Interaction of PLP with GFP-MAL2 in the Human Oligodendroglial Cell Line HOG

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Abstract

The velocity of the nerve impulse conduction of vertebrates relies on the myelin sheath, an electrically insulating layer that surrounds axons in both the central and peripheral nervous systems. Oligodendrocytes are the myelin-producing glial cells in the central nervous system. A deeper understanding of the molecular basis of myelination and, specifically, of the transport of myelin proteins, will contribute to the search of the aetiology of many dysmyelinating and demyelinating diseases, including multiple sclerosis. Recent investigations suggest that proteolipid protein (PLP), the major myelin protein, could reach myelin sheath by an indirect transport pathway, that is, a transcytotic route via the plasma membrane of the cell body. If PLP transport relies on a transcytotic process, it is reasonable to consider that this myelin protein could be associated with MAL2, a raft protein essential for transcytosis. In this study, carried out with the human oligodendrocytic cell line HOG, we show that PLP colocalized with green fluorescent protein (GFP)-MAL2 after internalization from the plasma membrane. In addition, both immunoprecipitation and immunofluorescence assays, indicated the existence of an interaction between GFP-MAL2 and PLP. Finally, ultrastructural studies demonstrated colocalization of GFP-MAL2 and PLP in vesicles and tubulovesicular structures. Taken together, these results prove for the first time the interaction of PLP and MAL2 in oligodendrocytic cells, supporting the transcytotic model of PLP transport previously suggested.

Introduction

The myelin sheath is an electrically insulating layer that surrounds axons in both the central and peripheral nervous systems. Oligodendrocytes (OLs) are the glial cells that produce myelin in the central nervous system (CNS) [1,2]. The presence of myelin sheath and its discontinuities, the nodes of Ranvier, allows saltatory conduction of action potential. In the absence of myelin, the velocity of nerve impulse conduction would be pathologically slow. To form the myelin sheath, OLs wrap their processes - extensions of the plasma membrane- around the axons [3], giving rise to different membrane domains and subdomains [4]. The various subdomains of OLs plasma membrane are not separated, as it occurs with basolateral and apical domains of epithelial polarized cells. Nevertheless, the myelin composition is drastically different from that of the plasma membrane of the cell body since, similar to the apical membrane of epithelial cells, myelin sheath is rich in glycosphingolipids (GSLs) and cholesterol [5]. Therefore, although myelinating OLs do not polarize segregating typical apical and basolateral surface subdomains, they can be considered as polarized cells [6].

The formation of the myelin sheath in the CNS is a highly complex process which involves the synthesis, transport, and target of large amounts of membrane proteins and lipids by OLs [7].

During OLs differentiation, several proteins and lipids segregate to form the myelin sheath. In spite of myelin composition, typical of the apical plasma membrane of polarized cells, studies in vitro showed that myelin sheet biogenesis has features of basolateral traffic. In this regard, vesicular stomatitis virus G protein (VSV-G), a basolateral marker, accumulated in the myelin sheet, whereas influenza virus hemagglutinin (HA), an apical marker, accumulated in the plasma membrane of the cell body, suggesting that the myelin membrane is the target of a basolateral-type pathway [8,9].

PLP, the major myelin protein, is an integral membrane protein with four transmembrane domains. PLP and DM20, a smaller isoform generated by alternative splicing, are the most abundant proteins in the CNS myelin, comprising the 50% of total myelin proteins [1]. PLP has been associated with the low-density CHAPS-insoluble membrane fraction in cultured OLs [10], although integration of PLP into different membrane domains is a dynamic process that depends on the trafficking stage. OLs lacking PLP are still capable of myelinating axons, although physical stability of myelin decreases, since PLP is responsible for the compaction of myelin sheaths [11]. Mutations of the PLP gene cause dysmyelinating diseases in man and animals, such as Pelizaeus-Merzbacher disease, an X-linked recessive leukodystrophy [12,13].

Crucial points on PLP traffic have still to be elucidated regarding its transport to the myelin sheath. After its synthesis in
of PLP in GFP-MAL2/HOG cells

1. Overexpression of MAL2 does not modify localization of PLP in GFP-MAL2/HOG cells

In a previous study carried out by our group, we performed the characterization of MAL2 in oligodendrocytic cells [26]. Now, we address the relationship of PLP with MAL2 proteolipid. In an initial phase, for that purpose, we used the cell line HOG stably transfected with GFP-MAL2, a construct encoding a chimera consisting of GFP fused to the amino-terminal end of MAL2. To study the localization of endogenous PLP in this GFP-MAL2 overexpressing cell line, we performed double-label indirect immunofluorescence analysis with anti-PLP monoclonal antibody and P239 anti-MAL2 antibody (Figure 1A) or PLP antibody alone (Figure 1B). Results showed that localization of PLP in GFP-MAL2/HOG cells was similar to that observed in HOG cells. Thus, cultivating of cells under differentiation conditions had a similar effect on HOG and GFP-MAL2/HOG cell lines. In both lines, endogenous and exogenous MAL2 were up-regulated when cultured in differentiation medium (DM) and culturing of both cell lines in differentiation conditions resulted in an increase of PLP detection. In addition, in both cell lines PLP was detected in the plasma membrane, in vesicles scattered throughout the cytoplasm and in myelin-like sheets. In some differentiated cells the presence of MAL2 and GFP-MAL2-positive vesicles containing PLP located near the plasma membrane was also observed (Figure 1).

2. Partial localization of PLP in LEs/Lys

To investigate whether, in our system, PLP is stored in le endosomes/lysosomes (LEs/Lys) after its internalization into the cells in the absence of neurons, as it occurs in other oligodendrocytic cells [27], GFP-MAL2/HOG cells cultured in DM at 37 °C for 2 days were fixed and processed for confocal double-label indirect immunofluorescence analysis with anti-PLP monoclonal and anti-LAMP1 polyclonal antibodies. Results showed colocalization of both PLP and LAMP1 in some vesicles (Figure 2), although most of PLP did not localized to LAMP1-positive structures. In GFP-MAL2/HOG cells, only a small fraction of PLP is detected in LEs/Lys, suggesting that accumulation of PLP in these endosomes does not take place in this system.

3. PLP interacts with GFP-MAL2 in GFP-MAL2/HOG cells

To study the relationship between PLP and GFP-MAL2, GFP-MAL2/HOG cells cultured in DM at 37 °C for 2 days were fixed and processed for confocal double-label indirect immunofluorescence analysis with anti-PLP monoclonal antibody. Colocalization of PLP and GFP-MAL2 was observed in numerous vesicles throughout the cytoplasm and the myelin-like sheets (Figure 3A). Moreover, GFP-MAL2-positive vesicles containing PLP could also be found close to the plasma membrane in some cells and in the extracellular medium (Figure 3B), suggesting that GFP-MAL2-positive vesicles containing PLP could be released outside the cell.

Previous studies have demonstrated that exit of cargo from the ARE/SAC and further transport to the apical surface is reduced at 18 °C [23,28]. Thus, transcytosing molecules accumulate in apical recycling endosomes under such conditions. To improve conditions of colocalization of PLP with GFP-MAL2, we cultured GFP-MAL2/HOG cells in DM at 37 °C for 2 days, and then, at 18 °C for 120 min. Thereafter, cells were fixed and processed for confocal double-label indirect immunofluorescence analysis with an anti-PLP monoclonal antibody (Figure 4A). As results showed, under these conditions there is an increase in the number of cells exhibiting colocalization of PLP with GFP-MAL2 in a pericentrosomal compartment.

It has been shown that CD59, a transcytotic glycosylphosphatidylinositol (GPI)–anchored protein, accumulated in the ARE/SAC of HepG2 cells and colocalized with MAL2 at 18 °C [19]. In GFP-MAL2/HOG cells, CD59 colocalized at 18 °C with GFP-MAL2 in the GFP-MAL2-positive compartment [26]. Accordingly, to assess the colocalization of PLP with CD59, we cultured GFP-MAL2/HOG cells in DM for 2 days and, then, at 18 °C for 120 min. Thereafter, cells were fixed and processed for confocal double-label indirect immunofluorescence analysis with an anti-PLP polyclonal antibody and Alexa Fluor
647-conjugated secondary anti–mouse antibody to detect anti–CD59 primary antibodies. Confocal images showed colocalization of GFP-MAL2, PLP and CD59 scattered in small regions of the GFP-MAL2-positive compartment (Figure 4B).

To determine whether PLP and GFP-MAL2 are in fact interacting in our system, we conducted co-immunoprecipitation studies by using a monoclonal anti-PLP or a polyclonal anti-GFP antibody. Controls included preimmune mouse or rabbit serum, the supernatants recovered after the centrifugation of the immunoprecipitate, a control without primary antibody and the whole cell lysates. SDS-PAGE analysis of the extracts immunoprecipitated with anti-PLP monoclonal antibody showed that a fraction of GFP-MAL2 coimmunoprecipitated with PLP (Figure 5), suggesting that at least a fraction of PLP is interacting with GFP-MAL2. As shown in Figure 5, the reverse assay confirmed the interaction, finding PLP in extracts immunoprecipitated with anti-GFP polyclonal antibody.

4. PLP colocalized with GFP-MAL2 after its internalization from the plasma membrane

Several studies suggest that PLP could reach the myelin sheath by indirect transport, that is, via the plasma membrane of the cell body. Therefore, if PLP travels by transcytosis, internalization from this domain will occur. After its internalization from the cell body (Figure 1A and B), it is expected to colocalize with GFP-MAL2, which is localized in the myelin sheath by indirect transport. Indeed, confocal images showed colocalization of GFP-MAL2, PLP and CD59 scattered in small regions of the GFP-MAL2-positive compartment (Figure 4B).
body plasma membrane, PLP would travel to the myelin sheath probably passing through other compartments involved in transcytosis, such as the MAL2-positive compartment. To study the intracellular transport of PLP from the cell surface, we performed a cell surface biotinylation assay. Cell surface proteins of GFP-MAL2/HOG cells were biotinylated at 4°C using a Sulfo-NHS ester of biotin. After biotinylation, cells were cultured in DM either for 1 hour at 4°C, or at 37°C for 20, 40, 60 and 150 min. Cells were then fixed and treated for triple-labelled indirect immunofluorescence analysis with an anti-PLP monoclonal antibody and streptavidin-Alexa Fluor 555, to visualize biotinylated proteins. Our results showed that endocytosed PLP and GFP-MAL2 colocalized in vesicles of cells incubated for 150 min (Figure 6). Vesicles positive for PLP and GFP-MAL2 were also observed in cells incubated even for 20 min, although at a lesser extent.

5. PLP and GFP-MAL2 colocalized with actin cytoskeleton

We have previously shown that GFP-MAL2/HOG cells cultured in DM exhibited a GFP-MAL2-positive tubular reticulum arising from the peri-centrosomal region and extending towards processes [26]. Since it has been demonstrated that MAL2-positive vesicular carriers associate with actin filaments during transcytosis [29], we decided to investigate whether the GFP-MAL2-positive tubular reticulum and the vesicles positive for GFP-MAL2 were also associated with the actin cytoskeleton. In addition, we investigated whether the vesicles containing GFP-MAL2 and PLP were also associated with actin filaments. To do that, GFP-MAL2/HOG cells cultured in DM at 37°C for 2 days were fixed and processed for confocal triple-labelled indirect immunofluorescence analysis with anti-PLP monoclonal antibody and Phalloidin-TRITC to stain F-actin. Our results showed that a high proportion of GFP-MAL2-positive vesicles contained F-actin (Figure 7A). In addition, colocalization of PLP and F-actin was also observed in some GFP-MAL2-positive vesicles (Figure 7B). Herein, we also found that GFP-MAL2-positive tubular reticulum colocalized with F-actin (Figure 7C). In addition, several PLP-positive vesicles were also associated with this tubular reticulum (Figure 7C).

6. Electronmicroscopy (EM) immunocolocalization of PLP and GFP-MAL2

Finally, to confirm colocalization of PLP with GFP-MAL2 and to determine the ultrastructure of the organelles positive for these two proteins, we performed double-labelled immunoelectron microscopy on ultra thin cryosections. GFP-MAL2/HOG cells cultured at 37°C in DM for 2 days were processed for ultra thin cryosectioning and immuno-labeling with a rabbit polyclonal anti-GFP antibody and a mouse monoclonal anti PLP antibody. Anti-GFP and anti-PLP were detected with 10nm protein A-gold and 15nm anti-mouse-gold, respectively. Colocalization of PLP and GFP-MAL2 was confirmed in vesicles located near the plasma membrane (Figure 8A); in scattered cytoplasmic vesicles (Figure 8B); in tubulovesicles near the nucleus (Figure 8C) and in a tubulovesicular membrane-bound compartment (Figure 8D).

Discussion

In a previous work [26], we characterized a MAL2-positive compartment in the oligodendrocytic cell lines Oli-neu and HOG, showing that, after differentiation, these cells up-regulate the expression of MAL2, which is accumulated in an intracellular compartment exhibiting some of the main features of the ARE/SAC, such as colocalization with Rab11a and CD59, peri-centrosomal location, tubulovesicular morphology, sensitivity to disruption of the microtubule cytoskeleton with nocodazole and lack of internalized transferrin. These results suggested that the MAL2-positive compartment in oligodendrocytic cells could be a structure analogous to the ARE/SAC seen in epithelial cells, and might therefore have an important role in the traffic of myelin proteins during oligodendrocytic polarization. Given its crucial role in transcytosis, it is reasonable to hypothesise about an involvement of MAL2 in the traffic of certain myelin proteins. This possibility prompted us to study the relationship between myelin proteins and GFP-MAL2 in our system.

Previous reports have provided data suggesting that PLP could reach myelin sheath by an indirect pathway, that is, a transcytotic route similar to that seen for many apical proteins in epithelial cells, via the basolateral surface [6,10,14,27]. Motivated by this

Figure 2. Localization of PLP and LEs/Lys in GFP-MAL2/HOG cells. Cells cultured in DM at 37°C for 2 days were fixed and processed for confocal double-labelled indirect immunofluorescence analysis with anti-PLP monoclonal and anti-LAMP1 polyclonal antibodies. Monoclonal and polyclonal antibodies were detected using Alexa Fluor 647 and Alexa Fluor 555 secondary antibodies respectively. Positive vesicles for both PLP and LAMP1 can be seen (arrows), although most of PLP signal does not localize to LAMP1-positive Les/Ls. Images correspond to a confocal slice of 0.8 μm. The squares show enlarged images. Scale bar = 10 μm. doi:10.1371/journal.pone.0019388.g002
hypothesis we decided to investigate the relationship between endogenous PLP and MAL2 in our oligodendrocytic cells using the human HOG cell line. It is known that mature OLs form “myelin-like” membranes in vitro [5,30]. Likewise, the HOG cell line develops myelin-like sheets when cultured under differentiation conditions and, correspondingly, these structures accumulate myelin proteins such as MAL [26] and, as we show herein, PLP. To improve MAL2 detection, we stably transfected HOG cells with GFP-MAL2, a construction already proven to be a useful tool to study MAL2 in other epithelial [20,29] and oligodendrocytic [26] systems. Accordingly, our results show that overexpression of MAL2 does not modify significantly the localization of PLP in GFP-MAL2/HOG cells, since, in both cell lines, this proteolipid was detected in the plasma membrane, in vesicles scattered throughout the cytoplasm and in myelin-like sheets. In addition, culturing both HOG and GFP-MAL2/HOG in differentiating conditions resulted in an increase of PLP detection.

It has been demonstrated in oligodendroglial cells that, after its endocytosis, PLP is targeted to LEs/Lys, where, in absence of neuronal signals, it is stored [27,31]. A cAMP-dependent neuronal signal triggers the retrograde transport of PLP from LEs/Lys to the surface of oligodendrocytes. In contrast with those results, in our system PLP showed a slight colocalization with LEs/Lys, since most of PLP could not be detected in LAMP1-positive endosomes. Why accumulation of PLP in Lys does not take place in our system will have to be ascertained. It seems clear that the use of diverse cell systems in the different studies, with their particular features and maturation stages, is a key element probably involved...
Figure 4. Colocalization of PLP and GFP-MAL2 in GFP-MAL2/HOG cells cultured at 18°C. Cells were cultured in DM at 37°C for 2 days, and subsequently, at 18°C for 2 h. After that, cells were fixed and processed for confocal double-label indirect immunofluorescence analysis with anti-PLP monoclonal antibody and Alexa Fluor 555 secondary antibody. A. PLP colocalizes with GFP-MAL2 in pericentrosomal compartments (squares). Image corresponds to the projection of two 0.8 μm confocal planes. Panels B and C are enlargements of the squares shown in panel A, corresponding to confocal slices of 0.8 μm. Vertical x-z sections taken at the plane indicated by the line in the corresponding horizontal section are also shown. Nuclei were stained with TO-PRO-3. D. Cells were cultured in DM at 37°C for 2 days, and then, at 18°C for 2 h. After that, cells were fixed and processed for confocal triple-labelled indirect immunofluorescence analysis with anti-PLP polyclonal and anti-CD59 monoclonal antibodies. Monoclonal anti-CD59 antibody was detected using an Alexa Fluor 647 secondary antibody. Colocalization of PLP, CD59 and GFP-MAL2 can be seen in a compartment located at the pericentrosomal region (arrow). Cell periphery is outlined. Images correspond to a confocal slice of 0.8 μm. Scale bars = 5 μm.

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or affecting the distinct behaviour of PLP during its transit to the myelin sheets.

After its synthesis, PLP is transported to the Golgi by vesicular traffic. Nevertheless, the way by which PLP reaches the myelin sheath remains to be defined. In this regard, several studies suggest that PLP could reach the myelin sheath indirectly via the plasma membrane of the cell body rather than directly from the Golgi. Therefore, if PLP travels by transcytosis, after its transport from the Golgi to the cell body plasma membrane, internalization from this domain might take place. Accordingly to this hypothesis, it has been observed that PLP present in the cell body plasma membrane is internalized and stored into LEs/Lys in absence of neuronal signals [27]. After its internalization from the cell body plasma membrane, PLP would travel to the myelin sheath possibly passing through other compartments involved in transcytosis. Therefore, after its endocytosis from this domain, it would be also feasible to find PLP associated to the MAL2-positive compartment, given that MAL2 is a raft protein essential for transcytosis in other systems. By means of confocal microscopy, we have demonstrated colocalization of PLP and GFP-MAL2 in numerous vesicles present in the cytoplasm and the myelin-like sheets of cells cultured at 37°C. In addition, culturing of cells at 18°C increased the number of cells exhibiting colocalization of PLP with GFP-MAL2.

To analyse the intracellular transport of PLP from the cell surface, we performed cell surface biotinylation. Using this method, we observed internalization of PLP from the plasma membrane and its further colocalization with GFP-MAL2, suggesting transcytosis of PLP also in our system. Some of these internalized vesicles positive for both GFP-MAL2 and PLP were localized to a pericentrosomal region, compatible with the oligodendrocytic GFP-MAL2-positive compartment analogous to the apical ARE/SAC of epithelial cells. Double-labelled immunoelectron microscopy on ultra thin cryosections also indicates a feasible interaction of PLP with GFP-MAL2. Thus, we observed colocalization of both proteins in vesicles close to the plasma membrane; in scattered cytoplasmic vesicles; in tubulovesicles near the nucleus and in a tubulovesicular membrane-bound compartment.

Since the association of MAL2-positive vesicles with actin filaments during transcytosis has been demonstrated [29], we decided to investigate whether GFP-MAL2-positive vesicles containing PLP were also associated to actin cytoskeleton. Previously, we had observed the presence of a GFP-MAL2-positive tubular reticulum in cells cultured in DM at 37°C [26]. This structure arose from the peri-centrosomal region and extended towards processes. Our present results show a large percentage of GFP-MAL2-positive vesicles containing F-actin. Moreover, colocalization of PLP and F-actin was also observed in some GFP-MAL2-positive vesicles and GFP-MAL2-positive tubular reticulum colocalized with F-actin. In addition, several PLP-positive vesicles were associated with this tubular reticulum, suggesting that actin cytoskeleton could be involved in the MAL2-mediated transcytosis of PLP in this system, as it does in other epithelial cells.

An intriguing observation was the presence of GFP-MAL2-positive vesicles containing PLP with an approximated diameter of 1.5 μm close to the plasma membrane and in the extracellular medium of GFP-MAL2/HOG cells, suggesting that these GFP-MAL2-positive vesicles containing PLP could even be released outside the cell. Similar extracellular vesicles containing PLP have also been observed in HOG cells. Large membrane vesicles, up to 1 μm of diameter, secreted by shedding from the plasma membrane, have been described in many cell types [32]. These microvesicles (MVs), different from the exosomes released upon exocytosis of multivesicular bodies, participate in important biological processes, perform a recognized role in the process of communication between cells and have significant physiological and pathological roles [33], such as inflammation, tumor progression or horizontal transfer of proteins and RNA. It has been demonstrated that cultured OLs secrete exosomes carrying PLP and other myelin proteins [34,35], but no large MVs containing PLP had been described before in cultured OLs. However, MVs derived from oligodendrocytes have been found in the cerebrospinal fluid of multiple sclerosis patients, which highlights their pathological significance [36]. Further studies will have to determine the nature of these extracellular PLP-containing vesicles observed in our study. Despite the fact that their size is slightly larger than those of extracellular vesicles described in other cell types, due to the heterogeneity in MVs size [37], we would like to suggest that these structures could be MVs, and may be involved in cell-to-cell communication.

It has been previously demonstrated that OLs cultured in the absence of neurons express myelin genes and form myelin-like membrane sheets [30]. HOG cells have been also proved to form
myelin-like membrane sheets which, in addition, accumulate myelin proteins such as MAL [26] and PLP. These membrane sheets have been previously described in several studies with primary and mixed cultures [38,39]. Therefore, we consider that the system used in our study, oligodendroglial human HOG cell line, is a reliable system to carry out polarization studies in oligodendrocytic differentiated cells. However, further studies, with mixed primary cultures and in vivo systems, will be useful to corroborate these results which, anyway, are consistent with previous observations.

In conclusion, our study shows that in oligodendrocytic cells HOG, PLP and GFP-MAL2 colocalize after internalization from the plasma membrane, co-immunoprecipitate and, finally, colocalize at ultrastructural level, suggesting an association between PLP and GFP-MAL2 in this system, providing new insights into transcytotic transport of PLP to the myelin sheath.

**Materials and Methods**

**Antibodies and reagents**

Anti-PLP mouse monoclonal antibody MAB388, and Horse-radish peroxidase-conjugated secondary anti-IgG antibodies were from Millipore (Billerica, MA, USA). Anti-PLP goat polyclonal antibody sc-18529 was from Santa Cruz Biotechnology (California, USA). Mouse monoclonal antibody to CD59 (MEM-43/5) was kindly provided by Dr V. Horejsi (Institute of Molecular Biology, Prague, Czech Republic).
Genetics, Prague, Czech Republic). P239 anti-MAL2 rabbit polyclonal antibody was kindly provided by Dr M.A. Alonso (CBMSO, Madrid, Spain). Anti-LAMP1 rabbit polyclonal antibody Ra-lamp1 was kindly provided by Dr S. Carlsson (University of Umeå, Sweden). Sulfo-NHS-LC-Biotin was purchased from Thermo Scientific. Anti-GFP rabbit polyclonal serum A6455, TO-PRO-3 iodide 642/661, Alexa Fluor 488-, Alexa Fluor 647- and Alexa Fluor 594-conjugated secondary antibodies and Streptavidin-Alexa Fluor 555 were obtained from Molecular Probes (Eugene, OR, USA). Fetal bovine serum (FBS), insulin, triiodothyronine (T3), apo-transferrin, sodium selenite, putrescine, dibutyryl cyclic AMP (dbcAMP), protease inhibitor cocktail, Protein A-Sepharose, Phalloidin-TRITC and isobutyl-3,3-methyl xanthine (IBMX) were purchased from Sigma Chemical Co. (St. Louis, MO, USA). Mowiol was from Calbiochem (Merck Chemicals, Germany).

Cell lines

The human HOG cell line, established from a surgically removed human oligodendroglioma [30] was kindly provided by Dr. A. T. Campagnoni (University of California, UCLA, USA). Cells were cultured on Petri dishes in growth medium (GM) containing Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS), penicillin (50 U/ml), streptomycin (50 μg/ml) and antibiotics (50 μg/ml) at 37°C in an atmosphere of 5% CO2.

GFP-MAL2/HOG cells consisted of HOG cells stably transfected with a chimera consisting of green fluorescent protein (GFP) fused to the amino-terminal end of MAL2 (GFP-MAL2) [20,26]. To induce maturation, cells were cultured in differentiation medium (DM) [26], containing DMEM without FBS, supplemented with antibiotics and 50 μg/ml apo-transferrin, 0.5 mg/l insulin, 30 nM triiodothyronine (T3), 30 nM sodium selenite and 16.1 mg/l putrescine. HOG cells cultured in this medium were also treated with 0.5 mM dbcAMP and IBMX (a phosphodiesterases inhibitor that reduces degradation of dbcAMP) at a final concentration of 0.5 nM.

Immunoprecipitation and immunoblot analysis

GFP-MAL2/HOG cells monolayers were washed in cold PBS, incubated 20 min in 1 ml of cold lysis buffer (50 mM Tris–HCl pH 6.8, 150 mM NaCl, 1% Triton X-100) supplemented with 1 mM phenylmethylsulfonyl fluoride (PMSF) and Complete Mini Protease Inhibitor Cocktail, and scraped in lysis buffer to collect the lysate. Cell lysates were incubated with monoclonal anti-PLP or polyclonal anti-GFP antibodies for 2 h at 4°C with gentle mixing. As control, preimmune rabbit or mouse serum was used. A sample without primary antibody was also included. Antibody–antigen complexes were captured by incubation with protein A-Sepharose for 2 h at 4°C. Sepharose beads were collected by centrifugation at 14,000 r.p.m. for 2 min at 4°C. The supernatants were reserved for controls. Sepharose beads were washed twice in lysis buffer by centrifugation at 14,000 r.p.m. for 2 min at 4°C and, finally, immunoprecipitates were eluted by boiling in SDS-sample buffer. Samples were then subjected to SDS-PAGE in 12% acrylamide gels under reducing conditions and trans-
ferred to Immobilon-P membranes (Millipore). After blocking with 5% nonfat dry milk, 0.05% Tween 20 in PBS, blots were incubated for 1 hr at room temperature with monoclonal anti-PLP or polyclonal anti-GFP primary antibodies. After several washes with 0.05% Tween 20 in PBS, blots were incubated for 1 hr with secondary antibodies coupled to horseradish peroxidase, washed extensively, and developed using an enhanced chemiluminescence Western blotting kit (ECL, Amersham, Little Chalfont, UK).

Immunofluorescence microscopy
Cells grown on glass coverslips were fixed in 4% paraformaldehyde for 20 min, rinsed with PBS and treated with 20 mM glycine for 5 min to quench aldehyde groups. Cells were then permeabilized with 0.2% Triton X-100, rinsed and incubated with 3% bovine serum albumin in PBS for 30 min. For double-labelled immunofluorescence analysis, cells were incubated for 1 hr with the appropriate primary antibodies coupled to horseradish peroxidase, washed extensively, and developed using an enhanced chemiluminescence Western blotting kit (ECL, Amersham, Little Chalfont, UK).

Figure 8. EM immunocolocalization of PLP and GFP-MAL2. Cryosections of GFP-MAL2/HOG cells cultured in DM at 37°C were stained with a rabbit polyclonal anti-GFP antibody and a mouse monoclonal anti-PLP antibody. Anti-GFP and anti-PLP were detected with 10 nm protein A-gold and 15 nm anti-mouse-gold, respectively. Colocalization of PLP and GFP-MAL2 was observed in vesicles close to the plasma membrane (A); in scattered cytoplasmic vesicles (B); in tubulovesicles near the nucleus (C) and in tubulovesicular membrane-bound compartments (D). PM, plasma membrane; N, nucleus. Arrows point to 15 nm-gold particles. Scale bars = 100 nm. doi:10.1371/journal.pone.0019388.g008

Surface protein biotinylation assay
For labelling of cell surface proteins, GFP-MAL2/HOG cells grown on glass coverslips and cultured in DM for 2 days, were washed with ice-cold DMEM and incubated for 1 hr with 0.5 mg/ml of Sulfo-NHS-LC-Biotin (Thermo Scientific) in DMEM at 4°C. Then, cells were washed with 50 mM glycine in PBS to quench free biotin. After that, cells were cultured in DM either for 1 h at 4°C, or at 37°C for 20, 40, 60 and 150 min. Finally, cells were fixed and treated for immunofluorescence analysis with monoclonal anti-PLP antibody and streptavidin-Alexa Fluor 555 (Invitrogen) to visualize biotinylated proteins.

Immunogold-labelling EM
GFP-MAL2/HOG cells cultured at 37°C in DM for 2 days were fixed in 4% paraformaldehyde in 0.1 M sodium phosphate buffer, pH 7.4, at 4°C for 2 hour. After that, cells were incubated overnight with 8% paraformaldehyde in 0.1 M sodium phosphate buffer, pH 7.4, at 4°C. Fixed cells were washed in PBS containing...
20 mM glycine and embedded in 12% gelatine, infiltrated with 2.3 M sucrose and frozen in liquid nitrogen as described [31]. Cryosections (approximately 70 nm thick) were stained with a rabbit polyclonal anti-GFP antibody and a mouse monoclonal anti-PLP antibody. Primary antibodies were detected with 10 nm protein A-gold (EM Lab, Utrecht University, the Netherlands), to detect anti-GFP, and 15 nm anti-mouse-gold (British BioCell, Cardiff, UK), to detect anti-PLP. Cryosections were examined with a JEM1010 transmission EM (Jeol, Tokyo, Japan).

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Author Contributions

Conceived and designed the experiments: RB-M, JAL-G. Performed the experiments: RB-M, MP-H. Analyzed the data: RB-M, JAL-G. Wrote the paper: RB-M. Performed immunoelectron microscopy: MTR. Discussed data and edited the manuscript: RB-M, AA, FM, JAL-G.

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