Myo-inositol transport into endothelial cells derived from nervous system microvessels

Peter E. Kollros1,2, Gary W. Goldstein1,2,* and A. Lorris Betz1,3
University of Michigan Medical Center, Departments of 1Pediatrics, 2Neurology and 3Surgery, Ann Arbor, MI (U.S.A.)
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Myo-inositol, the precursor in the biosynthesis of inositol phospholipids and inositol phosphates, is found in many tissues at concentrations well above its concentration in the plasma, but the highest concentrations are found in the central nervous system and the neuroretina. We describe an active, sodium gradient-dependent transport of myo-inositol into cultured endothelial cells derived from bovine retinal microvessels. Transport is inhibited by cytochalasin B, and phloridzin > phloretin. Mannitol, sorbitol, and fructose do not inhibit uptake, but α-galactose inhibits uptake > L-glucose > D-glucose. The apparent $K_m$ of this transport system is 311 ± 47 (S.D.) μM and the apparent $V_{max}$ is 40.8 ± 2.8 (S.D.) pmol·mg protein $^{-1}$·min $^{-1}$. This transport system may be a key in the maintenance of high tissue concentrations as it could concentrate myo-inositol from the plasma into the extracellular spaces of the eye and central nervous system.

INTRODUCTION

Myo-inositol is an important precursor to the biosynthesis of inositol phospholipids and inositol phosphates, compounds which play an important role in signal transduction9,19. In mammals, myo-inositol is present in plasma at concentrations ranging from 25 to 100 μM12,14,17. Intracellular concentrations of myo-inositol are several-fold higher than in the circulation, with brain and neuroretinal tissue having the highest concentrations2,17,18,21. Although high intracellular concentrations could be maintained either by active myo-inositol transport or by synthesis from glucose, transport appears to be the primary mechanism for two reasons. First, the activity of the rate limiting enzyme in the synthesis of myo-inositol, inositol 1-phosphate synthetase (EC.5.5.1.4), is low in most tissues20. Second, elevated plasma glucose concentrations are associated with lowered rather than elevated myo-inositol tissue concentrations8,10,13,15,18.

Cerebrospinal fluid (CSF) and ocular vitreous humor concentrations of myo-inositol are also higher than plasma levels17,18,23. Transport of this compound from blood to brain in vivo and by the isolated choroid plexus in vitro has been reported24. However, active transport systems capable of maintaining myo-inositol concentration gradients between plasma and CSF or vitreous humor have not been previously described in endothelial cells that form the blood–brain and blood–retinal barriers. In this report, we characterize the transport of myo-inositol into cultured endothelial cells derived from bovine retinal microvessels. Retinal microvessels display similar structural and functional properties to brain microvessels, and endothelial cells in each form blood–tissue barriers. We show here that myo-inositol is transported by an active transport process dependent on the transmembrane sodium gradient and inhibited by glucose, galactose, phloridzin, phloretin and cytochalasin B.

MATERIALS AND METHODS

Cell acquisition and culture

Retinal endothelial cells were obtained by a modification of methods previously described6. Bovine eyes were obtained from the slaughterhouse. Retinas were dissected, placed in ice-cold M-199 medium supplemented with penicillin and streptomycin, and rinsed 3 times. The retinas were strained on a 118 μm nylon mesh and extensively washed. The trapped tissue was washed and then homogenized in a glass tube with a loose fitting Teflon pestle (0.25 mm clearance) at 400 rpm for 20 up and down strokes. The homogenate was centrifuged for 5 min at 1000 g. The pellet was resuspended in M-199 and gently agitated for 45–60 min. The digest was then layered over 5 ml of Ficol-Paque and centrifuged at 1250 g for 20 min. The interface cells and the pellet were transferred to a T25 flask coated with gelatin and incubated with minimal essential medium with l-valine replaced by d-valine (MEM-d-Val) for 16–24 h. The second...
day the medium was changed to MEM-p-Val supplemented with 16 U/ml heparin and 50 μg/ml of endothelial growth supplement (ECGS). The cells were grown and passaged at confluency with splits of 1 to 4. Cells used in these experiments were in the 6th through 14th passage, tested positive for low density lipoprotein (LDL) uptake and displayed the typical endothelial cell morphology on phase contrast microscopy.

**Transport experiments**

Cells were grown to confluency in 24-well (1.6 cm diameter) culture plates coated with fibronectin. The cells were then washed and preincubated for 12 min in either an isoinitol-free HEPES buffered Dulbecco’s medium, a HEPES buffered Earle’s balanced salt solution (EBS), or sodium-free EBS with choline replacing sodium. The appropriate additives except for the radioactive tracers were included in the preincubation solutions. In the cases where ouabain or 2,4-dinitrophenol (2,4-DNP) and iodoacetic acid were used, the preincubation period was extended to 30 min. The preincubation solution was aspirated off the cells and the incubation solution was added. At the end of the incubation, the experiment was stopped by the addition of 2 ml of ice-cold phosphate-buffered saline (PBS), pH 7.4. The cells were then quickly washed 4 more times over 20 s with ice-cold PBS. After the final wash, 1 N NaOH was added to each well. The cells were dissolved overnight at room temperature and then aliquots were taken for liquid scintillation counting and for total protein determination. When quantitation of lipid and inositol was undertaken, methanol was added to each well immediately following the final PBS wash.

During the incubation, the medium or buffer solutions contained [14C]sucrose as a marker of the extracellular space. By measuring the 14C disintegrations per minute (DPMs) in each sample, the contribution to the total H DPMs from extracellular H was determined and the intracellular label was calculated by subtracting the extracellular DPMs of H from the total H DPMs.

**Biochemical characterization of incorporated label**

We employed a modification of the method of Berridge to characterize the products of inositol incorporation. Immediately following the final wash, methanol was added to the cell layer, which was then scraped and both the cell layer and supernatant were transferred to a glass tube. The well was washed with methanol and the wash was combined with the sample to make a total methanol volume of 1 ml. To this, 0.8 ml of chloroform-HCl (200:1) was added, followed by 2 ml of 0.5 mM myo-inositol dissolved in water. The sample was then vortexed and centrifuged at 5 g for 5 min. The aqueous layer was separated and an aliquot of the organic layer was counted as an indication of myo-inositol incorporation into lipid. To the aqueous layer, 0.5 ml of a 50% slurry of Dowex AGI-X8 was added and mixed with the sample for 5 min. The sample was allowed to settle and the supernatant separated. The resin pellet was then washed twice with 5 mM myo-inositol solution. The supernatants and aqueous layer were pooled, frozen and lyophilized. The resin was counted using a liquid scintillation counter to quantitate myo-inositol incorporation into inositol phosphates. The lyophilate was resuspended in 1 ml of water and an aliquot counted to determine the amount of free myo-inositol. To assure the purity of the free myo-inositol, representative samples were then run through a desalting column of 1.5 × 0.6 cm consisting of Dowex 1-X8 (acetate form) and 1.5 cm × 0.6 cm amylite CG-50 (‘H form) with the Dowex placed above the Amberlite. The sample was eluted with an excess of deionized distilled water. Recovery of label from the column was 100%. The effluent was frozen, lyophilized, and reconstituted in a small volume for paper chromatography. The sample and standards were run in two solvent systems, n-butanol:ethanol:water (2:2:1) and n-butanol:pyridine:water (6:4:3) on Whatman 1 paper. In both systems the sample co-migrated as a single peak with the [3H]myo-inositol standard.

**Water space determination**

The intracellular water space of the cultured cells was measured using 3-O-methyl-glucose. In this experiment, [3H]-O-methyl-glucose was added to the cells and incubated for 45 min along with [14C]sucrose as a marker of extracellular water space. Since 3-O-methyl-glucose equilibrates with the cells during this period of time, the intracellular water space was calculated as the difference between the 3-O-methyl-glucose space and the sucrose space. The water space value was determined to be 7.4 ± 1.2 (S.D.) μl/mg of protein (n = 4).

**Materials**

Cell culture medium, fetal calf serum, calf serum, penicillin, and streptomycin were purchased from GIBCO Laboratories, Grand Island, NY. ECGS was purchased from Collaborative Research, Lexington, MA. [3H]myo-inositol and [14C]sucrose were obtained from American Radiolabeled Chemicals, St. Louis, MO. [3H]-O-methyl-glucose was obtained from New England Nuclear, Boston, MA. Bovine immunoglobulin and Coomassie brilliant blue were acquired from Bio-Rad, Richmond, CA. Acetylated low density lipoprotein labeled with 1,1'-dioctadecyl-1,3,3,3'-tetra-methyl-indo-carbocyanine perchlorate was acquired from Biomedical Technologies, Stoughton, MA. All other chemicals were purchased from Sigma, St. Louis, MO. Tissue culture plates and flasks were obtained from Costar, Cambridge, MA.

**RESULTS**

**Temperature dependence of inositol transport and accumulation**

Fig. 1 shows the time course over 80 min for uptake of myo-inositol by cultured endothelial cells derived from retinal microvessels. The data are expressed as a ratio of the cellular concentration to the culture medium concentration of 3H, calculated using the measured value of 7.4 μl/mg protein for the intracellular water space. Over this time course the uptake is linear and temperature dependent. Values greater than one show that there is accumulation of label from the medium into the cells so the intracellular concentration is greater than the medium concentration. This accumulation could be the result of either the active transport of myo-inositol or the metab-
Fig. 1. Time course of myo-inositol uptake and incorporation into cultured retinal endothelial cells. Each point represents the mean of 5 replicates and the error bars are S.D. The data are presented as a concentration ratio of the intracellular concentration to the extracellular concentration. The extracellular ambient myo-inositol concentration was 6 μM and water space of cells measured 7.4 μl/mg protein.

Fig. 2. Uptake and incorporation of myo-inositol into intracellular myo-inositol, inositol phospholipids, and inositol phosphates. Each value represents the mean of 4 replicates with error bars showing the S.D. The ambient myo-inositol concentration was 0.76 μM.

Fig. 3. Effect of sodium concentration on myo-inositol transport. Each sample represents the mean of 6 replicates with the error bars representing S.D. The ambient extracellular myo-inositol concentration was 2.4 μM.

TABLE I
Effect of sodium replacement, ouabain, and energy metabolism inhibitors on myo-inositol transport

<table>
<thead>
<tr>
<th>Condition</th>
<th>fmol·mg protein⁻¹·min⁻¹</th>
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<tbody>
<tr>
<td>Control buffer</td>
<td>67.7 ± 1.4</td>
</tr>
<tr>
<td>Choline buffer</td>
<td>14.0 ± 1.0**</td>
</tr>
<tr>
<td>Ouabain</td>
<td>32.6 ± 2.1**</td>
</tr>
<tr>
<td>2,4-DNP + iodoacetic acid</td>
<td>34.9 ± 1.4**</td>
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**P < 0.005 compared to 150 mM Na⁺.

Metabolism of inositol

Cellular uptake of myo-inositol and incorporation of myo-inositol into lipid and inositol phosphates over 21.5 h is shown in Fig. 2. The accumulation of labeled myo-inositol remains linear for the entire period. At early time points, virtually all of the accumulated label is in the free myo-inositol fraction. With time, an increasing amount of label is incorporated into lipid and then into inositol phosphates. To assess the purity of the myo-inositol fraction, samples were desalted and subjected to paper chromatography in 2 solvent systems. ³H from the myo-inositol fraction co-migrated as a single peak with the myo-inositol standard. Taken together, these experiments demonstrate that metabolism does not account for the accumulation of intracellular myo-inositol. Therefore, myo-inositol is actively transported into these cells. In order to minimize the effect of metabolism of [³H]myo-inositol during long incubations, the remainder of the experiments were confined to 150 min or less.

Sodium gradient dependence of myo-inositol transport

Active transport in mammalian cells is accomplished either by directly using high-energy phosphorylated compounds or by deriving energy from the sodium gradient across the cell membrane through co-transport. Table I shows the results from an experiment measuring myo-inositol uptake in the presence of ouabain, 2,4-DNP and iodoacetic acid, or in sodium-free EBS. The greatest inhibition of transport occurred in the sodium-free EBS. Inhibition of Na⁺,K⁺-ATPase with ouabain, or disrup-
Fig. 4. Kinetics of myo-inositol transport into cultured retinal endothelial cells. The solid squares represent the sodium-dependent myo-inositol transport. These values were calculated by subtracting the measured sodium-independent transport (open triangles) from the total myo-inositol transport (not shown). The inset shows the portion of the Lineweaver-Burk transformation for data points closest to the Y-axis. No weighting of points was performed.

Inhibition of energy metabolism with 2,4-DNP and iodoacetic acid, also reduced myo-inositol transport, although not to as great an extent as did sodium replacement. Fig. 3 shows the velocity of inositol transport as a function of the extracellular sodium concentration. The velocity of myo-inositol transport increases with increasing extracellular sodium, suggesting that the rate of transport is dependent upon the sodium gradient.

**Kinetic parameters of inositol transport**

The initial velocity of myo-inositol transport as a function of culture medium myo-inositol concentration is shown in Fig. 4. The portion of the transport which is sodium independent (as measured in a sodium-free choline buffer) is much less than the portion which is sodium dependent. The apparent $K_m$ for the sodium-dependent uptake was $311 \pm 47$ (S.D.) μM and the $V_{max}$ was $40.8 \pm 2.8$ (S.D.) pmol-mg protein$^{-1}$-min$^{-1}$.

**Specificity of the inositol transport system**

Inositol uptake was finally characterized by measuring transport in the presence of various hexoses, hexitols, and inhibitors of the glucose transport system. The results are shown in Table II. Cold myo-inositol is a very potent inhibitor of radioactive myo-inositol transport, but the other two tested hexitols, mannitol and sorbitol, did not significantly inhibit myo-inositol transport. Fructose did not inhibit transport, but D-galactose, D-glucose and L-glucose all significantly inhibited myo-inositol transport. Finally, phloretin, phloridzin, and cytochalasin B inhibited myo-inositol transport, but the order of potency is different than that for the inhibition of glucose transport, with phloridzin being a more potent inhibitor of myo-inositol transport than phloretin$^4$.

Fig. 5 shows the increasing inhibition of myo-inositol transport with increasing concentrations of glucose. It is important to note that even at physiologic glucose and myo-inositol concentrations, the transport of myo-inositol is inhibited to a measurable extent.

**DISCUSSION**

Endothelial cells derived from retinal microvessels were chosen for this study because they form the blood–tissue barrier at the retinal capillary and prevent many molecules found in the plasma from entering the retina. The function and structure of the blood–retinal barrier is very similar to that of the blood–brain
The eye and brain, these barriers also tend to buffer tissues from swings in plasma concentrations of other molecules, such as amino acids and glucose. Specialized transport systems exist at blood–tissue barriers to permit certain molecules to pass from the circulation into tissues. The major transport system for myo-inositol uptake into cultured bovine retinal microvessel endothelial cells is a saturable, carrier-mediated, active process deriving its energy from the sodium gradient. Myo-inositol is concentrated in these cells to levels well above the concentrations present in the cell culture medium. The apparent \( K_m \) of 311 ± 47 (S.D.) \( \mu M \), found in vitro suggests that the rate of transport of myo-inositol across the barrier in vivo is dependent upon the plasma myo-inositol concentration. Uptake is inhibited by replacement of sodium with choline or by addition of ouabain; both observations indicate sodium-dependent transport. Finally, uptake is inhibited by cytochalasin B, phloretin, phloridzin, \( \alpha \)-glucose, \( \beta \)-glucose, galactose, but not by fructose, mannitol or sorbitol added to the cell culture medium. The order of potency of the transport inhibitors, the sodium-dependence, and the inhibition by \( \beta \)-glucose permit us to conclude that myo-inositol is not transported by means of the major glucose facilitated-diffusion transport system\(^4\). Even under physiologic concentrations of glucose and myo-inositol, there is measurable inhibition of transport by glucose. Since our kinetic analysis was performed in the presence of 5 mM \( \alpha \)-glucose, the calculated Michaelis–Menten constant represents an apparent \( K_m \).

The transport of myo-inositol into neural cells in culture has also been studied. Cultured rat Schwann cells have a saturable sodium-dependent transport system\(^20\). As in endothelial cells, transport is inhibited by glucose, galactose, phloridzin and phloretin. In this system, the \( K_m \) was 30 \( \mu M \). In mouse neuroblastoma cells grown in culture, Yorek et al.\(^29\) found two saturable sodium-dependent transport systems, a high-affinity transport system with a \( K_m \) value of 12.4 \( \mu M \) and a \( V_{\text{max}} \) of 14.1 pmol-mg protein\(^{-1}\)min\(^{-1}\) and a low-affinity system with a \( K_m \) of 433 \( \mu M \) and a \( V_{\text{max}} \) of 215 pmol-mg protein\(^{-1}\)min\(^{-1}\). Myo-inositol transport into neuroblastoma cells was inhibited by glucose and by sorbitol\(^{20}\). In doing these inhibitor studies, the cells were preincubated with inhibitor for two weeks in the presence of sorbitol and in 30 mM glucose. The inhibition by 30 mM glucose had both competitive and non-competitive components, with the non-competitive inhibition being greater. The long preincubation time may account for the differences in the nature of the glucose and sorbitol inhibition these authors observed compared to other investigators.

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In the central nervous system, the tissue concentrations of myo-inositol are tightly controlled\(^1\). Normal cerebral and cerebellar myo-inositol concentrations are maintained even in the presence of dietary restriction\(^1\). Spector described a 3 tier system of myo-inositol homeostasis in brain\(^22\). The first level is regulated by renal excretion and intestinal absorption. The second level is transported into the central nervous system by the choroid plexus. The third level is by carrier-mediated transport from interstitial fluid into brain cells.

The active transport of myo-inositol by retinal microvessel endothelial cells at the blood–retinal barrier may be fundamental to its high concentration in the retina. The retinal capillary is similar to the structure of the cerebral capillary in terms of tight junctions and specialized transport systems which protect neural tissue from fluctuations in the concentrations of plasma components\(^9\). Both the brain and the retina have tissue myo-inositol concentrations that are much higher than plasma. Concentrations of myo-inositol in the vitreous humor and the CSF probably approximate the concentrations in retinal and brain interstitial fluid because there are no barriers to diffusion between these fluid spaces for small polar molecules\(^5\). These fluids have a myo-inositol concentration greater than the plasma but less than the concentrations in the adjacent tissues\(^17,18\). Since myo-inositol is concentrated from the plasma into the extra-cellular fluid of the eye and brain, we presume that the transport system we describe into cultured retinal endothelial cells in vivo is located on the luminal border of the capillaries. Also implicit in this model is a mechanism for release of myo-inositol at the anti-luminal border which allows myo-inositol to flow out of the cell into the interstitial fluid. For the brain, myo-inositol transport may occur both at the capillary endothelium into interstitial fluid, and at the choroid plexus into spinal fluid. For the eye, myo-inositol transport would occur at the capillary endothelial cell into the retinal interstitial fluid. The source of myo-inositol in the vitreous humor is not known, but transport by the ciliary body into the aqueous humor with subsequent diffusion into the vitreous, and transport by the pigmented retinal epithelium may supplement diffusion into the vitreous from retinal interstitial fluid. Retinal and cerebral parenchymal cells further concentrate interstitial myo-inositol. In effect, there is a serial concentration of myo-inositol, first across the blood–tissue barrier, and second into the parenchymal cells of the retina and the brain. Coupling the myo-inositol pumps in series may explain how these two tissues maintain high concentrations of myo-inositol.

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