ランブル鞭毛虫及び小形アメーバの感染は2004年度の調査結果と同様、広範に確認された。施設Iに関しては、男性施設利用者の約46%が大腸アメーバに感染していることが判明した。これまでの調査から、対象施設の利用者の部屋やフロアー単位での集団感染だけでなく、施設から施設への腸管寄生原虫感染の伝播の可能性も示唆されていることから、他の腸管寄生原虫を含め、赤痢アメーバの施設内集団感染実態調査を引き続き実施する必要性を認識した。

2)ジアルジアによる施設内感染の分子生物学的な解析:今回新たに設計したプライマーを用いた nested PCR による遺伝子型の決定方法により、特に糞便中のジアルジア数が少ない場合に特に有効であることが明らかとなった。ジアルジア感染者の糞便には、糞便を採取した日により糞便中のシスト数が大きく異なることを考えると、この nested PCR による遺伝子型の決定法が後のジアルジアによる施設内集団感染においても感染経路の解明に一役を担えることが期待できる。

F. 健康危険情報 なし

G. 研究発表

- 1. 論文発表
- 1) 鈴木 淳、村田理恵、小林正規、柳川義勢、 竹内 勤:知的障害者更正施設における赤痢ア メーバ等腸管寄生原虫の感染実態調査、日本臨 床寄生虫学会誌(印刷中)

2. 学会発表

1) 鈴木 淳、村田理恵、小林正規、柳川義勢、 竹内 勤: 知的障害者更正施設における赤痢ア メーバ等腸管寄生原虫の感染実態調査、第16 回日本臨床寄生虫学会(2005)

H. 知的財産権の出願・登録状況

1. 特許取得 なし

2. 実用新案登録 なし

3. その他 なし



AXENIC CULTIVATION OF ENTAMOEBA DISPAR IN NEWLY DESIGNED YEAST EXTRACT-IRON-GLUCONIC ACID-DIHYRDOXYACETONE-SERUM MEDIUM

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ABSTRACT: Yeast extract—iron—gluconic acid—dihyrdoxyacetone-serum medium that allows axenic cultivation of *Entamoeba dispar* was designed based on casein-free yeast extract-iron-serum (YI-S) medium, and the usefulness of the medium was assessed. The main differences from YI-S medium are replacement of glucose by gluconic acid, addition of dihydroxyacetone and D-galacturonic acid monohydrate, and sterilization by filtration. This medium promoted the axenic growth of 5 strains of *E. dispar* (2 strains of nonhuman primate isolates and 3 strains of human isolates). In addition, to clarify the biological basis for the growth of *E. dispar* in this medium, analyses of relevant enzymes on the glycolytic pathway of the amoebae as well as of the protozoans that are the best culture supplement for amoebae are being performed.

After axenic cultivation of *Entamoeba dispar* (strain: SAW760RRcloneAR) was reported (Clark, 1995) in casein-free yeast extract–iron–serum (YI-S) medium (Diamond et al., 1995) supplemented with gastric mucin, the efficacy of this axenic culture system was assessed for other *E. dispar* strains. However, despite its utility, the only *E. dispar* strain established as an axenic amoeba in YI-S medium thus far is SAW760RR clone A.

On the other hand, when a monoxenic culture system for *E. dispar* in biosate-cysteine-starch-iron-serum (BCSI-S) medium with *Pseudomonas aeruginosa* was developed (Kobayashi et al., 1998), a significant growth-promoting effect of dihydroxyacetone (DHA) was observed. DHA is a ketotriose and as a sugar source for *E. dispar* is thought to be directly metabolized to DHA phosphate (DHAP), which is an intermediary metabolite in the Embden-Meyerhof-Parnas glycolytic pathway.

Another significant result was that during characterization of the glycolytic pathway of Crithidia fasciculata, an effective culture associate of E. dispar in BI-S-33 medium (Diamond et al., 1978), approximately 35 times as much glucose-6-phosphate dehydrogenase (G6PDH) activity as that of Trypanosoma cruzi (Tulahuen strain) was detected. Trypanosoma cruzi can be used as a growth-promoting supplement for monoxenic culture of E. histolytica. However, it does not promote the growth of E. dispar, whereas C. fasciculata does (S. Kobayashi, unpubl.). G6PDH activity has not been detected in axenically grown E. histolytica (Reeves, 1972) or E. dispar (S. Kobayashi, unpubl.). G6PDH is an essential enzyme not only in the pentose-phosphate pathway but also in the Entner-Doudoroff pathway, which starts with G6P. G6PDH catalyzes the transformation of G6P to 6-phosphogluconate (6PG) via 6-phosphogluconolactone, and we focused attention on the glycolytic pathway after 6PG in E. dispar and examined the reactions after 6PG. Both the pentose-phosphate and Entner-Doudoroff glycolytic pathways can branch from 6PG. The activity of 6PG dehydratase (EC 4.2.1.12.) (Gottschalk and Bender, 1982; Nguyen and Schiller, 1989), which is one of the enzymes in the Entner-

Doudoroff pathway, was detected in the lysates of both E. histolytica (HM-1:IMSS clone 6) and E. dispar (CYNO 16:TPC), and its activity in E. dispar was 2.87 times greater than in E. histolytica. By contrast, no 6PG dehydrogenase activity in the pentose-phosphate pathway was detected in either lysate (S. Kobayashi, unpubl.). On the other hand, DHA, which has a growth-promoting effect, is thought to be metabolized to G3P. through DHAP, which is converted to glyceraldehyde-3-phosphate (G3P) by triose phosphate isomerase (EC 5.3.1.1.). Thus, if viable C. fasciculata actually supplies a growth-promoting substance like DHA, DHA may be preferable to viable C. fasciculata as a culture ingredient. On the basis of this information, we hypothesized the existence of some failure in the upstream portion of the glycolytic pathway in E. dispar, before DHAP or G3P production from glucose in YI-S medium. We, therefore, attempted to design an axenic culture medium for E. dispar by modifying the ingredients of YI-S medium, and we tested its usefulness.

MATERIALS AND METHODS

Reagents

All chemicals used in this study were of the highest quality commercially available unless otherwise stated.

Entamoeba dispar isolates

Five strains of *E. dispar* were subjected to a trial of axenic cultivation in this study. Two strains (CYNO 09:TPC and CYNO 16:TPC) from nonhuman primates were isolated in Japan (Kobayashi et al., 1998) in Robinson's medium (Robinson, 1968), and 2 strains (AS 2 IR and AS 16 IR) were isolated from humans in Iran. An *E. dispar* strain, SAW 1734R clone AR (SAW1734RclAR), was used as the reference strain. The xenic *E. dispar* strains were transferred to monoxenic culture medium with viable *C. fasciculata* and antibiotics (as in the classic approach Diamond used for *E. histolytica*, Diamond, 1983) in BI-S-33 medium. The *C. fasciculata* was then replaced with antibiotic-sensitive *P. aeruginosa* in BCSI-S medium because *P. aeruginosa* is removed easily by antibiotics and antibiotic-resistant *C. fasciculata* usually survive in test medium for axenic culture even after 2–3 subcultures, and it promotes the growth of *E. dispar*.

Yeast-iron-gluconic acid-dihydroxyacetone-serum medium

Yeast-iron-gluconic acid-dihydroxyacetone (YIGADHA) broth was prepared by first dissolving the following ingredients in 880 ml of water distilled and purified with a Milli-Q column (Millipore Co., Billerica, Massachusetts): K₂HPO₄, 1 g; KH₂PO₄, 0.6 g; NaCl, 2.0 g; yeast extract (BBL, Becton Dickinson Co., Cockeysville, Maryland), 30 g; gluconic acid, 5.0 g; DHA (Sigma Chemical Co., St. Louis, Missouri), 1.0 g; p-galacturonic acid monohydrate, 0.2 g; L-cysteine hydrochloride, 1.0 g; ascorbic acid, 0.2 g; and ferric ammonium citrate (brown), 22.8 mg.

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To complete the YIGADHA-serum (YIGADHA-S) medium, 2 ml of vitamin mixture #18 prepared according to the instructions for PDM-805 medium (Diamond and Cunnick, 1991), and 16 ml of heat-inactivated bovine serum was aseptically added to 88 ml of the YIGADHA broth. A 5-ml volume of the complete medium was then dispensed into 13- × 100-mm, screw-capped borosilicate glass culture tubes (Asahi Techno Glass Co., Chuo-ku, Tokyo, Japan).

Establishment of the culture of *Entamoeba dispar* with sterilized Crithidia fasciculata

After removing *P. aeruginosa* by washing and addition of antibiotics (polymyxin B, 130 units/ml), trophozoites of the 5 strains of *E. dispar* were transferred to YIGADHA-S medium with sterilized *C. fasciculata* (2–3 million/ml, ReF-1:PRR, ATCC no. 50083) by fixation with 10% (v/v) formalin as described in the literature (Clark, 1995). All 5 strains of *E. dispar* adapted to the culture at 35.5 C in the YIGADHA-S medium with the associate within 2 wk after 3–4 whole-medium changes by centrifugation (275 g, 4 min) and eventually started to grow continuously. When the same growth-promoting effect was later found with autoclaved *C. fasciculata* in Hanks' solution as with the formalin-fixed *C. fasciculata*, we switched the method of sterilizing *C. fasciculata* to autoclaving (121 C, 15 min) because of its simplicity.

When the amoebae were subcultured, 0.6-0.8 ml of the amoeba suspension $(2.5 \times 10^4 \text{ to } 5 \times 10^4/\text{ml})$ was usually transferred to the fresh medium, after the cultures were chilled for 5 min in an ice-water bath. They were then inverted several times to disperse the amoebae. Subcultures were successfully prepared at intervals of 3 and 4 days.

Growth kinetics of the amoebae

At 24-hr intervals for 96 hr, the number of amoebae in 5 μ l of the homogeneous amoeba suspension diluted with known volume of YIG-ADHA-S medium were counted under a microscope after chilling the culture tubes for 5 min in an ice-water bath. The data were then converted to number per milliliter.

Zymodeme analysis and polymerase chain reaction

Zymodeme analysis (Sargeaunt, 1988) and polymerase chain reaction (PCR) analysis (Tachibana et al., 1991; Cheng et al., 1993) of the amoeba isolates were performed to characterize the amoebae grown in the axenic culture.

RESULTS

YIGADHA-S medium was designed, and the culture associates tested could be removed with no loss of *E. dispar* viability as a result of searching for the improved basic medium and condition for axenic cultivation of *E. dispar*, based on the casein-free YI-S medium.

YI-S medium was therefore modified as follows. Glucose was replaced by GA, DHA and D-galacturonic acid monohydrate were added because of the significant growth-promoting effects under the axenic culture conditions, and the medium was sterilized by filtration and adjusted to pH 6.5 with NaOH.

As a result of these modifications, trophozoites of *E. dispar* started to grow continuously in the absence of viable *C. fasciculata* or *P. aeruginosa* for the first time, although supplementation with the formalin-fixed organisms was still necessary for growth.

The growth-promoting ability of *C. fasciculata* was also maintained in this YIGADHA-S culture system even after it was autoclaved (121 C, 15 min) in Hanks' solution. There was no difference between the growth-promoting effect of autoclaved *C. fasciculata* and of formalin-fixed organisms. Because

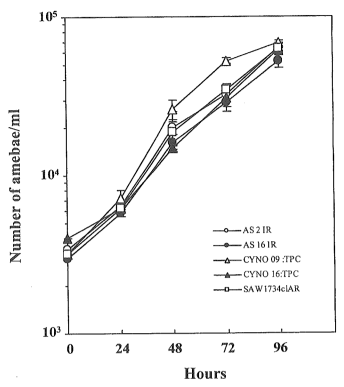


FIGURE 1. Growth kinetics of 5 strains of *Entamoeba dispar* in the 4-yr passaged cultures in YIGADHA-S medium supplemented with autoclaved *Crithidia fasciculata*. Mean numbers of amoebae in duplicate cultures are plotted.

the procedure was easier, the amoebae were maintained in the medium with autoclaved *C. fasciculata*. The growth kinetics of the 5 strains of *E. dispar* in this culture system are shown in Figure 1.

After 3 mo of maintenance in this culture system, 1 strain of *E. dispar* (CYNO 16:TPC) started to grow without any supplement, e.g., without autoclaved *C. fasciculata*. However, addition of 6PG (Sigma P-7877, 67 µg/ml), an intermediary metabolite in the Entner–Doudoroff pathway (Fig. 3), was necessary to maintain this strain without the autoclaved organisms.

After culturing with autoclaved *C. fasciculata* for 1 yr and 4 yr, respectively, other *E. dispar* strains started to grow axenically in the YIGADHA-S medium without the addition of any culture associates. The growth kinetics of 5 strains of axenically grown *E. dispar* are shown in Figure 2. The clear differences in growth kinetics between monoxenic culture (Fig. 1) and axenic culture (Fig. 2), except for the CYNO 09:TPC strain, suggest the existence of some other growth-promoting factor in autoclaved *C. fasciculata*.

The CYNO 16:TPC and CYNO 09:TPC strains were usually subcultured by transferring 1 ml of amoeba suspension to fresh YIGADHA-S medium, and the AS 16 IR, AS 2 IR, and SAW1734RclAR strains were subcultured by transferring 1 ml of amoeba suspension concentrated to an adequate density for subculture (3 \times 10⁴ to 6 \times 10⁴ amoebae/ml) by centrifugation (275 g, for 4 min).

The zymodemes of the 5 strains grown in both the monoxenic and axenic YIGADHA-S media were all judged to be type I. All 10 DNA samples of the 5 strains used in the PCR



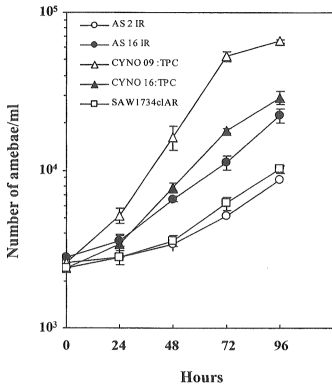


FIGURE 2. Growth kinetics of 5 strains of axenically grown *Entamoeba dispar* in 2-yr (CYNO 09: TPC), 3-yr (CYNO 16:TPC), and 2-mo (AS 2 IR, AS 16 IR, and SAW 1734RclAR) passaged cultures in YIGADHA-S axenic medium. Mean numbers of amoebae in duplicate cultures are plotted.

analyses described above were amplified with *E. dispar*–specific primers alone; the *E. histolytica*–specific primers did not elicit any DNA amplification. These findings are summarized in Table I.

DISCUSSION

We have previously reported a monoxenic culture system for *E. dispar* in BCSI-S medium in which glucose was replaced with starch and to which sterilized *C. fasciculata* were added after heat treatment at 56 C for 30 min followed by incubation in 1% hydrogen peroxide at 4 C for 24 hr (Kobayashi et al., 1998). The sterilized parasite appeared to be metabolically in-

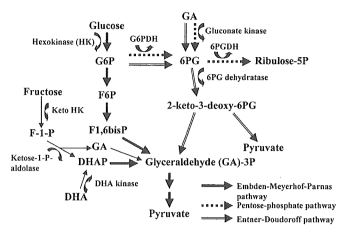


FIGURE 3. Metabolic map of 3 glycolytic pathways.

active as judged by the nuclear magnetic resonance spectra. Although the growth-promoting effect of *C. fasciculata* was partially lost as a result of this sterilization step, the minimum essential effect persisted, which made it possible to establish a long-term monoxenic culture of *E. dispar* with metabolically inactive *C. fasciculata*. In this study, we succeeded in producing an axenic culture system of *E. dispar* by using YIGADHA-medium designed on the basis of YI-S medium from the monoxenic culture of the amoeba with sterilized *C. fasciculata*.

During attempts to improve the culture system, it was found that by replacing some kinds of sugar from starch, the growth of E. dispar was promoted significantly in the BCSI-S monoxenic culture system. First, a marginal growth-promoting effect of fructose was found, although its effect was insufficient for axenic cultivation of E. dispar, and a more useful compound, DHA, was used. The concentration (0.1%) of DHA required for the growth of E. dispar in YIGADHA-S medium was very critical: DHA concentrations below 0.05% and greater than 0.15% failed to effectively promote the growth of E. dispar. However, despite the clear effect of DHA and fructose on the growth of E. dispar, the results of our biochemical assays for phosphorylation of DHA by DHA kinase, ketohexokinase, and ketose-1-phosphate aldolase, the enzymes involved in the conversion of fructose or DHA to DHAP (Fig. 3), showed no significant activity of these enzymes in the crude lysates of axenic E. dispar (CYNO 16:TPC) or E. histolytica (HM-1:IMSS clone

TABLE I. Zymodeme and PCR analyses of 5 strains of Entamoeba dispar.

Strain	Xenic (date)	Zymo	PCR†		
		Monoxenic (date)	Axenic (date)	Monoxenic	Axenic
SAW 1734RclAR	Z-III‡ (March 1985)	Z-I (January 2000)	Z-I (February 2003)	E. dispar	E. dispar
AS 2 IR	Z-I (June 1998)	Z-I (January 2000)	Z-I (February 2003)	E. dispar	E. dispar
AS 16 IR	Z-I (June 1998)	Z-I (January 2000)	Z-I (February 2003)	E. dispar	E. dispar
CYNO 09:TPC	Z-I (April 1992)	Z-I (January 2000)	Z-I (February 2003)	E. dispar	E. dispar
CYNO 16:TPC	Z-III (April 1992)	Z-I (January 2000)	Z-I (February 2003)	E. dispar	E. dispar

^{*} Zymodemes type I and III are classified as E. dispar.

[†] PCR analysis using 2 sets of oligonucleotide primers each (p11 plus p12 and p13 plus p14, respectively) for amplification of the DNAs of E. histolytica and E. dispar. The zymodeme and PCR analyses were performed around the same time.

‡ Data cited from Mirelman et al. (1986).

6) (data not shown). The mechanism of growth-promoting effect of DHA and fructose is not yet understood.

Pimenta et al. (2002) demonstrated an interesting morphological characteristic of E. dispar by electron microscopy: some concomitant bacteria cells were free and viable in the cytoplasm of E. dispar, without being surrounded by a distinct phagosome membrane. This suggests that some bacterial species can survive in the cytoplasm and exist in a symbiotic relationship with E. dispar. If the Entner-Doudoroff pathway does indeed function in glycolysis in E. dispar, as reported in E. histolytica under xenic conditions (Hilker and White, 1959), the reactions upstream in the Embden-Meyerhof-Parnas pathway (Fig. 3) including a reaction regulating the transformation of fructose-6phosphate to fructose-1,6-bisphosphate or vice versa, usually catalyzed by 6-phosphofructokinase (6-PFK) and fructose-bisphosphatase (EC 3.1.3.11.) and affecting the both the glycolysis and glycogenesis pathways, may not function well in E. dispar. The regulatory reactions in E. histolytica are well known to be regulated by a single unique enzyme (ppi-dependent 6-PFK; EC 2.7.1.90.), and the reaction is reversible and has no apparent regulatory function (Reeves et al., 1974, 1976).

The clear growth-promoting effect of autoclaved *C. fasciculata* (or *P. aeruginosa*) indicated that they contain as yet unidentified heat-stable growth-promoting substances for *E. dispar*. If the substances can be identified, YIGADHA-S medium will be improved. Further analyses of the axenically grown *E. dispar* will enable us to further elucidate the biological properties of *E. dispar* that differ from those of pathogenic *E. histolytica*.

ACKNOWLEDGMENTS

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GROWTH-PROMOTING EFFECT ON IRON-SULFUR PROTEINS ON AXENIC CULTURES OF ENTAMOEBA DISPAR

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Summary:

A growth-promoting factor (GPF) that promotes the growth of Entamoeba dispar under axenic culture conditions was found in fractions of mitochondria (Mi), hydrogenosomes (Hg) and chloroplasts (Cp) obtained from cells of six different protozoan, mammalian and plant species. We were able to extract the GPF from the Cp-rich leaf cells of a plant (spiderwort: Commelina communis L.) in an acetone-soluble fraction as a complex of chlorophyll with low molecular weight proteins (molecular weight [MW] approximately 4,600). We also found that on treatment with 0.6 % complexes of 2-mercapthoethanol (2ME), complexes of chlorophyll-a with iron-sulphur (Fe-S) proteins (e.g., ferredoxins [Fd] from spinach and Clostridium pasteurianum) and noncomplex rubredoxin (Rd) from C. pasteurianum have a growth-promoting effect on E. dispar. These findings suggest that E dispar may lack a sufficient quantity of some essential components of FeS proteins, such as Fe-S center.

KEY WORDS: growth, mitochondria, hydrogenosomes, chloroplasts, iron-sulphur protein. *Estamocha dispar*

Résumé : Les effets d'accélération de croissance des protéines ffr-soi fir dans la cillitre axénique d'*Etamoera dispar*

Des facteurs d'accélération de croissance (Growth promoting factor: GPF) favorisant le développement d'Entamoeba dispar ont été détectés dans la composition de la mitochondrie, de l'hydrogénosome et du chloroplaste isolés à partir de six sortes de cellules issues de protozoaires, mammiferes et plantes. De plus, les GPF de cellules de mésophile contenant une grande quantité de chloroplaste végétal (Commelina communis L.) ont pu être extraits comme une substance composée de protéines de faible masse moléculaire (≈ 4600) et de chloroplastes dans la composition acétone-soluble. À partir de ces résultats, nous avons préalablement pu découvrir des effets d'accélération de croissance dans le corps composé de protéines fer-soufre lépinard et ferrédoxine de Clostridium pasteurianum) et de chlorophylle a traité au 2-mercaptoéthanol 0,6 % et du corps simple rubrédoxine de C. pasteurianum. Ces observations ont suggéré qu'une composante essentielle formant la protéine fer-soufre d'E. dispar (comme noyau fer-soufre) semblait être insuffisante.

MOTS CLÉS : croissance, mitochondrie, hydrogénosome, chloroplaste, protéine lersoulre, Entamoeba dispar.

INTRODUCTION

mamoeba dispar grows well under xenic and monoxenic culture conditions along with enteric bacteria such as *Escherichia coli* or anaerobic bacteria such as *Fusobacterium symbiosum* (*Clostridium symbiosum* ATCC 14940) (Robinson, 1968; Diamond, 1982; Vargas *et al.*, 1990). However, even after the introduction of the axenic yeast extract-iron serum (YI-S) medium for *E. dispar* (Diamond *et al.*, 1995; Clark, 1995) the axenic cultivation of *E. dispar* remains difficult. Compared with *E. dispar*, pathogenic *E. bistolytica*, which is closely related to *E. dispar*, easily adapts to the axenic culture medium (TYI-S-33) (Diamond *et*

al., 1978). Additionally, Entamoeba histolytica is capable of invading the mucosa of the large intestine. We have also developed a new yeast extract-iron-gluconic acid-dihydroxyacetone-serum medium (YIGADHA-S) (Kobayashi et al., 2005) based on the YI-S medium and on the results of an investigation on bacterial metabolic products, and have succeeded in culturing five strains under axenic conditions. However, despite using the YIGADHA-S culture system, the axenic growth of four of five strains of E. dispar was found to be very poor, with the exception of one primate-derived strain (CYNO 09: TPC) isolated from a cynomolgus monkey. A further search for useful growth promoting factors (GPFs) revealed that autoclaved (121°C, 15 minutes) bacteria and more than 20 types of protozoan, mammalian and plant cells containing mitochondria (Mt), hydrogenosomes (Hg) and chloroplasts (Cp) have a significant growth-promoting effect on E. dispar. However, the degree of these growth-promoting effects differs among the GPFs. In the present study, we were able to extract GPF from the Cp-rich leaf cells of a plant (spiderwort: Commelina communis L.) in an acetonesoluble fraction as a chlorophyll complex with low mole-

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cular weight proteins. Significant amounts of iron (Fe), sulphur (S) and molybdenum (Mo) atoms were found in the protein-chlorophyll complex fraction along with the magnesium (Mg) atom of chlorophyll.

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Based on these findings, we hypothesized that some common components of Fe-S proteins present in Mt, Hg, Cp and bacteria support the growth of *E. dispar*. This is because in *E. dispar*, an essential redox Fe-S protein [ferredoxin (Fd)] is expected to be involved in energy metabolism, such as oxidative decarboxylation of pyruvate to acetyl-coenzyme A (acetyl-CoA), which is similar to that found in *E. bistolytica* (McLaughlin & Aley, 1985).

MATERIALS AND METHODS

E. DISPAR ISOLATE

n this study, one human-derived strain of axenically grown E. dispar (AS 16 IR) (Kobayashi *et al.*, 2005) was used to determine the growth promoting effect of Fe-S proteins.

AXENIC CULTIVATION OF E. DISPAR

The YIGADHA-S medium (Kobayashi *et al.*, 2005) containing 15 % heat-inactivated bovine serum was used as the axenic cultivation medium for *E. dispar*.

DONOR CELLS OF MT, HG, CP AND MITOSOMES

The Mt donors are as follows: protozoan parasites: i) Crithidia fasciculata (ReF-1, PRR strain, ATCC 50083); ii) Trypanosoma cruzi (Tulahuen strain); iii) Leishmania major (MHOM/SU/73/5-ASKH strain, ATCC 50155); and iv) Acanthamoeba castellanii (ATCC 30011). v) The vertebrate cells used as Mt donors were from the mouse lymphoblast cell line (P388D1, ATCC CCL-46). The Hg donors are as follows: protozoan parasites: vi) Trichomonas vaginalis (KO-11 strain) (Qi et al., 1995) and vii) Trichomonas foetus (Okamoto et al., 1998). viii) Leaves of the spiderwort (Commelina communis L.) were used as the Cp donors. Consisting amitochondrial protozoan parasite mitosomes that are mitochondrial remnant organelles (Tovar et al., 2003) was used as the mitosome donor, ix) Giardia intestinalis (syn. lamblia), (Portland-I strain, ATCC 30888). The bacterium used was Pseudomonas aeruginosa (PA:KEIO strain) (Kobayashi et al., 1998).

PREPARATION OF CELLS

C. fasciculata and *T. cruzi* were axenically cultured in liver infusion tryptose (LIT) medium (*Gutteridge et al.*, 1969) supplemented with 10 % foetal bovine serum (FBS). After cultivation for three and seven days res-

pectively, 2×10^7 cells of each parasite were harvested by centrifugation (650 g \times 8 minutes). The cells of each parasite were washed three times by centrifugation with 10 mM phosphate-buffered saline (PBS) (pH 7.4), and each pellet was suspended in approximately eight times its volume of 50 mM Tris-HCl (pH 7.4). L. major was axenically cultured in Schneider's medium (Gibco™) supplemented with 15 % FBS at 26°C for four days and harvested by centrifugation (650 g \times 8 minutes). A. castellanii was axenically cultured in peptone-yeast glucose (PYG) medium (Rowbotham, 1983; ATCC media formulations No. 712) for four days and harvested by centrifugation (650 g \times 4 minutes). The P388D1 cell line was cultured in RPMI 1640 medium (Nissui Pharmaceutical Co., Taito-ku, Tokyo, Japan) supplemented with 10 % FBS for four days and harvested by centrifugation (125 g \times 4 minutes). T. vaginalis and T. foetus were axenically cultured for three days in BI-S-33 medium in which the peptone components of TYI-S-33 medium (Diamond et al., 1978) are substituted by Biosate (BBL, Becton Dickinson Co., Cockeysville, Maryland, USA), and harvested by centrifugation (275 g × 4 minutes). The Cp-rich fraction from the wild spiderwort (C. communis L.) was isolated. The leaves of wild spiderwort were picked from the private field of Keio University (Tokyo) during the flower season from June to August (2000-2003). Fresh green spiderwort leaves were stripped from the stems, washed with tap water and then with distilled water. The leaves were ground with serum-free RPMI 1640 medium (RPMI) in an earthenware mortar by using a wooden pestle. The resulting leaf cell suspension was filtered through a double thickness gauze to remove the residue. The cell suspension was then washed three times with RPMI by centrifugation (440 g × 10 minutes). P. aeruginosa was cultured in Biosate-Iron broth (Diamond et al., 1978) for three days and harvested by centrifugation (650 g \times 4 minutes). G. intestinalis was axenically cultured in modified BI-S-33 medium (Keister, 1983) for three days and harvested by centrifugation (275 g \times 4 minutes).

PREPARATION OF MT, HG AND CP FRACTIONS

Mt, Hg and Cp rich fractions were prepared as described by Hogeboom (1955), Opperdoes *et al.* (1984) and Gorham (1955), respectively. Since the methodology for mitosome isolation has not been established, intact cells of *G. intestinalis* were tested for their growth-promoting effect without preparing the mitosome rich fraction. After washing the fractions with RPMI once by centrifugation (440 g \times 10 minutes), each fraction was suspended in 2 ml of RPMI. In order to test for their growth-promoting effect under axenic culture conditions, the fractions were sterilized by autoclaving (121° C, 15 minutes), and 0.2 ml of each suspension was added into the YIGADHA-S medium (5 ml). The

medium was then inoculated with a 0.8 ml suspension of *E. dispar* (AS16IR strain = AS 16 IR) (final density: 200-2,700 amoebae/ml).

NUPAGE® NOVEX 4-12 % BIS-TRIS GRADIENT PEPTIDE GEL ELECTROPHORESIS AND PREPARATIVE SODIUM DODECYL SULFATE-POLYACRYLAMIDE GEL ELECTROPHORESIS (SDS-PAGE)

The lyophilized cell fraction of spiderwort was extracted with 20 %, 40 %, 80 % and 100 % acetone solutions (20 mg/ml) for 10 minutes at 4° C. After centrifugation (17,800 g × 10 minutes, 4° C) of each acetone extract, the supernatant was collected. The supernatants that were extracted with 80 % and 100 % acetone solutions were diluted with distilled water to obtain an acetone concentration of 50 %. Each supernatant was then concentrated by evaporation by using a SpeedVac system (SVC 100; Savant Instruments, Inc., Farmingdale, New York, USA) until evaporated to dryness. The molecular weight of the proteins in each extract was determined by NuPAGE® Novex 4-12 % Bis-Tris gradient peptide (Invitrogen Corp., Carlsbad, California, USA) gel electrophoresis.

The bands of complexes of low molecular weight protein with chlorophyll that were detected in the supernatants of 80 % and 100 % acetone extracts were isolated by preparative SDS-PAGE on a 12 % gel in a Mini Prep Cell (Bio-Rad Laboratories, Inc., Hercules, California, USA) electrophoresis unit. SDS-PAGE was performed according to the procedure described by Laemmli (1970). For NuPAGE® (Novex 4-12 % gradient peptide gel = Novex 4-12 %) each dried and concentrated supernatant of the acetone extract was redissolved in the sample buffer with 50 mM dithiothreitol and heated at 70°C for 10 minutes to reduce the disulfide bonds. For SDS-PAGE (12 % gel) each supernatant was redissolved in the sample buffer with 1 % (v/v) 2-mercaptoethanol (2ME) and boiled for four minutes. Following NuPAGE®, the gel was stained with 2D-silverstain II (Daiichi Pure Chemicals Co. Ltd., Chuo-ku, Tokyo, Japan).

Assay of the acetone extracts of spiderwort leaf cells for a GPF

One millilitre supernatants (obtained after centrifugation; 17,800 g × 10 minutes, 4° C) of the 20 %, 40 %, 80 % and 100 % acetone extracts of the lyophilized cell fraction of spiderwort (20 mg/ml) was dialyzed three times against 200-250 volumes of distilled water for 18 hours by using a Spectra/Por® 3 (3500 MWCO) dialysis membrane for 18 h; the volume of each extract was adjusted to 2.4 ml. Each extract was sterilized by filtration (Sartorius membrane filter; 0.2-nm pore size), and 0.2 ml of each extract was added to the YIGADHA-S medium to test the growth-promoting effect of the extract on *E. dispar* (AS 16 IR).

Analysis using a scanning electron microscopy (SEM)/energy dispersive X-ray analyser (EDX)-integrated analysis system

Following the preparative SDS-PAGE, the complexes of chlorophyll with low molecular weight proteins in the 80 % and 100 % acetone-soluble fractions were dialyzed in the same manner as described above. After lyophilization, the dialyzed complexes were examined using an SEM and EDX-integrated analysis system SEM-EDX III Type N/H (Hitachi Science Systems, Ltd., Hitachinaka, Ibaragi, Japan) in the Nihonbashi laboratory, Hitachi High-Technologies Co., Tokyo, Japan.

PREPARATION OF CHLOROPHYLL-A COMPLEXES WITH PURIFIED IRON-SULPHUR PROTEINS

Purified Fd from spinach (1 mg/ml; Sigma F-3013) and C. pasteurianum (1 mg/ml; Sigma F-7629) and purified rubredoxin (Rd) from *C. pasteurianum* (15.7 mg/ml; Sigma R-2512) were dissolved in 0.6 % 2ME/H2O (20 mg of Fd or 31.4 mg of Rd/4 ml of 0.6 % 2ME), and the Fd and Rd solutions were dialyzed in the same manner as described above. Purified chlorophyll-a (Sigma C-5753) was dissolved in acetone (25 mg/25 ml acetone), and after adding 4 ml of distilled water, the solution was dialyzed in the same manner as described above. The Fd and Rd solutions were mixed with dialyzed chlorophyll-a and incubated for 10 minutes at 25°C. The solutions of chlorophyll-a complexes were then sterilized by filtration, and 0.2 ml of each solution was added to the YIGADHA-S medium to test their growthpromoting effect on E. dispar AS 16 IR.

STATISTICAL ANALYSIS OF GROWTH-PROMOTING EFFECT ON E. DISPAR

All experiments for testing growth-promoting effect were repeated at least twice. The data of each experiment were subjected to one-way analysis of variance (ANOVA) with Dunnett's multiple comparison post hoc test; the levels of statistical significance were taken as p < 0.05 and p < 0.01.

RESULTS

the Mt, Hg and Cp isolated from eight different types of cells were tested to determine whether they promoted the growth of axenically grown *E. dispar* AS 16 IR. The intact cells of *G. intestinalis* and *P. aeruginosa* were also tested for the same. A growth-promoting effect was apparently observed for every fraction except for the fraction of intact cells of *G. intestinalis*. Apparently, the intact cells of *E. histolytica* (HM-1: IMSS clone 6 strain) containing mitosomes (Leon-Avila & Tovar, 2004) also did not produce a

growth-promoting effect (data not shown). Although the growth-promoting effect of the two fractions of Mt from T. cruzi and the Hg fraction from T. vaginalis retained the statistically significant difference by Dunnett's test (p < 0.05), the growth-promoting effect of the other six fractions containing Mt, Hg and Cp was significantly different (p < 0.01 or p < 0.05). In particular, the effect of the Cp fraction from spiderwort exceeded that of the intact cells fraction of P. aeruginosa as shown in Figure 1. Thus, it was concluded that some of the Mt, Hg and Cp fractions, which were obtained under appropriate conditions as well as the intact bacterial cells contain a GPF.

Acetone extraction was used to obtain a water-soluble GPF from Cp-rich leaf cells of some plant species (e.g., spiderwort, cherry and morning glory), which have sufficiently strong leaf-cell membranes that can withstand the leaf crushing and cell isolation process. Figure 2 shows the results obtained on testing the 0 %, 20 %, 40 %, 80 % and 100 % acetone-soluble fractions from lyophilized Cp-rich leaf cells of spiderwort for a growth-promoting effect. The results showed that the 20 % and 80 % acetone-soluble fractions had a stronger growth-promoting effect than the other acetone-soluble fractions (0 %, 40 % and 100 %). The growth-promoting effects of the 20 % and 80 % acetone-soluble fractions were statistically significant at p < 0.01 (Dunnett's test);

however, the effects of the 0 %, 40 % and 100 % acetone-soluble fractions were retained at p < 0.05. The silverstain NuPAGE® analysis of these acetone-soluble fractions vielded a distinct dense protein band (MW 4600) that was common to the 20 %, 40 %, 80 % and 100 % acetone-soluble fractions. The intensity of the bands of the 20 % and 80 % acetone-soluble fractions were stronger than the hands of the other acetone-soluble fractions (40 % and 100 %) (Fig. 3). These results indicated that the efficacy of the growth-promoting effect was correlated with the intensity of the low molecular weight protein band (Figs 2, 3). The band of the complex of the chlorophyll with low molecular weight proteins (green colour) present in both the 80 % and 100 % acetone-soluble fractions was obtained by SDSpreparative electrophoresis; both the fractions showed a growth-promoting effect on E. dispar AS 16 IR (data not shown). The result showed that the band of the complex of chlorophyll with low molecular weight proteins, which was present in the acetone-soluble fractions, contained a GPF. The reduction in the intensity of the low molecular weight protein bands obtained by using the 40 % acetone-soluble fraction was considered to be caused by the incomplete solubility of chlorophyll. Some protein parts are believed to be precipitated together with the insoluble chlorophyll when extraction is performed with an intermediate concen-

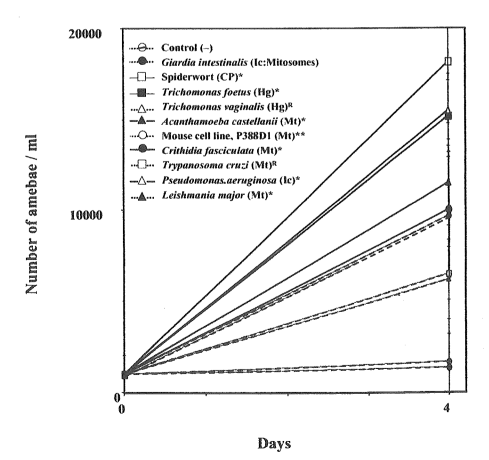


Fig. 1. – The effect of intact cells (Ic) and the fractions of mitochondria (Mt), chloroplasts (Cp) and hydrogenosomes (Hg) from ten types of cells, including bacterial, mammalian, plant and protozoan cells, on the growth of *Entamoeba dispar*.

The cellular components were sterilized by autoclaving at 121° C for 15 minutes. The growth kinetics of the *E. dispar* AS 16 IR strain in the YIGADHA-S medium are shown (mean numbers of amoebae in duplicate cultures are plotted). *, ***: the mean of the growth-kinetic level was significantly higher than that of the control (*p < 0.01 and **p < 0.05 by Dunnett's test). R: the significant difference was retained by Dunnett's test (p < 0.05).

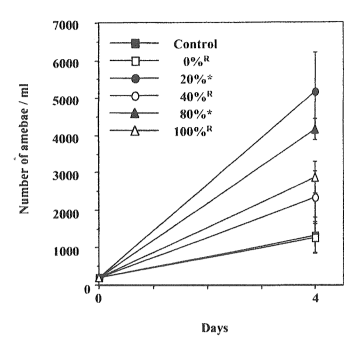


Fig. 2. – Effect of soluble fractions of spiderwort Cp-rich leaf cells extracted with five different concentrations of acetone (0 %, 20 %, 40 %, 80 % and 100 %) on the growth of $\it E. dispar.$

The growth kinetics of the *E. dispar* AS 16 IR strain in the YIGADHA-S medium are shown (mean numbers of amoebae in duplicate cultures are plotted). *: the mean of the growth-kinetic level was significantly higher than that of the control (*p < 0.01 by Dunnett's test). R: the significant difference was retained by Dunnett's test (p < 0.05).

tration of acetone such as 40 %. As compared to an 80 % acetone concentration, the 100 % concentration of acetone was considered to be slightly severe for the extraction of the low molecular weight proteins without

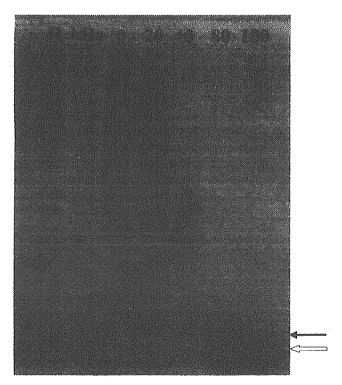


Fig. 3. – NuPAGE® (Novex 4-12 %) of five acetone-soluble fractions of spiderwort Cp-rich leaf cells extracted with different concentrations of acetone (0 %, 20 %, 40 %, 80 % and 100 %).

Molecular mass markers (M) are shown on the left. ← : acetone-soluble low molecular weight protein bands were isolated from the 20 %, 40 %, 80 % and 100 % acetone-soluble fractions. ←: broad green band of chlorophyll.

decreasing the activity of a GPF. As a result, the intensity of the low molecular weight protein bands obtained by using the 40 % and 100 % acetone-soluble fraction was believed to be reduced.

The SEM/EDX-integrated analysis system revealed that the low molecular weight protein hand contained Fe. S and Mo atoms along with the Mg atom of chlorophyll (Figs 4, 5). We believed that Fd present in Cp is a type of Fe-S protein containing Fe and S atoms. In order to confirm the growth-promoting effect of Fd, we examined the effects of purified spinach Fd (MW 12,000), C. pasteurianum Fd (MW 6,000) and purified C. pasteurianum Rd (MW 19,000) on the growth of E. dispar AS 16 IR. The analysis revealed that 0.6 % 2ME-treated C. pasteurianum Rd and each of the watersoluble chlorophyll-a complexes with 0.6 % 2ME-treated spinach Fd and C. pasteurianum Fd have a growthpromoting effect on E. dispar AS 16 IR with a statistically significant difference (p < 0.01) (Fig. 6). The noncomplexed Fd from spinach and from C. pasteurianum treated with 0.6 % 2ME also had a slight growth-promoting effect when compared with the growth kinetics of the control with or without chlorophyll-a; however, these growth-promoting effects were retained (p <

These results suggest that the GPF in the acetone-soluble fraction of spiderwort Cp-rich leaf cells is the complex of the chlorophyll-a with a component of an Fe-S protein, such as an inorganic Fe-S centre, released by the breakage of disulfide bonds due to treatment with 2ME. The Fe-S redox proteins are commonly found in enteric bacteria (e.g., Fd), mitochondria (e.g., Fe-S proteins in complex I and II), chloroplasts (e.g., Fd) and hydrogenosomes (e.g., Fd); however, there are diffe-

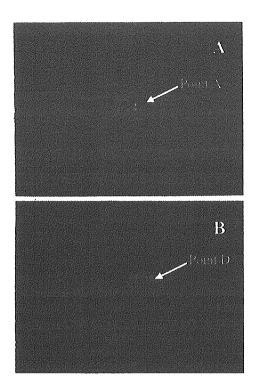


Fig. 4. – Scanning electron microscope (SEM) images of crystals observed in complexes of chlorophyll with low molecular weight proteins present in the 80 % (A) and 100 % (B) acetone-soluble fractions

By using an SEM/energy dispersive X-ray (EDX) analyzer, Fe atoms were detected at point A (A) and S and Mo atoms were detected at point D (B).

rences in the chemical structure of their Fe-S centers. In *E. dispar*, Fd is considered as an essential redox protein involved in energy metabolism, similar to that in *E. histolytica*, although neither species contains Mt. These findings suggest that *E. dispar* grown under axenic culture conditions may lack a sufficient quantity of some essential component of the Fe-S proteins (e.g., Fe-S centre).

The growth-promoting effect of the GPF was tested on three other strains of *E. dispar* (SAW 1734R clone AR, AS 2 IR and CYNO 16:TPC), (Koyabashi *et al.*, 2005); it was confirmed that the GPF produced the same effect on their growth (data not shown).

DISCUSSION

reviously, we designed the YIGADHA-S medium (Kobayashi *et al.*, 2005) for axenic cultivation of *E. dispar*. It contains dihydroxyacetone (DHA) that has a significant growth-promoting effect on *E. dispar*. DHA is a ketotriose and functions as a sugar source for *E. dispar*; therefore, it is considered to be directly metabolized to DHA phosphate, which is an intermediary metabolite in the Embden-Meyerhof-Parnas gly-

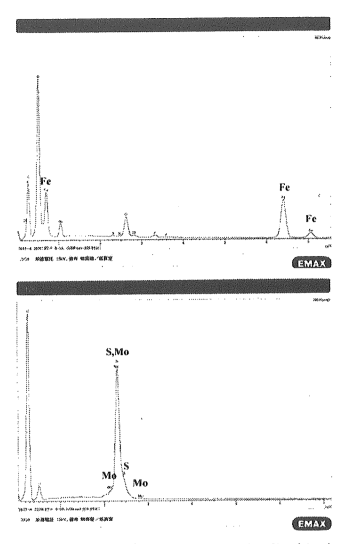


Fig. 5. – Specific peaks of the Fe atom at point A (Fig. 3) and S and Mo atoms at point D (Fig. 3) detected in the complexes of chlorophyll with low molecular weight proteins in the 80 % (A) and 100 % (B) acetone-soluble fractions by using an SEM/EDX analyzer.

colytic pathway. However, the growth of *E. dispar* was poor despite the presence of DHA.

In the present study, a GPF was detected in the autoclaved Mt, Hg and Cp fractions and in intact bacteria and was demonstrated to show a growth-promoting effect on E. dispar. However, the efficacy of each GPF from these organelles varied under different conditions. Prior to the present study, detection of a GPF from trophozoites of E. bistolytica (HM-1:IMSS clone 6 strain) was attempted. Since E. histolytica is genetically closely related to E. dispar, it contains an abundance of Fe-S proteins, such as Fd, which are only stable under anaerobic conditions, and it contains mitochondrial remnant mitosomes (Leon-Avila & Tovar, 2004). However, neither the intact cells nor any extract from the cells could promote the growth of E. dispar AS 16 IR (data not shown). One reason for the failure to extract a GPF from E. bistolytica trophozoites may be the fragility of

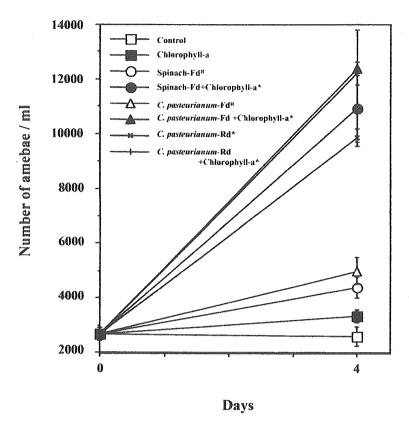


Fig. 6. - Effect of chlorophyll-a (-), 0.6 % 2ME-treated spinach ferredoxin (Fd) (-0-), Clostridium pasteurianum Fd (-0-) and C. pasteurianum rubredoxin (Rd) (-8-), and complexes of chlorophyll a with spinach Fd (-0-), C. pasteurianum Fd (-1-) and C. pasteurianum Rd (-1-) on the growth of E. dispar. and control (-1-).

The growth kinetics of the *E. dispar* AS 16 IR strain in the YIGADHA-S medium are shown (mean numbers of amoebae in duplicate cultures are plotted). *: the mean of the growth-kinetic level was significantly higher than that of the control (*p < 0.01 by Dunnett's test). R: the significant difference was retained by Dunnett's test (p < 0.05).

the cell membrane and organelles (e.g.,mitosomes) similar to that of the spinach leaf cells. The GPF was assumed to be composed of a heat-stable substance and some other common simple inorganic substances. It was feasible to isolate the GPF from acetone-soluble fractions of Cp-rich leaf cells as bands of low molecular weight proteins complexed with chlorophyll containing Fe, S and Mo atoms.

The results of this study indicate that the protein bands may be derived from Fe-S (e.g., Fd) or Mo-Fe-S proteins le.g., molybdoferredoxin, which is a common component of nitrogenase (present in rhizobium species)]. It appears that the GPF requires to form a complex with chlorophyll because the growth-promoting effects of purified Fd from spinach, C. pasteurianum and E. bistolytica strain HM-1: IMSS clone 6 (the Fd consists of 4Fe-4S centre) on E. dispar AS 16 IR were not remarkable (data not shown). We found that the complex of chlorophyll-a with 0.6 % 2ME-treated Fd containing a 2Fe-2S centre from spinach and that containing 4Fe-4S centre from C. pasteurianum have a growth-promoting effect on E. dispar AS 16 IR. This effect was also observed with 0.6 % 2ME-treated Rd, which has the most simple Fe-S centre (1 Fe and 4 cysteine-residues without inorganic S), without chlorophyll-a. It is not clearly understood why complex formation must occur between chlorophyll-a and the components (Fe-S centres and cysteine residues) of bacterial and plant Fd to produce a growth-promoting

effect is not clearly understood. However, it is considered that the complex of chlorophyll-a with the Fd component (MW greater than 100,000 by gel permeation chromatography) may be efficiently taken up by amoeba cells as a compact mass by a process, such as pinocytosis, and the complex may be degraded until an appropriate size of the Fe-S compound is obtained. We speculate that the Fe-S compounds are utilized as components of some essential Fe-S proteins such as amoebic Fd, which is an essential redox protein involved in oxidative decarboxylation of pyruvate to acetyl CoA in E. dispar, this is similar to that reported in E. histolytica (McLaughlin & Aley, 1985). It is reported that the amitochondrial protozoan parasite E. bistolytica, which is closely related to E. dispar, contains numerous fermentation enzymes that are similar to bacterial enzymes, including the pyruvate Fd oxidoreductase, Fd and alcohol dehydrogenase E, derived from concomitant enteric hacterial DNA by horizontal transfer (Rosenthal et al., 1997).

We succeeded in developing a more stable axenic cultivation system by adding the complex of chlorophyll with a component of Fe-S protein (e.g., Fd from spiderwort) treated with 2ME to the YIGADHA-S medium. We achieved an improved axenic cultivation system *for E. dispar* by using cellular components of protozoan, mammalian and plant cells as well as intact bacteria. However, we recognize that it is difficult to develop an axenic cultivation system for *E. dispar* that is at par

with the well-established cultivation system of *E. bistolytica* for use in biological studies such as biochemical and immunological studies, etc.

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SHORT COMMUNICATION

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Entamoeba histolytica and Entamoeba dispar infections in cynomolgus monkeys imported into Japan for research

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Abstract Three hundred and three stool samples of cynomolgus monkeys (Macaca fascicularis) imported from China and the Philippines were examined for Entamoeba histolytica/Entamoeba dispar infections. Microscopy detected E. histolytica/E. dispar cysts in 41 samples. Positive rates were higher in the monkeys from China (37.5%) than in the monkeys from the Philippines (3.7%). PCR analysis of 25 samples successfully cultured from the cysts demonstrated that 24 were E. dispar, one of the samples from China was E. histolytica. The one sample was also identified as E. histolytica by an antigen detection kit, although the monkey was asymptomatic and serology was negative. To our knowledge, this is the first report of E. histolytica isolation from cynomolgus monkeys based on the discrimination between E. histolytica and E. dispar.

Introduction

Amoebiasis, caused by infection with *Entamoeba histolytica*, is one of the most important parasitic diseases of

humans. In addition to symptomatic cases such as hemorrhagic colitis and liver abscesses, asymptomatic infections, in which only cysts are passed in the feces, also exist. Therefore, discrimination between *E. histolytica* and the morphologically indistinguishable but non-pathogenic *E. dispar* is requisite (WHO 1997).

Amoebiasis has also been reported in captive and in wild-trapped non-human primates (Amyx et al. 1978; Beaver et al. 1988). However, recent studies have demonstrated the prevalence of E. dispar infections, but not E. histolytica, in 17 species of captive non-human primates (Smith and Meerovitch 1985), baboons (Jackson et al. 1990), Japanese macaques (Rivera and Kanbara 1999), seven species of captive Old World Macaca monkeys (Tachibana et al. 2001) and chimpanzees (Tachibana et al. 2000). In Japan, cynomolgus monkeys (Macaca fascicularis) have been imported for experimental use in medical research. However, the prevalence of E. histolytica/E. dispar infections, based on discrimination between the two species, is unknown. In the present study, we surveyed imported monkeys for E. histolytica and E. dispar infections.

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Materials and methods

Stool samples were obtained from 215 cynomolgus monkeys from the Philippines by five different shipments and from 88 cynomolgus monkeys from China by four different shipments from 2000 April to 2002 June. Microscopic observation of trichrome-stained stool smears was performed to detect *E. histolytica/E. dispar* cysts. Stools with cysts were cultured xenically in Robinson's medium (Robinson 1968). After 3 days of cultivation, trophozoites were collected by centrifugation using a Percoll-gradient, as described by Tachibana et al. (1990). Genomic DNA of the trophozoites was extracted by a single-tube PCR kit (Takara) and then subjected to PCR amplification using primer sets specific for *E. histolytica* (p11 plus p12) and for *E. dispar* (p13 plus p14), as described by Tachibana et al. (1991). An antigen-capture

ELISA for *E. histolytica* was performed with the *E. histolytica* II kit (TechLab), using cultured trophozoites. Serodiagnosis for *E. histolytica*-infection was performed by an indirect immunofluorescent test, using Amoeba-Spot IF (bioMerieux).

Results

The results of the microscopic and PCR tests are summarized in Table 1. E. histolytica/E. dispar cysts were detected in 13.5% (41/303) of the stools. The positive rates varied among the different shipments and countries from 2.3 to 66.7%. The cyst-positive rate in the shipments from China, 37.5%, was much higher than that from the Philippines, 3.7%. When the 41 cyst positive samples were cultured in Robinson's medium, 25 samples were grown successfully. The main cause of the failure of culture in 16 samples was an outgrowth of Blastocystis hominis trophozoites. PCR analysis of the 25 samples revealed that 24 were E. dispar and one was E. histolytica. No mixed infections were found.

The trophozoites judged as *E. histolytica* by PCR showed a positive OD value of 1.78 in antigen detection ELISA, whereas all the other trophozoites, judged as *E dispar*, had negative OD values of less than 0.05. In the serological tests, none of the monkeys was scored as positive to *E. histolytica*, including the monkey with the *E. histolytica*-infection. The *E. histolytica*-positive monkey was judged to be asymptomatic.

Discussion

Recently, in Japan, *E. dispar* infections, but not *E. histolytica*, were reported in 43% of *M. fuscata* (Rivera and Kanbara 1999); in 66% of captive *Macaca* monkeys (Tachibana et al. 2001); and in 56% of chimpanzees (Tachibana et al. 2000). The dominance of *E. dispar* infections observed in the present study

accords with these previous reports. However, PCR analysis in this study was done from cultured parasites and not directly from fecal samples. Recently, it has been shown that culture in particular underestimates *E. histolytica* infection (Blessmann et al. 2002). Therefore, we cannot exclude the possibility that the prevalence of *E. histolytica* might be biased in this study. The difference of positive rates between China and the Philippines may depend on the hygienic managements of the monkey colonies or may reflect different positive rates in wild macaques.

In the present study, one isolate was identified as E. histolytica based on PCR analysis of the peroxiredoxin gene and antigenicity of the surface lectin (Haque et al. 2000). It might be essential to discriminate the isolate with a closely related parasite, such as Entamoeba chattoni or Entamoeba moshkovskii. Since the cysts in stool smears had four nuclei, infection with E. chattoni was ruled out. In addition, the possibility of E. moshkovskii could be ruled out because the PCR analysis did not amplify peroxiredoxin genes of the parasite (Tachibana et al. 1991; Cheng et al. 2004). To date, a limited number of E. histolytica isolates from non-human primates has been reported, that is, in one Japanese macaque (Tachibana et al. 1990) and in three species of old world and three species of new world monkeys (Verweij et al. 2003). To our knowledge, this is the first report of the isolation of E. histolytica from cynomolgus monkeys based on the discrimination between E. histolytica and E. dispar.

Although the monkey infected with *E. histolytica* was asymptomatic and did not have a positive serology to *E. histolytica*, asymptomatic cyst passers can become symptomatic under immunosuppressive conditions. Furthermore, the *E. histolytica* cysts in the stool of infected monkeys represent a zoonotic hazard to the caretakers. Tests to differentiate between *E. histolytica* and *E. dispar*, followed by successful treatment to exclude *E. histolytica* from the monkeys, are essential for the safe use of monkeys in experiments.

Table 1 Detection of E. histolytica/E. dispar cysts in feces of cynomolgus monkeys and differentiation of both species by PCR analysis of cultured trophozoites

Country Philippines	Shipment	Number of monkeys	Number of positives by microscopy (%)		Number of successful cultures	Number of PCR positives	
			micros	copy (70)	cultures	E. histolytica	E. dispar
			2	(5.1)	1	0	1
* *	2	44	2	(4.5)	1	0	1
	3	44	2	(4.5)	2	0	2
	4	44	1	(2.3)	1	0	1
	5	44	1	(2.3)	1	0	1
Subtotal		215	8	(3.7)	6	0	6
China	1	6	4	(66.7)	3	0	3
	2	50	20	(40.0)	7	0	7
	3	7	2	(28.6)	2	1	1
	4	25	7	(28.0)	7	0	7
Subtotal		88	33	(37.5)	19	1	18
Total		303	41	(13.5)	25	1	24

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ERRATUM

Koichi Koyama

Dendritic cell expansion occurs in mesenteric lymph nodes of B10.BR mice infected with the murine nematode parasite Trichuris muris

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Unfortunately, Table 1 was published with errors. The correct Table 1 is given here.

Table 1 Kinetics of CD11c+ B220- cells in MLNs of Trichuris muris-infected B10.BR mice

Days p.i.	Percentage of positive staining cells ^a				Total cells isolated (×10 ⁷) ^b
	CD11c ⁺ B220 ⁻	CD4 ⁺	CD8+	B220 ⁺	
Uninfected 14 20 25 32	1.0 ± 0.2^{c} 0.9 ± 0.2 $2.0 \pm 0.2^{**}$ 1.0 ± 0.3 0.9 ± 0.1	39.2 ± 1.4 39.1 ± 2.5 30.7 ± 3.0** 31.8 ± 3.2*** 36.7 ± 3.3	23.4 ± 2.5 21.6 ± 2.0 $18.0 \pm 1.3**$ $18.6 \pm 2.3*$ 20.1 ± 2.2	32.0 ± 2.9 35.3 ± 4.9 44.3 ± 2.3** 42.9 ± 4.1*** 38.6 ± 4.7*	2.70 ± 0.77 $4.05 \pm 0.79*$ $6.40 \pm 1.18**$ $4.55 \pm 0.65**$ $4.76 \pm 1.34*$

^aMLNCs were prepared from the MLNs of uninfected or Trichuris muris-infected B10.BR mice on days 14, 20, 25, and 32 p.i. MLNCs were stained with PE-anti-CD11c and FITC-anti-B220 MoAbs, or PE-anti-CD4 and FITC-anti-CD8 MoAbs. Stained cells were then analyzed using a FACScan bTotal numbers of mononuclear cells isolated from the MLNs of uninfected and infected mice

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^cResults represent the mean ± SD for five mice at each time-point and are representative of three independent experiments that gave the

similar results *P < 0.05; **P < 0.01; ***P < 0.001 compared with uninfected controls





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Entamoeba invadens: cysteine protease inhibitors block excystation and metacystic development

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Abstract

We examined the effects of six cysteine protease inhibitors on the excystation and metacystic development of Entamoeba invadens. Excystation, which was assessed by counting the number of metacystic amoebae after the induction of excystation, was inhibited by the cysteine protease inhibitors Z-Phe-Ala-DMK and E-64d in a concentration-dependent manner during incubation compared to the controls. Neither inhibitor had a significant effect on cyst viability; thus, their inhibitory effects were not due to the toxic effect on cysts. Metacystic development, when determined by the number of nuclei in amoeba, was also inhibited by these protease inhibitors, because the percentage of 4-nucleate amoebae was higher than in the controls on Day 3 of incubation. Although other cysteine protease inhibitors, Z-Phe-Phe-DMK, E-64, ALLM, and cathepsin inhibitor III, had a weak or little effect on the excystation, they inhibited cysteine protease activity in the lysates of E. invadens cysts. Broad bands with gelatinase activity of metacystic amoebae, as well as cysts and trophozoites, were detected in the gelatin substrate gel electrophores and were inhibited by Z-Phe-Ala-DMK. There was a difference in the protease composition between cysts and trophozoites, and the protease composition of metacystic amoebae changed from cyst-type to trophozoite-type during development. These results strongly suggest that cysteine proteases contribute to the excystation and metacystic development of E. invadens, which leads to successful infection.

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Index Descriptors and Abbreviations: Entamoeba invadens; Protozoa; Cysteine protease; Excystation; Metacystic development; DMSO, dimethyl sulfoxide; DTT, dithiothreitol; SDS-PAGE, sodium dodecyl sulfate-polyacrylamide gel electrophoresis

1. Introduction

Cysteine proteases are not only important virulence factors, but also play a role in the growth and differentiation in many protozoan parasites (McKerrow, 1989; Rosenthal, 1999; Sajid and McKerrow, 2002). This is also true of *Entamoeba histolytica*, because there is accumulating evidence for the potential role of cysteine proteases in the pathogenesis of invasive amebiasis as well as in the growth of the parasites (reviewed by Que and

Reed, 2000). In *Entamoeba invadens*, which has been used as a model of encystation and excystation of *E. histolytica*, specific cysteine protease inhibitors significantly reduced the efficiency of encystation, although the effect of inhibition was secondary through decreased trophozoite multiplication (Sharma et al., 1996). However, no studies on the role of cysteine proteases in the excystation and metacystic development of *Entamoeba* have so far been reported.

Excystation and metacystic development are necessary for *Entamoeba* infection, and their processes have been described for *E. histolytica* (Cleveland and Sanders, 1930; Dobell, 1928). Since *E. histolytica* does not encyst

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57)

efficiently in axenic culture, E. invadens, a reptilian parasite, has been commonly accepted as a model for the study of encystation and excystation (Eichinger, 1997; López-Romero and Villagómez-Castro, 1993). Excystation is the process through which the whole organism escapes from the cyst through a minute perforation in the cyst wall. Metacystic development is the process in which a hatched metacystic amoeba with four nuclei divides to produce eight amoebulae, which grow to become trophozoites (Cleveland and Sanders, 1930; Dobell, 1928; Geiman and Ratcliffe, 1936). The transfer of E. invadens cysts in an encystation medium to a growth medium induces in vitro excystation (Garcia-Zapien et al., 1995; Makioka et al., 2002; McConnachie, 1955; Rengpien and Bailey, 1975). In this study, we examined the effect of cysteine protease inhibitors on the excystation and metacystic development of E. invadens. Here, we report that cysteine proteases contribute to these processes of E. invadens.

2. Materials and methods

Trophozoites of the IP-1 strain of E. invadens were cultured in an axenic growth medium, BI-S-33 (Diamond et al., 1978), at 26 °C. To obtain cysts, trophozoites $(5 \times 10^5 \text{ cells/ml})$ were transferred to an encystation medium called 47% LG (LG is BI without glucose; Sanchez et al., 1994). After three days of incubation, the percentage of encystation reached 80% on average. The cells were harvested and treated with 0.05% sarkosyl (Sigma Chemical, St. Louis, MO) to destroy the trophozoites (Sanchez et al., 1994). The remaining cysts were washed with phosphate-buffered saline, counted, and then suspended in a growth medium. The viability of the cysts was determined by trypan blue dye exclusion, and the number of nuclei per cyst was determined after staining with modified Kohn's stain (Kumagai et al., 2001). Cyst preparation included 30% dead or denatured cysts and 70% viable cysts, where 4-nucleate cysts are 30% and 1to 3-nucleate cysts are 70%. For the experiments on the excystation and metacystic development of E. invadens, duplicate cultures of 5×10^5 cysts/ml were incubated with inhibitors for three days. Metacystic amoebae were counted in a hemocytometer on Days 1 and 3, and their viability was determined by trypan blue dye exclusion. Viable metacystic amoebae and cysts were clearly distinguished as light yellow and light blue in color, respectively. The former was also identified by positive motility. The cysteine protease inhibitors used in this study, purchased from Calbiochem (San Diego, CA), are listed in Table 1. These inhibitors were previously used in cultures of human fibroblasts infected with Toxoplasma gondii to examine their effect on the intracellular development of parasites (Shaw et al., 2002). All of the inhibitors were dissolved in dimethyl sulfoxide (DMSO).

Table 1
Cysteine protease inhibitors used in the present study

Inhibitor	Specificity		
Cysteine protease inhibitors			
Z-Phe-Ala-DMK	Cysteine proteases		
Z-Phe-Phe-DMK	Cysteine proteases		
E-64	Cysteine proteases		
E-64d	Cysteine proteases		
Calpain inhibitor 2 (ALLM)	Ca ²⁺ -dependent cysteine proteases		
Cathepsin (cysteine) protease inhibitor	•		
Cathepsin inhibitor III	Cathepsin proteases		

Abbreviations: Z-Phe-Ala-DMK, Z-Phe-Ala-diazomethylketone; Z-Phe-Phe-DMK, Z-Phe-Phe-diazomethylketone; E-64d, (2S, 3S)-trans-epoxysuccinyl-L-leucylamido-3-methylbutane ethylester; E-64, N-[N-(L-3trans-carboxirane-2-carbonyl)-L-leucyl]-agmatine; calpain inhibitor 2, N-acetyl-leu-leu-methioninal (ALLM); and cathepsin inhibitor III, Z-Phe-Gly-NHO-Bz-pOMe.

The control cultures received the same volume of DMSO.

Metacystic development was determined by the number of nuclei per amoeba. The cells were harvested on Days 1 and 3 in cultures with or without inhibitors and stained with modified Kohn's stain. The number of nuclei per amoeba was determined by the double-counting of least 100 amoebae.

For the assay of cysteine protease activity, cysts $(2\times10^7/\text{ml})$ were harvested, washed, and subjected to three freeze–thaw cycles in a phosphate-buffered saline. After centrifugation, the supernatants were obtained as lysates. Protease activity was quantified by the cleavage of synthetic peptide substrate Z-Arg-Arg-AMC (benzyloxycarbonyl-arginine-arginine-4-amino-7-methyl-coumarin; Sigma) as previously described (Keene et al., 1986), and recorded as the initial velocity of the cleavage of the fluorescent 4-amino-7-methylcoumarin group/5 μ l lysate. The lysates were preincubated for 15 min at room temperature with 10 and 50 μ M each of the cysteine protease inhibitors as described above.

Protease gel activity was assessed by gelatin substrate gel electrophoresis as previously described (Keene et al., 1986). The cysts and trophozoites were solubilized using a Laemmli sample buffer (Laemmli, 1970) without a reducing agent, and the supernatants after centrifugation were used. To obtain metacystic amoebae with 4-nuclei and 1-nucleus separately, the cysts were transferred to the growth medium with or without 10 µg/ml aphidicolin (Sigma). The cultures with aphidicolin on Day 1 contained a higher percentage of 4-nucleate amoebae, while the cultures without the drug on Day 3 included that of 1-nucleate amoebae (Makioka et al., 2003). Metacystic amoebae in both cultures were lysed by treatment with a small volume of 0.05% sarkosyl, which had no effect on the cysts. The supernatants after centrifugation were then treated with the Laemmli sample buffer. In certain experiments, trophozoites were treated similarly as for metacystic amoebae. SDS-PAGE was conducted in non-



reducing conditions on 10% gels that had been copolymerized with 0.1% gelatin: equivalent to 5×10^4 loaded per lane. After electrophoresis, the gels was washed for 1 h in 2.5% Triton X-100 to remove SDS, rinsed twice in distilled water, and incubated in 100 mM Tris-HCl (pH 7.4) buffer containing 5 mM EDTA and 2 mM DTT with or without 1 mM Z-Phe-Ala-DMK for 12–18 h at 37 °C. After staining with Coomassie blue and several cycles of destaining, the gelatinase activity was detected as clear bands on the Coomassie blue-stained background of the control gels. All of the experiments of this study were performed at least three times and similar results were obtained. Therefore, the data presented in the results are representative.

3. Results

The effect of cysteine protease inhibitors on the number of metacystic amoebae of *E. invadens* after the transfer of cysts to a growth medium is shown in Fig. 1. Among the inhibitors tested, two cysteine protease inhibitors, Z-Phe-Ala-DMK and E-64d, were effective. For this reason, only the results of these inhibitors are shown. The number of metacystic amoebae in cultures with $10\,\mu\text{M}$ Z-Phe-Ala-DMK during incubation was comparable to the controls, whereas it significantly decreased in cultures with 50 and $100\,\mu\text{M}$ Z-Phe-Ala-DMK compared to the controls. Similarly, metacystic amoebae decreased in number during incubation in cultures with more than $50\,\mu\text{M}$ E-64d. The effects of cysteine protease inhibitors on cyst viability are shown in Fig. 2. The number of viable cysts in the control cultures

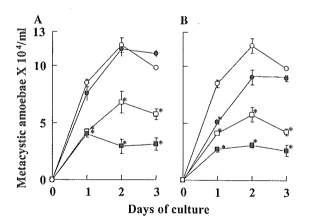


Fig. 1. Effect of cysteine protease inhibitors on the number of metacystic amoebae of *Entamoeba invadens*. Cysts were transferred to a growth medium containing various concentrations of cysteine protease inhibitors Z-Phe-Ala-DMK (A) and E-64d (B). The mean numbers \pm SE of metacystic amoebae for the duplicate cultures are plotted (each asterisk indicates P < 0.05). Concentrations of 0, 10, 50, and 100 μM are indicated by the white circles, black circles, white squares, and black squares, respectively.

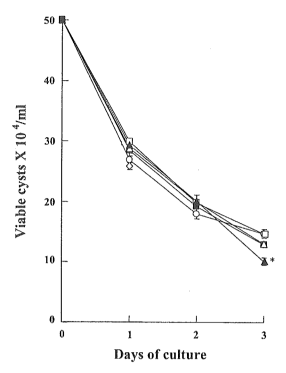


Fig. 2. Effect of cysteine protease inhibitors on the cyst viability of E. invadens in the growth medium. The experimental conditions were the same as those for Fig. 1. The mean numbers \pm SE of viable cysts for the duplicate cultures are plotted (each asterisk indicates P < 0.05). Control (white circles), $50 \,\mu\text{M}$ Z-Phe-Ala-DMK (white squares), $100 \,\mu\text{M}$ Z-Phe-Ala-DMK (black squares), $50 \,\mu\text{M}$ E-64d (white triangles), and $100 \,\mu\text{M}$ E-64d (black triangles).

decreased during incubation. It is considered that most immature cysts contained in culture degenerate or die during incubation. The number of viable cysts in cultures containing 50 and $100\,\mu\text{M}$ Z-Phe-Ala-DMK or E-64d during incubation was comparable to or greater than that of the controls, except for $100\,\mu\text{M}$ E-64d on Day 3.

The effects of cysteine protease inhibitors on metacystic development were examined by counting the number of nuclei per cell. As shown in Fig. 3, 9% of the metacystic amoebae were 4-nucleate on Day 1 of incubation in the control cultures, whereas 29 and 34% of the amoebae were in cultures with 100 µM each of Z-Phe-Ala-DMK and E-64d, respectively. The percentage of 4-nucleate amoebae in the control cultures then decreased to 3% on Day 3, following the increased percentages of 1- to 3-nucleate amoebae. In contrast, the percentage of 4-nucleate amoebae in cultures with Z-Phe-Ala-DMK and E-64d was still 19 and 27% on Day 3, respectively, suggesting the inhibition of metacystic development due to these inhibitors.

Since there was a difference in the inhibitory effect on excystation among the cysteine protease inhibitors used, we examined the effects of these inhibitors on cysteine protease activity in cyst lysates. As shown in Fig. 4,