compression myelopathy. One month after G-CSF administration, mean recovery rate of JOA score was 29.1%. In contrast, it was 1.1% in the control group at 1 month after initial treatment. In addition, we observed that both motor power and pain sensation scores significantly increased in the G-CSF group compared with the control group at 1 month after treatment. No surgical treatment was performed in patients of either group during the month after G-CSF administration or initial treatment, and they were equally provided conservative treatment such as bed rest. Thus, the present results strongly suggest that G-CSF administration exhibited a neuroprotective effect for the injured spinal cord in patients with worsening symptoms of thoracic myelopathy and improved the myelopathy.

To the best of our knowledge, there has been no other medical treatment that has provided reliable evidence for improvement of thoracic myelopathy. This study provides evidence that G-CSF neuroprotective therapy may be useful as a medical treatment of patients with worsening symptoms of thoracic compression myelopathy. The G-CSF therapy may be especially useful for patients in whom the treatment of complications other than myelopathy needs to be given priority and thus requires a long waiting period before surgery.

In our present trial, no severe side effects occurred. Thus, we suggest that the dose (10  $\mu$ g/kg per d), duration (5 consecutive days), and route (intravenous administration) of G-CSF administration used in this study are principally safe for the treatment of patients with thoracic myelopathy.

The biggest limitation of this study was that the trial was performed as an open-labeled study and the selection of patients to the G-CSF group and the control group was not randomized. We cannot deny the possibility that a placebo effect of injection may participate in the improvement of neurological symptoms. To increase the level of evidence, in the next stage the study design should be a randomized, double-blind placebo-controlled study. By conducting a phase IIb clinical trial in a large number of patients with the study design described earlier, we will be able to reach a better conclusion regarding the effectiveness of G-CSF neuroprotective therapy for patients with worsening symptoms of thoracic compression myelopathy.

# > Key Points

- ☐ A multicenter prospective controlled clinical trial was performed to confirm the feasibility of G-CSF administration for patients with worsening symptoms of thoracic myelopathy.
- For 10 patients with progressive myelopathy, G-CSF (10 μg/kg per day) was intravenously administered for 5 consecutive days.
- ☐ The administration of G-CSF caused neurological recovery in the patients.

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# Accumulation of Activated Invariant Natural Killer T Cells in the Tumor Microenvironment after $\alpha$ -Galactosylceramide-Pulsed Antigen Presenting Cells

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#### Abstract

*Purpose* The intravenous administration of α-Galactosylceramide (α-GalCer)-pulsed antigen presenting cells (APCs) is well tolerated and the increased IFN- $\gamma$  producing cells in the peripheral blood after the treatment appeared to be associated with prolonged survival. An exploratory study protocol was designed with the preoperative administration of α-GalCerpulsed APCs to clarify the mechanisms of these findings, while especially focusing on the precise tumor site.

*Methods* Patients with operable advanced lung cancer received an intravenous injection of  $\alpha$ -GalCer-pulsed APCs before surgery. The resected lung and tumor infiltrating lymphocytes (TILs) as well as peripheral blood mononuclear cells

were collected and the invariant NKT (iNKT) cell-specific immune responses were analyzed.

Results Four patients completed the study protocol. We observed a significant increase in iNKT cell numbers in the TILs and augmented IFN- $\gamma$  production by the  $\alpha$ -GalCer-stimulated TILs.

Conclusion The administration of  $\alpha$ -GalCer-pulsed APCs successfully induced the dramatic infiltration and activation of iNKT cells in the tumor microenvironment.

**Keywords** Invariant NKT cell · antigen presenting cell · immunotherapy · tumor infiltrating lymphocyte · non-small cell lung cancer

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#### Introduction

Vα24 invariant natural killer T (Vα24 iNKT) cells are a unique innate lymphocyte subpopulation characterized by the expression of a canonical invariant T cell receptor with a specific  $\alpha$ -chain gene rearrangement (Vα24-Jα18) and pairing mostly with a Vβ11 β-chain in human. Synthetic glycolipid,  $\alpha$ -Galactosylceramide ( $\alpha$ -GalCer) is a mouse and human iNKT cell ligand, presented by a monomorphic class I-like antigen presenting molecule CD1d [1–3]. Ligand activated iNKT cells exhibit both direct and indirect potent anti-tumor activity.

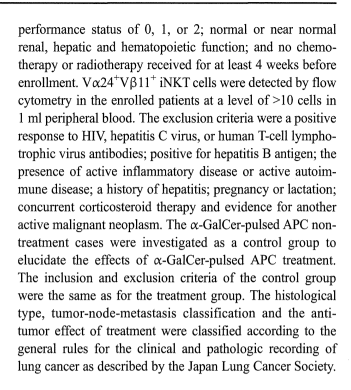
Patients with malignant diseases show either a decreased number or functionally impaired  $V\alpha 24$  iNKT cells in human peripheral blood mononuclear cells (PBMCs) [4-9]. Head and neck cancer patients with poor circulating iNKT cell number show significantly worse clinical outcomes, suggesting an important contribution of iNKT cells to anti-tumor responses [10]. In addition, the ability to produce IFN- $\gamma$  from circulating iNKT cells in cancer patients is preserved even though the absolute number of iNKT cells decreases, and thus, residual iNKT cells might still have a good competence to exert anti-tumor responses. Therefore, the expansion and activation of these cells in vivo may be therapeutically meaningful in patients with severely decreased or functionally deficient Vα24 iNKT cells. Clinical studies of α-GalCerpulsed antigen presenting cells (APC)s have been conducted to recover a functionally sufficient number of  $V\alpha 24$  iNKT cells [11–14]. A phase I/II study ofα-GalCer-pulsed APCs in patients with advanced or recurrent non-small cell lung cancer (NSCLC) found that the treatment elicits  $V\alpha 24$  iNKT cell-dependent immune responses, which are correlated with prolonged overall survival time [13]. The mechanisms that underlie this positive clinical outcome are still unclear.

The current clinical trial focused on the iNKT cell-specific immunological responses in the tumor microenvironments to investigate further anti-tumor mechanisms of  $V\alpha24$  iNKT cells after  $\alpha\textsc{-}GalCer\textsc{-}pulsed$  APC treatment. Therefore, in this exploratory study, the preoperative administration of  $\alpha\textsc{-}GalCer\textsc{-}pulsed$  APCs was performed to clarify the iNKT cell specific immune responses at the tumor site more precisely. The results indicated that  $\alpha\textsc{-}GalCer\textsc{-}pulsed$  APCs successfully induced the activation of tumor infiltrating  $V\alpha24$  iNKT cells in the lung.

#### Material and Methods

# Patient Eligibility Criteria

The study included patients between 20 and 80 years of age, with a diagnosis of clinical stage IIB or IIIA NSCLC that was to be treated surgically. Further inclusion criteria were a



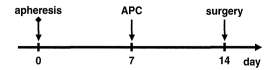
#### Clinical Protocol and Study Design

The study was carried out in the Department of Chest Surgery, Chiba University Hospital, Japan, according to the standards of Good Clinical Practice for Trials on Medicinal Products in Japan. The protocol was approved by the Institutional Ethics Committee (No. 1972). In addition, this trial underwent ad hoc reviews by the Chiba University Quality Assurance Committee on Cell Therapy.

The study design is illustrated in Fig. 1. Written informed consent was obtained from all of the patients before undergoing a screening evaluation to determine eligibility. Clinical and laboratory assessments were conducted once a week, including of a complete physical examination and standard laboratory values. Any adverse events and changes in laboratory values were graded according to the National Cancer Institute Common Toxicity Criteria version 4.0.

# Preparation of APCs from Peripheral Blood

All procedures were carried out according to the Good Manufacturing Practice standards. Eligible patients underwent peripheral blood leukapheresis (COBE Spectra, Gambro BCT,



**Fig. 1** Study design of α-GalCer-pulsed APC administration. The patients received α-GalCer-pulsed APCs. The timing for both apheresis and α-GalCer-pulsed APC administration are shown. APC, α-GalCer-pulsed APC administration



Inc., Lakewood, CO) and PBMCs were collected and further separated by density gradient centrifugation (OptiPrep, Nycomed Amersham, Oslo, Norway). Thereafter, whole PBMCs were cultured with GM-CSF and IL-2, as previously described [11, 15]. Briefly, PBMCs were washed three times and resuspended in AIM-V (Invitrogen Corp., Carlsbad, CA) with 800 units/ml of human granulocyte macrophage colonystimulating factor (GeneTech Co., Ltd., China) and 100 Japanese reference units per milliliter of recombinant human IL-2 (Imunace, Shionogi, Osaka, Japan). The cultured cells were pulsed with 100 ng/ml of specific ligand, α-GalCer (KRN7000; Kirin Brewery, Gunma, Japan) on the day before administration. Whole cells were harvested after 7 days of cultivation, washed 3 times and resuspended in 100 ml of 2.5 % albumin in saline. The patients received an intravenous injection of the cultured cells once (Fig. 1). The criteria for  $\alpha$ -GalCer-pulsed APC administration included a negative bacterial culture 48 h before APC injection, cell viability >70 % and an endotoxin test 48 h before APC injection with a result <0.7 Ehrlich units/ml. The patients were injected with  $1 \times 10^9$  cells/m<sup>2</sup>/injection of APCs.

#### Phenotype Evaluation of APCs

The phenotypes of  $\alpha$ -GalCer-pulsed APCs were determined using a FACSCalibur flow cytometer (BD biosciences). The monoclonal antibodies (mAb) used were FITC-labeled anti-HLA-DR, CD83, CD14; phycoerythrin-labeled anti-CD86, CD1d; and allophycocyanin-labeled anti-CD11c, CD40 (Becton Dickinson, San Diego, CA). Isotype-matched control mAbs were used as negative controls.

Preparation of Tumor Infiltrating Lymphocytes, Tumor Cells, Normal Lung Mononuclear Cells and Lymph Nodes Mononuclear Cells

Fresh tumor tissue specimens were obtained from the surgical specimens and the tissue was cut into small pieces with scissors. The tissue specimen was placed in a flask with a mixture of 0.1 mg/ml DNase type I, 1 mg/ml collagenase type IV and 0.5 mg/ml hyaluronidase type V (all from Sigma, St. Louis, MO) in RPMI 1640 and stirred at room temperature for 1 h. The resultant cell suspension was washed in HBSS and subjected to two-layered (75 and 100 %) Ficoll-Hypaque discontinuous density gradient centrifugation at 1200 g for 20 min. The cells from the 100 % interface and 75 % interface were used as tumor infiltrating lymphocytes (TILs) and tumor cells, respectively. Normal lung tissue and lymph nodes were excised from the surgical specimen, cut with scissors in RPMI 1640 containing enzymes and passed through a gauze filter. The resultant cell suspension was washed in HBSS and subjected to Ficoll-Hypaque gradient centrifugation. The interface was collected and used as either a normal lung or lymph node.

## Immunological Monitoring

PBMC samples were obtained at least twice before APC administration and 1 week after APC injection.

Flow Cytometric Analysis of  $V\alpha 24^{+}V\beta 11^{+}$  Inkt Cells in the Peripheral Blood and TILs

The cell concentrations of  $V\alpha24^+V\beta11^+$  iNKT cells in PBMCs, TILs and mononuclear cells from normal lung tissue or lymph node were assessed by flow cytometry. Mononuclear cells were three-color stained with FITC-conjugated anti–T-cell receptor (TCR)  $V\alpha24$  mAb (C15; Immunotech, Marseilles, France), phycoerythrin-conjugated anti-TCR  $V\beta11$  mAb (C21, Immunotech) and APC-conjugated anti-CD3 mAb (UCTH1; BD Bioscience). The stained cells were subjected to flow cytometry and the percentages of  $V\alpha24^+V\beta11^+CD3^+$  cells among mononuclear cells were calculated. Thereafter, the number of iNKT cells (counts/ml) was estimated based on the PBMC counts.

#### Single-Cell Enzyme-Linked Immunospot Assay

PBMCs, TILs and cells from normal lung tissue or lymph nodes were washed 3 times with PBS and then were stored in liquid nitrogen until use. IFN- $\gamma$ -secreting cells were assayed in 96-well filtration plates (Millipore, Bedford, MA) coated with mouse anti-human IFN- $\gamma$  (10 µg/ml; Mabtech, Nacka Strand, Sweden). The cells (5×10<sup>5</sup> per well) were incubated for 16 h with or without  $\alpha$ -GalCer (100 ng/ml) in 10%FCS containing RPMI. Phorbol 12-myristate 13-acetate (10 µg/ml) plus ionomycin (10 nmol/l) was used as a positive control. After culture, the plates were washed and incubated with biotinylated anti-IFN- $\gamma$  (1 µg/ml; Mabtech). Spot-forming cells were quantified by microscopy.

# Quantitative Real Time PCR of $V\alpha 24$ Invariant TCR and CD1d Expression

Total RNA was extracted from the tumors, normal lung tissue and lymph nodes using TRIzol Reagent (Sigma Aldrich) and reverse transcribed using Superscript II RT (Invitrogen Life Technologies) and oligo (dT12–18) primers (Invitrogen Life Technologies). The primers specific for the constant region of TCR  $\alpha$  chain (C $\alpha$ ) (sense, CGCCTTCAA CAACAGCATTA; antisense, ACCAGCTTGACATCA CAGGA), TCR V $\alpha$ 24 (sense, GCAAAGCTCTCT GCACATCA; antisense, CCAGGGTTGAGCCTCTGTC), CD1d (sense,gtcagggaagtcggaactga; antisense, atcctgagacatggcacacc) were used with 5  $\mu$ g of sample cDNA and



amplified with Taq polymerase (Promega). Quantitative real-time PCR was performed using real-time Taq-Man technology and an ABI PRISM 7000 sequence detector (Applied Biosystems, Foster City, CA). The expression was normalized using the  $C\alpha$  signal for  $V\alpha24$  and GAPDH for CD1d.

#### Statistical Methods

Statistical analyses were performed using Student's *t*-test.

#### Results

#### **Patient Characteristics**

A total of 4 patients met the inclusion criteria and were enrolled in the study. The patient characteristics are summarized in Table I. The study included one patient with adenocarcinoma and three patients with squamous cell carcinoma. Two patients were stage IIb and two were stage IIIa primary lung cancer. No patients had received any previous treatments.

In addition, a total of 6 patients who had not received  $\alpha$ -GalCer-pulsed APC injection were enrolled as the control group. Fresh tumor tissue, normal lung tissue and lymph nodes were excised from the surgical specimens. The patient characteristics of the control group are also listed in Table I.

# Phenotypes of αGalCer-Pulsed APCs

The phenotypes of  $\alpha$ GalCer-pulsed APCs prepared for administration were analyzed by flow cytometry. All profiles for each patient are shown in Fig. 2. The percentages of HLA-DR<sup>+</sup>, CD11c<sup>+</sup>, CD86<sup>+</sup>, CD40<sup>+</sup>, CD83<sup>+</sup> and CD1d<sup>+</sup> cells were determined by the overtone subtraction test using the population comparison platform in the FlowJo software package. More than 50 % of the cultured cells were HLA-DR<sup>+</sup> cells, 10 % to 50 % were CD11c<sup>+</sup> cells and 50 % to

80 % were CD86<sup>+</sup> cells. Interestingly, the majority of the cultured cells were CD3<sup>+</sup> T cells or CD56<sup>+</sup>CD3<sup>-</sup> NK cells, indicating the expression of HLA-DR<sup>+</sup>, CD11c<sup>+</sup> or CD86<sup>+</sup> on human T cells or NK cells (data not shown). Some variations were observed in the expression of CD83 (23.9–55.4 %), CD40 (8.5–17.1 %) and CD1d (23.1–69.9 %; Fig. 2).

#### Adverse Events

No serous (grade >2) toxicity or severe side effects were observed in any patients.

Immunological Monitoring of PBMCs and Resected Specimens

Immunological assays were conducted for all patients. The frequency of peripheral blood  $V\alpha24$  iNKT cells in all patients was measured by FACS. Figure 3 shows that two patients (cases 002 and 004) showed an increased number of circulating  $V\alpha24$  iNKT cells after the  $\alpha$ -GalCer-pulsed APC administration. No clear relationship was found between the number of circulating  $V\alpha24$  iNKT cells and the  $\alpha$ -GalCer-pulsed APC administration in the remaining two patients (cases 001 and 003).

Immunological assays were also performed for TILs and mononuclear cells (MNC)s from normal lung and lymph node tissues. TILs from all 4 cases in the  $\alpha$ -GalCer-pulsed APC administration group contained a high percentage of V $\alpha$ 24 iNKT cells in comparison to the normal lung MNCs (TILs; 1.86 %, 0.32 %, 0.15 % and 0.39 % vs. lung MNCs; 0.031 %, 0.013 %, 0.003 % and 0.01 %, Fig. 4a). The frequency of V $\alpha$ 24 iNKT cells in the TILs in case 001 was 60 times higher than the normal lung MNCs. Though the content of V $\alpha$ 24 iNKT in the normal lung MNCs was extremely low in case 003, the V $\alpha$ 24 iNKT cells were found to have accumulated in the TILs. The average percentage of V $\alpha$ 24 iNKT cells in the TILs was 50 times higher than that

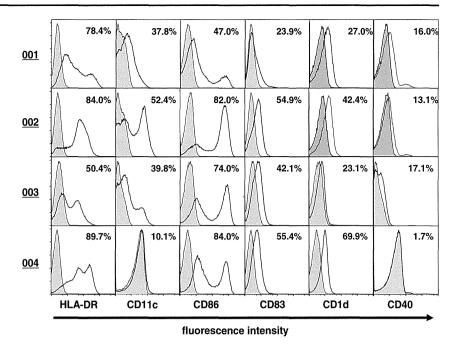
**Table I** Patient characteristics of  $\alpha$ -GalCer-pulsed APC group and control group

Case	Treata	Age/Sex	Histology	c-stage	Operation method
001	APC <sup>b</sup>	75/M	$Ad^d$	T2N1M0 (stage IIB)	Lobectomy+LND <sup>g</sup>
002	APC	76/M	$\mathrm{Sq}^\mathrm{e}$	T2N1M0 (stage IIB)	Lobectomy+LND
003	APC	74/M	Sq	T1N2M0 (stage IIIA)	Lobectomy+LND
004	APC	68/M	Sq	T3N1M0 (stage IIIA)	Pneumonectomy+LND
c-01	cont <sup>c</sup>	71 M	Sq	T2N1M0 (stage IIB)	Lobectomy+LND
c-02	cont	55 M	large <sup>f</sup>	T2N1M0 (stage IIB)	Lobectomy+LND
c-03	cont	70 M	Sq	T3N0M0 (stage IIB)	Lobectomy+LND
c-04	cont	72 M	Sq	T3N1M0 (stage IIIA)	Bilobectomy+LND
c-05	cont	56/M	Sq	T2N1M0 (stage IIB)	Lobectomy+LND
c-06	cont	63/M	Ad	T2N2M0 (stage IIIA)	Lobectomy+LND

<sup>a</sup>Treat, Treatment; <sup>b</sup>APC, α-GalCer-pulsed APC administration; <sup>c</sup> cont, control; <sup>d</sup>Ad, Adenocarcinoma; <sup>e</sup> Sq, Squamous cell carcinoma; <sup>f</sup> large, large cell carcinoma; <sup>g</sup> LND, Lymph Node dissection



Fig. 2 Flow cytometric analysis of  $\alpha$ -GalCer-pulsed APCs. The expression levels of HLA-DR, CD11c, CD86, CD83, CD1d and CD40 were assessed by flow cytometry. Shaded areas: background staining with an iso-type control. Solid lines: staining profiles of the indicated molecules. Values represent the percentages of positive cells



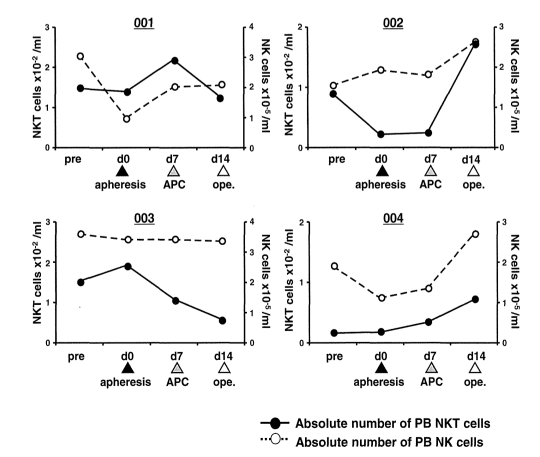
in normal lung MNCs. The  $V\alpha24$  iNKT cell frequency in the draining lymph nodes of each case was almost the same as that in the normal lung MNCs (Fig. 4a).

The proportion of V $\alpha$ 24 iNKT cells in the control group showed a relatively high percentage of TILs in comparison to the normal lung MNCs (TILs; 0.031 %, 0.058 %, 0.13 %, 0.47 %, 0.18 % and 0.12 % vs. lung MNCs; 0.034 %, 0.011 %, 0.004 %, 0.039 %, 0.014 % and 0.02 %,

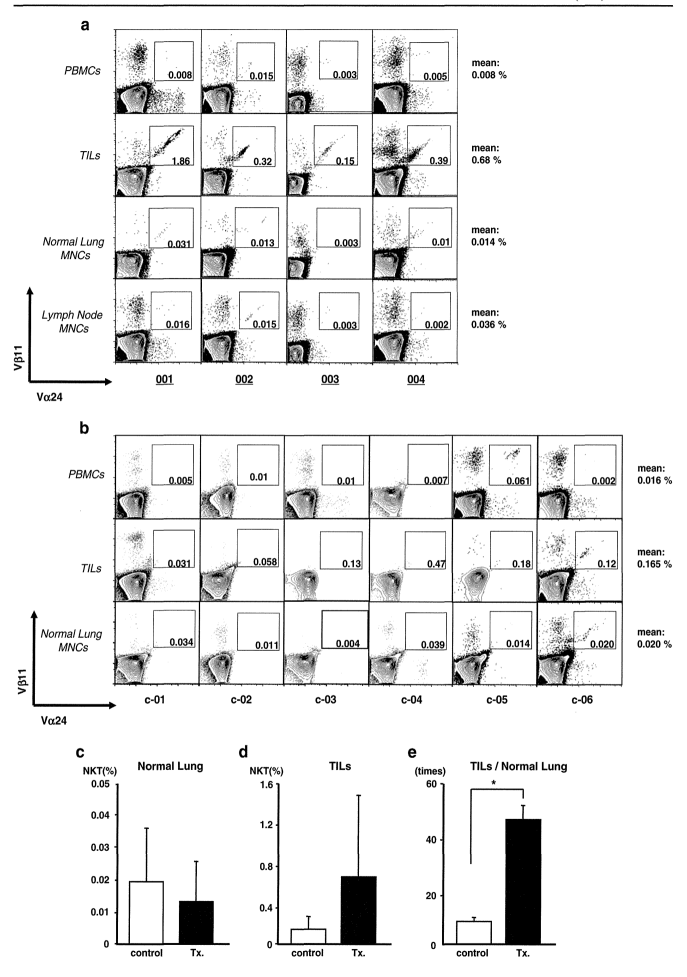
Fig. 4b). The average percentage of  $V\alpha24$  iNKT cells in the TILs was only 8 times higher than that in the normal lung MNCs.

Normal lung MNCs in the control group demonstrated a trend toward a higher  $V\alpha24$  iNKT cell rate in comparison to the treatment group (Fig. 4c). On the other hand, the proportion of  $V\alpha24$  iNKT cells in TILs tended to increase in the  $\alpha$ -GalCer-pulsed APC injected group in comparison to the

Fig. 3 Immunological monitoring of PBMCs of patients with α-GalCer-pulsed APC administration. The absolute number of peripheral blood iNKT cells  $(V\alpha 24^{+}V\beta 11^{+} \text{ cells})$  and NK cells (CD56<sup>+</sup>CD3<sup>-</sup> cells). Flow cytometric analysis and automated full blood counts (Chiba University Hospital) indicated the absolute number of Vα24 iNKT cells and NK cells. APC, α-GalCer-pulsed APC administration; ope., operation



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▼ Fig. 4 The frequencies of Vα24 iNKT cells in TILs and mononuclear cells from normal lung and lymph node tissues. a, The proportion of Vα24 iNKT cells (Vα24<sup>+</sup>Vβ11<sup>+</sup> cells) in PBMCs on day 14, TILs, normal lung MNCs and lymph node MNCs in theα-GalCer-pulsed APC administration group were assessed by flow cytometry. The lymph node MNCs column depicts one representative MNC profile in the draining lymph nodes including hilar (#10, 11 and 12) and mediastinal (#1, 3, 4 and 7) nodes. b, The proportion of Vα24 iNKT cells in PBMCs, TILs and normal lung MNCs in the control group were assessed by flow cytometry. c-d, The comparison between the Vα24 iNKT cell contents in normal lung MNCs c and TILs d in the α-GalCer-pulsed APC treatment group and the control group. e The TIL/Normal Lung MNC ratio of Vα24 iNKT cell proportion. control, control group; Tx, α-GalCer-pulsed APC administration group; \*p=0.0008

control group (mean percentage, 0.68 % and 0.165 %, Fig. 4d). The V $\alpha$ 24 iNKT cell ratio of TILs/normal lung in the  $\alpha$ -GalCer-pulsed APC administration group was significantly higher than that of the control group (p=0.0008, Fig. 4e).

The number of IFN-γ-producing cells after restimulation with α-GalCer in vitro was concurrently monitored in PBMCs, TILs, normal lung MNCs and lymph node MNCs using an ELISPOT assay. An analysis of the PBMCs and resected specimen showed the highest value of α-GalCerresponsive IFN-γ-producing cell number in the TILs of the  $\alpha$ -GalCer-pulsed APC treated group (Fig. 5a). The absolute number of α-GalCer-responsive IFN-γ-producing cells in the TILs was apparently high in cases 001 and 002, whereas a relatively low value was seen in cases 003 and 004. This observation was not detected with the use of control group specimens (Fig. 5b). Together with the results in Fig. 4, the administration of α-GalCer-pulsed APCs induced the mobilization of endogenous  $V\alpha 24$  iNKT cells into the primary site of the lung cancer and augmented the IFN-γ-producing ability of tumor infiltrating  $V\alpha 24$  iNKT cells.

In addition, the number of IFN- $\gamma$ -producing cells in PBMCs was determined after restimulation with  $\alpha$ -GalCer

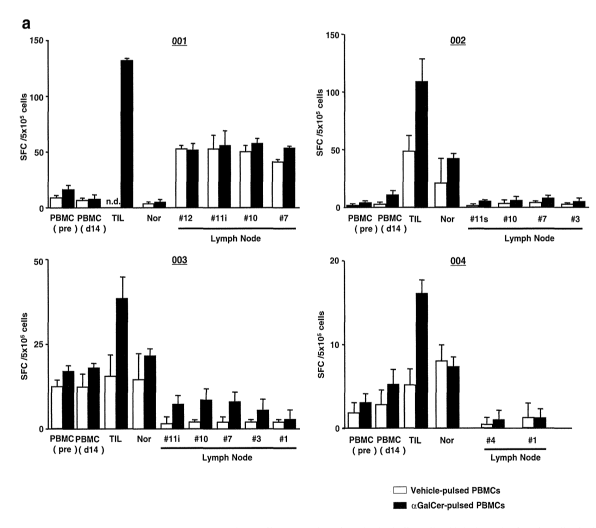
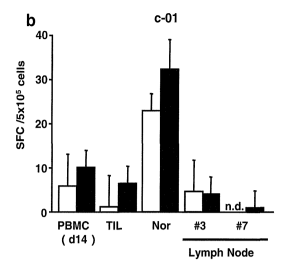
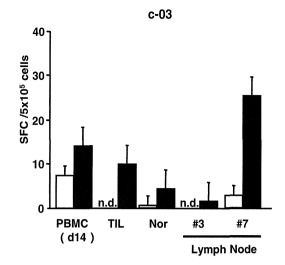


Fig. 5 Detection of α-GalCer-reactive IFN- $\gamma$ -producing cells by enzyme-linked immunospot assay. a Cryopreserved PBMCs, TILs, normal lung MNCs and lymph node MNCs of the α-GalCer-pulsed APC treated group were thawed and cultured overnight with either α-GalCer or vehicle. The presence of IFN- $\gamma$ -producing cells was quantified by an enzyme-linked immunospot assay. The resected draining

lymph nodes including hilar (#10, 11 and 12) and mediastinal (#1, 3, 4 and 7) nodes are shown. Spot number of IFN- $\gamma$  with standard deviation for triplicate culture of 4 cases are shown. **b** Spot-forming cell number in the control group. SFC, Spot Forming Cell; pre, pretreatment; d14, day 14; Nor, normal lung MNCs; n.d., not done; #11 s, lymph node #11 superior; #11i, lymph node #11 inferior







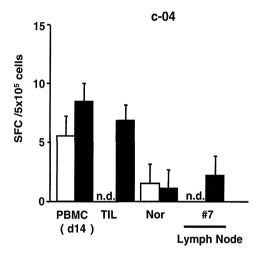


Fig. 5 (continued)

in vitro. In Fig. 5a, the number of IFN- $\gamma$  producing cells in PMBC increased 14 days after treatment in case 002 and 004, indicating that global NKT cell activation in these patients.

The mRNA Expression Level of  $V\alpha 24$  Inkt Cell Receptor and CD1d

Quantitative RT-PCR was performed to further confirm the increase in the number of V $\alpha$ 24 iNKT cells. The mRNA from the primary tumor, normal lung tissue and lymph node samples was obtained from the  $\alpha$ -GalCer-pulsed APC treated patients and the relative expression level of V $\alpha$ 24 invariant TCR and the constant region of TCR (C $\alpha$ ) mRNA were evaluated. The mRNA for V $\alpha$ 24 TCR was highly expressed in the tumor (Fig. 6a). The V $\alpha$ 24-J $\alpha$ 18 invariant TCR mRNA could not be detected in the normal lung tissue since the lung parenchyma included mainly alveolar epithelial cells and only a low copy number of C $\alpha$  mRNA was detected. The relative gene expression of CD1d was also evaluated by quantitative RT-PCR. CD1d expression was ascertained both in normal lung tissue and tumor tissue in all

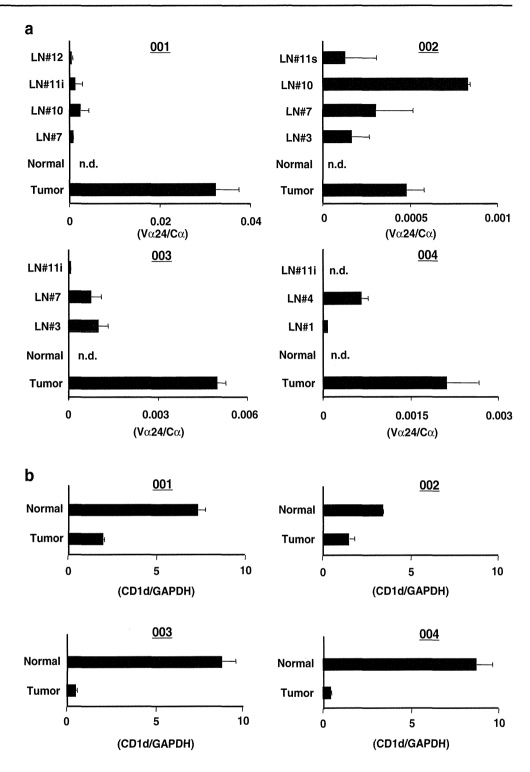
4 cases and the expression level appeared to be higher in normal lung tissue (Fig. 6b).

#### Discussion

The major aim of this study was to investigate the  $V\alpha24$  iNKT cell-specific immune responses in the primary tumor site after the intravenous injection of  $\alpha$ -GalCer-pulsed APCs in patients with advanced NSCLC. The development of these immunotherapeutic approaches with the administration of  $\alpha$ -GalCer-pulsed APCs requires a thorough knowledge of local immune responses.  $V\alpha24$  iNKT cells accumulate in lung cancer lesions [5], as observed in this report (Fig. 4e). A significant increase in the tumor infiltrating  $V\alpha24$  iNKT cell population was detected after the administration of  $\alpha$ -GalCer-pulsed APCs in comparison to the non-injected control group. This observation is quite reasonable for the iNKT cell-targeted therapy aimed at activation of  $V\alpha24$  iNKT cells in the tumor located site in vivo. The activation status of CTLs, rather than just the existence



Fig. 6 The relative mRNA expression of the  $V\alpha 24^+$  TCR in tumor, normal lungs and lymph nodes. Cancer tissue, non-cancerous lung tissue and lymph nodes were obtained from α-GalCer-pulsed APC treated patients. a The expression level of Vα24 TCR mRNA in each sample was analyzed by quantitative RT-PCR. Each mRNA was quantified by the standard curve method and copy numbers of Vα24 TCR were normalized by the copy number of the constant region of the TCR  $\alpha$  chain ( $C\alpha$  mRNA. **b** The expression level of CD1d mRNA in each sample was analyzed by quantitative RT-PCR. Each mRNA was quantified by the standard curve method and copy numbers of CD1d were normalized by the copy number of the GAPDH mRNA. LN, lymph node; Normal, normal lung tissue; Tumor, tumor tissue; n.d., not detected; #11 s, lymph node #11 superior; #11i, lymph node #11 inferior



of CTLs, has great prognostic significance [16–18]. Therefore, IFN- $\gamma$ -producing cells were also monitored in PBMCs, TILs and MNCs from normal lung tissue and lymph nodes using an ELISPOT assay. After starting the protocol, standard preoperative chemo-radiotherapy was introduced to treat locally advanced NSCLC, and such a change of treatment strategy hindered the entry of patients into this protocol. In spite of the limited number of patients analyzed, the results obtained by ELISPOT assay indicated that tumor infiltrating mononuclear cells had augmented IFN- $\gamma$ 

producing capacity, which may have a positive impact on the tumor microenvironment.

Tumor infiltrating lymphocytes are found in a variety of cancers and they are thought to be a result of a host immune response directed against tumor cells. Several reports have shown that the presence of large numbers of tumor-infiltrating CD8<sup>+</sup> T cells are associated with a favorable prognosis in esophageal carcinoma [18, 19], colorectal cancer [20, 21], ovarian cancer [22], and pancreatic carcinoma [23], while the infiltration of CD4<sup>+</sup> T cells that posses regulatory function,



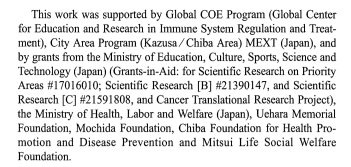
such as Foxp3<sup>+</sup> regulatory T cells (Treg) are associated with the poor prognosis of ovarian cancer [24, 25]. The balance between CD8<sup>+</sup> CTLs and Tregs in tumors is critical for disease progression and survival [24, 26, 27]. These diverse results indicate that the functional roles of TILs are complicated and uncertain and the effects of TILs might vary with the type and stage of cancers. In addition to CD8<sup>+</sup> cytotoxic T cells, tumor infiltrating  $V\alpha24$  iNKT cells have been reported to be a positive prognostic factor for colorectal carcinoma [28]. The current results indicated that the injection of  $\alpha$ -GalCer-pulsed APCs could induce the accumulation of  $V\alpha24$  iNKT cells in TILs, which would therefore lead to a good prognosis after a complete surgical resection.

Although a complete surgical resection is regarded as the optimal treatment for NSCLC, only around 25 % of NSCLC are suitable for potentially curative resection. Despite optimal surgical management, the 5-year survival rate of resected NSCLC ranges between 85.9 % for pathological stage Ia and 41 % for pathological stage IIIa [29]. Approximately 50 % or more of patients with NSCLC who undergo surgery experience relapse due to the existence of microscopic lesions that could not be detected by preoperative screening. Recently, adjuvant chemotherapy given after surgery has been shown to improve survival [30-32]. A meta-analysis suggested that cisplatin-based adjuvant chemotherapy could yield an absolute overall survival advantage of 5 % at 5 years [33]. At the same time, chemotherapeutic agents often show severe toxic effects and it was reported that in patients with early-stage disease have deleterious effects on long-term survival. This emphasizes the importance of development of less-invasive preoperative or postoperative therapy to suppress the growth of micrometastases. Therefore, immune cells for tumor surveillance, such as NK and iNKT cells, which possess anti-tumor activity, should be beneficial and post-surgical adjuvant immunotherapy by the use of these cells may be favorable since the residual tumor is quite small after a complete resection.

#### **Conclusions**

 $\alpha\text{-}GalCer\text{-}pulsed$  APC administration successfully induced the dramatic infiltration and activation of  $V\alpha24$  iNKT cells in the tumor lesion. This report is the first clinical trial of  $V\alpha24$  iNKT cell targeted immunotherapy that shows a functional  $V\alpha24$  iNKT cell accumulation in the tumor microenvironment. These results encourage the further development of immunotherapy aimed at the activation of endogenous  $V\alpha24$  iNKT cells in the lung.

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**Conflict of interest** The authors declare that they have no conflict of interest

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# 薬理と治療(JPT)

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● パネルディスカッション3/ これからの時代を担う IRB の機能と責務-中央 IRB(共同 IRB を含む)の普及に向けて-/講演 4

# 中央 IRB 等への移行過程で生じた課題と その解決に向けた取り組み

花岡英紀 青柳玲子 松本和彦 吉澤弘久 小池竜司 大学病院臨床試験アライアンス推進室および中央 IRB 検討ワーキンググループ

> **』 ፪●**፮ **፮**፮፮፮ ライフサイエンス出版

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【禁 無断転載・複製】

第11回CRC と臨床試験のあり方を考える会議 2011 in 岡山/ 2011年9月25日・岡山コンベンションセンター

# ● パネルディスカッション3/

これからの時代を担う IRB の機能と責務ー中央 IRB (共同 IRB を含む) の普及に向けて一/講演 4

# 中央 IRB 等への移行過程で生じた課題と その解決に向けた取り組み

花 岡 英 紀<sup>1)</sup> 青 柳 玲 子<sup>1)</sup> 松 本 和 彦<sup>2)</sup> 吉 澤 弘 久<sup>3)</sup> 小 池 竜 司<sup>4)</sup> 大学病院臨床試験アライアンス推進室および中央 IRB 検討ワーキンググループ

大学病院臨床試験アライアンスではいわゆる中央IRBのあり方について検討を行うとともに、千葉大学、信州大学、 新潟大学による中央IRBの審議を2010年7月より試験的に一試験について開始した。本稿では大学病院臨床試験 アライアンスで実施している中央IRB検討ワーキンググループ(以下WG)およびその上部組織である推進室で の議論、さらに欧州の中央IRBのあり方をふまえたIRBのあるべき姿と、実際の取り組みについて述べるもので ある。また、この内容については、筆者らの私見にもとづいており、各大学でのコンセンサスを得られた段階に至っ ていない。

#### はじめに

近年、治験の活性化の中で、中央IRBに向けた取り組みの必要性が謳われている。これは、多くの場合、治験を開始するにあたり、それぞれのIRBへの申請に関して多くの時間と労力を必要とするため、その業務に当たる費用が高額となり、治験依頼者への負担が増しているという事実がある。また、一方で、経験の乏しいIRBにおいて複雑な治験の審議をすることについて問題視する事実もある。このような状況のなか、GCPでは2度の改訂を経ているにもかかわらず、いまだ中央IRBの活動は活発ではない。平成23年6月には日本医師会治験促進センターによる治験等適正化作業班による提言がなされた。作業班では、中央IRBの設置とIRB事務局と治験事務局の外部委託による効率化も検討された。

大学病院臨床試験アライアンスは、国際共同試験をわが国で主導することを目的として関東信越7大学において先進的な取り組みを治験において実施してきた。そのなかで中央IRBについては、はたして実施にあたりどのような課題があるか具体的に検討

が必要との結論に至り、推進室の下部組織として信州大学を担当校とし各大学の担当者からなるWGを構成し議論を重ねてきた。

## WGの検討内容について

そもそも中央IRBの設置の前提としていくつかの 疑問がある。治験を実施する各施設個別の状況を知 らない中央IRBにおいて、①迅速性と効率性と、科 学性と倫理性は共に成り立つにはどのようにしたら いいのか、②被験者を保護するという原則はどのよ うにしたら確保されるのか、という2課題はWGの 検討以前においてそもそも重要な問題であり、IRB の審議の根幹を成すものである。中央IRBで十分な 議論ができる体制を維持すること、さらにそこでは、 その施設固有の問題点を被験者保護という立場から 話し合える状況にする必要があると考えている。

一方で、③審査手続きの共通化、④中央IRBの設置場所、⑤具体的な審議の方法、という3課題については、技術的な側面が強い課題であり、解決可能な課題とした。

## 欧米の中央IRBのあり方について

海外には中央IRBがすでに多く設置され、活用さ

<sup>1)</sup> 千葉大学 2) 信州大学 3) 新潟大学 4) 東京医科歯科大学

英国	3 機関の承認が必要
	1.NHS National Health Service 行政区ごとの local Research Ethical
	Committee (REC)
	2.Main REC
	3.MHRA (Medicines and healthcare products Regulatory Agency)
フランス	被験者保護委員会 CCP
	被験者保護に関する事項
	臨床試験の <u>プロトコルデザイン</u> に関する事項
	施設に関する事項
	医療製品保険安全局 AFSSAPS
	試験薬等の品質と安全性に関する事項
	被験者の安全性に関する事項(適格基準、投与量、被験者のモニタリング。
	対照薬)
米国	All committees are registered with the Office for Human Research Protections
	(OHRP)
	FDA, pharmaceutical companies, contract research organizations, as well as
	AAHRPP perform regular inspections of our processes.
	An FDA inspection in July 2009 resulted in no findings.
	CERTIFICATION: Staff and board members have earned certifications, such
	as Certified IRB Professional (CIP), Certified Clinical Research Professional
	(CCRP) and Certified IRB Manager (CIM) .

# 図1 欧米の中央 IRB に関する大学病院臨床試験アライアンスによる訪問調査

		IRB 審議資料	
IRB 開催日	千葉大	新潟大	信州大
2010/7/20	新規実施の適否	新規実施の適否 (治験参加カード)	新規実施の適否 (治験参加カード)
2010/9/21	実施計画書 2.0 版 同意説明文書	実施計画書 2.0 版	実施計画書 2.0 版
2010/10/18		同意説明文書	
2010/11/15	実施計画書 2.1 版 同意説明文書 監査手順書 2.0 版	実施計画書 2.1 版 同意説明文書 監査手順書 2.0 版	実施計画書 2.1 版 同意説明文書 監査手顧書 2.0 版
2011/2/21	院内の重篤有害事象	-devil	west.
2011/3/22	継続審査、安全性情報	継続審査、安全性情報	継続審査、安全性情報
2011/4/18	実施計画書 2.2 版 治験薬概要書 2.0 版 同意説明文書	実施計画書 2.2 版 治験薬概要書 2.0 版	実統計画會 2.2 版 治験薬概要書 2.0 版
2011/5/16	同意説明文書	同意説明文書	同意説明文書
2011/9/20	実施計画書 2.3 版 同意説明文書 緊急逸脱報告書	実施計画書 2.3 版 同意説明文書	実施計画書 2.3 版 同意説明文書

# 図3 3大学から千葉大学中央 IRB へ提出され審議された内容

れている国々があるが、その設置の基準や役割はさまざまである。海外の状況(⑥)を検討するためにわれわれは実際に規制当局や医療機関を訪問し、調査を行った(図1)。このなかで、施設固有の問題点を審議するローカルIRBと計画書の審議をする中央IRBがそれぞれ業務を分担し審議を行っていることを知った。さらに、規制当局との連携も重要なるであった。また、米国ではコマーシャルIRBが数多く設置されているが、それぞれのIRBではその質の高さと独立性を示すための取り組み(認定の取得、業務内容の公開、FDAの調査を受けた時期の公開)を行っていることが明らかになったが、一方で施設固有の問題点を審議することについては難しい状況であることが判明した。

# 医師主導治験の中央IRBについて

2010年7月千葉大学医学部附属病院治験審査委員

千葉大学. 新潟大学. 信州大学で実施中の中央IRB体制 7 | | 1) 治験実施の適否についての審査 中央IRB (千葉大IRB) にて審議 (施設固有事項含む) ③④ 2) 治験の継続についての審査 (重篤な有害事象発生時など) ①②③④(信州) 億州大 ···· 1) 施設問右直項 ③資料を提出し 聖香依頼 2) 有害事象報告書 信州大学が必要と判断した場合 ①審查結果通知 ①審查依賴! ②審查結果通知 ③専門IRBの審査結果 \*信州大学では病院側の判断で 必要ならば専門IRBの意見を 聴くことができる。その審認 結果を中央IRBに報告する。 その審査 信州大IRB:専門IRB 千葉大IRB:中央IRB

図2 WGで検討された中央 IRB のフロー

会では、信州大学医学部附属病院長および新潟大学 医歯薬総合病院長から千葉大学医学部附属病院長へ の医師主導治験の審議依頼を受け審議を開始した。 これは国立大学病院としては初めての取り組みであ る。この審議を実施するにあたり3大学ではWGで 検討された中央IRBの審議のフロー(図2)をもと にそれぞれ手順書の改訂(⑦)契約書の作成とこれ に伴う審査費用の算定(⑧)を行った。ここでは申 請元の大学のIRBを「施設で発生した重篤有害事象 を審議する専門IRB」と位置づけている。(⑨)3大学 から中央IRB事務局へ提出される書類は内容がすべ て同一でないことより中央IRBで承認された内容や 時期が異なっている(図3)。このことは治験事務 局と中央IRB事務局との独立性を意味し、推進する 立場とこれを監督する立場のぞれぞれの事務局の独 立性が確保された状況となっている。(⑩) 医師会の 作業班の提案する各大学の治験事務局を廃止し中央 に委託することについては、作業の効率性に加え、 技術的に千葉大学の事務局が各大学の業務を引き受 けることは困難との結論に至った。

#### まとめ

およそ10の課題について検討を行いながら,具体的に中央IRBを運用するに至った。規制当局との連携については今までまったく議論をされていな内容である。つまり,その本質において単なる効率化ではなく真の意味での中央IRBの役割を考えることが重要であり,欧米での取り組みを十分考慮のうえ,進めて行く必要がある。大学病院臨床試験アライアンスではわが国を代表するグループとして欧米からも信頼される中央IRB体制の構築を実施する。

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# <sup>[原著]</sup> 脊髄障害性疼痛に対する顆粒球コロニー 刺激因子(G-CSF)の治療効果

加 藤 啓 Ш 崎 正 志 大 河 昭 彦 佐久間 毅 本 光 子 高 橋 宏 橋 宏 林 浩 III辺 紬 藤 崇 之 矢 丈 雄 内 友 規 領 由 古 Ш 田 宫 下 智 大 萬納寺 誓 X 染 谷 幸 男 鎌 田 尊 X 紀1) 修 将 行 池 田 橋 本 井 上 雅 俊 花 岡 英 国府田 正 雄 高 橋 和 久

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# 要旨

脊髄障害性疼痛を有する13例を対象として、顆粒球コロニー刺激因子(G-CSF)を用いた神経保護療法の臨床試験を施行した。対象症例は、脊髄症の急性増悪と関連する疼痛を認めた 6 例(脊髄症急性増悪群)および脊髄症の後遺症と考えられる疼痛を認めた 7 例(脊髄症後遺症群)の二群に分けられた。G-CSFを 5  $\mu$ g/kg/日×5日間(1 例)、 $10\mu$ g/kg/日×5日間(11例)、 $10\mu$ g/kg/日×3日間(1 例)の投与量・期間で点滴静注投与し、投与前後での疼痛の変化を Visual analogue scale (VAS) にて評価した。11例で投与後に疼痛が軽減し、2 例では完全に消失した。疼痛軽減効果は全例、投与開始から 1 週間以内で認められた。脊髄症後遺症群では無効例が 2 例存在した。 VASは、脊髄症急性増悪群で投与前平均63.3mmから投与後 1 週間で平均25.0mm(P<0.05)に、脊髄症後遺症群で投与前平均67.1mmから投与後 1 週間で平均44.3mm(P<0.05)に減少した。投与後 6 ヵ月間の観察期間中に 7 例で疼痛軽減効果の減弱が見られた。効果の減弱は投与後 3 ヵ月時に 4 例、6 ヵ月時に 3 例で認められた。今回の検討から、G-CSFが脊髄障害性疼痛に対して疼痛軽減効果を有することが示唆された。

Key words: 脊髓障害性疼痛,顆粒球コロニー刺激因子,神経保護療法

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# I. 緒 言

脊髄障害性疼痛とは、2009年以降に本邦で使用 されるようになった新しい概念である。厳密な定 義は定められてはいないが、ヘルニア・脊柱靭帯 骨化症などによる圧迫性脊髄症や、急性脊髄損傷 などの脊髄障害に起因すると考えられる多彩な痛 みを総称している[1,2]。侵害受容性疼痛に比べ 神経障害性疼痛の占める割合が大きいのが特徴で あり、通常では痛みを起こさない程度の刺激が強 い痛みを引き起こすアロディニア (異痛症)や. 障害脊髄高位より尾側の締め付けられるような自 発痛などに代表される。非ステロイド性抗炎症薬 (NSAIDs) の効果は低く、最近では末梢神経障 害性疼痛で得られたエビデンスをもとに、プレガ バリンなどの抗てんかん薬や一部の抗うつ薬など の有効性が解明されてきたが[3-5],多くの症例 で必ずしも十分な除痛効果が得られるわけではな い。病態が十分に解明されておらず、症例によっ て薬剤に対する反応性が異なるため、治療に難渋 することがしばしばである。本邦では平成22年度 に厚生労働省により脊髄障害性疼痛症候群が研究 奨励分野に指定され、研究が進められている。

我々は、2008年から急性期脊髄損傷患者およ び圧迫性脊髄症急性増悪患者に対する顆粒球コ ロニー刺激因子(Granulocyte-colony stimulating factor: G-CSF) を用いた神経保護療法の臨床試 験を進めている[6-9]。G-CSFは血球系に作用す る増殖因子であり、顆粒球系細胞の分化・増殖・ 生存促進などの作用を有する[10]。本邦ではがん 化学療法による好中球減少症や、末梢血幹細胞移 植時の造血幹細胞の末梢血への動員などの目的で 臨床使用されている。中枢神経系においては、骨 髄細胞を脳・脊髄中へ動員する作用や[11], 脳卒 中モデルに対する神経保護作用などが報告されて おり[12],海外では脳梗塞に対する臨床試験が報 告されている[13]。以上より、脊髄損傷に対して もG-CSFが治療効果を発揮しうる可能性が想定 されたため、われわれはマウスおよびラットの 脊髄損傷モデルを用いてG-CSFの有効性および その作用機序につき報告してきた[14-18]。その 結果. G-CSF投与により後肢運動機能が有意に 改善した。その機序に関しては、①G-CSFによ

り動員された骨髄由来幹細胞が脊髄損傷部に生 着する,②直接的に神経細胞死を抑制する,③ Oligodendrocyteの細胞死を抑制し、髄鞘を保護 する、4炎症性サイトカイン (TNF- $\alpha$ , IL-1 $\beta$ ) 発現を抑制する,⑤血管新生を促進する,などが 想定された[14-18]。これらの根拠から、慢性圧 迫性脊髄症の急性増悪に対してもG-CSFが神経 保護作用を有する可能性が示唆されたため、圧迫 性脊髄症急性増悪患者に対する治療薬としての G-CSFの安全性・有効性を確認するため臨床試 験を開始した[6,7]。結果として, G-CSFの安全 性が確認され、全例で神経症状の改善が認められ た。さらに、予期せぬ効果として、G-CSF投与後 に脊髄障害に起因する疼痛が軽減した例を少なか らず経験した[19]。このことからG-CSFが脊髄 障害性疼痛に対して何らかの疼痛軽減効果を有す ることが示唆された。今回は、臨床試験例におけ るG-CSF投与後の脊髄障害性疼痛の変化を検討 した。

# Ⅱ. 方 法

2009年1月から2010年2月の期間に、脊髄障害 に関連する疼痛を認めた16例に対し、G-CSFを 用いた神経保護療法の臨床試験を施行した。今回 は、その中で投与から6ヵ月以上経過観察しえた 13例を対象として検討した(表1)。症例1~6 では、後縦靭帯骨化症(OPLL)および黄色靱帯 骨化症 (OYL) による圧迫性脊髄症の急性増悪 (日本整形外科学会頚髄症治療判定基準 [JOAス コア]が直近の1ヵ月で2点以上悪化)に対し てG-CSFの投与が行われた。この6例での脊髄 障害性疼痛は、脊髄症の急性増悪との関連が強く 示唆されたため、脊髄症急性増悪群として分類し 検討を進めた。G-CSFの投与量・期間は、症例 1で5μg/kg/日×5日間,症例2~6で10μg/ kg/日×5日間の点滴静注であった。6例全例で、 G-CSF投与後1.5~8週で脊髄症に対する後方除 圧固定術または椎弓切除術が行われた。

症例7では、頚椎症性脊髄症に伴うと考えられる疼痛が持続しており、症例8~13では、脊髄症に対する手術の術後に脊髄関連の疼痛が遺残していた。したがって、症例7~13を脊髄症後遺症群

表 1 脊髄障害性疼痛に対する G-CSF 投与症例

症例 No.	年齢 (歳) /性別	診断	脊髄症 急性 増悪	責任 高位	G-CSF 投与量 (µg/kg/日)	G-CSF 投与 期間	手術術式 a [範囲]	G-CSF 投与 ~手術
	, imp		- E / C/	n distribut di spognogia ja ja kanana menden distribut di berikti sekti sekti sekti sekti sekti sekti sekti sek	(μg/ ng/ μ/	(日)	otorio minimi in in 1888 il 1990 paga paga paga paga paga paga paga pag	(週)
1	68/男	後縦靱帯骨化症	+	T4/5	5	5	後方除圧固定 [T1-8]	1.5
2	46/男	後縦靱帯骨化症	+	T6/7	10	5	後方除圧固定 [T2-11]	8
3	75/男	後縦靭帯骨化症	+	C4/5	10	5	後方除圧固定 [C2-7]	7
4	64/男	後縦靭帯骨化症	+	C4/5	10	5	後方除圧固定 [C2-7]	2
5	32/男	後縦靱帯骨化症	+	T8/9	10	5	後方除圧固定 [T7-12]	4
6	67/男	黄色靭帯骨化症	+	T11/12	10	5	椎弓切除 [T10-12]	5
7	73/男	頚椎症性脊髄症		C4/5	10	5		><=>400000000000000000000000000000000000
8	58/男	後縦靭帯骨化症術後		C5/6	10	5	one-politic	·
9	72/男	脊髓円錐症候群術後	_	T12/L1	10	5		-
10	71/男	後縦靭帯骨化症術後		C5/6	10	5	destin	none.
11	78/男	黄色靱帯骨化症術後	Name of the last o	T10/11	10	5	NEWSON-	and the same of th
12	70/女	後縦靭帯骨化症術後	_	C4/5	10	5	(disprime	
13	38/女	胸髄腫瘍術後		T11/12	10	3	-torona-	4004

a: G-CSF 投与後の手術,T: Thorachic vertebrae (胸椎)/C: Cervical vertebrae (頚椎)

と分類して検討を進めた。G-CSFの投与量・期間は,症例  $7 \sim 12$ で $10\mu g/kg/H \times 5$  日間の点滴静注であった。症例13でも同一条件でG-CSFの投与を開始したが,3 日間の投与後に尿路感染に伴う発熱が生じたため,投与が中止となった。

疼痛部位と障害脊髄の位置関係から,脊髄障害性疼痛を2型に分類した。すなわち,障害髄節高位の痛みおよび障害髄節から2髄節以内の領域の痛みをat-level pain、髄節より3髄節以上尾側の領域の痛みをbelow-level painと定義した。疼痛の程度は,Visual analogue scale(VAS: $0\sim100$ mm)を用いて評価した。脊髄症の評価には,JOAスコア(頚髄症: $0\sim17$ 点,胸髄症: $0\sim11$ 点)を用いた。上下肢の筋力に評価には,American Spinal Injury Association(ASIA)運動スコア( $0\sim100$ 点),上下肢体幹の感覚の評価にはASIA 痛覚スコア( $0\sim112$ 点)を用いた。

統計学的評価にはフィッシャーの正確確率検定とウィルコクソンの符号付順位検定を用い, P<0.05を有意差ありとした。

# Ⅲ. 結果

脊髄症急性増悪群 6 例における疼痛は、at-level painが 4 例、below-level painが 2 例であり、有痛期間は $0.2 \sim 4$ 年(平均1.5年)であった(表2)。脊髄症後遺症群 7 例における疼痛は、at-level painが 3 例、below-level painが 4 例であり、有痛期間は  $1 \sim 27$ 年(平均10.9年)であった(表2)。脊髄症後遺症群における有痛期間が、有意に長かった(P < 0.01)。

脊髄症急性増悪群では6例全例で10mm以上のVASの低下が認められ、投与後1週間以内に除痛効果が得られていた(表3A)。完全に疼痛が消失した例も1例(症例5)存在した。VASは投与前平均63.3mmから投与後1週間で平均25.0mmに減少した(P<0.05)。除痛効果は3例で投与後3ヵ月時に減弱が見られ、1例(症例3)では投与直前の疼痛に戻った。投与後6ヵ月の時点でのVASは平均33.3mmであった(P<0.05)。

脊髄症後遺症群では7例中5例で10mm以上の VASの低下が認められた(表3B)。完全に疼痛 が消失した例も1例(症例11)認められた。無

表 2 脊髄障害性疼痛の性状

症例 No.	疼痛の分類	疼痛の部位	疼痛の性質	有痛期間 (年)
1	below-level	両下肢全体	しびれ感の強い痛み	0.7
2	at-level	胸背部	絞め付けられるような痛み	0.8
3	below-level	両下腿	しびれ感の強い痛み	3
4	at-level	右手	しびれ感の強い痛み	0.3
5	at-level	背部 右胸部	しびれ感の強い痛み 刺すような痛み	4
6	at-level	両鼠径部	しびれ感の強い痛み	0.2
7	below-level	両足底部	しびれ感の強い痛み	13
8	at-level	左手指	しびれ感の強い痛み	1
9	below-level	両足底部	刺すような痛み	3
10	at-level	両手指 両下肢全体	しびれ感の強い痛み	3
11	below-level	両大腿内側	しびれ感の強い痛み	19
12	at-level	両前腕以下	灼熱感を伴う痛み	27
13	below-level	左下肢全体	しびれ感の強い痛み	10

表3A G-CSF投与後のVASの変化(脊髄症急性増悪群)

,it had			VAS	VAL AT A PARA DESCRIPTION OF THE PARA DESCRIPTION OF T	VAS減少值a				
症例 - No.	投与 直前	1週	1ヵ月	3ヵ月	6カ月	1週	1ヵ月	3ヵ月	6カ月
1	50	25	25	25	25	25	25	25	25
2	60	30	30	30	20	30	30	30	40
3	50	30	30	50	50	20	20	0	0
4	50	25	25	45	45	25	25	5	5
5	80	0	0	0	0	80	80	80	80
6	90	40	40	60	60	50	50	30	30
平均	63.3	25.0*	25.0*	35.0*	33.3*	38.3	38.3	28.3	30.0
± SD	± 17.5	±13.4	$\pm 13.4$	$\pm 21.4$	$\pm 22.3$	$\pm 22.9$	$\pm 22.9$	$\pm 28.4$	$\pm 28.8$

VAS:  $0 \sim 100$ mm

表 3 B G-CSF 投与後の VAS の変化 (脊髄症後遺症群)

.i mi			VAS	4.,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	VAS減少值 a				
症例 No.	投与 直前	1 週	1ヵ月	3ヵ月	6ヵ月	1週	1ヵ月	3ヵ月	6ヵ月
7	70	70	70	70	70	0	0	0	0
8	50	40	40	40	50	10	10	10	0
9	90	90	90	90	90	0	0	0	0
10	80	50	50	50	50	30	30	30	30
11	60	0	0	60	60	60	60	0	0
12	60	20	20	20	30	40	40	40	30
13	60	40	40	40	60	20	20	20	0
平均	67.1	44,3*	44.3*	52.9	58.6	22.9	22.9	14.3	8.6
± SD	±13.8	± 29.9	$\pm 29.9$	$\pm 22.9$	$\pm 18.6$	$\pm 22.1$	$\pm 22.1$	$\pm 16.2$	$\pm 14.6$

VAS:  $0 \sim 100$ mm

a: 投与直前のVASからの減少値

<sup>\*:</sup> 投与直前に比し有意に減少(P<0.05)

a: 投与直前のVASからの減少値

<sup>\*:</sup> 投与直前に比し有意に減少(P<0.05)

改善率 (%) JOAスコア(点) 症例 No. 投与1ヵ月前 投与直前 1ヵ月後 6ヵ月後 1カ月後 6ヵ月後 1 8.5/11 1.5/11 6/11 8/11 47.4 68.4 2 7.5/115.5/116.5/1110/11 18.2 81.8 3 16/17 8.5/17 14.5/17 14.5/17 70.6 70.6 4 14/17 9.5/1714.5/17 14.5/17 66.7 66.7 5 6/11 6/11 28.6 28.6 4/11 6/11 6 35.7 6/11 4/11 6.5/11 6.5/1135.7 平均 44.5\* 58.6\*  $\pm$  SD  $\pm 21.0$  $\pm 21.3$ 

表4A G-CSF投与後のJOAスコアの変化(脊髄症急性増悪群)

JOA スコア: 日本整形外科学会頚髄症治療判定基準(頚髄症:  $0 \sim 17$ 点、胸髄症:  $0 \sim 11$ 点)\*: 脊髄症後遺症群(表 4 B)と比べ有意に改善(P < 0.01)

症例		JOAスコラ		改善率 (%)		
No.	投与1ヵ月前	投与直前	1ヵ月後	6ヵ月後	1ヵ月後	6ヵ月後
7	13/17	13/17	13/17	13/17	0	0
8	5/11	5/11	6/11	6/11	16.7	16.7
9	5.5/11	5.5/11	5.5/11	5.5/11	0	0
10	4.5/17	4.5/17	8/17	8/17	28	28
11	4/11	4/11	6/11	6/11	28.6	28.6
12	11/17	11/17	11/17	11/17	0	0
13	3/11	3/11	3/11	3/11	0	0
平均	244444444	**********			10.5	10.5
± SD					± 13.6	± 13.6

表4B G-CSF投与後のJOAスコアの変化(脊髄症後遺症群)

JOAスコア: 日本整形外科学会頚髄症治療判定基準(頚髄症: 0~17点,胸髄症: 0~11点)

効例が 2 例(症例 7 , 9 ) あり,両者とも below-level pain の例であった。有効例では全例,1 週間以内に除痛効果が得られていた。 VAS は投与前平均67.1mmから投与後 1 週間で平均44.3mmに減少した(P<0.05)。一方,除痛効果は 1 例で投与後 3 ヵ月時に,3 例で投与 6 ヵ月時に減弱が見られ,経過観察期間中に 3 例で投与直前の疼痛に戻った。投与後 3 ヵ月の時点での VAS は平均52.9mmと再度上昇していた。

JOAスコアは脊髄症急性増悪群の6例全例でG-CSF投与後に増加が見られた(表4A)。投与後1ヵ月および6ヵ月時の改善率はそれぞれ平均44.5%,58.6%であった。急性増悪前と比較しても遜色ない程度まで回復していた。脊髄症後遺症群では7例中3例でJOAスコアの増加を認めたが、投与後1ヵ月および6ヵ月時の改善率はいずれも平均10.5%に過ぎなかった(表4B)。除痛効果のなかった2例(症例7,9)では、JOAスコ

アの改善は得られていなかった。

脊髄症急性増悪群のASIAスコアは、運動、痛 覚ともに投与後1ヵ月および6ヵ月時で、投与 直前と比べ有意に改善を認めた(表5A)。一方、 脊髄症後遺症群のASIAスコアは改善を認めな かった(表5B)。無効例の2例(症例7,9)では、 運動、痛覚スコアともに、増加点数は0であった。

末梢血中の白血球数は投与開始後1日目から有意に増加した(表6)が、投与終了後速やかに減少し、1週間の時点では投与前の値に戻っていた。白血球の分画中、顆粒球が著明に増加していたが、単球の増加も有意であった。CRPも有意に増加していたが、軽度の増加であり臨床的に問題となる値ではなかった。リンパ球数は投与前後で変化を認めなかった。投与期間中、および投与後に重篤な有害事象の発生は認めなかった。