Oligodendroglia Development in Cell Culture as Monitored with a Monoclonal Antibody

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A new marker for young oligodendrocytes has been identified by a monoclonal antibody (mOg-1, IgM isotype) prepared from cerebellar plasma membrane stimulated mouse lymphocytes. mOg-1 reactive cells in the mouse cerebellum first appear at day 19 of gestation. Future white matter layers of fixed sections of neonatal rat cerebellum were labeled with mOg-1. Although EM analysis has shown cell-surface binding by presumptive oligodendroglia in neonatal cerebellum, the antibody does not bind to compact myelin. In cell cultures prepared from 6-d-old mice, 11% of the cells bound mOg-1 after 3 d in culture, but up to 5% of the cells bound mOg-1 after 2 weeks in culture. Of these same Og-1-positive cells, 69% bound anti-galactocerebrosides and 65% bound anti-myelin basic protein. After a week in culture Og-1-positive cells often produced lamellated sheets extending a millimeter over the polylysine substratum in the absence of normal myelin formation. mOg-1 recognizes a cell-surface determinant distinct from well-characterized oligodendroglial molecules (galactocerebrosides, sulfatide and myelin basic protein) that is expressed early in oligodendrocyte development. The antibody has been used to follow the maturation of oligodendrocytes in cultures of both normal and jimpy mouse cerebellum.

The nervous system is a complex network derived from multiple associations between different cell types. The limited number of distinct cell types and the availability of neurological mutants with defective cerebellar histogenesis make the mouse cerebellum an attractive model for studying cell-cell interactions. The development of antibodies toward neural cell-surface molecules could be instrumental in establishing the role of specific determinants in histogenesis. We have used the hybridoma technology of Kohler and Milstein (1975, 1976) to prepare monoclonal antibodies that might be employed to identify neural cell types in culture, analyze cell lineage relationships, and investigate cell interactions in reaggregate cultures. Sheep cerebellar plasma membranes were used to immunize BALB/c mice. The resulting hybridomas were tested for production of monoclonal antibodies cross-reactive with cerebellar cultures prepared from mice.

In this report, we describe a monoclonal antibody (mOg-1) reactive with the surface of young oligodendrocytes, prior to the appearance of galactocerebroside on these cultured cells. This antibody has been used to monitor oligodendrocyte maturation in cerebellar cultures. A preliminary report of these studies has previously appeared (Seeds and Collins, 1983).

Materials and Methods

Animals

All mice were obtained from Jackson Laboratories, (Bar Harbor, ME) and maintained as a breeding colony. C57Bl/6 and jimpy mice were used for cell culture, while BALB/c mice were used for immunizations. Sprague-Dawley rats were a gift from Dr. Robert Lasher. Lamb brains were purchased from Colorado Lamb Co. (Denver, CO).

Cell suspensions and cultures

The cerebellum was removed from fetal or postnatal mice of various ages and dispersed into a single-cell suspension, as described previously (Krystosek and Seeds, 1981). Diced tissue was dissociated in a solution of 0.25% trypsin (hog pancreas, ICN Pharmaceuticals) and 0.6 mg % deoxyribonuclease (DNase I, bovine pancreas, Nutritional Biochemicals) in saline 1 (0.138 m NaCl, 5.4 mm KCl, 1.1 mm NaHPO4, 1.1 mm KH2PO4) containing 0.4% glucose and 0.01% CaCl2. The suspension was incubated for 12 min at 37°C, dispersed by gentle pipetting in basal Eagle's medium (BEM) with 10% fetal calf serum (FCS), and passed through a 200 μm Nylon screen. Surface cultures were prepared from the dissociated cells by pipetting them onto 18 mm glass coverslips previously coated with poly-(D-lysine) (Yavin and Yavin, 1977). The cultures were maintained at 37°C in a 5% CO2:95% air incubator. Schwann cell cultures were initiated by dissecting the root portions of dorsal root ganglia from 2 d postnatal mice and preparing cell suspensions as described by Ransou et al. (1977). Briefly, the tissue was treated with 0.25% collagenase at 37°C for 5 min, triturated with Sigma Cote-d Pasteur pipettes in 0.5 ml serum-free culture media containing 10 μg DNase, and washed and pipetted on coverslips as above. Clonal cell lines, including the methylcholanthrene-induced rat astrocytoma C6 (Benda et al., 1968) and mouse oligodendrogloma G0, (Zimmermann and Matier, 1948), as well as an ethylcholostere-activated mouse schwannoma TRM66 (Fields et al., 1975), were cultured on coverslips in Dulbecco's modification of Eagle's medium (DMEFM) with 5-10% fetal calf serum in an atmosphere of 10% CO2:90% air.

Monoclonal antibody production

BALB/c mice were immunized with cerebellar plasma membranes prepared as described by Stuhlfauth and Seeds (1983) from bovine cerebellum. We used 100-500 μg protein, determined by the method of Lowry et al. (1951), for each injection. Initially, 0.2-0.4 μl of a 1:1 suspension with complete Freund's adjuvant was injected subcutaneously. Subsequently, mice were injected intraperitoneally minus adjuvant at 2 week intervals (3-4 times). Spleen cells from one mouse were fused 4 d after the last injection with hypoxanthine/aminopterin/thymidine-sensitive myeloma cells P3-NS-1/Ag 4-1 (Kohler et al., 1976) at a ratio of 3:1 in the presence of 4% Merck PEG 4000 containing 15% dimethylsulfoxide by the procedure of Gallo et al. (1977). The fused cells were distributed into five 24-well Costar dishes preconditioned with 5 × 10⁶ peritoneal macrophages/well in complete HAT medium. The cultures were fed every 3-4 d.
Approximately 2 weeks after fusion, aliquots of media were removed for testing from these wells that showed substantial growth and concentrated 10-fold by precipitating in 45% saturated ammonium sulfate. Dialed supernatants were then tested by immunofluorescence with surface cultures. Antibody-producing hybridoma cells were cloned twice by the limiting dilution method of Oi and Herzenberg (1980).

Immunofluorescence

Indirect immunofluorescence labeling of surface cultures was used for testing immunized mouse sera and hybridoma supernatant samples, and for analysis of cell-type specificity of monoclonal antibodies. In addition to mOg-1, the following cell-type-specific antibodies were used: a rabbit anti serum raised against galactocerebroside (rGalC, #68R-1594) (Joffe and Rapport, 1963), a mouse monoclonal to galactocerebroside (mGalC) (Ranscht et al., 1982), a rabbit anti-glycerol-3-phosphate dehydrogenase (rG3PDH) (Leveille et al., 1980), a rabbit anti-glial fibrillary acidic protein (rGFAP) (Dahl and Bignami, 1973), a rabbit antivimentin (gift of Dr. R. Evans), a mouse monoclonal (IgG) anti-myelin basic protein (mMBP) (Hybritech, Inc., San Diego, CA), a rabbit antiseraum against neonatal mouse cerebellum (rCbl-1) (Seeds, 1975), and tetanus toxin and rabbit anti-tetanus toxin (Yavin et al., 1981). Cultured-on coverslips were incubated with 30 µl of diluted antibody, rinsed 5 times with saline containing 10 µM CaCl2 and 5 µM MgCl2, fixed for 5 min with 4% paraformaldehyde, washed, and incubated with fluorescein (FITC) or tetramethyl rhodamine (TRITC)-conjugated second antibodies.

For double indirect immunofluorescence with mOg-1 and rGalC, coverslips were incubated simultaneously with the 2 antibodies, washed, fixed, and incubated with a mixture of FITC-goat anti-mouse Ig (GAM) (Meloy Laboratories, Springfield, VA) and TRITC-got anti-rabbit Ig (GAR) (Cappel Laboratories, Malvern, PA).

A similar procedure was used for mouse GalC and mOg-1; however, TRITC-GAM-µ-chain (affinity-purified, TAGO, Inc., Burlingame, CA) and FITC-GAM-µ-chain (Cappel Laboratories, Malvern, PA) were used. For studies using rG3PDH, r-vimentin, or rGFAP, coverslips were incubated with mOg-1, washed, fixed, and treated with methanol for 5 min at -20°C to permeabilize the cell membrane. The cultures were next incubated with rG3PDH, r-vimentin, or rGFAP, and finally with a mixture containing FITC-GAM and TRITC-GAR. Double labeling with mOg-1 and mMBP on cerebellar cultures was done in a sequential fashion. Live cells were incubated with mOg-1, washed, fixed with 4% paraformaldehyde for 5 min, and permeabilized with 0.05% Triton X-100 for 2 min, incubated with mMBP, washed, and incubated with the FITC-GAM-µ-chain specific and TRITC-GAM-µ-chain specific. Finally, double labeling with mOg-1 and tetanus toxin (the generous gift of Drs. W. Habig and M. Hardegee) was performed by incubating the coverslip first with a mixture of the 2 markers, then with rabbit anti-tetanus toxin, fixing, incubating with TRITC-got anti-rabbit Ig (GAR) and with FITC-GAM. Coverslips were mounted face down in Gelvatol (Monsanto, St. Louis, MO) and examined with a Zeiss epifluorescence microscope.

Immunostaining and electron microscopy of fixed sections

Rats of various ages were sedated with 15% chloral hydrate, perfused through the left ventricle with cold 5% paraformaldehyde in 0.15 M Sorenson's PO4 buffer containing 0.08 M sucrose at pH 7.7. Brains were removed, dissected, and fixed further for 3.5 hr. Vibratome sections, 30 µm, were stained in a 24-well dish with mOg-1 at 4°C overnight, then incubated with 1 ml monospecific rabbit IgG (anti-mouse IgM for 60 min, and finally labeled with 1 ml HRP-GAR IgG (TAGO, Burlingame, CA). Peroxidase labeling was achieved by the addition of 0.5 ml of 0.05% DAB-HCl for 10 min, followed by 5.5 ml of 0.3% solution of H2O2 for 10 min.

Sections were prepared for electron microscopy by incubation with 1% OsO4, dehydration with 50, 70, 95, and 100% ethanol, 2-hydroxypropylmethacrylate, and lastly Epon 5020 mix with catalyst. The sections were mounted between Aclar fluorocarbon film, polymerized, and attached with epoxy glue to an Epon block. They were then thin-sectioned with a Philips 300 EM.

Autoradiography

Surface cultures on coverslips were exposed to [3H]-thymidine (2 µCi/ml) for 72 hr, washed, labeled by indirect immunofluorescence and air-dried. The coverslips were mounted faceup on slides and coated with Kodak NTB2 emulsion. After 3-4 d at 4°C, they were developed with Kodak Dektol, mounted, and examined with phase and fluorescence optics.

Complement-dependent cytotoxicity

Monolayer cultures of cerebellar cells from 6- to 8-d-old C57BL/6 mice were maintained in vitro for 1 d. Some of the coverslips were then removed, washed with BEM without serum, and incubated with monoclonal antibody and guinea pig complement (diluted 1:8) for 20 min at 37°C in a 5% CO2, 95% air incubator. All dilutions were made in BEM. The reaction was terminated by 2 gentle rinses with BEM containing 10% fetal calf serum. Cultures were immediately processed for immunocytochemical analysis or maintained in sterile conditions in BEM with 10% fetal calf serum.

Results

Cell-type specificity of mOg-1

Supernatants from mOg-1-producing clones were used to label cerebellar cultures of 6-8 d postnatal C57BL/6 mice grown for 4 d in vitro. Only cells with an oligodendrocyte-like morphology were stained (Fig. 1), and these constituted 1.1% of the total viable cells. This specificity for oligodendrocytes was confirmed by double-labeling studies that used mOg-1 with other antibodies specific to various neural cell types. Cells binding mOg-1 did not react with tetanus toxin (a neuronal marker), anti-Chbl-1 (a neuronal marker), or anti-GFAP (an astrocyte marker) (Fig. 2). The mOg-1-reactive cells did not bind rabbit anti-vimentin, a marker for mesenchymal intermediate filaments (Fig. 3). However, subsets of mOg-1-positive cells did bind mGalC, rG3PDH, and mMBP (Fig. 4). GalC, the major glycolipid in myelin (Nor-
Figure 2. Double-labeling with mOg-1 antibody and specific neuronal and astrocyte markers. Dissociated cerebellar cells (A) were incubated with tetanus toxin, rabbit anti-tetanus toxin (C') and mOg-1 (B). Another neuronal marker anti-Cbl-1 (F) was used with mOg-1 (E) to distinguish cerebellar cell populations (D). The astrocyte specific anti-GFAP (I) was added to cells (G) labeled with mOg-1 (H). Fluorescent conjugates were used to visualize these bound antibodies, as described in Materials and Methods. The mOg-1 antibody reacts with a cell population distinct from these neuronal and astrocytic cells.

The mOg-1 antibody, developed by Dubois and Autilio (1966), has been shown to be expressed only by oligodendrocytes in a variety of rat (Raff et al., 1978), human (Kennedy et al., 1980), and bovine (Lisak et al., 1981) CNS cultures; however, not all mOg-1 cells bind anti-GalC (Fig. 4C). G3PDH has also been used as an oligodendrocyte marker (Leveille et al., 1980). Although G3PDH enzyme appears to be localized primarily in oligodendrocytes, it is also present in lesser amounts in astroglial cell cultures (McCarthy and DeVellis, 1980). Myelin basic protein (MBP), an important component of myelin, can be detected in oligodendrocytes prior to myelination (Stemberger et al., 1978). In dissociated brain cell cultures, where myelination does not take place, oligodendrocytes are still able to express MBP (Bologa-Sandru et al., 1981). The MBP reactivity is confined to the more central structures of the Og-l-positive cells (Fig. 4, E, F). In additional studies, cells binding rabbit antibody to myelin associated glycoprotein also bound mOg-1 (results not shown).

Schwann cells, the PNS equivalent of oligodendrocytes, prepared from dorsal root ganglia, were incubated with mOg-1. Labeled cells exhibiting the bipolar spindle shape characteristic of Schwann cells are shown in Figure 5. Some dependence of mOg-1 labeling on length of time in culture was noted. After 4 d in culture this marker is apparently lost. Dorsal root ganglia neurons were negative at all times tested.

Clonal tumor cell lines of presumed glial and Schwann cell origin—the rat astrocytoma C6, the mouse oligodendroglioma G26, and schwannoma TRM6B—were also tested for immunolabeling with mOg-1. Even though we included G26, a presumptive oligodendroglioma that has been shown to synthesize sulfatide, a myelin component (Seeds and Marks, 1979), all 3 cell lines were negative for mOg-1 binding, when tested shortly after initiation or at confluency. If mOg-1 antigen is involved in the differentiated functions of myelinating cells, it is apparently lost in establishment of these cell continuous lines.

Localization of mOg-1 staining in brain sections

White matter was very heavily labeled at day 8 (Fig. 6A), and occasional cells subadjacent to the internal granular layer were labeled at day 3 (not shown). White matter and structures of the internal granular layer were labeled in fixed sections of the
adult rat cerebellum (not shown). White matter portions of other brain areas were also labeled with mOg-1. Antibody to GaIC showed a similar labeling of cerebellar sections. Cells deep in the granule layer and white matter showed surface labeling with mOg-1 at the EM level (Fig. 6B). Although morphology of these cells is inconclusive, their position, nuclear morphology, and electron opacity are suggestive of oligodendroglial cells. Other cells, identified as mature oligodendrocytes by the criteria of Privat (1975) and Skoff et al. (1976), were not stained with mOg-1. Neither were several examples of compact myelin. Similar staining distribution has been observed with frozen sections of mouse cerebellum.

**Developmental studies**

Cell suspensions were prepared from the cerebellum of fetal and postnatal mice of various ages. Binding of mOg-1 to 1 aliquot of the cells was determined after 4 hr by immunofluorescence. A second portion of cells was placed in culture and immunolabeled 24 hr later, when a distinctive morphology could be seen in cultures. The earliest time positive cells were seen in suspensions was fetal day 19, or in cultures prepared from an 18 d fetal mouse and maintained 1 d in culture. Thus, we conclude that Og-1 m first makes its appearance at fetal day 19. Cultures prepared from younger animals and maintained for longer periods of culture confirmed this finding. For example, cultures prepared from 15 d fetal mice contained Og-1-positive cells after 3–5 d in culture, suggesting that in terms of Og-1 expression, maturation in culture parallels that seen in vivo.

In cell cultures prepared from 6-d-old mice, 1.1% of the cells bound mOg-1 after 3 d in culture. After 2 weeks in culture, up to 5% of the cells bound mOg-1. This culture period was typified by a decrease in small uniaxial bipolar cells and by proliferation of astrocytes. Where these percentages are relative, the actual numbers of Og-1-positive cells were apparently increasing as judged by Og-1-positive cells/field. Though much of the increase may derive from maturation of Og-1-negative precursors, a portion may be related to proliferation of these precursor cells or possibly the Og-1-positive cells themselves. Cultures incubated for 72 hr with [3H]thymidine and immunolabeled with mOg-1 and FITC-GAM contained Og-1-positive cells that had incorporated thymidine (Fig. 7).

Since monoclonal antibody mOg-1 was found to be complement-fixing and to be IgM by Ouchterlony analysis using class-specific anti-mouse immunoglobulins, this problem was also investigated using complement-mediated lysis of Og-1-positive cells in culture. Cerebellar cultures prepared from 6 d postnatal mice were treated after 1 d in culture with mOg-1 and guinea pig complement. The complement control indicated that at the dilution used (1:6) the complement alone was not cytotoxic. However, when tested 3 hr after treatment, no intact cells labeled with mOg-1 could be found. After several days, Og-1-positive cells were again observed in the culture. This phenomenon was quantified. Coverslips were withdrawn from culture at 2 d intervals and the relative number of Og-1-positive cells determined (Fig. 8). The percentage of Og-1-positive cells in the control cultures rises steadily for the first 4 d after treatment. An apparent decrease was seen at day 6, but this may be due to the very fast proliferation of astrocytes during this period, since the value is calculated as the percentage of total cells. An increase was likewise seen in Og-1-positive cells 2 d after complement lysis in the presence or absence of 10^{-3} M cytosine arabinoside, indicating that a nongrowing precursor for Og-1 was present in the culture at the time of complement-dependent lysis. However, the increase by 4 d suggests that some division of Og-1 precursor cells may be required for maximal expression of Og-1 in these cerebellar cultures. These findings confirm the hypothesis that precursors of Og-1-positive cells exist in the cultures at day 1 and express the Og-1 antigen at later times.
positive cells were more likely to survive the switch to serum-free conditions than were some other cell types, but additional cerebellar cells were not directed toward oligodendrocyte differentiation. However, if the cells were switched to N2 after only 1 d of culture, similar cell numbers and a 2-fold increase in the percent mOg-1-positive cells compared to serum grown cultures were found at day 4; similar to the observation of Raff et al., (1983) with optic nerve cultures.

Some forms of oligodendroglial cells can produce “veil-like” membranous structures in culture (Sommer and Schachner, 1981; Szuchet and Stefansson, 1980). After a week in culture, Og-1-positive cells often produced lamellar sheets extending for millimeters over the polylysine substratum (Fig. 9). These cells most likely correlate with the “active oligodendroglial” cells described by Tennekoon et al. (1980), which, by other criteria, are classified as probable oligodendroblasts.
Og-1 in the jimpy mutant mouse

The X-linked recessive mutation, jimpy, of the mouse is characterized by the virtual absence of myelin in the CNS (Sidman et al., 1964). The biochemical nature of the defect in the jimpy is not known. Since Og-1 appears to be a marker for oligodendroglial development, its presence on cultured cells from jimpy cerebellum was examined (Table 2). Og-1 was present on cells from the jimpy cerebellum, although mOg-1-positive cells represented a smaller percentage (3.5 vs 5.0%) of the total cell population than in cultures of normal litters. A similar proportion of these Og-1-positive cells was also positive for mGal-C binding; however, the jimpy cells failed to bind myelin basic protein antibody even after 14 d of culture. A previous study (Bologa-Sandru et al., 1982) also found an absence of myelin basic protein in 14 d cultures of jimpy. Thus, Og-1 appears to be a marker for early development of oligodendroglia and is expressed prior to the genetic block in the jimpy mouse.

Discussion

In this report, we describe a monoclonal antibody (mOg-1) that is directed against a surface component expressed on oligodendroglial cells and Schwann cells. These cells are responsible for myelination in the CNS and PNS, respectively. Double-labeling experiments with known neural cell markers were used for this identification. Only the known oligodendroglial cell markers, GaIC, rG3PDH, and MBP were seen on the Og-1-positive population. Tetanus toxin, a neuronal marker, and anti-GFAP, an astrocyte marker, labeled other cells in the population.

We could not identify all Og-1-positive cells as oligodendrocytes since only 60–70% of them double-labeled with other oligodendroglial markers in culture. However, several lines of evidence suggest that the Og-1-positive cells that are negative for other markers may be less mature oligodendrocytes. Simple monitoring of the cultures with or without complement-mediated lysis of Og-1-positive cells initially present indicates that the Og-1 antigen itself is acquired with increasing time in culture.

Tissue sections, labeled with mOg-1 and observed by light and electron microscopy, support the conclusion that Og-1-positive cells are oligodendrocytes. Positively stained cells are located deep in the granular layer and in the white matter, where we would expect to see oligodendroglial cells. However, many cells identifiable as oligodendrocytes in adult tissue by morphological criteria are not labeled. Neither are several examples of mature myelin. This may be due to incomplete penetration of the antibodies into the tissue, but positive oligodendrocytes adjacent to negative ones are difficult to explain on this basis. It seems more likely that Og-1 is a cell-surface component of less mature oligodendrocytes perhaps involved in the earliest stages of myelination.

The time of appearance of Og-1 antigen in the developing mouse brain is also consistent with this interpretation. Og-1-positive cells were first seen in tissue from 19 d fetal mice. This is 2–3 d before the production of GaIC (Schachner and Willinger, 1979) and well before the onset of myelination. Abney et al. (1981) have shown that oligodendrocytes develop on the same schedule in dissociated cell cultures of embryonic rat brain as...
they do in vivo using GalC as a marker. Similarly, our experiments suggest the same phenomenon for Og-1. However, one must be careful in the interpretation of these results. Recent elegant studies by Raff et al. (1983) have shown that, in glial cell cultures derived from optic nerve, glial differentiation and expression of certain cell markers are influenced by the composition of the tissue culture medium. The effect of serum-free medium on cerebellar cells appears to be a precocious appearance of Og-1, with an overall reduction in total oligodendrocyte (Og-1-positive cells) numbers after 8 to 11 d compared to serum-containing cultures. This finding probably reflects a need for further cell division of the oligodendrocyte stem cells during culture, which is inhibited by N2 medium. In addition, experiments by others have shown that oligodendroglial differentiation in vitro may be modified by the number of neurons in the culture (Barbarese et al., 1981; Bologa et al., 1982; Mirsky et al., 1980). Although we never see 100% of the Og-1-positive cells labeled with other oligodendrocyte markers, this may reflect the fact that we are making a static analysis of a dynamic population. That is, precursor cells may be constantly dividing in culture, acquiring the early marker Og-1 and then developing further. At any point in time some cells would be only partway along this path.

Oligodendroglia and Schwann cells share a number of myelin-specific proteins and glycolipids, including GalC, sulfatide, and MBP (Mirsky et al., 1980). Schwann cells stop making these markers a few days after being placed in culture, while oligodendrocytes continue for extended periods. Expression of Og-1 antigen follows similar patterns. It is present on both Schwann cells and oligodendroglia but is lost from the surface of Schwann cells after 4 d in culture.

The fact that a large degree of variation in cellular shapes and morphological features exists within the oligodendroglia is commonly known, and we have observed this in the Og-1-positive population. Mori and Leblond (1970) were among the first to

Table 1. Immunolabeling of Og-1-positive cells with anti-GalC, anti-G3PDH, or anti-MBP

| Days in culture | GalC/Og-1 (%) | G3PDH/Og-1 (%) | MBP/Og-1 (%) |
|----------------|--------------|----------------|--------------|
| 4              | 54           | 100            | 27           |
| 7              | 67           | 100            | 46           |
| 14             | 69           | 100            | 65           |

Cerebellar cultures were prepared from 6 d postnatal C57Bl/6 mice and double-labeled after various times in culture. The percentage of Og-1-positive cells expressing known oligodendroglial markers is shown.

Table 2. Oligodendroglial development in jimpy cultures

| Animal          | Og-1 + (%) | GalC/Og-1 (%) | MBP/Og-1 (%) |
|-----------------|------------|---------------|--------------|
| Jimpy           | 3.5        | 58            | 0            |
| Normal littermates | 5          | 61            | 57           |

Cerebellar cultures were prepared from 6 d postnatal mice and double-labeled after 7 d in culture. The percentage of Og-1-positive (+) cells in the cultures is indicated, as well as the percentage showing both labels.
suggest this variation may depend on the degree of oligodendroglial maturation. Applying the structural guidelines of Skooff et al. (1976) and Privat (1975), it is apparent that many O1-positive cells are immature oligodendroglial cells. Both oligodendroblasts and many O1-positive cells have relatively large amounts of cytoplasm, which is not typical of fully differentiated oligodendrocytes.

Schachner et al. (1981) have described 4 oligodendroglial-specific antigens (O1-O4) detected by monoclonal antibodies. Cells expressing O1 and O2 antigen also express GalC. However, only a fraction of the O3- and O4-positive cells (64 and 47%, respectively) are GalC-positive. The O3- or O4-positive, GalC-negative cells express no other cell-type-specific markers, paralleling the situation we have observed with O1-1. Berg and Schachner (1981) further investigated the possibility that these cells were immature oligodendroglia by performing immunoelectron microscopy on monolayer cultures of fetal and early postnatal mouse cerebellum. They concluded that approximately 5% of the O-positive cells were fully mature oligodendroglia, while greater than 90% should be more appropriately labeled oligodendroblasts. Under their culture conditions, these putative oligodendroglial precursors never fully mature.

Our data suggest a similar phenomenon. O1 antibody labels oligodendroglia as well as immature oligodendroblasts. Only a fraction of these cells can be shown to mature in culture. However, both O3 and O4 are present at earlier developmental times than O1 (day 16 fetal vs day 19). In addition, O3 antibody internally labels astrocytes, and this is not seen with O1. These factors suggest that O1-1 antigen is probably not identical to either O3 or O4, but these markers may be present on related cell populations. Since both O4 and O1 are mouse IgMs, it is not possible to perform double labeling with these molecules or with the interesting monoclonal A2B5 (Eisenbarth et al., 1979); sequential binding of these antibodies and their fluorescent conjugates has not been satisfactory. Attempts to conjugate biotin directly to O1 have led to inactivation, presumably due to an active amino group in the recognition site.

Cerebellar cultures were occasionally fixed with methanol or chloroform: methanol (1:1) before the addition of antibodies directed against intracellular antigens during the double-labeling studies. If O1-1 was not bound and fixed with paraformaldehyde prior to this treatment, the binding pattern was lost or greatly diminished. This suggests that the organic fixation step was destroying antigenicity either through denaturation or possibly extraction of the antigen. Fixation with paraformaldehyde before binding had minimal effect.

A number of well-characterized surface components—including GalC and sulfatide, as well as related molecules such as glucocerebroside and ceramide—were tested for their ability to bind O1-1. In the first method, direct binding was measured as described by Ranscht et al. (1982). The second method involved adsorption of O1-1-containing supernatants diluted slightly less than the titer end point, by the various lipids being tested. The adsorbed antibody was then tested in the usual immunofluorescent assay system with cerebellar cultures, and, in one instance, quantitated with 125I-anti-mouse immunoglobulin as second antibody. Both techniques confirmed that O1-1 antigen is not GalC, sulfatide, or the related compounds. Additional studies showed that O1 was tryptic sensitive.

Although the O1-1 antigen is extractable from cell membranes in nionic detergents, O1-1 is irreversibly denatured by SDS, thus precluding its identification by immunoblotting techniques. However, using 35S to metabolically label cerebellar cell cultures, a labeled molecule of approximately 90,000 M, that binds to O1-1 has been isolated on affinity columns containing purified goat anti-mouse IgM immunoglobulin, and it appears to be the O1-1 antigen. Although the molecular weight is similar to that of myelin-associated glycoprotein (MAG) (Quarles et al., 1983), O1-1 does not react with rat or human MAG (A. Noronha, R. H. Quares, and N. W. Seeds, unpublished observations).

These studies have described a cell-surface antigen, O1-1, that appears early in oligodendrocyte development. Current studies are investigating the possible role of O1-1 in oligodendrocyte–axon recognition events and myelination in reaggregate (Seeds, 1971; Seeds and Haffke, 1978) brain cell cultures.

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Figure 9. Lamellar sheet formation by oligodendrocytes. Large lamellar extensions millimeters in length are visualized by mOg-1 binding and FITC-goat anti-mouse Ig in cerebellar cell cultures after 7 d in culture.
