Dendroaxonal Transcytosis of Transferrin in Cultured Hippocampal and Sympathetic Neurons

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Previous studies using overexpressed polymeric immunoglobulin receptor in cultured neurons have suggested that these cells may use a dendroaxonal transcytotic pathway (Ikonen et al., 1993; de Hoop et al., 1995). By using a combination of semiquantitative light microscopy, video microscopy, and a biochemical assay, we show that this pathway is used by the endogenous ligand transferrin (Tf) and its receptor. Labeled Tf added to fully mature hippocampal neurons changes the intracellular distribution of its receptor from preferentially dendritic shortly after addition to dendritic and axonal at longer times. Incubation of living neurons with (caged)FITC-Tf followed by uncaging in the dendrites results in the later appearance of fluorescence in the axon of the same cell. In "chambered" sympathetic neurons in culture, 125I-Tf or iron as 55Fe-Tf added to the cell body/dendrite chamber is recovered in the axonal chamber, showing that internalized ligand from the cell body–dendrite area is released at the axonal end. Finally, we show that excitatory neurotransmitters increase Tf receptor transcytosis, whereas inhibitory neurotransmitters reduce it. The dendritic uptake, transcytotic transport, and axonal release of physiologically active Tf demonstrated here could be envisioned for other trophic factors and therefore have important consequences for neuronal anterograde target maturation. Moreover, the changes in transcytosis after neurotransmitter addition may be important in the cellular responses that follow electrical activation.

Key words: transferrin; receptor; l-glutamate; hippocampal neurons; sympathetic neurons; transcytosis

In polarized cells the movement of molecules endocytosed at one plasma membrane domain followed by transport to the opposite is called transcytosis (for review, see Mostov, 1991; Nelson, 1992; Mostov and Cardone, 1995). The best-characterized transcytotic molecule is the polymeric immunoglobulin receptor (plgR). Newly synthesized receptors first are targeted to the basolateral surface of epithelial cells, where after binding to the ligand they undergo transport across cells to the apical surface where, after cleavage, ligand and receptor are released. At least three sorting events take place in this event. The first is sorting in the Golgi apparatus for basolateral delivery. The second is sorting in the plasma membrane into clathrin-coated pits for endocytosis and transport to basolateral early endosomes. The third is in the basolateral endosomes, where the receptor must be sorted into transcytotic vesicles and thus avoids transport to the degradative compartment or the recycling compartment or both. In epithelial and endothelial cells, transcytosis has been shown to occur for molecules such as immunoglobulins (Hunziker and Mellman, 1989; Sooranna and Contractor, 1991; Neutra et al., 1988), nerve growth factor (NGF) (Siminoski et al., 1986), epidermal growth factor (EGF) (Maratos-Flier et al., 1987; Brändli et al., 1991), insulin (King and Johnson, 1985), human gonadotropin hormone (hGC), (Ghinea et al., 1994), and transporter molecules such as LDL (Hashida et al., 1986) or transferrin (Tf) (Soda and Tavassoli, 1984; Friden et al., 1991; Cerneus et al., 1993). Neuronal cells, like epithelia, are highly polarized (for review, see Rodriguez-Boulan and Powell, 1992; Craig and Banker, 1994). Although neuronal transcellular transfer has been demonstrated for viruses, toxins, immunoglobulins, trophic factors, and fluid phase molecules (Dumas et al., 1979; Pickard and Silverman, 1981; Evinger and Erichsen, 1986; LaVail and Margolis, 1987; Kuypers and Ugolini, 1990; Ikonen et al., 1993; de Hoop et al., 1995; von Bartheld et al., 1996), the existence of a neuronal transcytotic pathway similar to that of epithelial cells (see above) is suggested primarily from studies with wheat germ agglutinin (WGA) (LaVail and Margolis, 1987) and the plgR (Ikonen et al., 1993; de Hoop et al., 1995). WGA injected into the aqueous chamber of the eye is taken up by endocytosis from the dendrites of the retinal ganglion cells, and after crossing the cells without a major concentration in the Golgi apparatus, it concentrates later in vesicular structures in the axons of the optic nerve. plgR, a transcytotic molecule of epithelia, when expressed in cultured hippocampal neurons is first delivered from the cell body to the dendritic surface, where after binding to its ligand is transported to dendritic early endosomes and later to vesicular structures in the axons.

In this work we have investigated the intracellular trafficking of Tf receptor (TfR) and its ligand Tf. Tf is the major iron-transporting protein in the vertebrate body. Iron (Fe3+) is a
universal cofactor for mitochondrial energy regeneration, and it supports the growth and differentiation of all cell types. In the CNS, iron is a key component of systems responsible for myelination and the synthesis of several neurotransmitters (Beard et al., 1993). Although an essential nutrient, iron is also a potent toxin because it is a powerful oxidant and must therefore be stringently regulated. Dysfunction in iron metabolism has been involved in brain disorders such as Parkinson’s and Alzheimer’s diseases (for review, see Faucheux et al., 1995; Loeffler et al., 1995). Iron reaches the brain using a TfR-mediated process that is not yet fully understood. It is taken up by brain Tf and transported into TfR-expressing cells. In the cytosol, iron is complexed with the iron-binding protein ferritin. Major sites of accumulation of iron in the brain are oligodendrocytes, located mainly in the basal ganglia. Studies comparing the differential localization of iron and TfRs in the brain led to the hypothesis that iron might be vectorially transported via neurons expressing high level of TfRs toward the iron storage zones (Hill et al., 1985). This hypothesis is supported by data showing that injected radioactive iron is found first in the receptor-rich regions and some weeks later in the storage zones (Morris et al., 1992). Axonal release of Tf has been observed in the peripheral nervous system, although neurons do not synthesize any Tf (Markelonis et al., 1985; Kiffmeyer et al., 1991). Although indicative of brain transcytosis of Tf, neither of these observations formally examine the question of neuronal transcytosis or the cell biology of this phenomenon.

Using three complementary approaches, quantitative immunofluorescence light microscopy and video microscopy of hippocampal neurons in culture and a biochemical assay of Tf/iron uptake and release in sympathetic neurons grown in “Campanot” chambers (Campenot, 1992), we show that Tf and its receptor undergo dendroaxonal transcytosis. Moreover, we show that transcytosis of the TfR is modulated by exogenous Tf and neurotransmitters.

**MATERIALS AND METHODS**

**Cells**

*Rat hippocampal neurons.* Hippocampal cells were prepared from 18-d-old rat embryos according to the method of Goslin and Banker (1991). Briefly, the hippocampi are dissociated by trypsin and mechanical treatments. Cells are plated onto poly-l-lysine-treated glass coverslips. After allowing attachment for 4–12 hr in medium (N2 medium) containing 10% horse serum, the coverslips are transferred to dishes containing a monolayer of glial cell in a serum-free medium (N2 medium). Proliferation of non-neuronal cells was prevented by adding 5 μM cytosine arabinoside (ARA-C) (Calbiochem, La Jolla, CA). All experiments were performed in cells kept 14–21 d in culture.

Sympathetic explants and compartmentalized culture. Superior cervical ganglia were dissected from 19- to 20-d-old rat embryos according to the method of Goslin and Banker (1991). The sympathetic ganglia were dissociated by trypsin and mechanical treatments. Cells are plated onto poly-l-lysine-treated glass coverslips. After allowing attachment for 4–12 hr in medium (N2 medium) containing 10% horse serum, the coverslips are transferred to dishes containing a monolayer of glial cell in a serum-free medium (N2 medium). Proliferation of non-neuronal cells was prevented by adding 5 μM cytosine arabinoside (ARA-C) (Calbiochem, La Jolla, CA). All experiments were performed in cells kept 14–21 d in culture.

**TfS and antibodies**

Human holo-Tf (Sigma, St. Louis, MO) was coupled to rhodamine (lissamine rhodamine B sulfonyl chloride, Molecular Probes, Eugene, OR) and CMNB-caged fluorescein SE (5-(((tuccimimidyl)oxy)carbonylethanol)amino)fluorescein 5-carboxymethyl-2-nitrobenzyloether; Molecular Probes) according to the manufacturer’s recommendation, with the following modifications: (1) the fluorochrome was added in five steps at 5 min intervals, and (2) the reaction was stopped after 1 hr in 1 mM glycine, pH 8.5. Coupled and free fluorochrome were separated on a PD10 column (Pharmacia, Piscataway, NJ) and eluted in a Na-HEPES, pH 7.2, 0.15 M NaCl buffer. Human apo-TF (Sigma) was coupled to 3Fe as described (Sterverding et al., 1995).

MAP2 rabbit polyclonal was provided by Javier Díez-Guerra, Centro Biología Molecular, Madrid Spain, and was used at 1:1000 dilution; monoclonal anti-(rat) TfR clone OX26 (PharMingen, San Diego, CA) was used at 1:100.

**Endocytosis experiments**

For the internalization of labeled Tf, Rh or (caged)FITC-cells were incubated for 2 hr at 37°C in culture N2 medium (Goslin and Banker, 1991), with or without Tf, and then incubated for different times in N2 medium (without TF) containing 300 nM of either Rh or (caged)FITC-labeled human Tf. The cells were then fixed and analyzed by immunofluorescence microscopy or mounted for fluorescence video microscopy (see below).

Activation of GABA and glutamate synapses

δt-glutamic acid (glutamate) and GABA (Sigma) were added at concentrations of 5 μM and 10 μM, respectively. Hippocampal neurons were incubated for 4 hr in the presence of Fe-Tf and then for 2 hr in Tf-free medium (starvation) or in Fe-Tf containing medium in the presence of glutamate or GABA, fixed, and stained using an anti-TfR antibody (see below).

**Immunofluorescence**

After Rh-Tf internalization, cells on coverslips were fixed in 4% para-formaldehyde in PBS for 30 min at room temperature followed by quenching of aldehyde groups with 50 mM NH₄Cl in PBS for 10 min, mounted, and analyzed by fluorescence microscopy. For the detection of the TfR under the different experimental conditions (see Results), cells were fixed as above and then permeabilized in PBS/5% blocking solution 0.05% saponine for 10 min at 37°C, incubated for 1 hr with the anti MAP2 and anti-TfR antibodies, washed three times and incubated for 1 hr with fluorochrome-coupled secondary antibodies, washed three times with permeabilizing buffer and once in PBS and once in water, and finally mounted in Dabco/Mowiol as described in Hémar et al. (1995).

**Image analysis**

A semiautomatic program was developed to detect, count, and localize vesicles. It runs on a Series 151 digital image processor (Imaging Technology, Bedford, MA) hosted by a SPARCstation 20 (SUN, Mountain View, CA). The program recognizes vesicles by applying a two-step strategy method (Olivo, 1992): (1) all possible vesicle candidates are detected by local maxima filtering; and (2) a measuring mask is applied at each such position, and real vesicles are selected according to geometrical criteria, such as size and spacing, and statistical criteria, such as intensity, average, and SD values against local background. Region masks defining dendritic and axonal processes are defined by thresholding the fluorescence images and retaining pixels with values above an interactively selected threshold or by interactively drawing an overlay over the phase contrast, and for each image the cross-sectional areas of axons and dendrites are measured (xax and xdd). Vesicles are finally counted and attributed to either the axonal (vxax) or the dendritic (vxdd) compartment by applying a logical AND operator between the vesicle and masks images. The results are presented as the mean ± SEM of the percentage of the number of axonal vesicles relative to the total number of vesicles, (vxax/sax)/(vxax/sax + vdd/dd) × 100. To determine statistical significances, the total number of Tf- or TfR-labeled vesicles in axons and dendrites (see Results for the number of cells analyzed for each experiment) under the different experimental conditions was subjected to Student’s t test analysis as described in Press et al. (1992).

**Video microscopy**

Hippocampal neurons grown at low density were deprived of Tf (starved), incubated for 20 min with (caged)FITC-Tf, rinsed, mounted
in a homemade video chamber containing normal culture medium
(with Tf) without phenol red, and observed with a 63× PlanNeofluar
lens in a Zeiss inverted microscope (Axiovert). Cells grown in relative
isolation with a large dendritic tree and an identifiable axon were
chosen by phase contrast microscopy and then illuminated for 1 sec
with the FITC excitation wavelength. This always resulted in lack of
emission. To uncage the (caged)FITC-Tf, a small region of the den-
drites was illuminated with the UV filter with the field aperture closed
to its minimum (this resulted in the excitation of an area of ~5 µm in
diameter). After UV excitation, uncaging was confirmed by excitation
with the FITC wavelength. The axon of the cell in which uncaging was
performed was then visualized in the FITC channel with a 1 sec
exposure every 5 min. All images were captured with an SIT camera
(Hamamatsu, Photonics Deutschland) and processed in a Power
Macintosh (9100) computer equipped with a Scion LG3 image grabber
(Scion Co.). Images were analyzed using National Institutes of Health
image processing software.

Electron microscopy

Electron microscopy was used to examine Tf-positive structures in the
cells. Neurons, grown on coverslips, were incubated with Tf–HRP for
25 min in serum-free media at 37°C, washed in serum-free media, and
fixed using 2.5% glutaraldehyde in 50 mM sodium cacodylate buffer, pH
7.35, for 30 min at room temperature. After they were fixed, neurons
were washed with cacodylate buffer and reacted with diaminobenzidine
to visualize the Tf–HRP (Parton et al., 1992). Neurons were subse-
sequently post-fixed in 2% OsO4 for 1 hr at 4°C and washed with dH2O.
Cells were selected using an inverted microscope and marked on
coverslips for en face sectioning subsequent to electron microscopy
processing. Cells on coverslips were stained with 0.5% uranyl acetate
and dehydrated through a graded series of ethanol. Epon was infiltr-
ated into neurons for several hours, and the plastic was polymerized
overnight at 60°C. Marked cells were cut out and glued onto blank
plastic stubs so that neurons could be cut parallel to their growing
surface. Sections with a gold interference color were cut, picked up on
formvar-coated grids, and viewed in a Zeiss EM10 electron micro-
scope operated at 60 kV, without section staining.

Immunoprecipitation of TfR

Newly synthesized proteins in sympathetic neurons grown in explants
were labeled with 200 µCi/ml of 35S-methionine for 6 hr in medium
containing 1:10 cold methionine (metabolic pulse). At the end of the
pulse, cells were chased for 16 hr in medium deprived of or containing
100 nM Tf. Then, cell body and axonal masses were separated with a
razor blade, the material was lysed, large aggregates were discarded
after centrifugation, and aliquots of supernatant were subjected to
trichloroacetic precipitate and used for scintillation counting. Equal
amounts of radioactivity were then subjected to immunoprecipitation
as described by De Strooper et al. (1995) using antibody against TfR.
OX-26. Immunoprecipitates were subjected to SDS-PAGE, and the gel
was dried and exposed to x-ray film. Autoradiograms were scanned and
radioactivity was quantitated using National Institutes of Health Image
software.

Transcytosis assay in chambered sympathetic neurons

3H-inulin (TRA 32A, Amersham, Arlington Heights, IL) (0.5 µCi/ml, 1–5
Ci/ml) 1:1000 in medium without Tf was first added in the central
chamber, and cells were incubated for 2 hr at 37°C. Then lateral and
central chambers milieu were collected and washed, and radioactivity was
counted. In nonleaking cultures (as assayed by the lack of 3H-inulin in the
lateral chambers), 300 nM 125I-Tf (labeled as described in Fracker and
Speeks, 1978) was diluted in medium without Tf and added to the central
chamber for 1 hr at 37°C. The central and lateral chambers were then
washed, and the medium in the lateral chambers was replaced by N2
medium without Tf and with 5 mM nitrotriacetic acid to prevent the
re-endocytosis of released Tf. The radioactive Tf endocyctosed in the first
hour was allowed to recycle or transcytose for 2 hr at 37°C, the medium
was collected, and radioactivity was counted. After washing, 96 nM 55Fe
was added to the central chambers of the same cultures, and the same
procedure was followed.

RESULTS

Rhodamine-Tf is redistributed from dendritic to axonal
intracellular compartments in rat hippocampal
neurons in culture

We first analyzed the intracellular distribution of Tf after different
times of internalization. Hippocampal neurons were
preincubated for 2 hr in Tf-free medium (Tf starvation),
and then rhodamine-labeled Tf (Rh-Tf) was added for 20 or 90
min at 37°C. The cells were fixed, and the distribution of the
labeled ligand to the axonal and dendritic territories was
determined and quantified. To define axons and dendrites,
the dendritic territory was labeled with the specific marker MAP2
(Caceres et al., 1984). Axons were then defined by phase
contrast microscopy and by the lack of MAP2 staining. The
results of these experiments are shown in Figure 1. After a
short time of endocytosis, Rh-Tf labeling appeared as small
dots present mainly in the dendrites (Fig. 1A, a–c). Axonal
labeling was evident after 90 min of endocytosis (Fig. 1A, d–f).
To determine the location of the ligand to either axons or
dendrites, we used a semiautomatic computer software (de-
scribed in Materials and Methods) that permitted us to com-
pare the relative amounts of fluorescent vesicles in axons and
dendrites from experiment to experiment. By using this pro-
gram, region masks of dendrite and axonal processes are first
defined. Because some axons may run along dendrites, only the
axons running in isolation were counted. Moreover, to avoid
overestimation of dendritic vesicles caused by vesicles present
in axons running parallel to dendrites, vesicles located on the
edge of the dendritic mask were not taken into account. Then
bright spots representing vesicles were counted and attributed
to either the axonal or dendritic domains, and their number
was normalized per surface unit (Fig. 1B). By this means,
diffuse surface fluorescence is not taken into account. The use
of this semiquantitative method revealed a 46% increase in the
percentage of axonal Tf-labeled vesicles after 90 min of incu-
bation compared with that after 20 min. In absolute values,
18.6 ± 2.8% of Tf-labeled vesicles were found in axons after 20
min of internalization, and 27.2 ± 3.2% were found after 90
min (Fig. 1C).

Although not dramatic, the change in distribution of Tf at
different times of internalization, from preferentially dendritic to
dendritic and axonal, suggests transcytosis. However, the use of
this method does not permit us to conclude whether the axonal Tf
after 90 min is originated in the dendritic/cell body region,
transcytosis, or attributable to a different kinetics of axonal
internalization.

Transcytosis of (caged)FITC-Tf in living
hippocampal neurons

To examine more directly the transcytotic movement of internal-
zated Tf, we then analyzed the distribution of (caged)FITC-Tf in
living hippocampal neurons. Caged compounds are not fluores-
cent on excitation at 490 nm unless first uncaged by excitation at
a wavelength under 360 nm (Mitchinson, 1989). Hippocampal
neurons were incubated for 30 min with (caged)FITC-labeled Tf.
After excess ligand washing, the cells were mounted for video
microscopy analysis in normal culture medium (see Materials and
Methods). Under phase contrast observation, a cell was chosen
and then illuminated first at the excitation wavelength of FITC.
As shown in Figure 2A, no fluorescein labeling was observed in
the entire neuron. Illumination at 360 nm (Hoechst filter) of two
small dendritic areas (~5 µm in diameter) resulted in the appear-
ance of fluorescent dots exclusively in the excited areas (Fig. 2C,D) corresponding to the internalized Tf. The axon of this cell, identified by conventional morphological criteria (thin and of uniform area), did not show any fluorescence immediately after uncaging of FITC in the dendrites (Fig. 2E). However, some labeled vesicles were evident 45 min later (Fig. 2F). Labeled structures were also found in the cell body (not shown). Because Tf was made fluorescent only in the dendrites, this result shows that ligand internalized from this surface can move to the axons, thus proving the existence of dendroaxonal transcytotic movement. Whether Tf transcytosed bound to its receptor was analyzed next.

Tf receptor is redistributed from the dendrites to the axons in the presence of ligand

The intracellular pathway of Tf and its receptor in nonpolarized cells is well known. Diferric-Tf (holo-Tf) binds to its receptor at neutral pH and is endocytosed. At the acidic pH of the endosomes, iron dissociates from Tf (apo-Tf), which still bound to the receptor is recycled back to the cell surface where it dissociates and becomes free for another round of binding and internalization (Dautry-Varsat et al., 1983; Klausner et al., 1983). Thus, inside the cell Tf and receptor remain together. Because in Tf-starved hippocampal neurons Tf is preferentially internalized from dendrites after short times of endocytosis (Fig. 1A, a–c) but then appears in the axons at longer times (Fig. 1A, d–f), a similar change in distribution would be expected for the TfR.

Hippocampal neurons were incubated for 2 hr in medium without Tf followed by incubation in 300 nM Rh-Tf for 20 min (a–c) and 90 min (d–f). a, d, Phase contrast; b, e, MAP2 labeling (dendritic marker); c, f, internalized Tf. After 20 min of internalization (c) Tf positive structures are preferentially dendritic (compare the distribution of dots in c with that of the MAP2-positive processes in b). After 90 min of internalization (f), numerous dots are present in processes (small arrows in f) negative for MAP2 (e). B, Example (based on the cell shown in A, a–c) of how the computer program identifies axons and dendrites and determines the distribution of Tf-positive (rhodamine) dots. The computer recognizes dendrites on the basis of MAP2 labeling (middle panel) and axons by phase contrast (left panel) and lack of MAP2 labeling, and then allocates and counts the Tf dots. C. With longer incubation times there is an increase in Tf-positive structures in axons.
Increased (Fig. 3A, d–f). Quantitative analysis shows that in the presence of Tf in the medium, the percentage of axonal TIR-positive vesicles increased 2.1-fold (29.4% at steady state vs 13.7% after starvation; \( p < 0.001 \)) (Fig. 3B). To prove that indeed the receptor present in axons had reached the axonal surface, cells maintained in Tf-containing medium were incubated for 20 min with Rh-Tf, and the percentage of internalized Tf in axons and dendrites was determined. In these cells, 36.6% of fluorescent dots were in axons (Fig. 3A, g–h; quantitation in Fig. 3B). This amount is significantly different (\( p < 0.05 \)) from the 18.6% observed after 2 hr starvation followed by 20 min internalization (see first part of Results). This result shows that the low axonal labeling observed after 20 min endocytosis of Rh-Tf in starved cells as compared with 90 min endocytosis was not caused by a kinetic difference of Tf endocytosis in axons and dendrites.

Altogether this last series of experiments suggests that both Tf and TIR follow a similar transcytotic route from dendrites to axons and that this transcytosis of TIR is stimulated by the binding of Tf. It is unlikely that the significant change in TIR distribution is caused by an increase in axonal mass, because we are using fully developed, synaptically active neurons that have little growth capacity and, even if they do grow, the time of incubation (2 hr) is too short to account for significant process elongation.

Transcytosed Tf is present in vesicles in the axons

To analyze the morphological appearance of Tf-containing structures in both axons and dendrites, we performed electron microscopy of cells maintained in Tf-containing medium incubated for 20 min with Tf coupled to HRP. HRP-positive structures were indeed found in axons and dendrites (Fig. 4). Vesicles and tubules filled with reaction product were seen in the dendrites. In the axons, labeled vesicles were observed along both axonal shafts (Fig. 4B) and varicosities (Fig. 4C). Although this experiment does not conclusively demonstrate that the axonal Tf-containing vesicles are indeed transcytotic, this could be inferred from the fact that the axonal Tf is in small vesicular structures and not in the large multivesicular body-like vesicles, the classic carriers of retrogradely transported material (Parton et al., 1992).

Use of Campenot chambers reveals that Tf and iron are taken up from the cell body/dendrites and released from the axons

We used dissociated sympathetic neurons grown in Campenot chambers to biochemically determine dendroaxonal transcytosis of Tf. This culture system consists of a central and two lateral chambers. In this compartmentalized culture, axons originating from neurons plated in the central chamber grow across a silicone grease layer barrier and enter a separate fluid environment within
Before using the compartmentalized system we determined whether in sympathetic neurons TfR changed distribution from preferentially cell body/dendritic to axonal in the presence of exogenous Tf (Figs. 1, 3). For this, we used sympathetic explants in which the central cell body mass can be easily separated from the peripheral halo of axons (Fig. 5A). 35S-methionine labeling of newly synthesized proteins followed by a chase in the presence or absence of exogenous Tf and immunoprecipitation with an anti-TfR antibody revealed a significant increase in the amount of TfR in the axonal mass on addition of exogenous Tf (Fig. 5B). This result implies that Tf in the medium changes the distribution of the receptor from preferentially cell body/dendritic to axonal also for sympathetic neurons.

Using the Campenot chambers we next analyzed whether physiologically active Fe-Tf also transcytosed. 125I-Tf was added to the central chamber for 1 hr at 37°C. After they were washed, the axons in the lateral chamber were incubated for 2 hr in culture medium without Tf and with nitrilotriacetic acid, a chelating agent that prevents Tf re-endocytosis after release at the cell surface, and radioactivity was measured in both the central and lateral chambers. Recovery of radioactive Tf in the central chamber reflects recycling, whereas in the lateral chambers it would reflect transcytosis. Figure 6B shows that Tf transcytosed from the central (cell body/dendrites) to the lateral (axonal) chamber. Recovery in the lateral chamber was ~7% of the total Tf in the central and lateral chambers. However, this is an underestimation of transcytosis, because only a small fraction of the axons of the cells are in the lateral chamber. Given the low abundance of TfR in axons in the absence of exogenous Tf, it is unlikely that the Tf released into the lateral milieu comes from Tf that is internalized in axons in the central chamber and anterogradely transported.

To rule out that the radioactivity in the lateral chambers was caused by leakage of medium from the central chamber, 3H-inulin was added to the central chambers of the same cultures, and the radioactivity was measured. Inulin was not found in significant amounts in the lateral chambers. The values for 125I-Tf and 3H-inulin are shown in Table 1.

To determine whether Tf transcytosed bound to iron, we analyzed the routing of 55Fe-Tf. 55Fe-Tf was added to the central chamber of cultured sympathetic neurons, and its appearance was measured in the lateral chamber. Figure 5B and Table 1 show that 55Fe appeared in the lateral chamber at 2 hr of chase. This result suggests that Tf and iron transcytose together. Although not
significant, the proportion of $^{55}$Fe radioactivity recovered in the axonal milieu is higher than that of Tf. This could indicate that the proportion of Fe-bound Tf in the axon (transcytosing Tf) is higher than that in the recycling pathway.

**Transcytosis is influenced by the addition of excitatory and inhibitory neurotransmitters**

In the previous sections we characterized the intracellular trafficking of Tf and TfR and showed data suggesting that transcytosis of the TfR is modulated by Tf. We next asked whether synaptic activity would also affect TfR trafficking. Synaptic activity changes ion fluxes on the postsynaptic cell by activation of neurotransmitter (NT)-receptors. Activation of the intracellular signaling cascades leads to changes in gene expression and phenotype modifications (Ginty et al., 1992). Although the effect of membrane depolarization in synaptic vesicle recycling is well characterized, it is not known whether changes in membrane potential also affect the intracellular trafficking of membrane receptors and ligands present on the dendritic membrane and dendritic milieu. This could have an important role in the plastic changes accompanying increased synaptic activity. To examine this question we added low concentrations of GABA ($10^-5M$) to hippocampal neurons and analyzed and quantitated the intracellular distribution of TfR (Fig. 7). As shown earlier, in cells starved from exogenous Tf, the percentage of TfR in axons is 13.7%. Addition of glutamate to the medium during the starvation period (2 hr) increases this to 24.1%. This difference is statistically significant ($p < 0.001$). A similar effect was found on addition of NMDA to the starvation medium (not shown). However, addition of glutamate to cells grown in the presence of exogenous Tf produced a nonsignificant change in axonal TfR ($p > 0.3$), suggesting that transcytosis is a saturable phenomenon and could be triggered either by increasing the concentration of ligand in the medium or by increasing membrane excitability under basal levels of ligands (Fig. 6), but the effects are not additive. Contrary to the effect of the excitatory agonists, activation of inhibitory receptors by GABA decreases axonal TfR. In Tf-starved cells (-Tf) glutamate addition induces a clear increase in the percentage of axonal TfR-positive structures ($p < 0.001$) (**), whereas GABA addition does not significantly change the number of axonal labeled organelles. In cells grown in the constant presence of Tf (+Tf) a 2 hr incubation with GABA decreases the percentage of axonal TfR ($p < 0.001$ *)). Glutamate addition does not produce a further increase (*).

**DISCUSSION**

Our understanding of transcytosis comes largely from studies in polarized epithelial cells expressing IgR (Mostov and Cardone, 1995). The first step in the transcytotic movement of immunoglobulins is the binding to specific high-affinity receptors on the plasma membrane followed by internalization in clathrin-coated vesicles that pinch off from the membrane and fuse with underlying endosomes. Some of the ligand thus internalized remains associated with the receptor, possibly in an endocytic subcompartment of higher pH than the classic endosome, is packaged into specific carrier vesicles, and is transported to the opposite plasma membrane where release of the ligand occurs. In this work we provide evidence that in polarized hippocampal and sympathetic neurons Tf and its receptor follow a similar "classic" transectotic pathway from the dendrites to the axons. First, exogenous Tf added to Tf-starved cells is taken up from the dendritic surface where the receptor is located (Figs. 1, 3). Second, the exogenous Tf fills submembranous early endosomes (Fig.

### Table 1. Transport of $^{125}$I transferrin and $^{55}$Fe-transferrin from the central to the lateral compartments in “chambered” sympathetic neurons

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Central chamber</th>
<th>Lateral chamber</th>
<th>Percent in the lateral chamber</th>
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<tbody>
<tr>
<td><strong>Experiment 1</strong></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>$^3$H-inuline 2 hr incubation</td>
<td>92456 dpm</td>
<td>576 dpm</td>
<td>0.62</td>
</tr>
<tr>
<td>$^{125}$I-TF 2 hr chase</td>
<td>35655 cpm</td>
<td>2553 cpm</td>
<td>6.7</td>
</tr>
<tr>
<td>$^{55}$Fe-Tf 2 hr chase</td>
<td>11400 dpm</td>
<td>864 dpm</td>
<td>7.0</td>
</tr>
<tr>
<td><strong>Experiment 2</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$^3$H-inuline 2 hr incubation</td>
<td>84964 dpm</td>
<td>1120 dpm</td>
<td>1.3</td>
</tr>
<tr>
<td>$^{125}$I-Tf 2 hr chase</td>
<td>42889 cpm</td>
<td>3497 cpm</td>
<td>7.5</td>
</tr>
<tr>
<td>$^{55}$Fe-Tf 2 hr chase</td>
<td>13672 dpm</td>
<td>3408 dpm</td>
<td>20.0</td>
</tr>
</tbody>
</table>

**Figure 7.** Histogram showing the effect of glutamate and GABA addition on the intracellular distribution of TfR. In Tf-starved cells (-Tf) glutamate addition induces a clear increase in the percentage of axonal TfR-positive structures ($p < 0.001$ (**)), whereas GABA addition does not significantly change the number of axonal labeled organelles. In cells grown in the constant presence of Tf (+Tf) a 2 hr incubation with GABA decreases the percentage of axonal TfR ($p < 0.001$ *). Glutamate addition does not produce a further increase (*).
4). Third, the Tf internalized in the dendrites moves to the axon in vesicular organelles (Fig. 2). Fourth, Fe-Tf internalized in the cell body/dendrite region reaches the axonal plasma membrane and is released to the axonal milieu as Fe-Tf (Fig. 5), showing efficient release of physiologically active ligand. Finally, the intracellular trafficking of Tf/TIR is increased by activation of glutamate (excitatory) receptors and decreased by activation of GABAergic (inhibitory) receptors (Fig. 6).

Our biochemical approach with the Campenot chambers reveals that 7% of Tf transcytoses. This number may be an underestimation, because the number of axes crossing the barrier into the cell body/dendrite-free chamber is far less than the number of cell bodies in the central chamber. Accurate quantitation of transcytosis would require a system in which endocytosis and recycling were measured only in those cell bodies that extend axons in the lateral chambers. No such system is available yet.

We observed that transcytosis of the TfR is stimulated by the presence of ligand; in its absence most receptors are dendritic. This is consistent with previous results using starved cells (Parton et al., 1992). The dendritic distribution of TfR found in non-starved hippocampal neurons (Cameron et al., 1991) might be explained by the presence of apo-Tf, and not holo-Tf, in the culture medium. Although the effect of Tf in the trafficking of the receptor is not very well documented, differences have been measured in the kinetics of endocytosis and recycling of the TfR in the absence and presence of Tf (Girones and Davis, 1989; Sainte-Marie et al., 1991). These authors have shown that Tf accelerates both endocytosis and, to a larger extent, recycling of the TfR, resulting in the highest proportion of TfR at the cell surface in the presence of Tf. Moreover, we could assume that the increase in transcytosis of TfR in the presence of ligand is controlled by a mechanism similar to that resulting in increased transcytosis of the pIgR in the presence of dimeric immunoglobulin A (dIgA) (for review, see Mostov and Cardone, 1995).

We also show that iron is transcytosed, suggesting that the axonally transported Tf is still bound to iron and therefore able to bind to receptors in the target cells on release. This shows that transcytosis of Tf may play a physiological role and that iron accumulation in certain brain regions may arise by this mechanism. Other trophic factors such as FGF and NFG could also act via transcytosis (Ferguson et al., 1990; von Bartheld et al., 1996).

The fact that iron is transcytosed raises the question of the transcytotic pathway in neurons. Under the low pH of endocytic compartments, iron dissociates from Tf (Dautry-Varsat et al., 1983). Therefore the axonally transported iron may be bound to Tf in a nonacidic compartment. Whether separate compartments exist in neurons for the sorting of Tf destined for transcytosis and recycling is not known. In epithelial cells, transcytotic molecules endocytosed from the basolateral surface join an apical recycling compartment in which sorting between basolaterally and apically targeted molecules occurs. In epithelial cells, Tf accumulates in this apical recycling compartment after long internalization times. Whether this is the case in neurons and whether Tf destined for transcytosis accumulates in an axonal “recycling” compartment is not known. In the case of differentiated neurons, we did not detect any “large” Tf-containing compartment in axons (Fig. 1), suggesting that a similar compartment does not exist. However, it is possible that the recycling compartment is present in the cell body. Another interesting and unresolved matter is the mechanism of iron/Tf release on the axonal surface. One can envision that ligand and receptor exposed to the extracellular milieu dissociate because of the low concentration of ligand in this environment, simply following the rules of concentration kinetics. This could explain the release of ligand in the lateral chambers of cultured sympathetic neurons. In situ, ligand would dissociate from the receptor in regions where adjacent cells express a large number of unoccupied receptors.

Our work shows that dendroaxonal transcytosis is used for Tf. This simple discovery opens numerous questions for both cell biology and neurobiology. From a cell biology perspective, it would be of interest to know whether the signals for transcytosis in the Tf receptor are the same as those for other transcytotic molecules (i.e., pIgR), where transcytosing and recycling Tfs are sorted, and what are the mechanisms of transport. For neurobiology, the challenge ahead is the physiological role of transcytosis both during neuronal development and in the mature brain. Our results on the increase of Tf transcytosis after activation of glutamate (excitatory) but not GABAergic (inhibitory) synapses would suggest that the changes that take place after increased synaptic activity might also be mediated by changes in uptake and trafficking of trophic factors. It is well known that anterograde effects in the nervous system, those from neurons to their target organs, are mediated by activation of neurotransmitter receptors on the postsynaptic cell (Comb et al., 1987; Ginty et al., 1992). The activation of neurotransmitter receptors leads to the stimulation of second messenger systems that transfer the signals to the nucleus where expression of specific genes is regulated. Our data show that NT-receptor activation also induces changes in uptake-trafficking of trophic factors normally present in the extracellular milieu. One possibility is that the electrical activity-induced increase in uptake-transcytosis of extracellular ligands in the postsynaptic cell plays a role in the appearance of phenotypic changes normally observed under such conditions. Alternatively, the increase in intracellular trafficking does not have a direct effect on the postsynaptic cell; rather, it may be due to a change in uptake-transcytosis after glutamate/NMDA activation taking place but on distant targets. Whether the increase observed in transcytosis after glutamate/NMDA activation is attributable to an increase in endocytosis or exclusively in transcytosis is not yet clear. Because activation of NMDA receptors by glutamate stimulates protein tyrosine phosphorylation (Bading and Greenberg, 1991) and phosphorylation increases TfR transcytosis in epithelia (Cardone et al., 1994), the increase in axonal TfR could be attributable to higher transcytosis. The data on synaptic activity and transcytosis may have an important physiological connotation; trophic factors present in the external milieu can be more or less efficiently used by the postsynaptic cell or by cells at distant places depending on the type of trans-synaptic activity. Our work paves the way for the study of the functional implications of transcytosis during neuronal development and for the long-term adaptive changes in the mature brain, events in which trans-synaptic regulation of gene expression is involved.

REFERENCES


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