対しては肝移植以外の治療法がないことから, 早期 PBC の診断を確実に行い,必要に応じて 治療を行うことが重要である.

文 献

- 1) 厚生労働省「難治性の肝・胆道疾患に関する調査研究」班編:原発性胆汁性肝硬変(PBC)の診療ガイド. 2010, 文光堂, 東京
- Jones, E.A. and Bergasa, N.V.: The pathogenesis and treatment of pruritis and fatigue in patients with PBC. Eur. J. Gastroenterol. Hepatol. 11; 623-631, 1999
- 3) Gershwin, M.E., Rowley, M., Davis, P.A., et al.: Molecular biology of e-oxo-acid dehydrogenase complexes and anti-mitochondrial antibodies. Prog. Liver Dis. 10; 47-61, 1992
- 4) Nakamura, M., Kondo, H., Mori, T., et al.: Anti-gp 210 and anti-centromere antibodies are different risk factors for the progression of primary biliary cirrhosis. Hepatology 45; 118-127, 2007
- 5) Nakanuma, Y., Zen, Y., Harada, K., et al.: Application of a new histological staging and grading system for primary biliary cirrhosis to liver biopsy specimens: Interobserver agreement. Pathol. Int. 60; 167-174, 2010
- 6) 厚生労働省難治性疾患克服研究事業「難治性の 肝・胆道疾患に関する調査研究」班 編:原発性胆 汁性肝硬変(PBC)の診療ガイドライン(2011年). 2011年3月
- 7) Bizzaro, N., Covini, G., Rosina, F., et al.: Overcoming a "probable" diagnosis in antimitochondrial antibody negative primary biliary cirrhosis: Study of 100 sera and review of the literature. Clinic Rev. Allerg. Immunol. 2010 Dec 29 [Epub ahead of print]
- 8) Chazouillères, O., Wendum, D., Serfaty, L., et al.: Primary biliary cirrhosis-autoimmune hepatitis overlap syndrome: clinical features and response to therapy. Hepatology 28; 296-301, 1998
- 9) Takuma, K., Kamisawa, T. and Igarashi, Y.:

- Autoimmune pancreatitis and IgG4-related sclerosing cholangitis. Curr. Opin. Rheumatol. 23; 80-87, 2011
- 10) Hennes, E. M., Zeniya, M., Czaja, A. J., et al.: Simplified criteria for the diagnosis of autoimmune hepatitis. Hepatology 48; 169–176, 2008

Summary

Diagnosis of Primary Biliary Cirrhosis

Kazuhide Yamamoto* and Yasuhiro Miyake*

Primary biliary cirrhosis (PBC), a chronic cholestatic liver disease, which is characterized by the destruction and disappearance of intrahepatic small bile ducts. Middle aged women are those primarily affected by PBC. Autoimmune mechanisms may be involved in the process of bile duct destruction. Long standing cholestasis accompanied with varying degrees of interface hepatitis may eventually develop into liver cirrhosis. Anti-mitochondrial antibodies (AMA), which are directed to the E2 component of the 2oxo acid dehydrogenase complex, are positive in more than 90% of PBC patients. They are a highly sensitive and specific marker for the diagnosis of PBC. PBC is diagnosed based on three major findings including 1) elevation of biliary enzymes such as ALP and γ -GTP. 2) positive AMA, and 3) characteristic pathological findings including CNSDC, granulomas, bile duct loss or chronic cholestasis. Diagnosis of PBC may be difficult in cases assosiated with AMAnegative PBC or PBC-autoimmune hepatitis (AIH) overlap syndrome. Clinical stages are classified into asymptomatic PBC (aPBC) and symptomatic PBC (sPBC). The latter is further classified into s1-PBC with itching, unaccompanied by jaundice and s2-PBC with jaundice.

Key words: primary biliary cirrhosis, anti-mitochondrial antibody, overlap syndrome, chronic non-suppurative destructive cholangitis

*Department of Gastroenterology and Hepatology, Okayama University Graduate School of Medicine, Dentistry and Pharmaceutical Science, 2-5-1 Shikata-cho, Kita-ku, Okayama city, Okayama 700-8558, Japan PSCとその類縁疾患、オーバーラップス:基礎

自己抗体からみたPSCおよび類縁疾患 (IgG4関連硬化性胆管炎を含む)の病態

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索引用語:自己抗体,原発性硬化性胆管炎,P-ANCA,IgG4関連硬化性胆管炎

] はじめに

原発性硬化性胆管炎(primary sclerosing cholangitis: PSC) は、進行性の慢性胆汁うっ 滞性肝疾患であり, 胆管壁の線維性肥厚によ る肝内外胆管の多発性・びまん性狭窄を特徴 とする.一方、PSC患者の全国調査では6% に膵炎の合併を認めており、一部の症例は 血清中IgG4の上昇を示している. よって, PSCと診断された症例の中にIgG4関連硬化 性胆管炎(IgG4-related sclerosing cholangitis: IgG4-SC)例が含まれている可能性が考えら れている. PSCでは、コルチコステロイドや ウルソデオキシコール酸などの薬物治療の 有効性は一般的に乏しく, 高頻度に潰瘍性 大腸炎(Ulcerative colitis: UC) など炎症性腸 疾患を合併する. IgG4-SCはIgG4関連疾患 の概念に含まれ、しばしば自己免疫性膵炎 (Autoimmune pancreatitis: AIP)に合併する. また、PSCと異なりコルチコステロイドに対 する治療反応性は良好であり、涙腺・唾液腺 炎や後腹膜線維症などを合併することがあ る. PSCとIgG4-SCは異なる臨床像を呈する が、両疾患とも自己免疫性肝疾患の範疇に含 まれ、血清中に種々の自己抗体が出現する.

本稿では、PSCおよびIgG4-SCにおける血清中自己抗体と病態との関連について述べる.

2 PSC (表1, 表2)

1. 抗核抗体

わが国において 2007年~2008年にかけて 実施された PSC の全国調査における抗核抗 体(anti-nuclear antibody: ANA) 陽性率は 37%であり 10 , 欧米からの報告と同程度の結果で あった. PSC 患者における ANA の蛍光パター ンとしては、homogeneous と speckled がそ れぞれ約半数ずつに認められたと報告されて いる 20 . しかし、ANA は PSC に特異的な自己 抗体ではなく、病態との関連についても不明 である.

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表1 PSC患者の血清中に出現する自己抗体

38 1 100 % HANDING TANK A SECTION				
抗体	陽性率			
ANA	37%			
P-ANCA	2.4%(本邦 ELISA)			
	60~93%(欧米 IIF)			
抗 catalase 抗体	$16 \sim 60\%$			
抗α-enolase抗体	$11 \sim 31\%$			
抗lactoferrin抗体	22~50%			
抗neutrophil elastase 抗体	18%			
抗 cathepsin G 抗体	5~35%			
抗BPI抗体	5~46%			
胆管上皮抗体	63 %			
抗カルジオリピン抗体	$27 \sim 66\%$			
抗亜硫酸オキシダーゼ抗体	56%			
抗甲状腺ペルオキシダーゼ抗体	16%			
リウマチ因子	15%			

表2 自己抗体とPSC患者の臨床像

抗体	関連する臨床像
P-ANCA	UC合併、肝内外の胆管に病変
抗 catalase 抗体	ALP高値, 重症例
抗lactoferrin抗体	UC合併
抗cathepsin G抗体	肝硬変
抗BPI抗体	肝硬変
抗カルジオリピン抗体	長期の罹患期間、ALP高値、γ-グ
	ロブリン高値、組織学的な進行度
抗亜硫酸オキシダーゼ抗体	UDCA投与で抗体価の低下

2. P-ANCA

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わが国のPSC患者における抗好中球細胞質ミエロペルオキシダーゼ抗体(perinuclear neutrophil antibody: p-ANCA) 陽性率は2.4%であり¹⁾, 欧米のPSC患者における陽性率60~93%³⁾に比べて著しく低いことが報告されている。これは、両者におけるp-ANCAの測定法の違いによるものと考えられている。P-ANCAの測定法には、間接免疫蛍光法(Indirect immuneofluorescence assay: IIF)と酵素免疫法(Enzyme-Linked ImmunoSorbent Assay: ELISA)の2通りが使用されており、

欧米では主にIIFでの測定が行われている.しかし、わが国では、マイクロプレート上に固相化された遺伝子組み換え蛋白であるmyeloperoxidaseに血清を反応させてANCAの検出を行うELISA法が保険診療の適応となっている.IIFでは好中球をエタノール固定したものを基質として使用するため、未知の抗原に反応するANCAが検出される可能性がある.以前から、PSCや潰瘍性大腸炎患者でIIFにより検出されるp-ANCAは、顕微鏡的多発血管炎で検出される典型的なp-ANCAの蛍光パターンと異なること

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から"atypical" p-ANCAとよばれてきた. Terjungらは 4 、PSC患者の血清中に出現する"atypical" p-ANCAの対応抗原が分子量50kDの核膜タンパクであることをみいだし、"atypical" p-ANCA陽性PSC患者の95%でこの核膜タンパクに対する自己抗体が検出されたと報告している。さらに、最近では、この分子量50kDの核膜タンパクが細菌に発現しているFtsZタンパクと分子相同性を持つ β -tubulin isotype 5であることが報告されている 5 .

P-ANCAは、PSC以外に炎症性腸疾患でも高率に検出されるが、UCやクローン病を合併したPSC患者では非合併例に比べてp-ANCA陽性率の高いこと $(67\sim88\% \text{ vs.}29\sim40\%)$ が報告されている $^{6,7)}$. また、p-ANCA陽性例では陰性例に比べて肝内胆管と肝外胆管の両方に病変を認める頻度が高いという報告もある 7 .

上記以外のP-ANCAの対応抗原としては、catalase $(16\sim60\%)$, α -enolase $(11\sim31\%)$, lactoferrin $(22\sim50\%)$, neutrophil elastase (18%), cathepsin G $(5\sim35\%)$, BPI (bactericidal/permeability-increasing protein) $(5\sim46\%)$ が報告されている $^{2,8\sim12}$). 抗 catalase 抗体陽性例には重症例が多く,抗 cathepsin G抗体と抗BPI抗体は肝硬変例で陽性率が高い、抗lactoferrin抗体陽性例は,UC合併例に多い。

3. 胆管上皮抗体

PSC患者の63%で血清中に胆管上皮抗体が出現すると報告されている¹³⁾. 血清中に胆管上皮抗体を認めるPSC患者の58%で肝組織中の胆管上皮にToll様受容体(Toll-like receptor: TLR) 4とTLR9が発現しており、TLR4やTLR9を発現した胆管上皮細胞をそれぞれのリガンドであるLPSやCpG-DNAで

刺激するとIL-1 β やIFN- γ , TNF- α , TGF- β の産生が亢進する $^{(4)}$. これらの結果より、PSCの病態形成における自然免疫の関与が推測されている.

4. 抗カルジオリピン抗体

抗カルジオリピン抗体は、抗リン脂質抗体症候群の診断や治療効果、再発の予知などに有用な自己抗体であるが、PSC患者でも27~66%の症例で陽性となる $^{2,15)}$. 抗カルジオリピン抗体陽性例では、PSCの罹患期間が長く、血清中ALP値が高いとされている。また、抗カルジオリピン抗体価は、血清中ALP値や γ -グロブリン値、組織学的な進行度(Ludwig分類)と有意に相関することも報告されている。

5. 抗SO抗体

亜硫酸オキシダーゼ(sulfite oxidase: SO)は、真核生物のミトコンドリアに存在する金属酵素であり、抗ミトコンドリア M4抗体の対応抗原とされている。抗SO抗体は、PBCやAIH患者に比べてPSC患者で有意に陽性率が高く、治療前のPSC患者における陽性率は56%と報告されている¹⁶⁾、一方、抗SO抗体陽性PSC患者にUDCAを投与すると、抗体価が低下する。

6. その他

抗甲状腺ペルオキシダーゼ抗体が16%, リウマチ因子が15%の患者で陽性になると 報告されているが²⁾、病態との関連は不明で ある.

3 IgG4-SC

IgG4-SC症例の74~80%でANAが陽性である。また、血清中IgG4の上昇(135 mg/dL以上)が診断に有用である^{17,18)}。特に、血清中IgG4の上昇は、PSC患者の9%にしか認めらなかったとの報告もあり¹⁹⁾、IgG4-SCと

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PSCの鑑別に有用なマーカーと考えられる. 一方、IgG4SC症例のほとんどがAIPとの関連を有している。AIPにおけるANA陽性率は58%であり、23%でリウマチ因子の出現、86%で血清中IgG4の上昇が認められる200. AIP患者において、ANA陽性またはIgG4の上昇のどちらか一方、さらにANA陽性またはIgG4の上昇またはリウマチ因子の出現のどれか1つを満たす症例の割合について検討すると、それぞれ95%と97%が該当すると報告されている。IgG4SCの治療薬としてはコルチコステロイドが第一選択であり治療反応性は良好であるが、治療経過に伴って血清中IgG4値は低下する.

最近. ランダムペプチドライブラリーを用 いた研究により、AIP患者の90%以上で血清 中に膵腺房細胞のubiquitin-protein ligase E3 component n-recognin 2 (UBR2) Ø 1186-1192 アミノ酸配列と高い分子相同性を有するヘリ コバクター・ピロリのplasminogen-binding protein (PBP)の298-304アミノ酸配列に対す る抗体が存在すると報告された20.また, AIP患者の血清中には、heat shock protein 10 (92 %) ♥amylase alpha-2A (100%), pancreatic secretory trypsin inhibitor (42%), carbonic anhydrase II (59 %) に対する自 己抗体が検出されることも報告されてい る^{22~25)}. なお, 抗 carbonic anhydrase II 抗体 については、PSC 12例における検討により 全例で血清中に検出できなかったとの報告が ある²⁶⁾. よって, 抗carbonic anhydrase II抗 体はPSCとIgG4-SCの鑑別に有用な可能性が あるが、今後の検討が必要である.

4 おわりに

PSCではp-ANCAが高率に陽性となるが、 病態との関連については不明な点が多い. また、PSCに特異的なバイオマーカーはみつかっておらず、診断や病勢の評価に有用なマーカーの発見が求められている。一方、IgG4-SCでは、血清中IgG4が疾患特異的バイオマーカーとして有用であることが明らかにされているが、IgG4と病態との関連についての検討が必要である。これらの課題を克服するためにも、今後はゲノミクスのみならずプロテオミクス、さらには糖鎖解析までを含めた幅広い分野での共同研究が必要と思われる。

京 献

- 1) 滝川 一: PSCの全国調査. 難治性の肝・胆道 疾患に関する調査研究. 平成20年度総括・分担 研究報告書 74-77, 2009
- 2) Angulo P, Peter JB, Gershwin ME et al: Serum autoantibodies in patients with primary sclerosing cholangitis. J Hepatol 32: 182–187, 2000
- Weismüller TJ, Wedemeyer J, Kubicka S et al: The challenges in primary sclerosing cholangitisaetiopathogenesis, autoimmunity, management and malignancy. J Hepatol 48: S38–57, 2008
- 4) Terjung B, Spengler U, Sauerbruch T et al:

 "Atypical p-ANCA" in IBD and hepatobiliary disorders react with a 50-kilodalton nuclear envelope protein of neutrophils and myeloid cell lines. Gastroenterology 119: 310–322, 2000
- 5) Terjung B, Söhne J, Lechtenberg B et al: p-ANCAs in autoimmune liver disorders recognise human beta-tubulin isotype 5 and cross-react with microbial protein FtsZ. Gut 59: 808–816, 2010
- 6) Seibold F, Weber P, Klein R et al: Clinical significance of antibodies against neutrophils in patients with inflammatory bowel disease and primary sclerosing cholangitis. Gut 33: 657–662, 1992
- Bansi DS, Fleming KA, Chapman RW: Importance of antineutrophil cytoplasmic antibodies in primary sclerosing cholangitis and ulcerative colitis: prevalence, titre, and IgG subclass. Gut 38: 384–389, 1996
- 8) Lindgren S, Nilsson S, Nässberger L et al : Antineutrophil cytoplasmic antibodies in patients

肝胆膵 62巻4号·2011年4月

754

- with chronic liver diseases: prevalence, antigen specificity and predictive value for diagnosis of autoimmune liver disease. Swedish Internal Medicine Liver Club (SILK) J Gastroenterol Hepatol 15: 437–442, 2000
- Roozendaal C, de Jong MA, van den Berg AP et al: Clinical significance of anti-neutrophil cytoplasmic antibodies (ANCA) in autoimmune liver diseases. J Hepatol 32: 734–741, 2000
- 10) Roozendaal C, Van Milligen de Wit AW, Haagsma EB et al: Antineutrophil cytoplasmic antibodies in primary sclerosing cholangitis: defined specificities may be associated with distinct clinical features. Am J Med 105: 393–399, 1998
- 11) Peen E, Almer S, Bodemar G et al: Antilactoferrin antibodies and other types of ANCA in ulcerative colitis, primary sclerosing cholangitis, and Crohn's disease. Gut 34: 56–62, 1993
- 12) Orth T, Kellner R, Diekmann O et al: Identification and characterization of autoantibodies against catalase and alpha-enolase in patients with primary sclerosing cholangitis. Clin Exp Immunol 112:507-515, 1998
- 13) Xu B, Broome U, Ericzon BG et al: High frequency of autoantibodies in patients with primary sclerosing cholangitis that bind biliary epithelial cells and induce expression of CD44 and production of interleukin 6. Gut 51: 120–127, 2002
- 14) Karrar A, Broomé U, Södergren T et al: Biliary epithelial cell antibodies link adaptive and innate immune responses in primary sclerosing cholangitis. Gastroenterology 132: 1504–1514, 2007
- 15) Zachou K, Liaskos C, Rigopoulou E et al: Presence of high avidity anticardiolipin antibodies in patients with autoimmune cholestatic liver diseases. Clin Immunol 119: 203–212, 2006
- 16) Preuss B, Berg C, Altenberend F et al: Demonstration of autoantibodies to recombinant human sulphite oxidase in patients with chronic liver disorders and analysis of their clinical relevance. Clin Exp Immunol 150: 312–321, 2007

- 17) Nakazawa T, Ohara H, Sano H et al: Clinical differences between primary sclerosing cholangitis and sclerosing cholangitis with autoimmune pancreatitis. Pancreas 30: 20–25, 2005
- 18) Nishino T, Oyama H, Hashimoto E et al: Clinicopathological differentiation between sclerosing cholangitis with autoimmune pancreatitis and primary sclerosing cholangitis. J Gastroenterol 42: 550–559, 2007
- 19) Mendes FD, Jorgensen R, Keach J et al: Elevated serum IgG4 concentration in patients with primary sclerosing cholangitis. Am J Gastroenterol 101: 2070–2075, 2006
- 20) 川 茂幸,藤永康成,入澤篤志,他:自己免疫 性膵炎と膵癌の鑑別のポイント. 膵臓 23:555– 569,2008
- 21) Frulloni L, Lunardi C, Simone R et al: Identification of a novel antibody associated with autoimmune pancreatitis. N Engl J Med 361: 2135–2142, 2009
- 22) Takizawa S, Endo T, Wanjia X et al: HSP 10 is a new autoantigen in both autoimmune pancreatitis and fulminant type 1 diabetes. Biochem Biophys Res Commun 386: 192–196, 2009
- 23) Endo T, Takizawa S, Tanaka S et al: Amylase alpha-2A autoantibodies: novel marker of autoimmune pancreatitis and fulminant type 1 diabetes. Diabetes 58: 732-737, 2009
- 24) Asada M, Nishio A, Uchida K et al: Identification of a novel autoantibody against pancreatic secretory trypsin inhibitor in patients with autoimmune pancreatitis. Pancreas 33: 20–26, 2006
- 25) Okazaki K, Uchida K, Ohana M et al : Autoimmune-related pancreatitis is associated with autoantibodies and a Th1/Th2-type cellular immune response. Gastroenterology 118: 573– 581, 2000
- 26) Gordon SC, Quattrociocchi-Longe TM, Khan BA et al : Antibodies to carbonic anhydrase in patients with immune cholangiopathies. Gastroenterology 108: 1802–1809, 1995

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Clinical and Experimental Immunology ORIGINAL ARTICLE

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Prolonged recurrence-free survival following OK432-stimulated dendritic cell transfer into hepatocellular carcinoma during transarterial embolization

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Summary

Despite curative locoregional treatments for hepatocellular carcinoma (HCC), tumour recurrence rates remain high. The current study was designed to assess the safety and bioactivity of infusion of dendritic cells (DCs) stimulated with OK432, a streptococcus-derived anti-cancer immunotherapeutic agent, into tumour tissues following transcatheter hepatic arterial embolization (TAE) treatment in patients with HCC. DCs were derived from peripheral blood monocytes of patients with hepatitis C virus-related cirrhosis and HCC in the presence of interleukin (IL)-4 and granulocyte-macrophage colony-stimulating factor and stimulated with 0.1 KE/ml OK432 for 2 days. Thirteen patients were administered with 5×10^6 of DCs through arterial catheter during the procedures of TAE treatment on day 7. The immunomodulatory effects and clinical responses were evaluated in comparison with a group of 22 historical controls treated with TAE but without DC transfer. OK432 stimulation of immature DCs promoted their maturation towards cells with activated phenotypes, high expression of a homing receptor, fairly well-preserved phagocytic capacity, greatly enhanced cytokine production and effective tumoricidal activity. Administration of OK432-stimulated DCs to patients was found to be feasible and safe. Kaplan-Meier analysis revealed prolonged recurrence-free survival of patients treated in this manner compared with the historical controls (P = 0.046, log-rank test). The bioactivity of the transferred DCs was reflected in higher serum concentrations of the cytokines IL-9, IL-15 and tumour necrosis factor-α and the chemokines CCL4 and CCL11. Collectively, this study suggests that a DC-based, active immunotherapeutic strategy in combination with locoregional treatments exerts beneficial anti-tumour effects against liver cancer.

Keywords: dendritic cells, hepatocellular carcinoma, immunotherapy, recurrence-free survival, transcatheter hepatic arterial embolization

Introduction

Many locoregional therapeutic approaches including surgical resection, radiofrequency ablation (RFA) and transcatheter hepatic arterial embolization (TAE) have been taken in the search for curative treatments of hepatocellular carcinoma (HCC). Despite these efforts, tumour recurrence rates remain high [1,2], probably because active hepatitis and cirrhosis in the surrounding non-tumour liver tissues causes de novo development of HCC [3,4]. One strategy to reduce tumour recurrence is to enhance anti-tumour immune responses that may induce sufficient inhibitory effects to prevent tumour cell growth and survival [5,6]. Dendritic cells (DCs) are the most potent type of antigen-presenting cells in the human body, and are involved in the regulation of both innate and adaptive immune responses [7]. DC-based immunotherapies are believed to contribute to the eradication of residual and recurrent tumour cells.

To enhance tumour antigen presentation to T lymphocytes, DCs have been transferred with major histocompatibility complex (MHC) class I and class II genes [8] and co-stimulatory molecules, e.g. CD40, CD80 and CD86 [9,10], and loaded with tumour-associated antigens, including tumour lysates, peptides and RNA transfection [11]. To induce natural killer (NK) and natural killer T (NK T) cell activation, DCs have been stimulated and modified to

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Table 1. Patient characteristics.

			Largest						
Patient no.	Gender	Age (years)	HLA	TNM stages	No. of tumours	tumour (mm)	Child-Pugh	KPS	Post-TAE Rx
1	М	60	A11 A33	III	5	35	В	100	RFA
2	M	57	A11 A24	III	1	21	В	100	RFA
3	M	57	A11 A31	III	2	39	В	100	RFA
4	M	77	A2 A24	III	2	35	Α	100	RFA
5	F	83	A11 A24	III	3	29	В	100	RFA
6	F	74	A2 A24	II	1	35	Α	100	RFA
7	F	72	A24 A33	III	3	41	В	100	RFA
8	F	65	A2 A11	II	4	12	В	100	RFA
9	M	71	A2 A11	II	4	16	Α	100	RFA
10	M	79	A11 A24	III	2	40	Α	100	RFA
11	M	71	A2 A24	II	1	28	A	100	RFA
12	M	56	A2 A26	III	2	25	В	100	RFA
13	M	64	A2 A33	III	2	37	В	100	RFA

M, male; F, female; TNM, tumour—node—metastasis; Child—Pugh, Child—Pugh classification; KPS, Karnofsky performance scores; TAE, transcatheter arterial embolization; Rx, treatment; HCC, hepatocellular carcinoma; HLA, human leucocyte antigen; RFA, percutaneous radiofrequency ablation.

produce larger amounts of cytokines, e.g. interleukin (IL)-12, IL-18 and type I interferons (IFNs)[10,12]. Furthermore, DC migration into secondary lymphoid organs could be induced by expression of chemokine genes, e.g. C-C chemokine receptor-7 (CCR7) [13], and by maturation using inflammatory cytokines [14], matrix metalloprotein-ases and Toll-like receptor (TLR) ligands [15].

DCs stimulated with OK432, a penicillin-inactivated and lyophilized preparation of Streptococcus pyrogenes, were suggested recently to produce large amounts of T helper type 1 (Th1) cytokines, including IL-12 and IFN-γ and enhance cytotoxic T lymphocyte activity compared to a standard mixture of cytokines [tumour necrosis factor- α (TNF- α), IL-1β, IL-6 and prostaglandin E₂ (PGE₂)] [16]. Furthermore, because OK432 modulates DC maturation through TLR-4 and the β_2 integrin system [16,17] and TLR-4-stimulated DCs can abrogate the activity of regulatory T cells [18], OK432-stimulated DCs may contribute to the induction of anti-tumour immune responses partly by reducing the activity of suppressor cells. Recently, in addition to the orchestration of immune responses, OK432-activated DCs have themselves been shown to mediate strong, specific cytotoxicity towards tumour cells via CD40/CD40 ligand interactions [19].

We have reported recently that combination therapy using TAE together with immature DC infusion is safe for patients with cirrhosis and HCC [20]. DCs were infused precisely into tumour tissues and contributed to the recruitment and activation of immune cells *in situ*. However, this approach by itself yielded limited anti-tumour effects due probably to insufficient stimulation of immature DCs (the preparation of which seems closely related to therapeutic outcome [21,22]). The current study was designed to assess the safety and bioactivity of OK432-stimulated DC infusion into tumour tissues following TAE treatment in patients with cirrhosis and HCC. In addition to documenting the safety of

this approach, we found that patients treated with OK432-stimulated DCs displayed unique cytokine and chemokine profiles and, most importantly, experienced prolonged recurrence-free survival.

Patients and methods

Patients

Inclusion criteria were a radiological diagnosis of primary HCC by computed tomography (CT) angiography, hepatitis C virus (HCV)-related HCC, a Karnofsky score of \geq 70%, an age of \geq 20 years, informed consent and the following normal baseline haematological parameters (within 1 week before DC administration): haemoglobin \geq 8·5 g/dl; white cell count \geq 2000/µl; platelet count \geq 50 000/µl; creatinine < 1·5 mg/dl and liver damage A or B [23].

Exclusion criteria included severe cardiac, renal, pulmonary, haematological or other systemic disease associated with a discontinuation risk; human immunodeficiency virus (HIV) infection; prior history of other malignancies; history of surgery, chemotherapy or radiation therapy within 4 weeks; immunological disorders including splenectomy and radiation to the spleen; corticosteroid or anti-histamine therapy; current lactation; pregnancy; history of organ transplantation; or difficulty in follow-up.

Thirteen patients (four women and nine men) presenting at Kanazawa University Hospital between March 2004 and June 2006 were enrolled into the study, with an age range from 56 to 83 years (Table 1). Patients with verified radiological diagnoses of HCC stage II or more were eligible and enrolled in this study. In addition, a group of 22 historical controls (nine women and 13 men) treated with TAE without DC administration between July 2000 and September 2007 was included in this study. All patients received RFA therapy to increase the locoregional effects 1 week later [24].

They underwent ultrasound, computed tomography (CT) scan or magnetic resonance imaging (MRI) of the abdomen about 1 month after treatment and at a minimum of once every 3 months thereafter, and tumour recurrences were followed for up to 360 days. The Institutional Review Board reviewed and approved the study protocol. This study complied with ethical standards outlined in the Declaration of Helsinki. Adverse events were monitored for 1 month after the DC infusion in terms of fever, vomiting, abdominal pain, encephalopathy, myalgia, ascites, gastrointestinal disorder, bleeding, hepatic abscess and autoimmune diseases.

Preparation and injection of autologous DCs

DCs were generated from blood monocyte precursors, as reported previously [25]. Briefly, peripheral blood mononuclear cells (PBMCs) were isolated by centrifugation in LymphoprepTM Tubes (Nycomed, Roskilde, Denmark). For generating DCs, PBMCs were plated in six-well tissue culture dishes (Costar, Cambridge, MA, USA) at 1.4×10^7 cells in 2 ml per well and allowed to adhere to plastic for 2 h. Adherent cells were cultured in serum-free media (GMP CellGro® DC Medium; CellGro, Manassas, VA, USA) with 50 ng/ml recombinant human IL-4 (GMP grade; CellGro®) and 100 ng/ml recombinant human granulocyte-macrophage colony-stimulating factor (GM-CSF) (GMP grade; Cell-Gro®) for 5 days to generate immature DC, and matured for a further 2 days in 0·1 KE/ml OK432 (Chugai Pharmaceuticals, Tokyo, Japan) to generate OK-DC. On day 7, the cells were harvested for injection, 5×10^6 cells were suspended in 5 ml normal saline containing 1% autologous plasma, mixed with absorbable gelatin sponge (Gelfoam; Pharmacia & Upjohn, Peapack, NJ, USA) and infused through an arterial catheter following Lipiodol (iodized oil) (Lipiodol Ultrafluide, Laboratoire Guerbet, Aulnay-Sous-Bois, France) injection during selective TAE therapy. Release criteria for DCs were viability > 80%, purity > 30%, negative Gram stain and endotoxin polymerase chain reaction (PCR) and negative in process cultures from samples sent 48 h before release. All products met all release criteria, and the DCs had a typical phenotype of CD14⁻ and human leucocyte antigen (HLA)-DR+.

Flow cytometry analysis

The DC preparation was assessed by staining with the following monoclonal antibodies for 30 min on ice: antilineage cocktail 1 (lin-1; CD3, CD14, CD16, CD19, CD20 and CD56)-fluorescein isothiocyanate (FITC), anti-HLA-DR-peridinin chlorophyll protein (PerCP) (L243), anti-CCR7-phycoerythrin (PE) (3D12) (BD PharMingen, San Diego, CA, USA), anti-CD80-PE (MAB104), anti-CD83-PE (HB15a) and anti-CD86-PE (HA5.2B7) (Beckman Coulter, Fullerton, CA, USA). Cells were analysed on a fluorescence activated cell sorter (FACS0CaliburTM flow cytometer. Data

analysis was performed with CELLQuestTM software (Becton Dickinson, San Jose, CA, USA).

DC phagocytosis

Immature DCs and OK432-stimulated DCs were incubated with 1 mg/ml FITC dextran (Sigma-Aldrich, St Louis, MO, USA) for 30 min at 37°C and the cells were washed three times in FACS buffer before cell acquisition using a FACS-CaliburTM cytometer. Control DCs (not incubated with FITC dextran) were acquired at the same time to allow background levels of fluorescence to be determined.

Enzyme-linked immunosorbent assay (ELISA)

DCs were seeded at 200 000 cells/ml, and supernatant collected after 48 h. IL-12p40 and IFN- γ were detected using matched paired antibodies (BD Pharmingen) following standard protocols.

Cytotoxicity assays

The ability of DCs to exert cytotoxicity was assessed in a standard ⁵¹Cr release assay [19]. We used the HCC cell lines Hep3B and PLC/PRF/5 [American Type Culture Collection (ATCC), Manassas, VA, USA] and a lymphoblastoid cell line T2 that expresses HLA-A*0201 (ATCC) as target cells. Target cells were labelled with ⁵¹Cr. In a 96-well plate, 2.5×10^3 target cells per well were incubated with DCs for 8 h at different effector/target (E/T) ratios in triplicate. Percentage of specific lysis was calculated as follows: (experimental release – spontaneous release)/(maximum release – spontaneous release) × 100. Spontaneous release was always < 20% of the total.

NK cell activity

NK cell cytotoxicity against K562 erythroleukemia target cells was measured by using ⁵¹Cr-release assay, according to previously published methods [26], with PBMCs obtained from the patients. All experiments were performed in triplicate. Percentage of cytotoxicity was calculated as follows: {[experimental counts per minute (cpm) – spontaneous cpm]/[total cpm – spontaneous cpm]} × 100.

Intracellular cytokine expression

Freshly isolated PBMCs were stimulated with 25 ng/ml phorbol 12-myristate 13-acetate (PMA; Sigma-Aldrich) and $1\,\mu g/ml$ ionomycin (Sigma-Aldrich) at 37°C in humidified 7% CO₂ for 4 h. To block cytokine secretion, brefeldin A (Sigma) [27] was added to a final concentration of $10\,\mu g/ml$. After addition of stimuli, the surface staining was performed with anti-CD4-PC5 (13B8·2), anti-CD8-PerCP (SK1) and anti-CD56-PC5 (N901) (Beckman

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Coulter). Subsequently, the cells were permeabilized, stained for intracellular IFN- γ and IL-4 using the FastImmuneTM system (BD Pharmingen), resuspended in phosphate-buffered saline (PBS) containing 1% paraformaldehyde (PFA), and analysed on a flow cytometer (\approx 10 000 gated events acquired per sample).

IFN-γ enzyme-linked immunospot (ELISPOT) assay

ELISPOT assays were performed as described previously with the following modifications [28-30]. HLA-A24 restricted peptide epitopes, squamous cell carcinoma antigen recognized by T cells 2 (SART2)899 (SYTRLFLIL), SART3₁₀₉ (VYDYNCHVDL), multi-drug resistance protein 3 (MRP3)₇₆₅ (VYSDADIFL), MRP3₅₀₃ (LYAWEPSFL), MRP3₆₉₂ (AYVPQQAWI), alpha-fetoprotein (AFP)₄₀₃ (KYIQESQAL), AFP434 (AYTKKAPQL), AFP357 (EYSRRHPQL), human telomerase reverse transcriptase (hTERT)₁₆₇ (AYQVCGPPL) (unpublished), hTERT461 (VYGFVRACL) and hTERT324 (VYAETKHFL) were used in this study. Negative controls consisted of an HIV envelope-derived peptide (HIVenv₅₈₄). Positive controls consisted of 10 ng/ml PMA (Sigma) or a CMV pp65-derived peptide (CMVpp65328). The coloured spots were counted with a KS ELISPOT Reader (Zeiss, Tokyo, Japan). The number of specific spots was determined by subtracting the number of spots in the absence of antigen from the number of spots in its presence. Responses were considered positive if more than 10 specific spots were detected and if the number of spots in the presence of antigen was at least twofold greater than the number of spots in the absence of antigen.

Cytokine and chemokine profiling

Serum cytokine and chemokine levels were measured using the Bioplex assay (Bio-Rad, Hercules, CA, USA). Briefly, frozen serum samples were thawed at room temperature, diluted 1:4 in sample diluents, and 50 µl aliquots of diluted sample were added in duplicate to the wells of a 96-well microtitre plate containing the coated beads for a validated panel of 27 human cytokines and chemokines (cytokine 27-plex antibody bead kit) according to the manufacturer's instructions. These included IL-1β, IL-1Ra, IL-2, IL-4, IL-5, IL-6, IL-7, IL-8, IL-9, IL-10, IL-12p70, IL-13, IL-15, IL-17, basic fibroblast growth factor (FGF), eotaxin, G-CSF, GM-CSF, IFN-y, interferon gamma-induced protein (IP)-10, monocyte chemoattractant protein (MCP)-1, MIP-1α, MIP-1β, platelet-derived growth factor (PDGF)-BB, regulated upon activation normal T cell-expressed and secreted (RANTES), TNF- α and vascular endothelial growth factor (VEGF). Eight standards (ranging from 2 to 32 000 pg/ml) were used to generate calibration curves for each cytokine. Data acquisition and analysis were performed using Bio-Plex Manager software version 4.1.1.

Arginase activity

Serum samples were tested for arginase activity by conversion of L-arginine to L-ornithine [31] using a kit supplied by the manufacturer (BioAssay Systems, Hayward, CA, USA). Briefly, sera were treated with a membrane filter (Millipore, Billerica, MA, USA) to remove urea, combined with the sample buffer in wells of a 96-well plate, and incubated at 37°C for 2 h. Subsequently, the urea reagent was added to stop the arginase reaction. The colour produced was read at 520 nm using a microtitre plate reader.

Statistical analysis

Results are expressed as means ± standard deviation (s.d.). Differences between groups were analysed for statistical significance by the Mann–Whitney *U*-test. Qualitative variables were compared by means of Fisher's exact test. The estimated probability of tumour recurrence-free survival was determined using the Kaplan–Meier method. The Mantel–Cox log-rank test was used to compare curves between groups. Any *P*-values less than 0.05 were considered statistically significant. All statistical tests were two-sided.

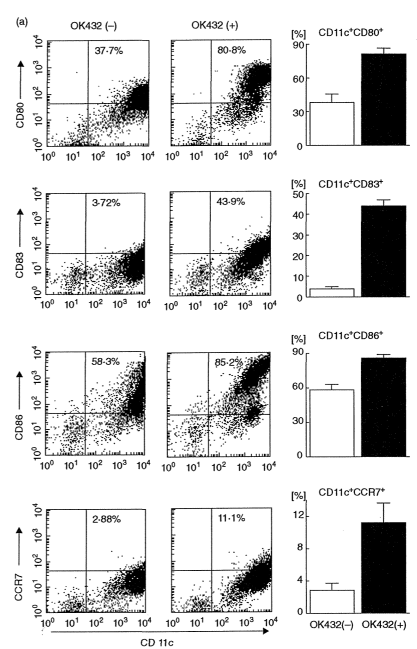
Results

Preparation of OK432-stimulated DCs

Adherent cells isolated from PBMCs of patients with cirrhosis and HCC (Table 1) were differentiated into DCs in the presence of IL-4 and GM-CSF. The cells were stimulated with 0.1 KE/ml OK432 for 3 days; $54.6 \pm 9.5\%$ (mean \pm s.d.; n = 13) of OK432-stimulated cells showed high levels of MHC class II (HLA-DR) and the absence of lineage markers including CD3, CD14, CD16, CD19, CD20 and CD56, in which $30.9 \pm 14.2\%$ were CD11c-positive (myeloid DC subset) and 14.8 ± 11.2 were CD123-positive (plasmacytoid DC subset), consistent with our previous observations [20]. As reported [32,33], greater proportions of the cells developed high levels of expression of the co-stimulatory molecules B7-1 (CD80) and B7-2 (CD86) and an activation marker (CD83) compared to DCs prepared without OK432 stimulation (Fig. 1a). Furthermore, the chemokine receptor CCR7 which leads to homing to lymph nodes [13,34] was also induced following OK432 stimulation.

To evaluate the endocytic and phagocytic ability of the OK432-stimulated cells, uptake of FITC-dextran was quantitated by flow cytometry (Fig. 1b). The cells showed lower levels of uptake due to maturation compared to DCs prepared without OK432 stimulation, while the OK432-stimulated cells derived from HCC patients preserved a moderate uptake capacity. As expected, the OK432-stimulated cells produced large amounts of cytokines IL-12 and IFN-γ (Fig. 1c). In addition, they displayed high cyto-

Fig. 1. Effects of OK432 stimulation on the properties of dendritic cells (DCs) generated from blood monocyte precursors in patients with cirrhosis and hepatocellular carcinoma (HCC) (n = 13). (a) Lineage cocktail 1 (lin 1⁻) human leucocyte antigen D-related (HLA-DR-) subsets with [OK432(+)] and without [OK432(-)] stimulation were analysed for surface expression of CD80, CD83, CD86 and CCR7. Dot plots of a representative case are shown in the left-hand panel. Mean percentages [±standard deviation (s.d.)] of positive cells are indicated in the right-hand panel. OK432 stimulation resulted in the expression of high levels of CD80, CD83, CD86 and CCR7 in the lin 1-human leucocyte antigen D-related (HLA-DR⁻) DC subset. (b) DC subsets with and without OK432 stimulation were incubated with fluorescein isothiocyanate (FITC) dextran for 30 min and the uptake was determined by flow cytometry. A representative analysis is shown in the upper panel. Mean fluorescence intensities (MFIs) (±s.d.) of the positive cells are indicated in the lower panel. OK432-stimulated cells showed lower levels of uptake due to maturation. (c) DC supernatants were harvested and the concentrations of interleukin (IL)-12 and interferon (IFN)-γ measured by enzyme-linked immunosorbent assay (ELISA). OK432-stimulated cells produced large amounts of the cytokines. The data indicate means \pm s.d. of the groups with and without the stimulation. All comparisons in (a-c) [OK432(+) versus OK432(-)] were statistically significant by the Mann-Whitney U-test (P < 0.005). (d) Tumoricidal activity of DCs assessed by incubation with 51Cr-labelled Hep3B, PLC/PRF/5 and T2 targets for 8 h at the indicated effector/target (E/T) cell ratios. OK432-stimulated cells displayed high cytotoxic activity against the target cells. The results are representative of the cases studied.



toxic activity against HCC cell lines (Hep3B and PLC/PRF/5) and a lymphoblastoid cell line (T2) although DCs without OK432 stimulation lysed none of the target cells to any great degree (Fig. 1d). Taken together, these results demonstrate that OK432 stimulation of IL-4 and GM-CSF-induced immature DCs derived from HCC patients promoted their maturation towards cells with activated phenotypes, high expression of a homing receptor, fairly well-preserved phagocytic capacity, greatly enhanced cytokine production and effective tumoricidal activity, consistent with previous observations [16,19].

Safety of OK432-stimulated DC administration

Prior to the administration of OK432-stimulated DCs to patients, the cells were confirmed to be safe in athymic nude mice to which 100-fold cell numbers/weight were injected subcutaneously (data not shown). Subsequently, OK432-stimulated DC administration was performed during TAE therapy in humans, in which DCs were mixed together with absorbable gelatin sponge (Gelfoam) and infused through an arterial catheter following iodized oil (Lipiodol) injection, as reported previously [20]. Adverse events were

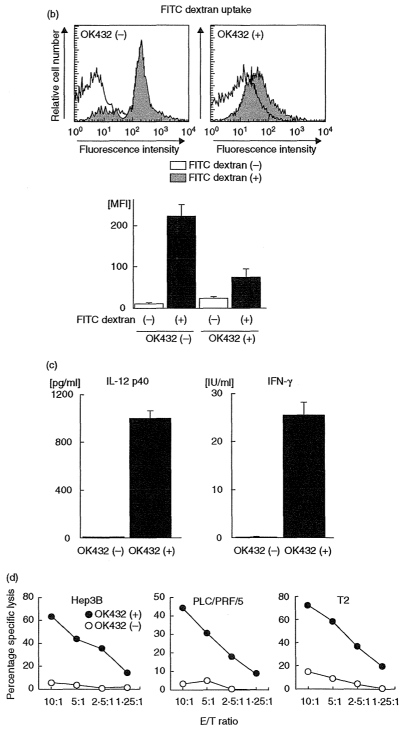


Fig. 1. Continued

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monitored clinically and biochemically after DC infusion (Table 2). A larger proportion (12 of 13) of the patients were complicated with high fever compared to those treated previously with immature DCs (five of 10) [20], due probably to the proinflammatory responses induced by OK432-stimulated DCs. However, there were no grades III or IV

National Cancer Institute Common Toxicity Criteria adverse events, including vomiting, abdominal pain, encephalopathy, myalgia, ascites, gastrointestinal disorders, bleeding, hepatic abscess or autoimmune diseases associated with DC infusion and TAE in this study. There was also no clinical or serological evidence of hepatic failure or autoimmune

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Table 2. Adverse events.

Patient	Fever		Abdominal		
no.	(days)	Vomiting	pain	Encephalopathy	Others†
1	2	No	No	No	No
2	2	No	No	No	No
3	1	No	No	No	No
4	3	No	No	No	No
5	3	No	No	No	No
6	4	No	No	No	No
7	10	No	No	No	No
8	No	No	No	No	No
9	2	No	No	No	No
10	1	No	No	No	No
11	2	No	No	No	No
12	2	No	No	No	No
13	1	No	No	No	No

[†]Other adverse events include myalgia, ascites, gastrointestinal disorder, bleeding, hepatic abscess and autoimmune diseases.

response in any patients. Thus, concurrent treatment with OK432-stimulated DC infusions can be performed safely at the same time as TAE in patients with cirrhosis and HCC.

Recurrence-free survival following DC infusion

A further objective of this study was to determine clinical response following DC infusion. A group of historical controls treated with TAE without DC administration was reviewed for this study (Table 3). The clinical characteristics including tumour burden and hepatic reserve were comparable between patients treated with TAE and OK432-stimulated DC transfer (n = 13) and those historical controls with TAE but without DC administration (n = 22). We com-

pared the recurrence-free survival between these patient groups. Kaplan–Meier analysis indicated that patients treated with TAE and OK432-stimulated DC transfer had prolonged recurrence-free survival compared with the historical controls that had been treated with TAE alone (recurrence rates 360 days after the treatments; two of 13 and 12 of 22, respectively; P=0.046, log-rank test) (Fig. 2). The results demonstrated that OK432-stimulated DC transfer during TAE therapy reduces tumour recurrence in HCC patients.

NK cell activity and intracellular cytokine responses in PBMCs

To assess systemic immunomodulatory effects of OK432-stimulated DC transfer, PBMCs were isolated 1 and 3 months after treatment and NK cell cytotoxicity against K562 erythroleukaemia target cells measured using the ^{51}Cr release assay (Fig. 3). The level of NK cell was unaltered following treatment. In addition, cytokine production capacity of lymphocyte subsets was quantitated by measuring intracellular IFN- γ and IL-4 using flow cytometry. There were also no significant changes in terms of cytokine production capacity in the CD4+, CD8+ and CD56+ subsets in the patients treated with OK432-stimulated DCs.

Immune responses to peptide epitopes derived from tumour antigens

To assess the effects on T cell responses to tumour antigens, PBMCs were obtained 4 weeks after DC infusion, pulsed with peptides derived from AFP, MRP3, SART2, SART3 and hTERT. IFN- γ production was then quantitated in an

Table 3. Clinical characteristics of patients treated with TAE + OK-DC and TAE alone.

	TAE + OK-DC	TAE	P
No. of patients	13	22	
Age (years)	68.2 ± 9.1	70.0 ± 7.6	n.s.†
Gender (M/F)	9/4	13/9	n.s.‡
White cell count ($\times 10^2/\mu l$)	34.4 ± 11.6	41.4 ± 18.9	n.s.†
Lymphocytes (×10²/μl)	10.4 ± 3.6	12.4 ± 4.7	n.s.†
Platelets (×10 ⁴ /µl)	11.5 ± 10.2	10.3 ± 5.8	n.s.†
Hepaplastin test (%)	64.6 ± 11.6	75.5 ± 24.3	n.s.†
ALT (IU/l)	56.7 ± 38.9	67.9 ± 44.6	n.s.†
Total bilirubin (mg/dl)	1.3 ± 0.7	1.1 ± 0.6	n.s.†
Albumin (g/dl)	3.4 ± 0.6	3.6 ± 0.4	n.s.†
Non-cancerous liver parenchyma (no.)			
Chronic hepatitis	0	8	
Cirrhosis (Child-Pugh A/B/C)	13 (5/8/0)	14 (6/8/0)	n.s.‡
TNM stages (I/II/III/IV-A/IV-B)	0/4/9/0/0	3/8/11/0/0	n.s.‡
No. of tumours	2.5 ± 1.3	1.9 ± 1.3	n.s.†
Largest tumour (mm)	30.2 ± 9.4	32.6 ± 15.2	n.s.†
AFP	204.8 ± 404.1	201.8 ± 544.2	n.s.†

Results are expressed as means \pm standard deviation. †Mann–Whitney U-test. †Fisher's exact test. TAE, transcatheter arterial embolization; OK-DC, OK432-stimulated dendritic cells; ALT, alanine transaminase; TNM, tumour–node–metastasis; AFP, alpha-fetoprotein; Child–Pugh, Child–Pugh classification; n.s., not significant.

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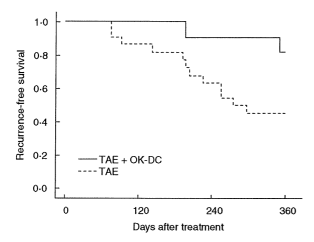
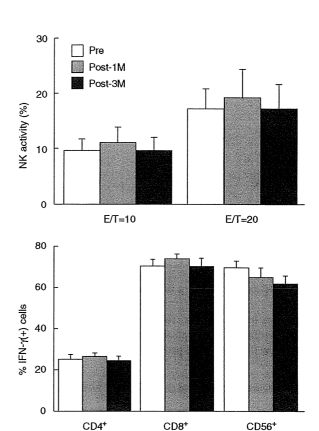


Fig. 2. Recurrence-free survival of patients treated with transcatheter hepatic arterial embolization (TAE) with [TAE + OK-stimulated dendritic cells (DC); n=13] and without (TAE: historical controls; n=22) OK432-stimulated DC administration. Time zero is the date of TAE. All patients underwent ultrasound, computed tomography (CT) scan or magnetic resonance imaging (MRI) of the abdomen about 1 month after treatment and at a minimum of once every 3 months thereafter. Kaplan–Meier analysis indicated that TAE + OK-DC treatment prolonged recurrence-free survival compared with the TAE-alone group (recurrence rates 360 days after the treatments; two of 13 and 12 of 22, respectively; P=0.046, log-rank test).

ELISPOT assay. Cells producing IFN- γ in response to stimulation with HLA-A24 [the most common HLA-A antigen (58·1%) in Japanese populations [35]]-restricted peptide epitopes derived from tumour antigens MRP3 and hTERT were induced in three of six HLA-A24-positive patients (numbers 2, 6 and 11) after treatment with TAE and OK432stimulated DCs (Fig. 4). To understand the immunological and clinical significance of the T lymphocyte responses, PBMCs obtained from the historical control patients who had been treated with TAE without DC administration were also evaluated by ELISPOT. Similarly, positive reactions were observed in four (numbers t8, t19, t20 and t22) of six HLA-A24-positive patients. These data indicate that T lymphocyte responses to HLA-A24 restricted peptide epitopes of tumour antigens were induced following the TAE therapy, but no additional responses were observed as a result of OK432stimulated DC transfer in the current study.

Serum levels of cytokines, chemokines and arginase activity

To screen for immunobiological responses induced following OK432-stimulated DC transfer, serum levels of cytokines and chemokines were measured simultaneously using the Bio-Plex multiplex suspension array system. The results were compared with the historical control patients treated with TAE without DC administration. Interestingly, serum con-



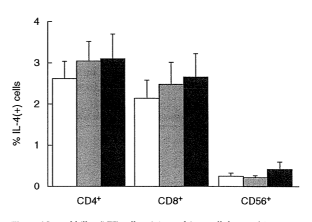
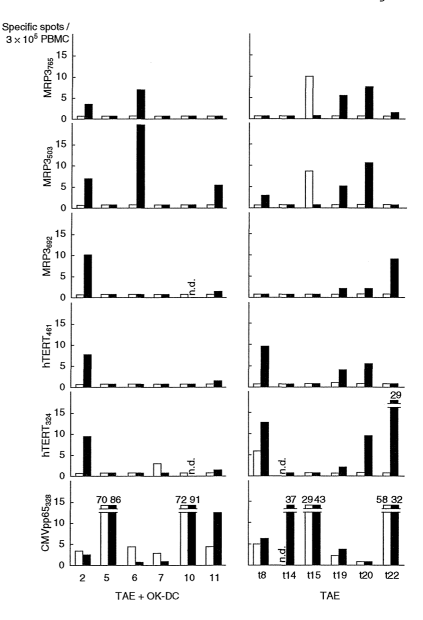


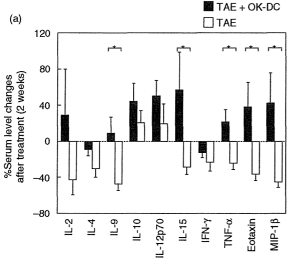
Fig. 3. Natural killer (NK) cell activity and intracellular cytokine production in peripheral blood mononuclear cells (PBMCs) of patients treated with OK432-stimulated dendritic cells (DCs) during transcatheter hepatic arterial embolization (TAE) therapy (n = 13). PBMCs were isolated before and 1 and 3 months after treatment and used for the analyses. Upper panel: NK cell cytotoxicity against K562 erythroleukaemia target cells was evaluated at the effector/target (E/T) cell ratios shown. NK cell activities were not changed following treatment. Middle and lower panels: PBMCs were stimulated with phorbol 12-myristate 13-acetate (PMA) and ionomycin, stained for CD4, CD8 and CD56 expression, permeabilized and stained for intracellular interferon (IFN)-γ and interleukin (IL)-4. Percentages of cytokine-positive cells were quantitated by flow cytometry. There were no significant changes in terms of cytokine production capacity in the CD4+, CD8+ and CD56+ subsets following the treatments. The data are given as means ± standard deviation of the groups.

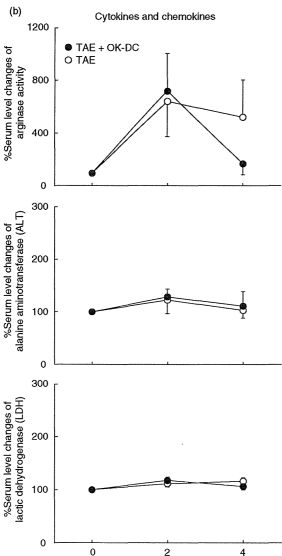
Fig. 4. Immune responses to human leucocyte antigen (HLA-DR⁻)-A24-restricted peptide epitopes derived from tumour antigens in HLA-A24-positive patients treated with OK432-stimulated DCs during transcatheter hepatic arterial embolization (TAE) therapy (numbers 2, 5, 6, 7, 10 and 11) and HLA-A24-positive historical controls treated with TAE without dendritic cell (DC) transfer (numbers t8, t14, t15, t19, t20 and t22). Peripheral blood mononuclear cells (PBMCs) were obtained before (open bars) and 1 month after the infusion (solid bars), pulsed with the peptides derived from squamous cell carcinoma antigen recognized by T cells 2 (SART2), SART3, multi-drug resistance protein 3 (MRP3), alpha-fetoprotein (AFP), human telomerase reverse transcriptase (hTERT) and interferon (IFN)-γ production was quantitated by enzyme-linked immunospot (ELISPOT). Negative controls consisted of a human immunodeficiency virus (HIV) envelope-derived peptide (HIVenv₅₈₄). Positive controls consisted of 10 ng/ml phorbol 12-myristate 13-acetate (PMA) or a cytomegalovirus (CMV) pp65-derived peptide (CMVpp65328). The number of specific spots was determined by subtracting the number of spots in the absence of antigen from the number of spots in its presence. T lymphocyte responses to the peptide epitopes were induced following TAE therapy, but no additional responses were observed after DC transfer. Numbers denote specific spots beyond the upper limit of y-axis; n.d., not determined.



centrations of IL-9, IL-15 and TNF- α were greatly increased after OK432-stimulated DC infusion, in contrast to their reduction following TAE treatment alone (Fig. 5a). Furthermore, the chemokines eotaxin (CCL11) and MIP-1 β (CCL4) were induced markedly after DC transfer, although they were also decreased after TAE alone. These data indicate that transfer of OK432-stimulated DC during TAE therapy induced unique immune responses that may be mediated by the cytokines IL-9, IL-15 and TNF- α and the chemokines eotaxin and MIP-1 β .

In addition, serum arginase activity was reported to reflect numbers of myeloid-derived suppressor cells (MDSCs) that may inhibit T lymphocyte responses in cancer patients [36]. Therefore, serum arginase activity was measured after OK432-stimulated DC infusion, and it was found that it was increased six- or sevenfold in patients treated with TAE. However, this increase was independent of the presence or absence of OK432-stimulated DC transfer (Fig. 5b). None the less, serum arginase activity was decreased again 4 weeks after treatment with both TAE and OK432-stimulated DC transfer but tended to be maintained at a high levels in patients treated with TAE without DC transfer. However, these differences did not reach statistical significance (P > 0.05). Because arginase activity is known to be relatively high in liver and HCC cells [37], the influence of tissue injury was assessed biochemically by measuring serum levels of ALT and LDH activities. We did not observe ALT or LDH elevation, indicating that the increase of arginase activity was not due to tissue damage following treatment. Collectively, these results demonstrate that infusion of OK432-stimulated





Weeks after treatment

Fig. 5. Cytokine and chemokine profiling and arginase activity in sera of patients treated with OK432-stimulated dendritic cells (DCs) during transcatheter hepatic arterial embolization (TAE) therapy (TAE + OK-DC; n = 13) and the historical controls treated with TAE without DC transfer (TAE; n = 22). (a) Serum samples were examined for their content of a validated panel of cytokines and chemokines. using the Bioplex assay. Percentage changes in serum levels 2 weeks after the treatments were calculated as follows: [(post-treatment level – pretreatment level)/pretreatment level] \times 100. The data are means \pm standard error of the mean (s.e.m.) of the groups. *P < 0.05 when compared by the Mann–Whitney U-test. (b) Serum samples were tested for arginase activity by conversion of L-arginine to L-ornithine, and for alanine aminotransferase (ALT) and lactic dehydrogenase (LDH) activities. While there was a trend for the arginase activity in the TAE + OK-DC group to decrease 4 weeks after treatment, the difference did not reach statistical significance (P > 0.05). Percentage changes in serum levels 2 weeks after the treatments were calculated as follows: [(post-treatment level pretreatment level)/pretreatment level] × 100. The data indicate means \pm s.e.m. of the groups.

DCs during TAE treatment may reduce the immunosuppressive activities of MDSCs, and assist in developing a favourable environment for the induction of anti-tumour immunity.

Discussion

Although many novel strategies, including immunotherapies, have been developed in an attempt to suppress tumour recurrence after curative treatments for HCC, recurrence rates and survival times have not been improved significantly [38]. In the current study, we first established that OK432stimulated DC administration during TAE therapy did not cause critical adverse events in patients with cirrhosis and HCC. Most importantly, DC transfer resulted in prolonged recurrence-free survival after combination therapy with TAE and OK432-stimulated DC administration. In terms of the immunomodulatory effects of DC transfer, although NK cell activity, intracellular cytokine production and T lymphocyte-mediated immune responses were not altered in PBMCs from treated patients, serum levels of IL-9, IL-15 and TNF- α and the chemokines eotaxin and MIP-1 β were enhanced markedly after DC transfer. In addition, serum levels of arginase activity were decreased following DC transfer. Collectively, this study demonstrated the feasibility, safety and beneficial anti-tumour effects of OK432stimulated DC infusion into tumour tissues for patients with cirrhosis and HCC, suggesting the ability of an active immunotherapeutic strategy to reduce tumour recurrence after locoregional treatment of HCC.

DCs were stimulated with OK432 prior to infusion into tumour tissues through an arterial catheter. OK432 was reported to activate DCs through its binding to TLR-2 and -4 [16,39] that can be used for cancer therapy [33]. The current results indicate that OK432 stimulation of immature DCs

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from HCC patients promoted their maturation processes while preserving antigen uptake capacity and enhancing tumoricidal activity, consistent with previous observations [16,19] and supporting the current strategy in which OK432-stimulated DCs were infused directly into tumour tissues. Because the tumoricidal activity of unstimulated DCs was not observed in in vitro experiments, OK432 stimulation obviously altered the cytotoxic properties of DCs. One of the mechanisms of DC killing was reported to be CD40/ CD40 ligand interaction [19]. Further studies are needed to determine the killing mechanisms of DCs derived from HCC patients in a direct [TNF, TNF-related apoptosis inducing ligand (TRAIL), Fas ligand, nitric oxide (NO) and perforin/ granzyme] and indirect (MHC-restricted) manner [40-43]. Although the main mechanism by which OK432-stimulated DCs prolonged the recurrence-free survival was not elucidated, the tumoricidal activity of mature DCs was implicated in in vivo enhancement of antigen presentation, co-stimulation and inflammatory cytokine production.

Very recent reports document injection of OK432stimulated DCs into patients with cancer of the gastrointestinal tract or pancreas [44,45], but their anti-tumour effects were not defined clearly. The current study shows for the first time that OK432-stimulated DCs induce beneficial antitumour responses when transferred into tumour tissues during TAE therapy. The anti-tumour responses may have been enhanced as a result of optimal activation of the DCs with OK432 or combining infusion of stimulated DCs with TAE therapy. Inappropriately activated DCs may be unable to generate sufficient numbers of properly activated effector T lymphocytes [46]. As shown in Fig. 1, all these alterations could contribute to the further enhancement of anti-tumour effects compared to those in our previous study with immature DCs [20]. Furthermore, the tumour cell deathpromoting therapies, e.g. chemotherapy [47] and TAE [48], can be expected to enhance the effects of therapeutic cancer vaccines by redressing the immunosuppressive tumour environment.

NK cell activity and intracellular cytokine responses in CD4+ and CD8+ T lymphocytes and CD56+ NK cell subsets in PBMCs were not changed significantly in patients treated with OK432-stimulated DCs. Furthermore, we did not observe tumour antigen-specific T lymphocyte responses associated clearly with DC administration. The data suggest therefore that the immune responses induced by the therapy applied here were not detectable systemically. Because cytotoxic T lymphocyte responses were enhanced in patients receiving $> 3 \times 10^7$ cells [49,50], the numbers of transferred OK432-stimulated DCs were apparently not sufficient to induce responses detectable in the peripheral blood, but were enough to exert beneficial anti-tumour effects. In addition, many studies have concluded that cytotoxic T lymphocyte responses rarely predict clinical outcomes of DC-based immunotherapies [51,52] and that in many cases, also including our own studies

[28,30], tumour-specific effector T lymphocytes co-exist with the tumours. Consistent with these observations, the current results suggest that cytotoxic T lymphocyte responses in PBMCs are not reliable predictors of beneficial anti-tumour effects in patients treated with the current OK432-stimulated DC strategy.

Serum levels of the cytokines IL-9, IL-15 and TNF-α and the chemokines eotaxin and MIP-1B were increased following OK432-stimulated DC transfer, but decreased after TAE therapy without DC administration. IL-9 and IL-15 belong to the cytokine receptor common gamma chain (γ_c; CD132) family, a member of the type I cytokine receptor family expressed on most lymphocyte populations [53]. IL-9 exerts pleiotropic activities on T and B lymphocytes, mast cells, monocytes and haematopoietic progenitors [54,55]. IL-15 and TNF-α are known to prime T lymphocytes and NK cells when secreted by DCs [56] and to induce anti-tumour immune responses [57]. Eotaxin is known to selectively recruit eosinophils also contributing to anti-tumour effects [58,59], and MIP-1ß is a chemoattractant for NK cells, monocytes and a variety of other immune cells [60]. In addition, serum levels of arginase tended to decrease after DC transfer. Because serum arginase activity reflects the numbers of MDSCs that inhibit T lymphocyte responses in cancer patients [36], the patients treated with OK432-stimulated DCs might have developed lower levels of suppressor cells. Collectively, the results suggest that infusion of OK432stimulated DCs may orchestrate the immune environment in the whole body that could enhance beneficial anti-tumour effects, although the precise molecular and cellular mechanisms associated with the actions of these cytokines and chemokines were not defined clearly in the current analysis.

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Disclosure

The authors have declared that no conflict of interest exists.

References

- 1 Omata M, Tateishi R, Yoshida H, Shiina S. Treatment of hepatocellular carcinoma by percutaneous tumor ablation methods: ethanol injection therapy and radiofrequency ablation. Gastroenterology 2004; 127:S159–66.
- 2 Belghiti J. Resection and liver transplantation for HCC. J Gastroenterol 2009; 44 (Suppl. 19):132–5.

- 3 Nakamoto Y, Guidotti LG, Kuhlen CV, Fowler P, Chisari FV. Immune pathogenesis of hepatocellular carcinoma. J Exp Med 1998; 188:341–50.
- 4 Ercolani G, Grazi GL, Ravaioli M et al. Liver resection for hepatocellular carcinoma on cirrhosis: univariate and multivariate analysis of risk factors for intrahepatic recurrence. Ann Surg 2003; 237:536–43.
- 5 Shankaran V, Ikeda H, Bruce AT et al. IFNgamma and lymphocytes prevent primary tumour development and shape tumour immunogenicity. Nature 2001; 410:1107–11.
- 6 Vulink A, Radford KJ, Melief C, Hart DN. Dendritic cells in cancer immunotherapy. Adv Cancer Res 2008; 99:363–407.
- 7 Banchereau J, Briere F, Caux C et al. Immunobiology of dendritic cells. Annu Rev Immunol 2000; 18:767–811.
- 8 Lemos MP, Esquivel F, Scott P, Laufer TM. MHC class II expression restricted to CD8alpha+ and CD11b+ dendritic cells is sufficient for control of Leishmania major. J Exp Med 2004; 199:725-30.
- 9 Ni K, O'Neill HC. The role of dendritic cells in T cell activation. Immunol Cell Biol 1997; 75:223–30.
- 10 Andrews DM, Andoniou CE, Scalzo AA et al. Cross-talk between dendritic cells and natural killer cells in viral infection. Mol Immunol 2005; 42:547–55.
- 11 Heiser A, Coleman D, Dannull J et al. Autologous dendritic cells transfected with prostate-specific antigen RNA stimulate CTL responses against metastatic prostate tumors. J Clin Invest 2002; 109:409–17.
- 12 Banchereau J, Steinman RM. Dendritic cells and the control of immunity. Nature 1998; 392:245–52.
- 13 Forster R, Schubel A, Breitfeld D et al. CCR7 coordinates the primary immune response by establishing functional microenvironments in secondary lymphoid organs. Cell 1999; 99:23-33.
- 14 MartIn-Fontecha A, Sebastiani S, Hopken UE et al. Regulation of dendritic cell migration to the draining lymph node: impact on T lymphocyte traffic and priming. J Exp Med 2003; 198:615–21.
- 15 Ratzinger G, Stoitzner P, Ebner S et al. Matrix metalloproteinases 9 and 2 are necessary for the migration of Langerhans cells and dermal dendritic cells from human and murine skin. J Immunol 2002; 168:4361-71.
- 16 Nakahara S, Tsunoda T, Baba T, Asabe S, Tahara H. Dendritic cells stimulated with a bacterial product, OK-432, efficiently induce cytotoxic T lymphocytes specific to tumor rejection peptide. Cancer Res 2003; 63:4112–18.
- 17 Okamoto M, Oshikawa T, Tano T et al. Mechanism of anticancer host response induced by OK-432, a streptococcal preparation, mediated by phagocytosis and Toll-like receptor 4 signaling. J Immunother 2006; 29:78–86.
- 18 Pasare C, Medzhitov R. Toll pathway-dependent blockade of CD4+CD25+ T cell-mediated suppression by dendritic cells. Science 2003; 299:1033-6.
- 19 Hill KS, Errington F, Steele LP et al. OK432-activated human dendritic cells kill tumor cells via CD40/CD40 ligand interactions. J Immunol 2008; 181:3108-15.
- 20 Nakamoto Y, Mizukoshi E, Tsuji H et al. Combined therapy of transcatheter hepatic arterial embolization with intratumoral dendritic cell infusion for hepatocellular carcinoma: clinical safety. Clin Exp Immunol 2007; 147:296–305.
- 21 Steinman RM, Banchereau J. Taking dendritic cells into medicine. Nature 2007; 449:419–26.

- 22 Tacken PJ, de Vries IJ, Torensma R, Figdor CG. Dendritic-cell immunotherapy: from ex vivo loading to in vivo targeting. Nat Rev Immunol 2007; 7:790–802.
- 23 Makuuchi M. General rules for the clinical and pathological study of primary liver cancer, 2nd edn. Tokyo: Kanehara & Co., Ltd, 2003.
- 24 Veltri A, Moretto P, Doriguzzi A, Pagano E, Carrara G, Gandini G. Radiofrequency thermal ablation (RFA) after transarterial chemoembolization (TACE) as a combined therapy for unresectable non-early hepatocellular carcinoma (HCC). Eur Radiol 2006; 16:661-9.
- 25 Dhodapkar MV, Steinman RM, Sapp M et al. Rapid generation of broad T-cell immunity in humans after a single injection of mature dendritic cells. J Clin Invest 1999; 104:173–80.
- 26 Orange JS, Brodeur SR, Jain A et al. Deficient natural killer cell cytotoxicity in patients with IKK-gamma/NEMO mutations. J Clin Invest 2002; 109:1501–9.
- 27 Klausner RD, Donaldson JG, Lippincott-Schwartz J. Brefeldin A: insights into the control of membrane traffic and organelle structure. J Cell Biol 1992; 116:1071–80.
- 28 Mizukoshi E, Nakamoto Y, Marukawa Y et al. Cytotoxic T cell responses to human telomerase reverse transcriptase in patients with hepatocellular carcinoma. Hepatology 2006; 43:1284–94.
- 29 Mizukoshi E, Nakamoto Y, Tsuji H, Yamashita T, Kaneko S. Identification of alpha-fetoprotein-derived peptides recognized by cytotoxic Tlymphocytes in HLA-A24+ patients with hepatocellular carcinoma. Int J Cancer 2006; 118:1194–204.
- 30 Mizukoshi E, Honda M, Arai K, Yamashita T, Nakamoto Y, Kaneko S. Expression of multidrug resistance-associated protein 3 and cytotoxic T cell responses in patients with hepatocellular carcinoma. J Hepatol 2008; 49:946–54.
- 31 Rodriguez PC, Quiceno DG, Zabaleta J et al. Arginase I production in the tumor microenvironment by mature myeloid cells inhibits T-cell receptor expression and antigen-specific T-cell responses. Cancer Res 2004; 64:5839–49.
- 32 Itoh T, Ueda Y, Okugawa K et al. Streptococcal preparation OK432 promotes functional maturation of human monocyte-derived dendritic cells. Cancer Immunol Immunother 2003; 52:207-14
- 33 Kuroki H, Morisaki T, Matsumoto K et al. Streptococcal preparation OK-432: a new maturation factor of monocyte-derived dendritic cells for clinical use. Cancer Immunol Immunother 2003; 52:561-8.
- 34 Gunn MD, Kyuwa S, Tam C et al. Mice lacking expression of secondary lymphoid organ chemokine have defects in lymphocyte homing and dendritic cell localization. J Exp Med 1999; 189:451–60
- 35 Imanishi T, Akaza T, Kimura A, Tokunaga K, Gojobori T. HLA 1991, Proceedings of the Eleventh International Histocompatibility Workshop and Conference. Tokyo: Oxford University Press, 1992.
- 36 Zea AH, Rodriguez PC, Atkins MB et al. Arginase-producing myeloid suppressor cells in renal cell carcinoma patients: a mechanism of tumor evasion. Cancer Res 2005; 65:3044–8.
- 37 Chrzanowska A, Krawczyk M, Baranczyk-Kuzma A. Changes in arginase isoenzymes pattern in human hepatocellular carcinoma. Biochem Biophys Res Commun 2008; 377:337–40.
- 38 Caldwell S, Park SH. The epidemiology of hepatocellular cancer: from the perspectives of public health problem to tumor biology. J Gastroenterol 2009; 44 (Suppl. 19):96–101.
- 39 Okamoto M, Oshikawa T, Tano T et al. Involvement of Toll-like

- receptor 4 signaling in interferon-gamma production and antitumor effect by streptococcal agent OK-432. J Natl Cancer Inst 2003; 95:316–26.
- 40 Liu S, Yu Y, Zhang M, Wang W, Cao X. The involvement of TNF-alpha-related apoptosis-inducing ligand in the enhanced cytotoxicity of IFN-beta-stimulated human dendritic cells to tumor cells. J Immunol 2001; 166:5407–15.
- 41 Lu G, Janjic BM, Janjic J, Whiteside TL, Storkus WJ, Vujanovic NL. Innate direct anticancer effector function of human immature dendritic cells. II. Role of TNF, lymphotoxin-alpha(1)beta(2), Fas ligand, and TNF-related apoptosis-inducing ligand. J Immunol 2002; 168:1831–9.
- 42 Nicolas A, Cathelin D, Larmonier N et al. Dendritic cells trigger tumor cell death by a nitric oxide-dependent mechanism. J Immunol 2007; 179:812–18.
- 43 Stary G, Bangert C, Tauber M, Strohal R, Kopp T, Stingl G. Tumoricidal activity of TLR7/8-activated inflammatory dendritic cells. J Exp Med 2007; 204:1441–51.
- 44 West E, Morgan R, Scott K et al. Clinical grade OK432-activated dendritic cells: in vitro characterization and tracking during intralymphatic delivery. J Immunother 2009; 32:66–78.
- 45 Hirooka Y, Itoh A, Kawashima H et al. A combination therapy of gemcitabine with immunotherapy for patients with inoperable locally advanced pancreatic cancer. Pancreas 2009; 38:e69–74.
- 46 Melief CJ. Cancer immunotherapy by dendritic cells. Immunity 2008; 29:372–83.
- 47 Zitvogel L, Apetoh L, Ghiringhelli F, Kroemer G. Immunological aspects of cancer chemotherapy. Nat Rev Immunol 2008; 8:59–73.
- 48 Ayaru L, Pereira SP, Alisa A et al. Unmasking of alpha-fetoprotein-specific CD4(+) T cell responses in hepatocellular carcinoma patients undergoing embolization. J Immunol 2007; 178:1914–22.
- 49 Thurner B, Haendle I, Roder C et al. Vaccination with mage-3A1 peptide-pulsed mature, monocyte-derived dendritic cells expands specific cytotoxic T cells and induces regression of some metastases in advanced stage IV melanoma. J Exp Med 1999; 190:1669–78.

- 50 Banchereau J, Palucka AK, Dhodapkar M et al. Immune and clinical responses in patients with metastatic melanoma to CD34(+) progenitor-derived dendritic cell vaccine. Cancer Res 2001; 61:6451-8.
- 51 Engell-Noerregaard L, Hansen TH, Andersen MH, Thor Straten P, Svane IM. Review of clinical studies on dendritic cell-based vaccination of patients with malignant melanoma: assessment of correlation between clinical response and vaccine parameters. Cancer Immunol Immunother 2009; 58:1–14.
- 52 Itoh K, Yamada A, Mine T, Noguchi M. Recent advances in cancer vaccines: an overview. Jpn J Clin Oncol 2009; 39:73–80.
- 53 Sugamura K, Asao H, Kondo M *et al.* The common gamma-chain for multiple cytokine receptors. Adv Immunol 1995; **59**:225–77.
- 54 Temann UA, Geba GP, Rankin JA, Flavell RA. Expression of interleukin 9 in the lungs of transgenic mice causes airway inflammation, mast cell hyperplasia, and bronchial hyperresponsiveness. J Exp Med 1998; 188:1307–20.
- 55 McMillan SJ, Bishop B, Townsend MJ, McKenzie AN, Lloyd CM. The absence of interleukin 9 does not affect the development of allergen-induced pulmonary inflammation nor airway hyperreactivity. J Exp Med 2002; 195:51–7.
- 56 de Saint-Vis B, Fugier-Vivier I, Massacrier C et al. The cytokine profile expressed by human dendritic cells is dependent on cell subtype and mode of activation. J Immunol 1998; 160:1666–76.
- 57 Shanmugham LN, Petrarca C, Frydas S et al. IL-15 an immunoregulatory and anti-cancer cytokine. Recent advances. J Exp Clin Cancer Res 2006; 25:529–36.
- 58 Kataoka S, Konishi Y, Nishio Y, Fujikawa-Adachi K, Tominaga A. Antitumor activity of eosinophils activated by IL-5 and eotaxin against hepatocellular carcinoma. DNA Cell Biol 2004; 23:549–60.
- 59 Simson L, Ellyard JI, Dent LA et al. Regulation of carcinogenesis by IL-5 and CCL11: a potential role for eosinophils in tumor immune surveillance. J Immunol 2007; 178:4222-9.
- 60 Bystry RS, Aluvihare V, Welch KA, Kallikourdis M, Betz AG. B cells and professional APCs recruit regulatory T cells via CCL4. Nat Immunol 2001; 2:1126-32.

Comparative Analysis of Various Tumor-Associated Antigen-Specific T-Cell Responses in Patients with Hepatocellular Carcinoma

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Many tumor-associated antigens (TAAs) recognized by cytotoxic T cells (CTLs) have been identified during the last two decades and some of them have been used in clinical trials. However, there are very few in the field of immunotherapy for hepatocellular carcinoma (HCC) because there have not been comparative data regarding CTL responses to various TAAs. In the present study, using 27 peptides derived from 14 different TAAs, we performed comparative analysis of various TAA-specific T-cell responses in 31 HCC patients to select useful antigens for immunotherapy and examined the factors that affect the immune responses to determine a strategy for more effective therapy. Twenty-four of 31 (77.4%) HCC patients showed positive responses to at least one TAA-derived peptide in enzymelinked immunospot assay. The TAAs consisting of cyclophilin B, squamous cell carcinoma antigen recognized by T cells (SART) 2, SART3, p53, multidrug resistance-associated protein (MRP) 3, alpha-fetoprotein (AFP) and human telomerase reverse transcriptase (hTERT) were frequently recognized by T cells and these TAA-derived peptides were capable of generating peptide-specific CTLs in HCC patients, which suggested that these TAAs are immunogenic. HCC treatments enhanced TAA-specific immune responses with an increased number of memory T cells and induced de novo T-cell responses to lymphocyte-specific protein tyrosine kinase, human epidermal growth factor receptor type 2, p53, and hTERT. Blocking cytotoxic T-lymphocyte antigen-4 (CTLA-4) resulted in unmasking of TAA-specific immune responses by changing cytokine and chemokine profiles of peripheral blood mononuclear cells stimulated by TAA-derived peptides. Conclusion: Cyclophilin B, SART2, SART3, p53, MRP3, AFP, and hTERT were immunogenic targets for HCC immunotherapy. TAA-specific immunotherapy combined with HCC treatments and anti-CTLA-4 antibody has the possibility to produce stronger tumor-specific immune responses. (HEPATOLOGY 2011;53:1206-1216)

epatocellular carcinoma (HCC) is the most common primary malignancy of the liver and becoming an important public health concern.^{1,2} Although many kinds of treatments have

Although many kinds of treatments have to prevent recurrence is necessary and the province of the province of

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MRP, multidrug resistance-associated protein; PBMC, peripheral blood

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mononuclear cell; TAA, tumor-associated antigen.

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been performed for HCC, their effects are limited because the recurrence rate of HCC is very high; therefore, the development of new therapeutic options to prevent recurrence is necessary.^{3,4}

To protect against recurrence, tumor antigen-specific immunotherapy is an attractive strategy. Many tumorassociated antigens (TAAs) and their epitopes recognized by cytotoxic T cells (CTLs) have been identified during the last two decades and some of them have been used in clinical trials for several cancers. The epitopes have been under investigation for the treatment of cancer, with major clinical responses in some trials. With regard to immunotherapy for HCC, few kinds of TAAs and their epitopes have been used and only clinical data of α-fetoprotein (AFP) have been reported. TAP: In human trials targeting AFP, it is possible to raise an AFP-specific T-cell response using AFP-derived peptides, but this has shown little