t1A produced by the hybridoma of the same name, deposited under accession number ECACC 96112640 for the manufacture of a reagent for "in vitro" and "in vivo" diagnosis of Psoriasis, substantially as herein described with reference to one or more of the Examples but 25 excluding comparative Examples.
16. A pharmaceutical composition for the treatment of Psoriasis which contains a monoclonal antibody which is the murine subclone ior t1A produced by hybridoma of the same name, deposited under accession number ECACC 96112640, substantially as herein described with reference to one or more of the Examples but excluding 30 comparative Examples.
17. Reagent for diagnosis of Psoriasis, substantially as herein described with reference to one or more of the Examples but excluding comparative Examples.



18. Reagent for diagnosis of Psoriasis which contains a monoclonal antibody recognising human CD6 which is the murine CD subclone ior t1A produced by hybridoma of the same name, deposited under accession number ECACC 96112640, substantially as herein described with reference to one or more of the Examples but 5 excluding comparative Examples.

19. A method of treatment of Psoriasis by administering to a subject in need thereof a therapeutically effective amount of a medicament including a monoclonal antibody recognising human CD6 which is the murine subclone ior t1A produced by the hybridoma of the same name, deposited under accession number ECACC 96112640 or a 10 humanised monoclonal antibody obtained therefrom, substantially as herein described with reference to one or more of the Examples but excluding comparative Examples.

DATED this 13th Day of June 2000

CENTRO DE INMUNOLOGIA MOLECULAR

15

Attorney: IVAN A. RAJKOVIC
Fellow Institute of Patent Attorneys of Australia
of BALDWIN SHELSTON WATERS

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A



FIGURE 1: Analysis for the modification by way of humanization of variable region of the heavy chain of ior t1A mAb.

														12
A	GLU	VAL	GLN	LLEU	VAL	GLU	SER	GLY	GLY	GLY	LEU	VAL		
B	GLU	VAL	GLN	LEU	VAL	GLU	SER	GLY	GLY	GLY	LEU	VAL		
C	GLU	VAL	GLN	LEU	VAL	GLU	SER	GLY	GLY	GLY	LEU	VAL		
														24
A	LYS	PRO	GLY	GLY	SER	LEU	LYS	LEU	SER	CYS	ALA	ALA		
B	GLN	PRO	GLY	GLY	SER	LEU	ARG	LEU	SER	CYS	ALA	ALA		
C	LYS	PRO	GLY	GLY	SER	LEU	LYS	LEU	SER	CYS	ALA	ALA		
														36
A	SER	<u>GLY</u>	<u>PHE</u>	<u>LYS</u>	<u>PHE</u>	<u>SER</u>	<u>ARG</u>	<u>TYR</u>	<u>ALA</u>	<u>MET</u>	<u>SER</u>	<u>TRP</u>		
B	SER	<u>GLY</u>	<u>PHE</u>	<u>THR</u>	<u>PHE</u>	<u>SER</u>	<u>ARG</u>	<u>TYR</u>	<u>ALA</u>	<u>MET</u>	<u>SER</u>	<u>TRP</u>		
C	SER	<u>GLY</u>	<u>PHE</u>	<u>LYS</u>	<u>PHE</u>	<u>SER</u>	<u>ARG</u>	<u>TYR</u>	<u>ALA</u>	<u>MET</u>	<u>SER</u>	<u>TRP</u>		
														48
A	VAL	ARG	GLN	THR	PRO	GLU	LYS	ARG	LEU	GLU	TRP	VAL		
B	VAL	ARG	GLN	ALA	PRO	GLY	LYS	GLY	LEU	GLU	TRP	VAL		
C	VAL	ARG	GLN	<u>ALA</u>	PRO	<u>GLY</u>	LYS	ARG	LEU	GLU	TRP	VAL		
														59
A	ALA	THR	ILE	SER	<u>SER</u>	<u>GLY</u>	<u>GLY</u>	<u>SER</u>	TYR	ILE	TYR	TYR		
B	SER													
C	ALA	THR	ILE	SER	<u>SER</u>	<u>GLY</u>	<u>GLY</u>	<u>SER</u>	TYR	ILE	TYR	TYR		
														71
A	PRO	ASP	SER	VAL	LYS	GLY	ARG	PHE	THR	ILE	SER	<u>ARG</u>		
B														
C	PRO	ASP	SER	VAL	LYS	GLY	ARG	PHE	THR	ILE	SER	<u>ARG</u>		
														82A
A	ASP	ASN	VAL	LYS	ASN	THR	LEU	TYR	LEU	GLN	MET	SER		
B	ASP	ASN	SER	LYS	ASN	THR	LEU	TYR	LEU	GLN	MET	ASN		
C	ASP	ASN	VAL	LYS	ASN	THR	LEU	TYR	LEU	GLN	MET	SER		
														92
A	SER	LEU	ARG	SER	GLU	ASP	THR	ALA	MET	TYR	TYR	CYS		
B	SER	LEU	ARG	ALA	GLU	ASP	THR	ALA	VAL	TYR	TYR	CYS		
C	SER	LEU	ARG	SER	GLU	ASP	THR	ALA	MET	TYR	TYR	CYS		
														102
A	ALA	<u>ARG</u>	ARG	ASP	TYR	ASP	LEU	ASP	TYR	PHE	ASP	SER		
B	ALA	<u>LYS</u>												
C	ALA	<u>ARG</u>	ARG	ASP	TYR	ASP	LEU	ASP	TYR	PHE	ASP	SER		
														113
A	TRP	GLY	GLN	GLY	THR	THR	LEU	THR	VAL	SER	SER			
B	TRP	GLY	GLN	GLY	THR	LEU	VAL	THR	VAL	SER	SER			
C	TRP	GLY	GLN	GLY	THR	LEU	VAL	THR	VAL	SER	SER			

FIGURE 2: Analysis for the modification by way of humanization of variable region of the light chain of 101 t1A mAb.

FIGURE 3

Table of the results of the expression of anti CD6 antigen in lymphocytes of the inflammatory infiltrate characteristic of the cutaneous lesions in patients with Psoriasis Vulgar.

Patient	CD6 Expression	Histological Diagnosis
1. IRM	+++	Psoriasis Vulgar
2. ZEAP	+++	Psoriasis Vulgar
3. DLH	+++	Psoriasis Vulgar
4. JMV	+++	Psoriasis Vulgar
5. MCGH	+++	Psoriasis Vulgar
6. DMMG	+++	Psoriasis Vulgar
7. ACP	+++	Psoriasis Vulgar
8. CCD	+++	Psoriasis Vulgar
9. MPC	+++	Psoriasis Vulgar
10. NRG	+++	Psoriasis Vulgar
11. MVMH	+++	Psoriasis Vulgar
12. IMVR	+++	Psoriasis Vulgar
13. MEGA	+++	Psoriasis Vulgar
14. CBT	+++	Psoriasis Vulgar

FIGURE 4

Evaluation of the clinical response to the topical treatment with the anti CD6 monoclonal antibody (ior t1) in patients with Psoriasis Vulgar after 84 days of follow up.

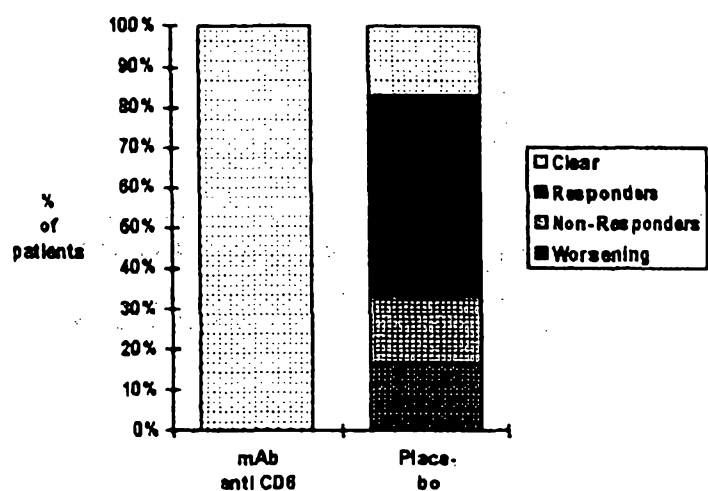


FIGURE 5

Clinical response of patients treated with a topical formulation containing 0.3, 1 or 3 mg of ior t1A mAb/gram of jelly evaluated by PASI

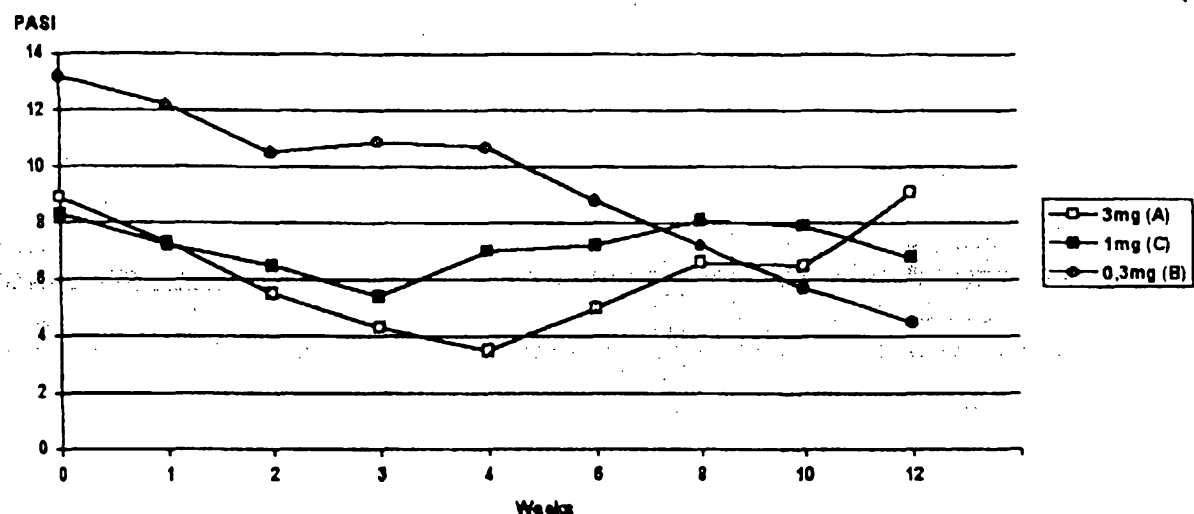
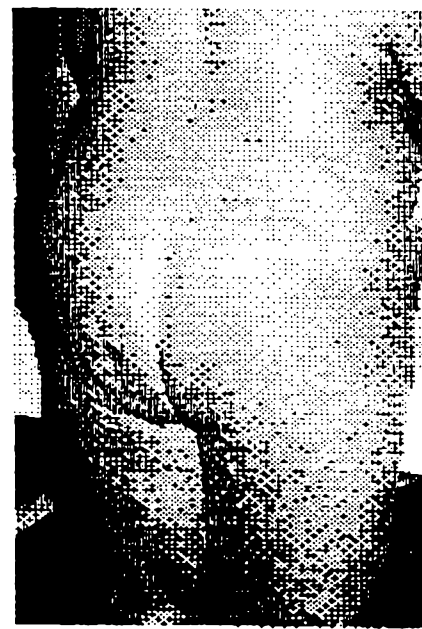
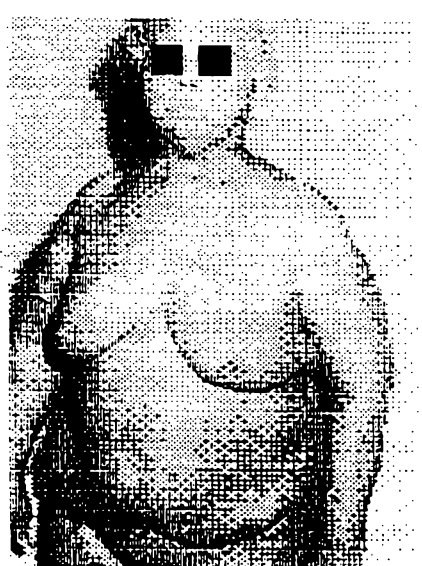
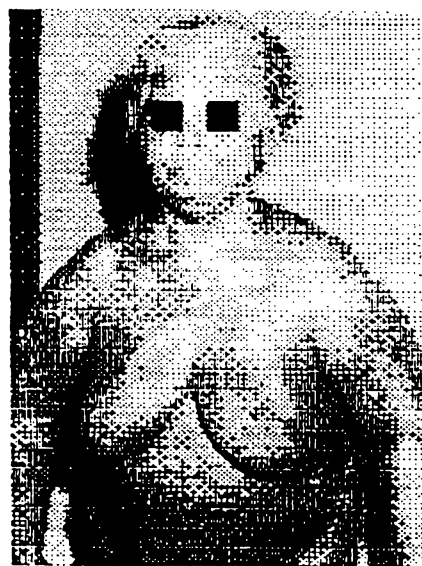


FIGURE 6

Clinical response of one patient with a severe form of generalized Psoriasis treated topically and endovenously with the anti CD6 monoclonal antibody



BASE
S
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FIGURE 7

