Thyroxine (T4) Transfer from Blood to Cerebrospinal Fluid in Sheep Isolated Perfused Choroid Plexus: Role of Multidrug Resistance-Associated Proteins and Organic Anion Transporting Polypeptides

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Thyroxine (T4) enters the brain either directly across the blood–brain barrier (BBB) or indirectly via the choroid plexus (CP), which forms the blood–cerebrospinal fluid barrier (B-CSF-B). In this study, using isolated perfused CP of the sheep by single-circulation paired tracer and steady-state techniques, T4 transport mechanisms from blood into lateral ventricle CP has been characterized as the first step in the transfer across the B-CSF-B. After removal of sheep brain, the CPs were perfused with 125I-T4 and 14C-mannitol. Unlabeled T4 was applied during single tracer technique to assess the mode of maximum uptake (Umax) and the net uptake (Unet) on the blood side of the CP. On the other hand, in order to characterize T4 protein transporters, steady-state extraction of 125I-T4 was measured in presence of different inhibitors such as probenecid, verapamil, BCH, or indomethacin. Increasing the concentration of unlabeled-T4 resulted in a significant reduction in Umax%, which was reflected by a complete inhibition of T4 uptake into CP. In fact, the obtained Unet% decreased as the concentration of unlabeled-T4 increased. The addition of probenecid caused a significant inhibition of T4 transport, in comparison to control, reflecting the presence of a carrier mediated process at the basolateral side of the CP and the involvement of multidrug resistance-associated proteins (MRPs: MRP1 and MRP4) and organic anion transporting polypeptides (Oatp1, Oatp2, and Oatp14). Moreover, verapamil, the P-glycoprotein (P-gp) substrate, resulted in ~34% decrease in the net extraction of T4, indicating that MDR1 contributes to T4 entry into CSF. Finally, inhibition in the net extraction of T4 caused by BCH or indomethacin suggests, respectively, a role for amino acid “L”

Abbreviations: THs, thyroid hormones; T4, thyroxine; T3, triiodothyronine; CSF, cerebrospinal fluid; CP, choroid plexus; BBB, blood–brain barrier; B-CSF-B, blood–cerebrospinal fluid barrier; CNS, central nervous system; ECF, extracellular fluid; ISF, interstitial fluid; BCH, β-2-aminobicyclo-(2,2,1)-heptane-2-carboxylic acid.
INTRODUCTION

Thyroid hormones (THs) are important regulators of normal growth and development in the central nervous system (CNS) and brain (1–4). Thyroxine (T4), a major type of lipophilic TH, is transported between blood and cerebrospinal fluid (CSF) in a restricted manner, which does not follow simple diffusion mechanism (5, 6). The absence of triiodothyronine (T3) and thyroxine (T4) hormones, such as in hypothyroidism, leads to serious damage in the brain and neuronal cells (7). Therefore, it has been suggested that the blood–brain barrier (BBB) and/or blood–CSF barrier (B-CSF-B) control thyroxine availability to the cerebral compartments (8). Indeed, thyroxine enters the CSF and brain parenchyma by two possible routes: either across the BBB, located at the level of cerebral capillary endothelium into brain extracellular fluid (ECF) and then by diffusion into the CSF (9), or via the B-CSF-B, formed by the choroid plexus (CP) epithelium (9, 10). However, the quantitative extent to which the BBB and B-CSF-B/CP contribute to T4 transport to the brain is poorly understood.

It was previously shown that the level of THs increases rapidly, within minutes of their intravenous (i.v.) injection, using an in vivo dog model (11). In fact, T4 cross instantly from blood into CSF through a carrier-mediated process. In addition, it was demonstrated that 1 h after i.v. injection of radiolabeled-T4, it accumulates to a large extent in the CP and gray matter, before any appearance in the white matter of the brain (12). This accumulation in the CPs, and the subsequent rise in the CSF levels, cannot be accounted for through a free diffusion mechanism from circulating plasma where THs are mostly found as protein bound. However, this increase is likely to occur through a carrier transport mechanism present at the blood side of the CP. Since the rate of T4 equilibrium into CSF is more rapid than that into brain, the CP might constitute another major pathway for the entry of these hormones into the CSF.

Thyroid hormone action at the cellular level depends primarily on the binding of T3 to its nuclear receptors (13), through type 2 deiodinases (D2), expressed in astrocytes. In fact, ~50% of intracellular T3a, active form of the hormone, derives from T4 already present within the brain. On the other hand, the remaining 50% of T3 depends on the entry of T4 from the circulation into the brain through various transporters that act across the BBB and CP. However, the transport mechanisms of THs into brain and the role of the CP transporters in this context are still poorly understood.

Using a rat model, a carrier transport mechanism was identified for T4 and T3 uptake at the BBB; however, their high accumulation by the CP was not investigated (14). Furthermore, it was shown that CP of the rat can accumulate T4 more rapidly than any other region in the brain (15). We have also revealed in an in vivo rabbit model that the distribution of T4 from CSF into the brain and CP is dependent on carrier-mediated transport mechanisms (16). In fact, the CP may potentially contribute to THs homeostasis in the brain ECF since the CSF secreted by the CP is in direct contact with the ventricular/sub-ventricular regions and the brain interstitial fluid (ISF). On the other hand, the BBB has been thought to be the major pathway for T3 and T4 entry into CNS ISF since its surface area is greater than that of the CP. Nevertheless, it was shown that the surface area of the CSF face of the CP may have a greater transport capacity, especially during early stages of brain growth and development (17).

Different transport mechanisms for the movement of THs from CP epithelial cells into CSF (15, 18) and from CSF into brain (19, 20) have been identified in earlier studies. However, only limited information is known about the initial uptake process from blood to CSF and then into CSF. Several limitations exist for the study of T4 uptake using in vitro and in vivo techniques. In fact, in vitro studies are complicated by the inability to gain access to the blood side of the CP while in vivo studies cannot distinguish the transport across the B-CSF-B/CP from that across the BBB. The current knowledge on how the brain regulates TH homeostasis is incomplete, and the role of B-CSF-B/CP is still not fully understood.

In this study, we have used an in situ-isolated perfused CP of the sheep which can selectively examine the B-CSF-B/CP, in complete separation from the BBB (19). Indeed, we have previously demonstrated, using this model, that 125I-T4 uptake at the blood face of the CP was mediated by both saturable and non-saturable uptake processes (19). Therefore, this study investigates the extraction of 125I-T4 at the basolateral (blood) side of the in situ perfused CP of the sheep, and the role of some protein transporters. Finally, the characteristics of T4 transport mechanisms were also examined using various drug inhibitors.

RESULTS

CSF Secretion Rate

The CSF secretion rate was measured in an in situ-isolated perfused CP of the sheep (Figure 1A). Experiments were stopped after 4 h of perfusion since an increase in the arterial pressure and a decrease in the CSF secretion rate were observed, clear indications of tissue deterioration. Results showed that the rate of CSF secretion remained constant during the 4 h of CP perfusion. In fact, the average secretion rate during 4 h was 132.1 ± 4.4 μl/min/g (n = 14, Figure 1B), consistent with previously published studies (21).
### Uptake of 125I-Labeled T4 Using Single Circulation Method

Mannitol was used as an extracellular marker, which allowed the measurement of thyroxine net cellular uptake, using the single-circulation paired tracer dilution technique at the basolateral side of the isolated perfused CP ([Figure 2A](#fig2A)). Indeed, mannitol can only diffuse from the vascular compartments across the fenestrated capillaries and is not taken up into the CP cells via any carrier-mediated process (10, 19). However, some mannitol diffuses across the CP via the paracellular route as the CP tight junctions are more permeable than those of the BBB. Comparison between the percentage recoveries of thyroxine versus that of mannitol ([Figure 2A](#fig2A)) enables the measurement of the net cellular uptake across the plexuses ([Figure 2B](#fig2B)) and hence corrects for any diffusion between the cells. Results showed that during the first 10 s of perfusion, the average maximum uptake of 125I-labeled T4 on the blood side of the CP was found to be 30% ([Figure 2B](#fig2B)).

The characteristics of basolateral transport of 125I-labeled T4 were then investigated, by measuring the maximum uptake ($U_{\text{max}}$) in 20 drops of perfusate and in less than 60 s time period ([Figure 3](#fig3)). Results showed that there was a significant decrease in the $U_{\text{max}}$% in presence of different concentrations of unlabeled-T4 ([Figure 3](#fig3)). Indeed, the $U_{\text{max}}$% fell from ~22%, when only trace levels of 125I-labeled T4 were present, to ~12% after the addition of 25 µM unlabeled-T4 ([Figure 3](#fig3)). In addition, a higher concentration of unlabeled-T4 (50 µM) caused a further significant reduction in the $U_{\text{max}}$%, indicating increased saturation of T4 carrier-mediated proteins. Moreover, complete saturation was achieved in presence of 100 µM of unlabeled T4 ([Figure 3](#fig3)).

Furthermore, data also showed that the obtained $U_{\text{net}}$% decreased as the concentration of unlabeled-T4 increased ([Table 1](#table1)). The inhibitory effect on $U_{\text{net}}$% ranged between ~45 and 71%, which was consistently lower than $U_{\text{max}}$, suggesting the presence of a significant amount of tracer backflux with time ([Table 1](#table1)).

In summary, increasing the concentration of unlabeled-T4, from 25 to 200 µM, resulted in a significant reduction in $U_{\text{max}}$%, which was reflected at various levels of inhibition. In fact, high concentrations of 100 and 200 µM of unlabeled-T4 caused a
complete inhibition of T₄ uptake into CP. Taken together, there was a significant decrease in the $U_{\text{net}}$% in presence of different concentrations of unlabeled-T₄, consistent with an increase in the inhibition level.

**Effect of Various Drugs on the Extraction of $^{125}$I-T₄, Using the Steady-state Method**

The characteristics of basolateral transport of thyroxine under the effect of various drugs were then investigated using the steady-state method. However, before evaluating the effect of each drug on the uptake of $^{125}$I-labeled T₄, steady-state extraction uptake of $^{125}$I-labeled T₄ was performed by collecting perfusates every 4 min, for a period of 1 h (Figure 4). In the steady-state method, the perfusion fluid contained 0.555 MBq of $^{125}$I-T₄ tracer and 2.77 MBq of $^{14}$C-mannitol extracellular marker in 100-ml perfusate. Results showed that steady-state extraction of $^{125}$I-labeled T₄ from the blood side was ~38% (Figure 4). In addition, the net extraction of $^{125}$I-T₄ reached ~16%, when the reference molecule mannitol was subtracted, indicating a role for protein transporters on the blood side of the tissue.

In order to understand the mechanism by which T₄ is transported from the blood to CSF across the CP, a number of drugs were used to target multidrug resistance-associated proteins (MRPs: MRP1 and MRP4), organic anion transporters and P-glycoprotein (P-gp). It has been suggested that these transporters might be involved in the uptake of thyroxine from blood into CP (20–23). The steady-state extraction of $^{125}$I-T₄ at the blood side (basolateral) of the perfused CP over 1 h was calculated relative to the extracellular marker $^{14}$C-mannitol. This allows to measure the extraction of $^{125}$I-T₄ in presence of drugs known to inhibit the efflux transporters, at the basolateral side of the CP, such as probenecid, verapamil, BCh, and indomethacin (24, 25). The net extraction of $^{125}$I-T₄ from the blood to CP, which represents the non-specific paracellular loss of T₄, was calculated by subtracting the extraction of $^{14}$C-mannitol from that of $^{125}$I-T₄ (Ex. $^{125}$I-T₄ − Ex. $^{14}$Cmannitol). It is important to note that each drug was applied at the same concentration at the blood and CSF sides of the CP (basolateral versus apical sides, respectively).

Data showed that probenecid caused the highest percentage change, in comparison to control. Indeed, after its addition, a ~45% significant inhibition was observed in the net extraction of $^{125}$I-T₄ ($p < 0.05$; Table 2). This suggests that probenecid has competed with the same Oatps for T₄ transport across the basolateral membrane of the CP. In fact, since all Oatps are probenecid sensitive (24), Oatps localized on the basolateral membrane of the CP (Oatp2 and Oatp14) (24, 26) as well as Oatp1 on the apical side (27) could mediate the transport of T₄ from the blood to the CSF.

Similarly, the addition of verapamil, the P-gp substrate, at a concentration of 10 µM resulted in a significant ~34% decrease in the net extraction of $^{125}$I-T₄ ($p < 0.05$; Table 2). However, the addition of the large neutral amino acid analog BCh, which is specific to “L” system, has produced a modest ~17% reduction in the net extraction of $^{125}$I-T₄, in comparison to control ($p < 0.05$; Table 2). Finally, indomethacin, an inhibitor of the organic anion transporter 1 (Oatp1), had a similar inhibitory effect to BCh ($p < 0.05$; Table 2). These results suggest that verapamil, BCh, and indomethacin reduce the net extraction of $^{125}$I-T₄ indicating

### Table 1 | The effect of different concentrations of unlabeled thyroxine (T₄) on the net uptake ($U_{\text{net}}$%) of radiolabeled $^{125}$I-T₄, from the blood side of the isolated perfused choroid plexuses of the sheep, using the paired tracer dilution technique.

| Concentration of unlabeled T₄ (µM) | $U_{\text{net}}$ (%) | Inhibition (%) |
|-----------------------------------|---------------------|--------------|
| Control (1.8 nM)                  | 17.0 ± 2.8          | n.d.         |
| 25 µM                             | 9.1 ± 1.9*          | 44.7         |
| 50 µM                             | 2.1 ± 0.2***        | 71.2         |
| 100 µM                            | 0***                | 100          |
| 200 µM                            | 0***                | 100          |

The inhibitory effect is represented in percentages (%). Values correspond to mean ± SEM, number of sheep n = 4 for each condition. *p < 0.05; **p < 0.001, in comparison to control.

**FIGURE 3 | Calculation of $U_{\text{max}}$ and $U_{\text{net}}$.** The inhibitory effect of different concentrations of unlabeled-T₄ on the calculated $U_{\text{max}}$% in the isolated perfused CP of the sheep, using the single-pass method. The maximum uptake of T₄ ($U_{\text{max}}$) is recorded when the maximum uptake of radioactivity has occurred. Results are expressed as the mean ± SEM. Statistical significance were determined using the Student’s t-test and shown as *p < 0.05, **p < 0.001, ***p < 0.0001.

**FIGURE 4 | Net extraction of T4 using the steady-state method.**

Steady-state extraction of $^{125}$I-labeled T₄, from the blood side was ~38%, whereas the net extraction reached ~16%, when the reference molecule mannitol was subtracted. Results are expressed as the mean ± SEM. Statistical significance were determined using the Student’s t-test and shown as *p < 0.05.
TABLE 2 | The effect of various drug inhibitors on the extraction of 125I-T4 from the blood side of the CPs.

| Extraction of 125I-T4 from blood to CP (%) | 125I-T4 | Mannitol | Net Ex | % of paired control |
|------------------------------------------|---------|----------|--------|-------------------|
| Mean control                             | 0.53 ± 0.03 | 0.32 ± 0.01 | 0.20 ± 0.02 | n.d. |
| PROB (1.0 mM)                             | 0.38 ± 0.05 | 0.24 ± 0.10 | 0.13 ± 0.05 * | 45.40 |
| VERAP (10 mM)                             | 0.45 ± 0.02 | 0.29 ± 0.01 | 0.16 ± 0.002 * | 34.30 |
| BCH (5.0 mM)                              | 0.49 ± 0.01 | 0.36 ± 0.02 | 0.13 ± 0.03 | 17.60 |
| INDO (1.0 mM)                             | 0.41 ± 0.06 | 0.28 ± 0.02 | 0.13 ± 0.04 * | 16.20 |

This group of experiments was paired experiments, each with its own control. PROB, probenecid; VERAP, verapamil; BCH, β-2-aminobicyclo-(2,2.1)-heptane-2-carboxylic acid; INDO, indomethacin; CP, choroid plexus. Values are mean ± SEM, n = 3–6, student paired t-test. *p < 0.05, in comparison to control.

DISCUSSION

The present study investigated the steady-state extraction of 125I-T4, transport at the basolateral (blood) side of isolated in situ-perfused CP of the sheep, in the presence of extracellular marker 14C-mannitol. Results demonstrated a carrier-mediated transport mechanism for T4, movement at the basolateral side of the CP involving various transporters. The following lines of evidence support the above statement: (1) the average secretion rate during the single pass technique was measured relative to mannitol, a passively distributed molecule. Therefore, the net steady-state extraction at both sides of the CP, mediating the uptake of T4. This is supported by a previous study in Xenopus oocytes, which showed that both transporters are multifunctional and involved in the transport of THs in the brain (39), retina, kidney, and liver (22). However, this does not exclude a role for Oatp1 or Oatp3 on the apical face of the CP epithelial cells, which are also probenecid sensitive (37, 40). Finally, Oatp14 (known as Oatp1c1) transporter has also been implicated in TH transport.

a role for P-gp, “L” system and Oatp1, respectively, in transporting T4 across the basolateral membrane, from the blood to CP.

During the control, the maximum uptake (Umax) of 125I-T4 in the single pass technique was measured relative to mannitol, a passively distributed molecule. Therefore, the net steady-state extraction reflects both uptake and efflux back to the blood, resulting in a lower extraction compared to Umax. Since Umax is an index of unidirectional uptake at the blood side of CP, our results suggest that the basolateral membrane has a high transfer rate for T4. The net uptake (Unet) was more consistent than Umax, which reflects a small backflux of T4 into the venous effluent. Our findings are consistent with a previous study showing that Umax for T4 was significantly inhibited at high concentrations of unlabeled-T4, exceeding those of physiological conditions (19).

In addition, a carrier transport mechanism for T3 and T4 has also been identified at the BBB in a rat model (14). Furthermore, our previous studies have shown that in an in vivo Ventriculo-Cisternal perfused rabbit model, a large accumulation of 125I-T4 in the CP was reduced by 80% in the presence of 200 µM of unlabeled-T4, and proved to be a component of saturation (16). Our results demonstrated that the entry of T4 from the blood into CP is partially mediated by a saturable process, since the uptake of T4 was markedly inhibited by excess of unlabeled-T4. This suggests that a carrier-mediated transporter is localized in the CP, used as a pathway for T4 entry from blood into the CSF compartment. Indeed, this confirms our hypothesis that carrier-mediated transport for T4 at the basolateral membrane of the CP may contribute to TH homeostasis in brain ECF. Although this study did not investigate the fate of T4 after its entry into the CP, T4 is known to bind to other proteins such as albumin (Alb), thyroid-binding globulin, or transthyretin (TTR). Finally, T4 could also be transported into the CSF space from the CP when complexed to TTR, or as a free hormone (28).

The existence of mechanisms regulating the transport of TH has been suggested in cerebrocortical neurons (29), astrocytes (30), glial cells (31), hepatocytes (32, 33), erythrocytes (34), and skeletal muscle (35). The cellular influx and efflux of THs are facilitated by transmembrane protein transporters; therefore, this study investigated the role of some of these transporters located at the basolateral side, using the isolated perfused CP. The steady state of T4 at the basolateral face was measured in presence of various inhibitors such as probenecid, verapamil, BCH, and indomethacin, which were added to the blood side of CP. In steady state, the CP secretes CSF into lateral ventricle, supporting the net flux of T4, from blood to CSF. Since unbound T4 concentration in the CSF (70 pM) is much higher than that of the plasma (20 pM), the net flux cannot be determined only by the rate of CSF secretion.

In order to characterize the systems implicated in the transport of 125I-T4, from basal to apical sides of the CP, the role of organic anion transporters were examined. Following the addition of probenecid, there was a significant inhibition in the net extraction of T4 after >30 min of perfusion. Reduction in the net uptake of 125I-T4 is indicative of probenecid inhibition to Oatp2, a sodium-independent transporter located at the basolateral side of the CP (36). In addition, Oatp2 is also localized at the abluminal and luminal sides of the brain capillary endothelial cells and is involved in transporting anions such as taurocholate, cholate, bile acids, estrogen conjugates, ouabain, and digoxin (37). In fact, taurocholate has been shown to have a similar effect to probenecid, providing additional evidence for the role of Oatp2 in the efflux of T4 from brain to the blood side (38). Moreover, our data suggest that Oatp2 and Oatp3 are localized on both sides of the CP, mediating the uptake of T4. This is supported by a previous study in Xenopus oocytes, which showed that both transporters are multifunctional and involved in the transport of THs in the brain (39), retina, kidney, and liver (22). However, this does not exclude a role for Oatp1 or Oatp3 on the apical face of the CP epithelial cells, which are also probenecid sensitive (37, 40). Finally, Oatp14 (known as Oatp1c1) transporter has also been implicated in TH transport.
described to play a role in the uptake of T4 at the basolateral side of CP epithelial cells (26), and it is important in transporting T1 at the BBB since it has a high Km for T1 (41). Indeed, Oatp14 has been shown to localize on the BBB and to contribute to T1 uptake into brain (41). In summary, our results demonstrated that various Oatps are involved in the uptake of T1 from the blood into CP, and that their role in the transport of T1 in previous work has been underestimated.

Following the addition of verapamil, a well-established substrate for P-gp (42), a marked reduction was observed in the net extraction of 125I-T1, suggesting that P-gp multidrug resistance MDR1 is involved in T1 transport from blood to the CP. This clearly indicates that verapamil interacts with P-gp resulting in a reduction in the amount of T4 recovered from the blood side, which may then cross into CSF through the apical membrane of the plexus. In fact, previous studies showed that MDR1 localizes sub-apically of the CP and confers an apical-to-basal transepithelial permeable barrier (43). In addition, verapamil has also been demonstrated to inhibit MDR/P-gp and to slow T3 efflux from rat hepatoma, cardiomyocytes, and fibroblasts (44, 45). This is in accordance with previous data showing that verapamil inhibited the efflux of Tn, reflecting an involvement of ABC transporter (44). Moreover, verapamil might also interact with MRPl located at the basolateral side and therefore prevents T1 from exiting the CP toward the blood. Furthermore, verapamil has also been shown to reduce the unidirectional transport of the anticancer drug vincristine, from basolateral to apical side of the brain capillary endothelial cells (42).

Few studies were performed on the role of “L” system transporter in the transport of T1 from the blood to CP. Following the addition of BCH, an amino acid analog, a significant inhibition was observed in the extraction of 125I-T1 at the basolateral face of the CP. This suggests that removal of T1 from CP to the blood side is mediated by the amino acid “L” system on the basolateral side. This would ultimately affect T1 action within tissue cells, leading subsequently to changes in the total concentration of T1, available in the CSF under normal physiological conditions. Previous studies have shown that cross-competition exists between BCH and THs in mouse neuroblastoma cells (46), also reported in the BBB (47). It has also been demonstrated that BCH caused a weak inhibition of T1 uptake at the basolateral side, confirming “L” system contribution of TH transport in isolated perfused CP (19). Finally, it is not known whether BCH would specifically displace THs from intracellular binding sites since it does not affect cytosol–nucleus movement of T1 in BeWo cells (48).

Following the addition of indomethacin, an established inhibitor of MRPl (49) and Oatp1 (50), the net extraction from blood to CP at the basolateral side was significantly inhibited. Our data suggest that MRPl and MRP4, located on the basolateral membrane of CPs (51), and Oatp1, located at the apical side of the CPs, are involved in mediating T1 transport from blood to CP. Therefore, it would be expected that T1 accumulated in the CP since it was not transported out into CSF, which tends to oppose any further entry of T1 from blood to CP.

Taken together, the inhibition in the uptake of 125I-T1 suggests the presence of a carrier mediated process at the basolateral side of the left ventricle choroid plexus (LVCP). The presence of this carrier at the B–CSF-B may contribute to T1 homeostasis in the brain ECF. In addition, the significant inhibition in 125I-T1 transport in presence of probenecid suggests the involvement of Oatp1, Oatp2, and Oatp14 in the transport of T1 into brain. The inhibitory effect of verapamil on the extraction of 125I-T1 from blood suggests that MDR1 contributes to 125I-T1 entry into CSF. Finally, inhibition in the net extraction of T1 caused by BCH or indomethacin suggests, respectively, a role for amino acid “L” system and MRPl in mediating T1 entry into CSF. Indeed, the carrier-mediated mechanism together with MDR1 and Oatps (Oatp1, Oatp2, Oatp14) might mediate a bidirectional transport of T1 from the circulating blood into the brain, playing a role in maintaining T1 concentration in the brain (Figure 5).

Several classes of TH transmembrane proteins belonging to the family of solute carrier (Slc) transporters have been identified such as Oatps and L-type amino acid transporters, which actively participate in the entry and exit of THs into and out of cells (52, 53). The apparent competition between the drugs that were used and the presumed T1 transporters, on either side of the CP, is indicative to the potential role of these transporters in T1 homeostasis into CP/CSF/brain. We hypothesize that the entry of T1 into CP tissue is not only driven by the lipid partitioning of the molecule into CP but also by a carrier-mediated transport mechanism. The drugs caused a reduction in this extraction leading to an inhibition of the transporters located on the blood side (namely, Oatp1, Oatp3, Oatp14, L system), preventing T1 entry into CP. Concurrently, since the latter transporters act in a bidirectional fashion, T1 transport toward the CSF cannot be ignored because apically localized transporters (namely, P-gp, Oatp1, “L” system) become activated, allowing T1 entry from CP into CSF (Figure 5). However, the role of efflux transporters MRPl and MRP4 is to function unidirectionally, from CP to blood, in order to prevent the accumulation of T4 into CP, by removing T1 from CP ECs. This will allow to maintain an uphill concentration gradient of T4 in blood, balancing T1 concentration in CP/CSF. Since P-gps and MRPs have been reported to transport different structurally and functionally unrelated toxic xenobiotics, natural product drugs, phospholipids, and conjugated compounds (54, 55), we propose that they might directly contribute to the B–CSF-B barrier to the entry of T1 in CSF/brain. Indeed, transporters such as Lat1, Mrp1, and Mrp4 were detected on the basolateral surface of LVCP ECs (51). When the extraction was measured from blood to CP, the influx of T1 into the CSF was mediated via P-gp, whereas Oatps (Oatp1, Oatp3, Oatp14) can function bi-directionally, allowing the influx and efflux of T1 movement into and out of CSF. Therefore, the involvement of the latter transporters in the net influx of T1 from blood to the CSF might explain why the free T1 concentration in the CSF is greater than that found in the plasma (11). On the other hand, various substrates were reported to compete and to be transported by Mrp1, including both organic anions and some cationic compounds. For instance, glucuronide conjugates, such as estradiol-17β-glucuronide (E217βG), and sulfate conjugates, such as estrone 3-sulfate, are among those preferred substrates (56). In addition, along with GSH, Mrp1 is capable of cotransporting certain cationic compounds such as the anti-cancer
drugs etoposide and vincristine (56). However, Mrp1 basolateral localization allows for efflux of substances from CSF into blood circulation (57). It is important to note that several classes of transporters such as Oatps, Na(+)/taurocholate co-transporting polypeptide, and amino acid transporters have been reported to transport TH (58).

In conclusion, the presence of transporters for cellular influx on the basolateral membrane of the CP would account for the efficient transcellular transport of T4 from blood to CSF, across the CP. The transport of T4 across the plasma membrane determines the intracellular concentration of the genomically active T3 (nuclear T3 receptor) which in turn depends on TH (T3 and T4) transport to the target cell and the activity of the different deiodinases.

**MATERIALS AND METHODS**

**Experimental Setup**

The method of isolated perfused lateral CP of the sheep was used in this study (18, 19). Briefly, sheep of either sex (Clun Forest strain) weighing 20–35 kg and aged 6–12 months old were used. They were anesthetized with intravenous (i.v.) injection of thiopentone sodium (20 mg kg⁻¹), heparinized (25,000 U, i.v.), and then exsanguinated. The brain was rapidly and carefully removed from the skull after all vessels and connections had been severed. The total number of animals used was 14 sheep. This study was approved by the ethical committee at the university.

**Cannulation of the Internal Carotid Arteries (ICAs)**

Both ICAs were cannulated on the base of the brain. Perfusion system was then started at 0.5–1.5 ml-min⁻¹ using a peristaltic pump (Watson-Marlows, UK). All other vessels in the circle of Willis were tied off in order to direct the perfusate into the anterior choroidal arteries supplying the lateral CP. The optic nerves were then sectioned allowing the brain to be removed from the skull cavity. The lateral ventricles were then opened, and the CPs were exposed and superfused with artificial cerebrospinal fluid (aCSF) and kept moist during the experiment. The venous outflow from both CPs was collected at a regular interval via a cannula inserted into the great vein of Galen.

**Perfusion Fluids**

After cannulation of ICAs, the CPs were perfused with mammalian Ringer solution containing 4.0 g·dl⁻¹ bovine serum albumin (Sigma Fraction V, UK). The composition of the perfusion fluids in the perfusate was (in millimolar): Na⁺ 145.8, K⁺ 5.4, Cl⁻ 119.7, HCO₃⁻ 25, HPO₄²⁻ 1.2, Ca²⁺ 2.35, Mg²⁺ 1.13, and glucose 5.0, in addition to 40% Dextran 70 in saline in order to maintain the colloid osmotic pressure in the absence of any protein (all compounds purchased from Sigma, UK). The perfusate was gassed with 95% O₂ and 5% CO₂, de-bubbled, pre-warmed to 37°C, and filtered with polymer wool before entering the plexus. During the experiment, the brain perfusion preparation was kept warm at 37°C in a water jacket, and by an external heat source. The composition of
perfusion fluids in the aCSF contained in (millimolar): Na+ 148, K+ 2.9, Cl− 135, HCO3⁻ 26, HPO4²⁻ 0.25, Ca²⁺ 2.5, Mg²⁺ 1.8, and glucose 5.0. The aCSF was pre-warmed to 37°C, gassed with 5% CO₂ in O₂, and its pH adjusted to 7.2 prior to reaching the CPs. The perfusion pressure and brain temperature were continuously monitored by a pressure transducer and digital probe thermistor (Edal, CD model, UK). Under these experimental conditions, the brain preparation was viable for at least 5 h. A loss in viability was indicated by a rise in arterial pressure and a fall in the venous outflow.

CSF Secretion Rate
This was determined from the difference in concentration of Evans blue albumin in arterial and venous perfusate samples. The concentration of the dye was determined using Unicam spectrophotometer at a wavelength of 625 nm. The secretion is given by Ki = Fv(V/A)−1 μl/min/g, where Fv = venous perfusion flow rate (μl/min/g wet weight), and V and A corresponds to venous and arterial spectrophotometer readings, respectively (59).

Paired Tracer Indicator Dilution Technique
This technique was first developed to study the transport of sugar and amino acids across the isolated perfused CP of the sheep (59). We performed this technique in order to measure the uptake of 125I-labeled T4 from a bolus injection during a single circulation of the CP. This uptake of T4 is proportionally related to the passage of a non-transported marker molecule, 14C-mannitol. The uptake of [125I]-T4 (both net and maximum) was measured under conditions in which the isolated lateral CPs were perfused with different concentrations (50–200 μM) of unlabeled T4. Under these conditions, the 100-μl bolus contained both isotopes (labeled-[125I]-T4 and 14C-mannitol) in addition to different concentrations (50–200 μM) of unlabeled T4.

Experimental Procedure
A 100-μl bolus Ringer solution (perfusate) containing 3 μCi 125I-T4 and 1 μCi 14C mannitol was injected into a calibrated sidearm in the perfusion circuit and then switched into either left or right CPs via closed system of taps. After 25 s, the dead space within the tubing was cleared and then a run of 20 sequential “one drop” samples of venous effluent were collected in ~60 s, followed by continuous collection of one final sample during 4 min in order to calculate the flow rate. This was considered as 1 cycle of 21 samples per run, followed by collection of a clearance sample during 10 min. The above cycle was repeated for at least 4 times (n = 4–6) accounting for a total of at least 84 samples per brain. A 3.5 ml of scintillation liquid Ultima Gold (Packard, UK) was added to each of the 20 drops collected, as well as to the samples of the injected bollus. The activities of 125I and 14C in the samples were then counted and analyzed.

Calculation of the Recovered Isotopes
After counting the samples, the recovered 125I and 14C in each of the 20 drops was then expressed as a percentage of the 125I and 14C injected in the 100-μl bolus (% of injectate recovered). The following equation was used to calculate the percentage uptake (U%) for each drop, based on the differences in recovery of the two isotopes, taken into account that for any given drop the recovery of 125I-T4 from the CP is far less than the recovery of 14C-mannitol.

\[ U\% = \frac{100\times \%14C\text{- mannitol recovered} - \%125I\text{- mannitol recovered}}{\%14C\text{- mannitol recovered}} \times 100. \]

The net uptake U_net over the whole run was calculated from the single drops and the final “4 min” collection samples, as follows:

\[ U_{net}\% = \frac{\Sigma 14C\text{- mannitol recovered} - \Sigma 125I\text{- T4 recovered}}{\Sigma 14C\text{- mannitol recovered}} \times 100, \]

where Σ is the sum of tracer recoveries for the whole run and the final “4-min” sample. Σ is expressed as percentage of the 14C or 125I originally injected.

Steady-state Extraction at the Basolateral Face
This technique measures the extraction of 125I-T4 from the blood into CP over 1–2 h and was previously described (59, 60). The mammalian Ringer solution contained 10 μCi-100 ml−1 125I-T4 (90 pmol·l⁻¹) and 40 μCi-100 ml⁻¹ 14C-mannitol as non-diffusible extracellular marker. The lateral CPs were perfused for 1 h until steady state has been achieved. The samples of arterial perfusate and venous effluent were collected at a regular intervals every 5 min, for a further 40 min, accounting for 8 samples per brain, for a total of 14 sheep. The tracer activities in 100-μl aliquots of arterial and venous samples were determined by liquid scintillation counting after addition of 3.5 ml of Ultima Gold (Packard, UK). The activities of both isotopes 125I and 14C were separated and converted to disintegration per minute (dpm); using internal stored quench curves on β-counter (LKB Rackbeta Spectral 1219, UK). The extractions of both 125I-T4 and 14C-mannitol at the blood side of the CP were calculated separately, using the equation below. The difference between the two extractions was considered as the cellular uptake of 125I-T4. Cellular uptake, also known as extraction (%) is

\[ \text{Extraction(%) = } \frac{F_v A^* - F_A V^*}{F_v A^*} \times 100, \]

where \( F_v = \) arterial flow rate (ml·min⁻¹·g⁻¹); \( F_A = \) venous flow rate (ml·min⁻¹·CPs wet weight); \( A^*, V^* = \) activity of the tracer 125I-T4 and 14C-mannitol in the arterial and venous effluent, respectively (dpm·ml⁻¹).

Statistics
All statistical calculations were performed using Microsoft Excel and GraphPad Prism version 5.0 (GraphPad Inc.). Results are expressed as the mean ± SEM. Statistical comparisons were performed using the Student's t-test in order to determine statistical significance at \( p < 0.05 \). Symbols indicate statistical difference: * \( p < 0.05 \), ** \( p < 0.001 \), *** \( p < 0.0001 \).

ETHICS STATEMENT
The Institutional Animal Care and Use Committee (IACUC) of the Lebanese University approved all experimental
procedures in this study. Surgical procedures were performed under deep anesthesia, and all animal experimental procedures were carried out in accordance with the guidelines of the Agriculture Ministry, which conform to the provisions of the Declaration of Helsinki (as revised in Brazil in 2013) and of the European Communities Council Directive (86/609/EEC).

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**AUTHOR CONTRIBUTIONS**

KZ and NK designed the study and performed experiments. NZ performed statistical analysis. MS, MH, WM, HH, and FK participated in data collection. KZ, FK, and NK analyzed data. KZ and NK wrote the manuscript. All the authors read and approved the final version of the manuscript.
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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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