

Figure 5.7. In vitro effect of nitric oxide (NO) generated from SIN-1 on expression of adhesion molecule, CD44, in LM-2 melanoma cell line. After LM-2 melanoma cells were incubated with various concentrations of SIN-1, NO generator for 4 hours, they were washed and stained with anti-CD44 antibody conjugated with FITC. Then their fluorescence intensity was measured by using Flow cytometer. Values are means \pm SD; significantly different from the culture with medium only (* $P < 0.05$).

lished in epidemiological and experimental studies. It is known that patients either with newly diagnosed cancers or with metastases or who are undergoing therapy have statistically significant less blood, urine, and hair Se than age- and sex-matched healthy controls. It seems that Se supplementation is important for cancer patients because Se has an ability to enhance various humoral and cellular immune responses.

The ability of C57BL/6J mice, maintained for 8 weeks on a Se-deficient (0.02 ppm Se), normal (0.20 ppm Se), or Se-supplemented (2.00 ppm Se) diet, to generate cytotoxic lymphocytes (CTL) and to destroy tumor cells was examined. Lymphocytes from mice fed the Se-supplemented diet had a greater ability to destroy tumor cells than those from mice fed the normal diet, whereas Se deficiency reduced the cytotoxicity (Roy et al. 1990). In addition, it is proposed that the enhancement of in vivo cytotoxicity of NK or CTL following Se supplementation is likely to act synergistically with tumor growth inhibition in the reduction of tumor incidence (Fig. 5.8) (Petrie et al. 1989). It seems that Se's direct effects on host defense cells to suppress or enhance their actions as well as effects on tumor cells occur in the promotion phase (Talcott et al. 1984).

Zinc is also an important nutrient for maintaining cellular immune functions. Zinc status was determined in patients with newly diagnosed squamous cell carcinoma of the oral cavity, oropharynx, larynx, and hypopharynx. In this case, patients with metastatic disease and with severe comorbidity were excluded. Results showed that approximately 50% of the subjects were zinc-deficient based on cellular zinc criteria and had decreased production of Th1 cytokines but not Th2 cytokines, decreased NK cell activity, and decreased

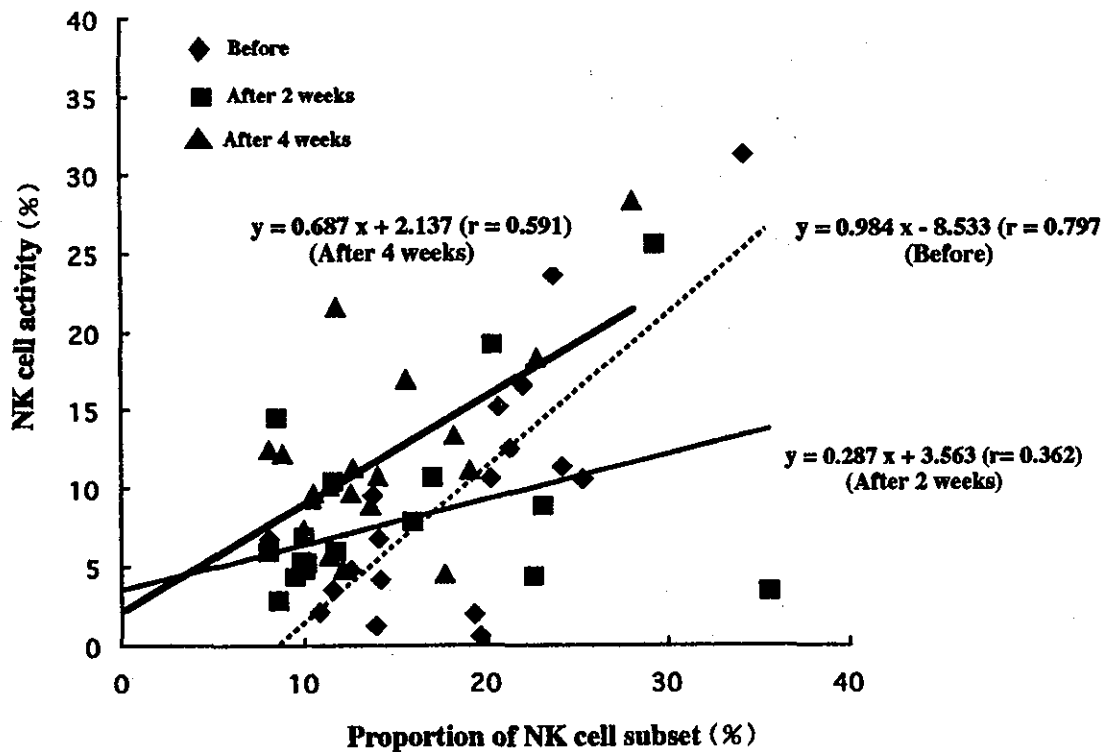


Figure 5.8. Changes in the activity and the proportion of natural killer (NK) cells in peripheral blood lymphocytes (PBL) of subjects with Aojiru drinking for 4 weeks. Nineteen female university students were selected as subjects (aged 20 to 22) in this study. They daily drank 90 ml of Aojiru juice for 4 weeks. At 2 and 4 weeks, peripheral blood was taken and their PBL were isolated. NK activity in the vertical axis and proportion of NK cell subset in the horizontal axis were plotted. As shown in this figure, NK activity of subjects with lower proportion of NK cell subset was largely enhanced following Aojiru drinking.

proportion of $CD4^+CD45RA^+$ cells in the peripheral blood (Prasad et al. 1998). Zinc concentration in polymorphonuclear cells was also decreased in the hospitalized subjects (Goode et al. 1991).

ROLE OF SOME FOODS IN CANCER PREVENTION AND IMMUNITY

Both epidemiological and animal studies have found anticarcinogenic potential in garlic and its constituent compounds. A review on a historical perspective on garlic and cancer was recently published (Milner, 2001). This review shows that water- and lipid-soluble allyl sulfur compounds have an ability to block experimentally induced tumors in a variety of sites including skin, breast, and colon, which mechanism is related to changes in DNA repair and immunocompetence. *In vitro* effects of garlic derivative (alliin) on both peripheral blood mononuclear cell (PBMC) proliferation and cytokine production induced by the mitogen were examined and increases in pokeweed mitogen (PWM)-induced lymphocyte proliferation, and interleukin (IL)-1 beta and tumor necrosis factor (TNF)- α productions were found (Salman et al. 1999). Lamm and Riggs (2001) have also reviewed the antitumor effect of garlic and described that the immune stimulation of garlic is able to

reduce the incidence of cancer. In addition, it has been also reported that aged garlic extract (AGE) significantly inhibits the growth of Sarcoma-180 (allogenic) and LL/2 lung carcinoma (syngenic) cells transplanted into mice and increases natural killer (NK) and killer activities of spleen cells in Sarcoma-180-bearing mice (Kyo et al. 2001).

In Japan, 40,000 households periodically purchase and drink Aojiru. Aojiru is named from its color and is a juice prepared from kale, which is a plant related to cabbage. The beneficial effect of Aojiru was investigated by measuring the activity of natural killer (NK) cells, which play an important role in protecting the body from bacteria and viral infections, and in excluding transformed cells and suppressing carcinogenesis (Cooley et al. 1999, Hirose and Kuroda 1999, and Baraz et al. 1999). NK activity of splenocytes from rats fed the freeze-dried Aojiru supplemented diet was about three times higher than that of control rats (Moriguchi and Muraga 2000).

To investigate its mechanism, the effect of Aojiru drinking on NK activity of peripheral blood lymphocytes of young female university students was examined. NK activity following Aojiru drinking for 4 weeks was significantly increased. As shown in Figure 5.8, the enhancement of NK activity is due not to increased proportion of NK cells but to increased activity of NK cell per se (Ogawa et al. 2001). Measuring cytokines in serum of these subjects, there was a significant increase of interleukin-2 (IL-2), inducing the activation of NK cells, after 4 weeks of Aojiru drinking (Fig. 5.9).

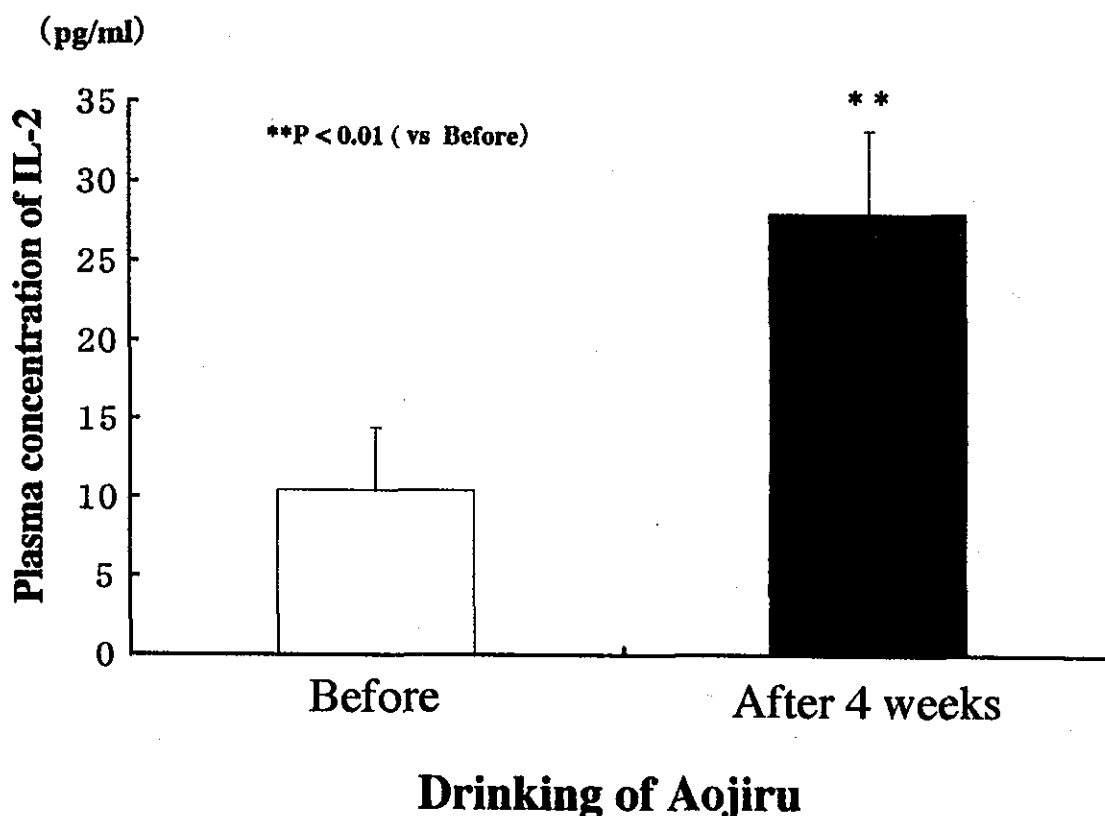


Figure 5.9. Plasma concentration of IL-2 in female university students following Aojiru drinking for 4 weeks. Values are means \pm SD ($n = 19$); significantly different from plasma concentration of IL-2 of subjects before Aojiru drinking (** $P < 0.01$).

These results suggest that the intake of some foods having immunoenhancing effect is beneficial for preventing carcinogenesis and maintaining health.

CONCLUSION

Nutritional supplementation is important for both maintenance of host immune function for perioperative cancer patients and the suppression of cancer incidence and promotion.

As described in this chapter, there are many nutrients having immunoenhancing effects. Some of them act directly to inhibit tumorigenesis and tumor growth. Even if cancer patients fall into the immunodeficient status following malnutrition, enteral or parenteral nutrition for supplying adequate nutrition can improve their nutritional status and immune functions and result in prolonging their survival time. In addition, to protect tumorigenesis in our body simply and with certainty, we have two options: one is not eating foods with possible carcinogens and the other is eating foods with immunoenhancing effects described in this chapter.

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ビタミンと免疫[†]

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はじめに

多くのビタミンが種々の栄養素の代謝の中で酵素の働きを助ける補酵素として重要な役割を担っており、それらビタミンの欠乏は細胞内での栄養素代謝の破綻をもたらし、その結果として免疫能を含む細胞機能の低下を誘導する。さらに、ビタミン欠乏は単独で起こることは稀であり、一般にたん白・エネルギー栄養不良 (PEM; Protein-energy malnutrition) に付随してみられる¹⁾ (表1)。また、入院患者においても単一あるいは2、3のビタミン欠乏が混在していることも知られている²⁾ (表2)。個々のビタミン欠乏時にはほとんどの免疫能の低下することが知られている³⁾ (表3)。本稿ではビタミンを脂溶性ビタミン (ビタミンA, D, EおよびK) と水溶性ビタミン (ビタミンB群とC) に分け、宿主免疫能への影響についてこれまでの研究成果について要約した。

1 脂溶性ビタミン

脂溶性ビタミンの中で免疫能との関連でこれまで多くの研究報告がみられるビタミンはAおよびEである。ビタミンDに関しては若干の報告はあるものの未だ十分ではない。さらに、ビタミンKに関しては免疫能に対する作用は低いと考えられている。

表1 たん白質・エネルギー栄養不良に伴うビタミン欠乏の発生頻度

ビタミン	重篤な栄養不良		中程度の栄養不良	
	調査数	欠乏 (%)	調査数	欠乏 (%)
ビタミンA	13/29	45	11/37	30
カロテン	32/33	97	25/32	78
葉酸	5/33	15	7/36	19
ビタミンC	0/20	0	0/19	0
ビタミンB ₁	12/28	43	18/27	67
ビタミンB ₂	5/31	16	3/24	13
ビタミンB ₆	12/34	35	8/42	19

表2 米国の入院患者にみられるビタミン欠乏の頻度

ビタミン	欠乏者の割合 (%)
ビタミンA	13
ビタミンE	12
葉酸	45
ビタミンC	12
ビタミンB ₁	31
ビタミンB ₂	12
ビタミンB ₆	27
ナイアシン	29
パントテン酸	15
ビタミンB ₁₂	10
ビオチン	1
2種の欠乏	38
3種の欠乏	14
4種の欠乏	6
5種の欠乏	10

[†] Vitamins and Immunity

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表3 種々のビタミン欠乏と免疫能 (抜粋)

ビタミン	免疫能の変化
ビタミンB ₆	・ジフテリア毒素に対する抗体産生の低下 ・SRBCに対する抗体産生細胞数の低下 ・リンパ球混合培養反応の低下
パントテン酸	・サルモネラ菌に対する抗体価の低下 ・SRBCに対する抗体産生細胞数の低下
ビタミンB ₁	・ラットにおけるヒト赤血球に対する抗体価の低下
ビタミンB ₂	・ラットおよびブタにおけるヒト赤血球に対する抗体価の低下
ビオチン	・ジフテリアに対する二次抗体価の低下
ビタミンB ₁₂	・TおよびB細胞数は正常 ・PHAに対するヒト末梢血リンパ球幼若化能の低下
ビタミンA	・ジフテリア毒素に対する抗体反応の低下 ・遅延型過敏反応の低下 ・末梢血T細胞数の低下とPHAに対する反応低下
ビタミンC	・ツ反に対する感受性発達を阻害 ・皮膚移植片の拒絶反応の低下 ・胸腺由来液性因子の産生低下

(1) ビタミンA およびカロテノイド

一般にビタミンA欠乏状態ではT細胞の成熟の場である胸腺の萎縮がみられ、細胞性ならびに体液性免疫ともに抑制され、細菌に対する易感染性や発ガン物質投与によるガン発生の増加が認められている^{4, 5)}。また、一方では高ビタミンA摂取により細胞性免疫や抗体産生の亢進することが見出されており、感染抵抗性の増大やマクロファージの殺腫瘍活性亢進を介した移植腫瘍に対する増殖抑制作用が報告されている^{6, 7)}。例えば、マウスの皮膚ガン発生に対する高ビタミンA摂取の効果をみた実験では、発生腫瘍数や腫瘍重量が著明に低下することを認めている(表4)。この機序としてマクロファージ数の増加や殺腫瘍活性亢進との関連が示唆されている。特にB₁₆メラノーマ細胞に対するマクロファージの殺腫瘍活性が食餌

中のビタミンA含量の増加に依存して高くなる傾向がみられている⁸⁾。また、マクロファージ活性化因子(MAF)を用いてマクロファージを活性化したところ、基礎食群のマクロファージでは約35%の殺腫瘍活性の上昇を認めた。これに対し、高ビタミンA食群では食餌中のビタミンA含量が増えるに伴い、逆にマクロファージのMAFに対する反応性は低下する傾向がみられている。このことは高ビタミンA食摂取によりT細胞が活性化され、活性化されたT細胞から産生されたMAFによりマクロファージが既に活性化されていることを示唆している。さらに、ビタミンAとの*in vitro*培養によってもラット肺胞マクロファージ(AM)の殺腫瘍活性が亢進されること⁹⁾や、遺伝的に胸腺が欠損したヌードマウスにおいても高ビタミンA食投与によりマクロファージ数の増加やオプソニン化羊赤血球(SRBC)に対する貪食能の有意な亢進を認めたこと¹⁰⁾から、ビタミンAによるマクロファージ機能の亢進が活性化Tリンパ球から産生されるサイトカインの一つであるMAFによるものと、Tリンパ球を介さず、ビタミンAそのものが直接作用する可能性があることが考えられる¹¹⁾。図1にマクロファージとNK細胞活性化の機序を要約した。換言すると、ビタミンAによるT細胞活性化を介して産生されるMAF、IL-2およびIFN- γ などのサイトカインによって二次的にマクロファージやNK細胞が活性化される系と、ビタミンAにより直接的に活性化される系とが存在する可能性が考えられる。しかし、ビタミンAの過剰摂取は皮膚の落屑、脱毛、筋肉痛や妊婦では胎児の奇形などの副作用をあらわすことが知られており¹²⁾、その摂取には注意を要する。一方、ビタミンAの前駆物質として知られる β -カロテンやその他のカロテノイドもまたビタミンAと同様に免疫賦活作用を有しており、しかもビタミンAとは異なり大量摂取によっても副作用はほとんど出現しないことが知られている¹³⁾。カロテノイド摂取により、好中球やマクロファージなどの食細胞機能の亢進、NK細胞活性の上昇、細胞障害性Tリンパ球機能の亢進などがみられ、これらによって移植腫瘍細胞の増殖抑制や発ガン抑

制が誘導されることが報告されている¹⁴⁾。

表4 マウス皮膚癌発生に対するビタミンAパルミチン酸 (RP) および13-cisレチノイン酸 (13cRA) の影響

実験群	食餌中RP又は13cRA含量 (IU/kg diet)	マウス当りの腫瘍数	マウス当りの腫瘍重量 (g)
基礎食	3,500	15.6±2.7	1.373±0.20
RP	60,000	14.3±2.6	0.20±0.04
	200,000	8.2±2.0	0.131±0.03
	700,000	3.4±1.2	0.007±0.001
13cRA	200,000	13.0±1.9	0.312±0.07
	700,000	14.8±3.0	0.049±0.01

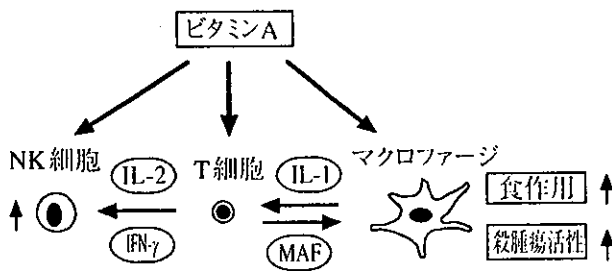


図1 ビタミンAによるマクロファージおよびNK細胞活性化の機序

(2) ビタミンE

ビタミンEは自然界にはトコフェロールと側鎖に二重結合を有するトコトリエノールとがあり、各々にクロマン環に結合したメチル基数の違いにより α -、 β -、 γ -および δ -の4つのタイプ、計8種の同族体が存在している。ビタミンEと言えば一般に α -トコフェロールのことを指し、抗酸化活性や体内含量が同族体の中では最も高いことが知られている。ビタミンE欠乏では免疫系だけでなく生殖機能を含め種々の組織・器官の機能と形態が損なわれる¹⁵⁾。免疫系では抗体産生能、リンパ球増殖能、好中球の走化性や殺菌能など広範囲にわたる免疫低下を誘導することが知られている。但し、動物実験ではビタミンE欠乏時に肺胞マクロファージ貪食能が亢進するという報告もある¹⁶⁾。一方、高ビタミンE食投与は体液性および細胞性免疫能を亢進し、生体防御能を高めること

が知られている¹⁷⁾。特に、細胞性免疫能の低下した高齢者では血中VLDLコレステロール当りのビタミンEレベルと末梢血リンパ球幼若化能との間に有意な正相関のあることが見出されている(図2)。さらに、ビタミンE補足により高齢者の低下した免疫能が改善されることが報告されている。この機序としてアラキドン酸から合成され、免疫抑制作用を有するPGE₂産生をビタミンEが抑制することとの関連が示唆されている。その他、ビタミンEには骨髄で産生された未熟T細胞が分化・成熟する場である胸腺の機能を高める作用のあることも見出されている¹⁸⁾。これらの作用の多くはビタミンEの抗酸化作用に帰するものであるが、同じく抗酸化作用を有する2-メルカプトエタノール(2-ME)投与ではビタミンE投与と同様の免疫能改善がみられないことから抗酸化作用それだけでは説明できないビタミンE固有の作用の存在が考えられている。

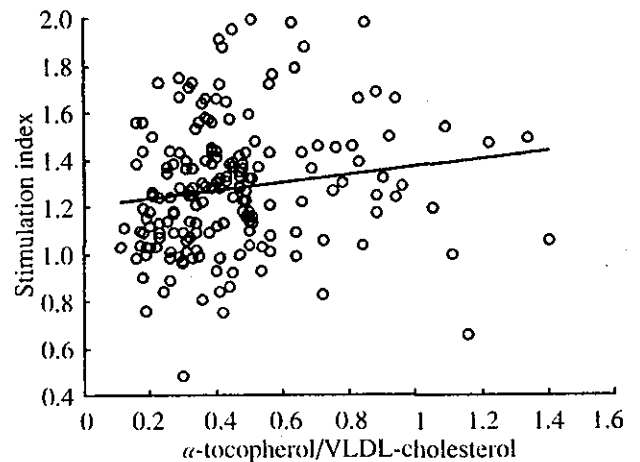


図2 末梢血リンパ球幼若化能と血中ビタミンE濃度との関係

(3) ビタミンD

ビタミンDはきのこや魚類に含まれるほかに、皮膚において紫外線によりプロビタミンDから合成される。このビタミンDが欠乏すると子どもではくる病、成人では骨軟化症を発症することが知られている。ビタミンDと免疫能との関連についての研究はこれまで比較的少なく、その中でくる病の子どもの好中球機能をみたものがある。ビタ

ミンD欠乏状態であるくる病においても好中球の殺菌能は保持されていたが、貪食活性は低下することが認められている¹⁹⁾。また、動物実験ではビタミンD欠乏により抗体産生能の低下することが見出されており、宿主免疫能を保持するうえで十分なビタミンD摂取が必要である。ビタミンDの補足効果については、ビタミンAとの併用によりイースト菌に対する貪食能が相乗的に充進されることも報告されている²⁰⁾。

II 水溶性ビタミン

(1) ビタミンB群

ビタミンB₁、B₂、B₆、B₁₂、葉酸およびパントテン酸などのビタミンB群と免疫能との関連についての研究は比較的少なく、しかもそのほとんどがこれら各ビタミンの欠乏時における免疫能について検討したものである。また、ヒトではこれらビタミン欠乏が単独で起こることはほとんどないことから、これまでの研究成果のほとんどが動物実験によるものである。いずれのビタミン欠乏においてもTおよびBリンパ球機能の低下が認められている。その中でも特にアミノ酸や核酸代謝に関与するビタミンB₆欠乏時にはTおよびBリンパ球数の減少をはじめとする顕著な免疫低下の生じることが知られている²¹⁾。いずれの場合も欠乏しているビタミンを補足することによりそれら機能が回復することから、ビタミンB群に関しては免疫を保持するうえで少なくとも欠乏にならない程度の摂取が望まれる。また、ビタミンB群の補足による免疫変化に関してはビタミンB₁およびB₆補足により食細胞の走化性が充進されたり²²⁾、ナイアシン補足によりヒト末梢血リンパ球やマウス脾臓リンパ球の増殖能が充進されること²³⁾が知られている。

(2) ビタミンC

ビタミンCは多くの動物ではグルコースから酵素的に合成されるが、ヒトをはじめとする霊長類やモルモットでは合成酵素が欠損しているために、体外より食事として経口的にビタミンCを摂取する必要がある。ビタミンCの欠乏症状としては壊

血病が知られている。ビタミンCが免疫の分野において脚光を浴びたのはノーベル賞を2度受賞したポーリング博士の「ビタミンC大量投与が風邪やガン発生を予防する」という発言に端を発する²⁴⁾。しかし、ビタミンCによる免疫賦活作用は図3に示すごとく、所要量(100mg/L)の10倍を越えるビタミンCを毎日連続して摂取している場合には末梢血リンパ球増殖能の充進が誘導されるが、一旦、ビタミンC摂取を止めると1週間後には充進していた免疫能が摂取前のレベルにまで戻り、さらに1日2gを越えるビタミンC摂取をしてもさらに高い免疫能の充進は誘導されない²⁵⁾。これら結果はビタミンC補足によって風邪等の予防を図るうえでのビタミンCの量や摂取方法に対して示唆を与えるものである。また、ビタミンCは抗酸化ビタミンの中で唯一の水溶性ビタミンであり、その作用の一つとして同じく有力な抗酸化ビタミンであるビタミンEが活性酸素等により酸化された場合、その還元作用によりもとの抗酸化能を有するビタミンEに回復することが見出され、

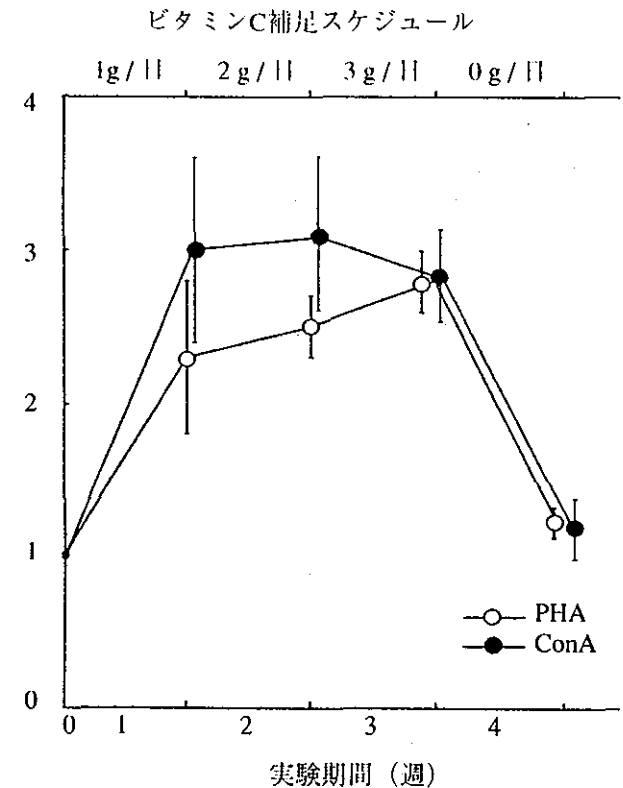


図3 末梢血リンパ球幼若化能に対するビタミンC大量投与の影響

注目されている²⁰⁾。このことは、ビタミンC補足による免疫賦活作用がビタミンC固有の作用だけ

ではなく、ビタミンEとの共同作用によって免疫能が亢進される可能性を支持するものである。

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Characterization of Corrinoid Compounds from a Japanese Black Tea (Batabata-cha) Fermented by Bacteria

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Characterization of Corrinoid Compounds from a Japanese Black Tea (Batabata-cha) Fermented by Bacteria

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A Japanese fermented black tea (Batabata-cha) contained a considerable amount of vitamin B₁₂ (456 ± 39 ng per 100 g dry tea leaves and 2.0 ± 0.3 ng per 100 mL of tea drink). A corrinoid compound was partially purified and characterized from the tea leaves. The patterns of the purified compound by the silica gel 60 thin-layer chromatography and C18 reversed phased high-performance liquid chromatography were identical to those of authentic vitamin B₁₂. When 20 week old vitamin B₁₂ deficient rats, which excreted substantial amounts (about 250 mg/day) of methylmalonic acid in urine as an index of vitamin B₁₂ deficiency, were fed the tea drink (50 mL/day, 1 ng of vitamin B₁₂) for 6 weeks, urinary methylmalonic acid excretion (169 ± 29 mg/day) of the tea drink-supplemented 26 week old rats decreased significantly relative to that (250 ± 32 mg/day) of the deficient rats. The results indicate that the vitamin B₁₂ found in the fermented black tea is bioavailable in mammals.

KEYWORDS: Vitamin B₁₂; cobalamin; corrinoid; tea; vitamin B₁₂ deficient rat; methylmalonic acid; hepatic vitamin B₁₂

INTRODUCTION

In manufacturing processes of black teas fermented by bacteria, such as Pu'erh tea, tea leaves are heat-treated with steam or roasting and then fermented with certain naturally occurring bacteria (1). Thus, they are completely different from the types of self-oxidized black teas (Keemun tea and Darjeeling tea, etc.). These black teas fermented by bacteria, which are found in some Asian countries, may contain various vitamins and/or biofactors synthesized by the concomitant bacteria.

Vitamin B₁₂ (B₁₂) is synthesized only in certain bacteria (2). Usual dietary sources of B₁₂ are known to be animal products but not plant products (3). If the fermented black teas contain considerable amounts of B₁₂, the black tea would contribute to human B₁₂ needs, especially for vegetarians.

Here, we describe the partial purification and characterization of a corrinoid compound from Japanese fermented black tea (Batabata-cha) leaves and also investigated the effect of feeding the tea drink on the B₁₂ status of B₁₂ deficient rats.

MATERIALS AND METHODS

Materials. Cyano-B₁₂ was obtained from Wako Pure Chemical Industries (Osaka, Japan). A B₁₂ assay medium for *Lactobacillus delbrueckii* subsp. *lactis* (formerly *Lactobacillus leichmannii*) ATCC7830

was obtained from Nissui (Tokyo, Japan). Silica gel 60 thin-layer chromatography (TLC) aluminum sheets were obtained from Merck (Darmstadt, Germany). All other reagents used were of the highest purity commercially available. Japanese fermented black tea (Batabata-cha) leaves and the tea drink were provided by Asahi, Ltd. (Toyama-city, Japan).

A UV-1600 UV-vis spectrophotometer (Shimadzu, Kyoto, Japan) was used for measuring the turbidity of the *L. delbrueckii* test culture in the microbiological method. A fully automated ACS 180 chemiluminescence B₁₂ analyzer (Chiron Diagnostics, East Walpole, MA) was used for the B₁₂ assay.

Extraction of Corrinoid Compounds for the Determination of B₁₂ Content of the Tea Leaves. The dried tea leaves (5 g) were powdered by a food mill and then suspended in 50 mL of 0.25 mol/L acetate buffer, pH 4.8, containing 0.2% (w/v) KCN as cyanation for stabilization. The total corrinoids were extracted from the suspension by boiling for 60 min at 98 °C in the dark. The suspension was centrifuged for 10 min at 5000g, and the supernatant was used for the B₁₂ assay.

Concentration of the Fermented Black Tea Drink. For the determination of B₁₂ in the tea drink, B₁₂ was concentrated with a Sep-Pak Vacc 20 cm³ (5 g) C18 cartridge (Waters Corp., Milford, MA). After the C18 cartridge was washed with 75% ethanol and equilibrated with distilled water, an aliquot (50 mL) of the drink was put on the cartridge. B₁₂ was eluted with 50 mL of 25% ethanol, and the eluate was evaporated to dryness under reduced pressure, dissolved in 1.0 mL of distilled water, and used for the B₁₂ assay.

Assay of B₁₂. B₁₂ was assayed by the microbiological method with *L. delbrueckii* subsp. *lactis* ATCC 7830 and a B₁₂ assay medium (Nissui) and by the chemiluminescence B₁₂ analyzer with intrinsic factor (IF) as described previously (4). The above B₁₂ extract and the concentrated drink were directly applied to the chemiluminescence

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analyzer. They were diluted with distilled water to a B₁₂ concentration range of 0.01–0.1 µg/L and used as samples for the microbiological method.

Purification of Corrinoid Compounds from the Fermented Black Tea. About 0.5 kg of the dried tea leaves was powdered by a model MX-X51-H food mill (National, Osaka, Japan), and suspended in 2 L of 0.25 mol/L acetate buffer, pH 4.8. KCN was added to the suspension at the final concentration of 10 mmol/L. Total B₁₂ was extracted from the suspension under the same conditions described above. The boiled suspension was centrifuged at 10 000g for 10 min. The supernatant was used for purification of corrinoid compounds. Amberlite XAD-4 resin (500 g) washed with 5 L of methanol and equilibrated with distilled water was added to the supernatant fraction and stirred for 3 h at room temperature in the dark. The resin suspension was passed through a glass funnel (Buchner type) with a type 25G1 glass filter (Iwaki, Tokyo, Japan), and the resin was washed with 5 L of distilled water. One liter of 80% methanol solution was added to the washed resin, and the suspension was stirred for 3 h at room temperature in the dark. The resin suspension was passed through the glass funnel. The 80% methanol eluant (about 1 L) containing corrinoid compounds was pooled and evaporated to a final volume of 20 mL under reduced pressure. The solution was put on a 24 mm × 70 mm column of Cosmosil 140C18-OPN (Nacalai Tesque, Kyoto, Japan), which was washed with 75% ethanol solution and equilibrated with distilled water. A corrinoid compound was eluted with 100 mL of a linear gradient (0–25%) of ethanol. The B₁₂ active fractions assayed by the microbiological method were pooled, evaporated to dryness under reduced pressure, and dissolved in a small amount of 70% 2-propanol solution containing 2.8% NH₄OH. The solution was put on a silica gel 60 TLC sheet and developed with 2-propanol/28% NH₄OH/water (7:1:2) as a solvent in the dark at room temperature. The dried TLC sheet was cut into small pieces (0.5 cm × 1.0 cm) with scissors. Corrinoid was extracted from the pieces in 70% 2-propanol solution containing 2.8% NH₄OH several times, and the extract was evaporated to dryness under reduced pressure, dissolved in 1.0 mL of distilled water, and used as samples for the microbiological B₁₂ assay. The B₁₂ active fractions assayed by the microbiological method were pooled, evaporated to dryness under reduced pressure, and dissolved in a small amount of distilled water. The concentrated solution was further purified by high-performance liquid chromatography (HPLC) using a Shimadzu HPLC apparatus consisting of a LC-6A pump, SPD-6A spectrophotometer, CTO-6A column oven, and C-R6A Chromatopac. The sample (100 µL) was put on a 150 mm × 4.6 mm i.d., 5 µm, Wakosil-II 5C18RS reversed phase HPLC column equilibrated with 20% methanol solution containing 1% acetic acid at 35 °C. The flow rate was 1 mL/min. The corrinoid compound was isocratically eluted with the same solution, monitored by measuring absorbance at 278 nm, and collected in 1 mL fractions. The B₁₂ active fractions assayed by the microbiological method were collected, concentrated, and used as a purified corrinoid compound.

Analytical TLC and HPLC of Corrinoid Compound Purified from the Black Tea. The concentrated solution (10 µL) of corrinoid compound purified from the tea leaves and authentic cyano-B₁₂ was spotted on the silica gel 60 TLC sheets and developed with solvents I [2-propanol/NH₄OH (28%)/water (7:1:2)] and II [1-butanol/2-propanol/water (10:7:10)] in the dark at room temperature. The TLC sheets were dried, and R_f values of the pink-colored spot of cyano-B₁₂ were determined.

The TLC sheets were also cut into small pieces (0.5 cm × 1.0 cm) with scissors. Corrinoids were extracted from the pieces in 80% (v/v) methanol several times, evaporated to dryness under reduced pressure, dissolved in 1.0 mL of distilled water, and used as samples for the microbiological B₁₂ assay.

In the case of HPLC, the concentrated solutions (10 µL) of the purified corrinoid compound and authentic cyano-B₁₂ were analyzed with the same reversed phase HPLC column as used for purification. The corrinoids were isocratically eluted with 20% methanol solution containing 1% acetic acid at 35 °C and monitored by measuring absorbance at 278 nm. The retention times of corrinoids were determined at a flow rate of 1 mL/min. The eluate from the HPLC column was collected, evaporated to dryness, dissolved in 1.0 mL of distilled water, and used as samples for the microbiological B₁₂ assay.

Animals and Experimental Diets. Fifteen male Wister rats (20 weeks old), born to 14 week old parents fed on a B₁₂ deficient diet for 8 weeks, were used. The B₁₂ deficient diet fed to the parents contained (g/kg diet): 400 soyabean protein (Fuji Oil Ltd, Osaka, Japan), 438 anhydrous glucose (Nacalai Tesque Ltd., Kyoto, Japan), 100 soyabean oil (Nacalai Tesque Ltd.), 50 salt mixture, 5 dl-methionine (Nacalai Tesque Ltd.), and 5 B₁₂-free vitamin mixture and 2 choline chloride (Nacalai Tesque Ltd.), as described previously (5). The 3 week old weanling rats were housed in individual metabolism cages at 24 °C in a room with a 12 h light–dark cycle. They were given free access to 16 g/day of the B₁₂ deficient diet and distilled water for 17 weeks. In the feeding experiments, the 20 week old B₁₂ deficient rats (four rats/group) were given free access to 16 g/day of the B₁₂ deficient diet and 50 mL of either distilled water, authentic cyano-B₁₂ solution (1 ng of B₁₂ per 50 mL), or the tea drink (1 ng of B₁₂ per 50 mL) for 6 weeks. All experimental procedures involving laboratory animals were approved by the Animal Care and Use Committee of Osaka Prefecture University.

Urinary Methylmalonic Acid Assay. The urine of the B₁₂ deficient, cyano-B₁₂-supplemented, and tea drink-supplemented rats was sampled for 24 h in individual metabolic cages at weeks 0, 1, 2, and 6 during the experiments. Urinary methylmalonic acid was assayed by HPLC as described previously (6).

Extraction of B₁₂ from Rat Liver. After food was withheld from the 26 week old rats overnight, the rats were killed by decapitation under diethyl ether anesthesia. Livers were washed with a chilled 9 g/L NaCl solution, weighed, and stored at –80 °C until analyzed. A portion (1 g) of the liver was cut into small pieces using a razor blade and homogenized in 10 times its volume of 10 mmol/L acetate buffer, pH 4.8. B₁₂ was extracted from the liver homogenate by boiling with KCN at acidic pH as described above and assayed by the microbiological assay.

Statistics. Statistical analysis was performed using GB-STAT5.4 (Dynamic Microsystems, Inc., Silver Spring, MD). One way and two way repeated measure analysis of variance (ANOVA) were used for assay of hepatic B₁₂ and urinary methylmalonic acid in the animal feeding test, respectively. When ANOVA results were significant, a posthoc two-tailed Student's *t*-test also was performed and considered significant at *P* < 0.05.

RESULTS AND DISCUSSION

B₁₂ Content Determination. The B₁₂ contents were determined by two methods, the IF-chemiluminescence and the microbiological methods. The B₁₂ contents of the dry tea leaves were 456 ± 39 ng and 368 ± 56 ng per 100 g dry weight by the IF-chemiluminescence method and the microbiological method, respectively. In the case of B₁₂, contents of the tea drink used for the feeding experiments were 2.0 ± 0.3 ng and 2.0 ± 0.8 ng per 100 mL by those two methods, respectively. Although B₁₂ contents were considerably lower in the black tea drink than in cow's milk (0.3 µg/100 g) (7), which greatly contributes to the B₁₂ intake of U.S. adult women (8), this is the first report on the occurrence of B₁₂ in tea leaves and their drinks.

Characterization of the Corrinoid Compound from the Black Tea. A corrinoid compound was purified from the tea leaves. The purified corrinoid compound and authentic cyano-B₁₂ were analyzed by TLC and HPLC. The R_f values for the purified compound were 0.64 and 0.22 on silica gel 60 TLC in solvents I and II, respectively. These values were identical to those for authentic cyano-B₁₂. The retention time of authentic B₁₂ by reversed phase HPLC was 9.4 min; it was also identical to that of the purified compound. These results strongly suggest that the compound purified from the fermented tea leaves is true B₁₂ but not corrinoid compounds inactive for humans. UV–vis spectroscopy and NMR spectroscopy could not be determined because a substantial amount of the purified compound was not obtained.

Feeding Test of the Tea Drink with the B₁₂ Status of B₁₂ Deficient Rats. To evaluate whether the B₁₂ found in the tea

Table 1. Effects of Feeding the Fermented Black Tea Drink on Urinary Methylmalonic Acid of B₁₂ Deficient Rats^a

groups	urinary methylmalonic acid (mg/day)		
	control	CN-B ₁₂ supplement	tea drink supplement
week 0	246 ± 51 ^a	253 ± 41 ^a	251 ± 35 ^a
week 1	246 ± 32 ^a	200 ± 25 ^a	181 ± 66 ^a
week 2	251 ± 104 ^a	218 ± 35 ^a	151 ± 50 ^b
week 6	250 ± 32 ^a	201 ± 28 ^a	169 ± 29 ^b

^a The 20 week old B₁₂ deficient rats (four rats/group) were given free access to 50 mL of either distilled water, the cyano-B₁₂ solution (1 ng/50 mL), or the tea drink (1 ng of B₁₂/50 mL) per day. ^{ab}The mean values with different superscript letters are significantly different; *P* < 0.05.

Table 2. Hepatic B₁₂ Contents of the 26 Week Old B₁₂ Deficient Rats Fed the CN-B₁₂ Solution and Tea Drink^a

groups	B ₁₂ contents (pg/g wet tissue)
control	746 ± 97 ^a
CN-B ₁₂ supplement	768 ± 129 ^a
tea drink supplement	1473 ± 252 ^b

^a The 20 week old B₁₂ deficient rats (four rats/group) were given free access to 50 mL of either distilled water, the cyano-B₁₂ solution (1 ng/50 mL), or the tea drink (1 ng of B₁₂/50 mL) per day. ^{ab}The mean values within a column with different superscript letters are significantly different; *P* < 0.01.

leaves is absorbed in the mammalian intestine and accumulated in the liver, feeding experiments of the tea drink to 20 week old B₁₂ deficient rats were conducted. There was no significant difference in the intakes of the diet and drink (water, cyano-B₁₂ solution, or the tea drink) among the rat groups during the experimental time course. When the 20 week old B₁₂ deficient rats, which excreted substantial amounts of methylmalonic acid (about 250 mg/day) in urine (as an index of B₁₂ deficiency), were given the tea drink (1 ng of B₁₂ per day) for 6 weeks, urinary methylmalonic acid excretion of the tea drink-supplemented 26 week old rats decreased significantly relative to the B₁₂ deficient (control) rats, but that of the cyano-B₁₂-supplemented rats did not (Table 1).

Although the rate of growth (61.6 ± 18.2 g) of the B₁₂ deficient rats given the tea drink had a tendency to be greater than that of the control (28.2 ± 7.4 g) and cyano-B₁₂-supplemented (20.6 ± 15.8 g) rats during the experiment, there was no significant difference in body weight among the rats fed the three experimental drinks after 6 weeks.

The hepatic B₁₂ contents were about 2-fold greater in the tea drink-supplemented rats than in both control and cyano-B₁₂-supplemented rats (Table 2); there was no significant difference in the hepatic B₁₂ contents between control and cyano-B₁₂-supplemented rats. Although the methylmalonic aciduria of the B₁₂ deficient rats could not be completely recovered by the 6 week feeding of the tea drink, the significant increase in the hepatic B₁₂ content of the tea drink-supplemented rats indicated that the feeding of the tea drink considerably improved B₁₂ status in the B₁₂ deficient rats.

Our previous study (9) has indicated that urinary levels of methylmalonic acid became undetectable in the B₁₂ deficient rats fed a cyano-B₁₂ (about 100 ng/day)-supplemented diet for 10 days. In this study, however, the cyano-B₁₂ (1 ng/day)-supplemented 26 week old rats did not show both significant recovery of methylmalonic aciduria and increase in hepatic B₁₂ content. Even the 26 week old B₁₂ deficient rats given the tea drink did not completely recover from methylmalonic aciduria. The results may be due to a lesser B₁₂ content (1 ng/day) of the administered authentic B₁₂ and tea drink. We did not use any

concentrated or purified compound from the tea leaves because of evaluation for bioavailability of B₁₂ found in the tea drink commercially available for humans.

Our preliminary experiments indicated that considerable amounts of the coenzyme B₁₂ (adenosylcobalamin and methylcobalamin) were found in the tea leaves and that the B₁₂ found in the tea drink existed as a free form (without binding to a macromolecular compound). These results suggest that the B₁₂ found in the tea drink would be assimilated more easily in the B₁₂ deficient rats than authentic cyano-B₁₂.

The results presented here indicate that the B₁₂ found in the Japanese black tea (Batabata-cha) fermented by bacteria is bioavailable in mammals. Although only 1–2 L of the fermented tea drink (20–40 ng of B₁₂) is not sufficient to satisfy the recommended dietary allowance (2.4 μg/day) for human adults, intakes of various B₁₂-containing plant foods [purple and green lavers (10), Chlorella tablets (11), and the fermented tea extract] would contribute to prevention of B₁₂ deficiency for vegetarians.

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Time Course of Changes in the Activity of Vitamin B₁₂-Dependent Methionine Synthase during Cell Growth of *Euglena gracilis* Z

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要 旨

ビタミンB₁₂給与および欠乏 *Euglena gracilis* Z の生育に伴うビタミンB₁₂依存性メチオニン合成酵素活性の変動を検討した。本酵素活性はビタミンB₁₂給与 *Euglena* 細胞で生育2日目の対数増殖初期で最大(2.1nmol/min/10⁶cells)となり、その後顕著に減少した。一方、ビタミンB₁₂欠乏 *Euglena* 細胞では生育期間を通して非常に低い活性(0.01-0.04nmol/min/10⁶cells)を示した。本酵素はビタミンB₁₂給与細胞の対数増殖初期にのみ高い活性が認められたため、対数増殖初期のビタミンB₁₂給与細胞を用いて本酵素の細胞内局在性の解明を試みた。その結果、ほとんどの本酵素活性が細胞質画分に回収されたが、単離・精製したミトコンドリアと葉緑体にも本酵素活性が検出され、これらオルガネラにおいても本酵素が存在し、メチオニン代謝に関与する可能性が示唆された。

Abstract

To verify physiological roles of vitamin B₁₂ (or cobalamin)-dependent methionine synthase in the vitamin B₁₂-requiring protozoan *Euglena gracilis* Z, the time course of changes in the enzyme activity during *Euglena* cell growth was studied. The enzyme activity of the vitamin B₁₂-sufficient *Euglena* cells reached a maximum (2.1nmol/min/10⁶cells) at the early logarithmic growth phase and significantly decreased thereafter. While that of the vitamin B₁₂-deficient cells was significantly low (0.01-0.04nmol/min/10⁶cells) during cell growth. Preliminary experiments indicated that most of the enzyme activity found in a homogenate of the cells grown for 2 days in the vitamin B₁₂-sufficient medium was recovered in the cytosolic fraction. The enzyme activity, however, was found in the percoll-purified mitochondria and chloroplasts, suggesting that the enzyme can function in methionine metabolism in the organelle.

Keywords : cobalamin, *Euglena gracilis* Z, methionine synthase, subcellular distribution, vitamin B₁₂

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1 目的

Euglena gracilis Zは紡錘形の形をした原生生物で、長さ約50 μ m、幅約10 μ mの大きさをしており、動物と植物の両方の特徴を有している¹⁾。

E. gracilis Zは生育にビタミンB₁₂ (B₁₂) を要求し²⁾、細胞内にB₁₂を活発に取込み・蓄積する²⁾ためB₁₂の生理機能解明のための研究材料として適している。

E. gracilis ZにはB₁₂依存性酵素としてN⁵-メチルテトラヒドロ葉酸とホモシステインからメチオニンの生成を触媒するメチオニン合成酵素 (EC 2.2.2.13)³⁾、アミノ酸や奇数鎖脂肪酸の異化経路でR-メチルマロニル-CoAからスクシニル-CoAの異性化反応を触媒するメチルマロニルCoAムターゼ (EC 5.4.99.2)⁴⁾、DNA合成に関与するリボヌクレオチドリダクターゼ (EC 1.17.4.2)⁵⁾が存在している。

B₁₂依存性メチオニン合成酵素は生物界に広く分布し、哺乳動物の各種細胞では細胞質に局在する酵素として知られている⁶⁾。一方*E. gracilis* Zにおいて本酵素は細胞質、ミトコンドリア、葉緑体の各オルガネラに分布することが報告³⁾されているが、*E. gracilis* Zの本酵素は非常に不安定なため詳細な酵素化学的性質や生理機能についての知見はない。そこで、*E. gracilis* ZにおけるB₁₂依存性メチオニン合成酵素の生理機能を解明する目的で生育に伴う本酵素活性の変動やB₁₂欠乏の影響について検討した。

2 実験方法

2-1 培養法

E. gracilis ZはKH培地¹⁾ (B₁₂を5 μ g/L含む)で光照射下 (30 μ E/m²/s) 27°Cで5日間振盪培養した。B₁₂欠乏細胞の調製は、KH培地で5日間前培養した*E. gracilis* Zの培養液1.0mLを無菌的にB₁₂を除去したKH培地150mLに接種し、同条件で5日間培養した。

2-2 細胞数の測定法

E. gracilis Zの細胞数は血球計算盤を用いて測定した。

2-3 粗酵素液の調製法

数日間生育させた*E. gracilis* Zの培養液 (1~10mL) を無菌的にサンプリングした後、3,000xg 10分間の遠心分離により*Euglena*細胞を集めた。細胞は蒸留水で2回洗浄後、10% (w/v) ショ糖を含む10mMリン酸カリウム緩衝液 (pH7.0) に懸濁後、超音波処理により破碎した。細胞破碎液は10,000xg 30分間遠心分離し、その上清を粗酵素液として実験に用いた。全ての操作は2(-)4°Cで行った。半角

2-4 ビタミンB₁₂依存性メチオニン合成酵素活性の測定法

本酵素活性の測定はHuangらの方法³⁾を改良して行った。また、Banerjeeら⁹⁾の半嫌氣的酵素活性測定法に準じて酵素反応液を調製した。酵素反応液の組成は、100mMリン酸カリウム緩衝液 (pH7.0)、152 μ M S-アデノシルメチオニン (シグマ社製)、50mM ヒドロキソB₁₂ (シグマ社製)、25mM アスコルビン酸、25mM ジチオスレイトール、500 μ M L-ホモシステイン (シグマ社製)、25 μ M N⁵-メチルテトラヒドロ葉酸 (シグマ社製)、粗酵素液とし全容量を1.0mLとした。酵素反応液は、N⁵-メチルテトラヒドロ葉酸を含まない反応液を予め調製し、37°C 5分間保温し、本酵素を還元的に活性化させた。

その後、基質N⁵-メチルテトラヒドロ葉酸を加え、37°C 10分間酵素反応を行った。酵素反応終了後、直ちに100°C 2分間の加熱処理により酵素反応を停止させた。酵素反応液は水中で5分間冷却した後、遠心分離により変性タンパク質を除去した。この遠心分離上清液をメンブレンフィルター (Millex-LH, 0.45 μ m, ミリポア社製) でろ過し、ろ過液10 μ Lを高速液体クロマトグラフィー (HPLC) の試料とした。

HPLC分析システムは島津社製SCL-10Avpシステムコントローラー, LC-10ADvp HPLCポンプ, DGU-12Aデガッサー, CTO-10Avpカラムオーブン, C-R6Aクロマトパック, 分光蛍光光度計RF-5000を用いた。HPLCの分析条件はTSK-GEL ODS-120A (4.6×250mm, 東ソー社製) カラムを用い, カラム温度30°C, 移動相7.0% (v/v) アセトニトリルを含む33mMリン酸カリウム緩衝液 (pH3.0) を用い, 流速0.5mL/minで分析した。酵素反応により生成したテトラヒドロ葉酸を励起波長290nm, 吸収波長356nmで測定した。また, 本酵素活性は対照の反応液 (予め100°C 5分間加熱処理した粗酵素液を用いて上述の反応液を調製し, 直ちに100°C 2分間の加熱処理を行った後, N⁵-メチルテトラヒドロ葉酸を添加した反応液) 中のテトラヒドロ葉酸量を差引き求めた。

2-5 タンパク質定量法

タンパク量はオボアルブミンを標準タンパク質としてバイオラッド社製プロテインアッセイ試薬を用いて定量した。

2-6 細胞分画法

生育2日目の培養液を3,000xg 10分間の遠心分離により *Euglena* の細胞を集めた。細胞は蒸留水で2回洗浄後, 0.33Mマンニトールを含む25mMグリシルグリシン-KOH緩衝液 (pH7.4) に懸濁後, 乳鉢・乳棒を用いて細胞を海砂と共に温和に摩砕することで細胞破碎液を調製した。細胞破碎液をガーゼでろ過し, 3,000xg 10分間の遠心分離上清画分を細胞抽出液とした。

細胞抽出液を5,000xg 10分間遠心分離し, 沈殿画分は0.33Mマンニトールを含む25mMグリシルグリシン-KOH緩衝液 (pH7.4) で懸濁し, 粗葉緑体とした。

上清画分はさらに11,000xg 10分間遠心分離し, 沈殿画分は0.25Mショ糖を含む25mMグリシルグリシン-KOH緩衝液 (pH7.4) で懸濁し粗ミトコンドリアとした。また, 上清画分は細胞質画分と

して実験に用いた。全ての操作は, 2-4°Cで行った。

粗ミトコンドリアおよび葉緑体はさらに Watanabeら¹⁰⁾ の方法でパーコール精製を行い無傷のミトコンドリアと葉緑体を調製した。酵素活性測定に際して各画分を超音波破碎し, 粗酵素液として使用した。

3 結果と考察

B₁₂ 給与および欠乏 *Euglena* 細胞の生育に伴う B₁₂ 依存性メチオニン合成酵素活性の変動を Fig. 1 に示す。B₁₂ 給与細胞では生育2日目で対数増殖初期となり, 生育4日目には定常期に達した。一方, B₁₂ 欠乏細胞では実験期間を通じて顕著な細胞増殖は確認できなかった。

B₁₂ 依存性メチオニン合成酵素活性は B₁₂ 給与細胞で生育2日目の対数増殖初期で最大 (2.1nmol/min/10⁶ cells) となり, その後顕著に減少した。この結果は, 対数増殖期の活発な細胞分裂に備えタンパク質合成に必須のメチオニンを供給するために本酵素が機能しているものと思われる。

一方, B₁₂ 欠乏細胞では生育期間を通して非常に低い酵素活性 (0.01-0.04nmol/min/10⁶ cells) を示した。この結果は, B₁₂ 欠乏により不安定なアポ酵素が増加するためと考えられ, 大腸菌¹¹⁾ や哺乳動物¹²⁾ で同様の結果が報告されている。

Isegawaら⁴⁾ は生育6日目の B₁₂ 欠乏 *Euglena* 細胞を用いて本酵素の細胞内局在性を検討し, 本酵素が細胞質, 葉緑体, ミトコンドリアにそれぞれ 68.9% (14.5pmol/min/mg protein), 18.4% (7.6pmol/min/mg protein), 9.5% (6.6pmol/min/mg protein) の割合で存在していることを報告している。しかし, 今回の実験結果から本酵素は B₁₂ 給与細胞の対数増殖初期にのみ高い活性が検出されたため, 対数増殖初期の B₁₂ 給与細胞を用いて本酵素の細胞内局在性を検討する必要がある。

B₁₂ 欠乏 *Euglena* 細胞を用いたトリプシン消化法によるオルガネラの分画法⁴⁾ が確立されているが, B₁₂ 給与細胞の細胞膜複合体は強固なため本