The Vacuole Binding to Mitochondria by VIG1
Contributes an Equal Inheritance of the Vacuoles in
Cyanidioschyzon merolae

Takayuki Fujiwara, Fumi Yagisawa, Mio Ohnuma, Yamato Yoshida,
Masaki Yoshida, Keiji Nishida, Osami Misumi, Haruko Kuroiwa and
Tsuneyoshi Kuroiwa*

Research Information Center for Extremophiles, Rikkyo University,
3–34–1 Nishiikebukuro, Toshimaku, Tokyo 171–8501, Japan

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Summary  Vacuoles function in endocytosis, storage and digestion of metabolites in eukaryotic
cells. They are inherited by the daughter cells. However, the mechanisms of vacuole inheritance
are poorly understood because the cells contain multiple vacuoles that behave randomly. Cyanidioschyzon
merolae cell has a minimum set of organelles. The vacuoles were equally inherited
by the daughter cells by binding to dividing mitochondria. The binding was mediated by VIG1.
However, the role of the binding in the vacuoles inheritance was poorly understood. We examined it
by inhibiting the binding cytochemically. The vacuoles, which were not bound to mitochondria, were
not equally inherited by the daughter cells. As the results, vacuole-less daughter cells were
generated. These results suggested that the binding contributed the equal inheritance of vacuoles and
ensured the permanence of vacuoles in daughter cells.

Key words  Lysosome, Mitochondria, Inter-organelle contact, Organelle inheritance, Red alga.

The inheritance of organelles is an essential feature of eukaryotic cell division. In mammals,
plants, and yeasts, lysosomes/vacuoles are inherited by the daughter cells (Bergeland et al. 2001,
Kutsuna et al. 2003, Han et al. 2003, Weisman 2006). However, the molecular mechanisms were
not established because the cells contain multiple vacuoles that behave randomly. Cyanidioschyzon
merolae cells offer many advantages for studies of vacuole inheritance. Each C. merolae cell has a
minimum set of organelles comprising 1 plastid, 1 mitochondrion and a few vacuoles. The
organelles are divided in order of a plastid, mitochondrion and a cell nucleus during the cell
division (Kuroiwa 1998). The vacuoles are inherited by daughter cells as binding with dividing
mitochondria (Misumi et al. 2005, Yagisawa et al. 2007, Fujiwara et al. 2010). These organelle
divisions can be highly synchronized by a light/dark cycle (Suzuki et al. 1994).

The transport and binding of vacuoles are not dependent on actin filaments and microtubules
(Takahashi et al. 1995, Matsuzaki et al. 2004, Yagisawa et al. 2007). Our previous study reported
that the coiled-coil protein VIG1 is essential for the transport and binding of vacuoles during the
inheritance in C. merolae (Fujiwara et al. 2010). VIG1 appears on the surface of free vacuoles in
the cytosol in the S phase, and then tethers the vacuoles to the mitochondria during the G2 and M
phases. The vacuoles are released from the mitochondrion in the daughter cells following VIG1
digestion in early G1 phase. By suppression of VIG1 by antisense RNA, migration of vacuoles was
disturbed (Fujiwara et al. 2010). However, a relationship between quantity and function of VIG1
during the processes of vacuole inheritance was not examined. Furthermore, it is thought that the
interaction between vacuoles and mitochondria played some kind of important role during the

* Corresponding author, e-mail: tsune@rikkyo.ne.jp
inheritance of vacuoles, but the role was poorly understood.

Cell cycle period-specific treatment with a translation inhibitor can suppress expression of VIG1 to various levels, since organelle divisions can be highly synchronized in *C. merolae*. In this study, we examine the function of VIG1 and the inheritance of vacuoles that are not migrated to mitochondria by the treatment of a translation inhibitor.

Materials and methods

**Synchronous culture and cycloheximide treatment**

*Cyanidioschyzon merolae* 10D-14 were synchronized according to the method of Suzuki et al. (1994). Cells were cultured in 2x Allen's medium at pH 2.3 (Allen 1959). Flasks were shaken under continuous light (40 W/m²) at 42°C. The cells were sub-cultured to 10⁷ cells/ml, and then synchronized by subjecting them to a 12 h light/12 h dark cycle at 42°C while the medium was aerated. For cycloheximide (chx) treatment, 0.5 M chx dissolved in ethanol was added to synchronized culture at 14 h at a dilution of 1 : 500. For the control experiment, ethanol was added in same condition. The cells were cultured in medium containing chx and/or ethanol until the cells were harvested for the immunoblotting and immunofluorescent microscopy analyses.

**Immunoblotting**

The antisera or antibodies against Dnm1, Dnm2 and VIG1, were previously prepared (Nishida et al. 2003, Miyagishima et al. 2003, Fujiwara et al. 2010). Immunoblotting was performed by conventional methods. *C. merolae* cells were harvested at each time from initiation of the synchronized culture and lysed in conventional SDS containing sample buffer. Thirty micrograms of proteins of each sample were separated by SDS-PAGE, and then were immunoblotted. The primary antisera or antibodies against Dnm1, Dnm2 and VIG1 were each used at a dilution of 1 : 1000. The secondary antibodies, alkaline phosphatase-conjugated goat anti-guinea pig or anti-rabbit IgG (Kirkegaard and Perry Laboratories) were each used at a dilution of 1 : 3000.

**Fluorescence microscopy**

Cells were harvested during the mitotic phase from synchronized cultures. After fixation and blocking of the cells (Nishida et al. 2004), VIG1 was stained with anti-VIG1 primary antibodies at a dilution of 1 : 200, and secondary goat anti-guinea pig IgG conjugated with Alexa 488 at a dilution of 1 : 100 (Fujiwara et al. 2010). Vacuoles were stained with anti-V-ATPase primary antibodies at a dilution of 1 : 100 and secondary goat anti-rabbit IgG conjugated with Alexa 555 at a dilution of 1 : 100 (both from Molecular Probes) (Yagisawa et al. 2009). Mitochondria were stained with anti-POR primary antisera at a dilution of 1 : 200, and secondary goat anti-rat IgG conjugated with Alexa 488 at a dilution of 1 : 300 (Fujiwara et al. 2010). Cells were observed using a fluorescence microscope (Olympus BX51) with a combination of narrow bandpass filter sets: BP470-490 BA510-550 (U-MNIBA2, Olympus) for Alexa-488, BP541-551 BA565-595 (XF37, Omega) for Alexa-555, and using a Hg arc lamp as a source of excitation light. Images were collected using a 3 charge-coupled device (3CCD) camera system, C7780-10 (Hamamatsu Photonics; Tokyo, Japan) and processed using the M8458-03 RCA-3CCD Photoshop plug-in software. The change of color channels and adjustment of fluorescence and merging of images were performed by Photoshop CS2 (Adobe Systems Inc., California, USA).

Results and discussion

The cell cycle of *C. merolae* was synchronized by 12 h light/12 h dark cycles. Immunofluorescent microscopy showed that behaviors of vacuoles and mitochondria during cell cycle (Fig.
The binding between vacuoles and mitochondria was inhibited by cycloheximide treatment targeting VIG1. (A) Phase-contrast and immunofluorescent images showing the inheritance of vacuoles. Vacuoles and mitochondria were shown in red and yellow, respectively. Ethanol was added as a control to chx experiment. (B) Schematic model showing the inheritance of vacuoles. C. merolae cell has 1 cell nucleus, 1 mitochondrion, 1 plastid and a few vacuoles. Vacuoles were located in the nucleus side of cytosol in the G1 phase, and then were transported to the mitochondrial surface by the G2 phase and continued to bind there during the M phase. Vacuoles were transported with divided mitochondria to daughter cells, and then vacuoles were released from the mitochondria in the early G1 of daughter cells. (C) Immunoblotting of Dnm1, Dnm2 and VIG1 during cell cycle in chx-treated cells. The phases of cell cycle and time from onset of synchronous culture were shown on the immunoblot. (D) Phase-contrast and immunofluorescence images showing expression of VIG1 and localization of vacuoles in chx experiment. VIG1 and vacuoles were shown in green and red, respectively. VIG1/Vc, immunofluorescence images of VIG1 overlaid with those of vacuoles. (E) Percentage of the migration of vacuoles in chx-affected cells. The x axis shows categories based on the percentage of vacuoles not migrated to the mitochondrial region in each mitotic cell. The y axis shows percent of cells falling in each category. In chx-affected cells, more than forty percent of vacuoles were not migrated to the mitochondrial region. Control: n=30; chx-affected: n=20. Bars indicate SD. M-meta, M-ana and M-telo shows metaphase, prophase and telophase in the M phase, respectively. Scale bars in A and D=2 μm.
1A). They were summarized in a schematic model (Fig. 1B). The vacuoles were distributed in the cytosol during the G1 phase and became bound to the dividing mitochondrion by the G2 phase. During the M phase, they were inherited by the daughter cells by binding to the dividing mitochondria. In the early G1 phase, the vacuoles were released from the mitochondrion and returned to the cytosol in the daughter cells (Fig. 1A and B).

In C. merolae, we already established an antisense-based suppression system. However, the inheritance of vacuoles that were not migrated to mitochondria could not be analyzed, because the cell cycle was not synchronized under the system. We tried, therefore, to inhibit the migration of vacuoles by period-specific inhibition of translation in the synchronous culture and examined the inheritance of vacuole including new daughter-cells after cell division.

To suppress VIG1, cycloheximide (chx), an inhibitor of translation, was added to synchronized cultures at 12, 14 or 16 h at final concentrations of 100 μM or 1 mM. The time point of 14 h and concentration of 1 mM were selected as the most efficient combination. To show selectivity for suppression of VIG1 in chx experiment, Dnm1 (dynamin for mitochondrial division constantly expresses during cell cycle) and Dnm2 (dynamin for plastid division mainly express during G2 and M phases) was used as controls. Immunoblotting showed that Dnm1 constantly expressed during cell cycle, Dnm2 and VIG1 mainly expressed in G2 and M phases in control experiment. In chx experiment, Dnm1 was not suppressed and Dnm2 was not completely suppressed, while VIG1 was almost suppressed (Fig. 1C). The cells harvested at 18 h were fixed and then immunostained. Immunofluorescence microscopy with antibodies against VIG1 and V-ATPase (Vacuole H^+-ATPase) showed that the localization of VIG1 and the behavior of vacuoles in chx experiment (Fig. 1D). In control cells, VIG1 was localized at the vacuoles migrated to the mitochondrial region while expression of VIG1 and the migration of vacuoles were suppressed in chx-affected cells (Fig. 1D). Chx did not inhibit the plastid division but the vacuole migration because Dnm2 was not suppressed by chx-treatment (Fig. 1C and D). Statistical data showed that more than half of the vacuoles were not migrated to the mitochondrial region in chx-affected cells (Fig. 1E).
migration of vacuoles was successfully inhibited by suppression of VIG1 in chx-treatment (Fig. 1D and E).

To examine period of VIG1 synthesis and relationship between quantity of VIG1 and the behavior of vacuoles, the 3 cells that varied in quantity of VIG1 were shown in a same field (Fig. 2A). Difference in VIG1 quantity was probably caused by the stage of the cell cycle in each cell when chx had been added, because cell cycle stages of individual cells slightly varied within control culture. Vacuoles had been migrated to the mitochondrial surface by the G2 phase in a control (Fig. 1A and B), but they did not migrate in chx-treated G2 cells that lacked VIG1 (Fig. 2A G2). In the prophase cell that VIG1 was weakly expressed, the some vacuoles failed to migrate to the mitochondrial region (Fig. 2A M-pro). In the prometaphase cell that VIG1 was substantially expressed, the vacuoles were completely migrated (Fig. 2A M-prometaphase). The G2 cell lacking VIG1 in chx experiment suggested that VIG1 had been synthesized 4 h before G2 phase. Therefore, the period of VIG1 synthesis was predicted to be S phase according to the examination of temporal relationship among organelle division cycle in the previous study (Imoto et al. 2010). Moreover, the percentage of vacuole on the mitochondrial region related to quantity of VIG1 between the prophase and prometaphase cells, suggesting that VIG1 was also involved in binding between vacuoles and mitochondria.

Furthermore, we examined the inheritance of vacuoles that failed to migrate such as the G2 cell. Immunofluorescent microscopy showed that the migration of vacuoles failed through the cell cycle without inhibiting mitochondrial and plastid division in chx-affected cells (Fig. 2B). In the telophase, the vacuoles, remaining in the cytosol, were not equally inherited by the daughter cells (Fig. 2B M-telo). As a result, vacuole-less daughter cells were generated (Fig. 2C). These results suggested that the binding between vacuoles and mitochondria by VIG1 contributed the equal inheritance of vacuoles, and ensured the permanence of vacuoles in daughter cells.

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