

Creatine and guanidinoacetate transport at blood-brain and blood-cerebrospinal fluid barriers

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Abstract While it was thought that most of cerebral creatine is of peripheral origin, AGAT and GAMT are well expressed in CNS where brain cells synthesize creatine. While the creatine transporter SLC6A8 is expressed by microcapillary endothelial cells (MCEC) at blood-brain barrier (BBB), it is absent from their surrounding astrocytes. This raised the concept that BBB has a limited permeability for peripheral creatine, and that the brain supplies a part of its creatine by endogenous synthesis. This review brings together the latest data on creatine and guanidinoacetate transport through BBB and blood-CSF barrier (BCSFB) with the clinical evidence of AGAT-, GAMT- and SLC6A8-deficient patients, in order to delineate a clearer view on the roles of BBB and BCSFB in the transport of creatine and guanidinoacetate between periphery and CNS, and on brain synthesis and transport of creatine. It shows that in physiological conditions, creatine is taken up by CNS from periphery through SLC6A8 at BBB, but in limited amounts, and that CNS also needs its own creatine synthesis. No uptake of guanidinoacetate from periphery occurs at BBB except under GAMT deficiency, but a net exit of guanidinoacetate seems to occur from CSF to blood at BCSFB, predominantly through the taurine transporter TauT.

Introduction

In the brain as in most tissues, the creatine (Cr) / phosphocreatine (PCr) / creatine kinase (CK) system plays essential roles to maintain the high energy levels necessary for CNS development and functions, through regeneration and buffering of ATP levels (Brosnan and Brosnan 2007; Wyss and Kaddurah-Daouk 2000). Recent works suggest that Cr in CNS may also act as true neurotransmitter and one of the main CNS osmolytes (Almeida et al. 2006; Bothwell et al. 2002). In mammals, half of Cr is obtained from diet, the other half being synthesized endogenously by a two-step mechanism involving arginine:glycine amidinotransferase (AGAT; EC 2.1.4.1; gene: *GATM*) and guanidinoacetate methyltransferase (GAMT; EC 2.1.1.2; gene: *GAMT*). Cr is distributed by blood to tissues and taken up by cells through a specific Cr transporter, SLC6A8 (gene: *SLC6A8*), also called CT1, CRT1, CRTR, CTR or CreaT (Wyss and Kaddurah-Daouk 2000).

The brain is protected from variations in the blood milieu by different barriers which tightly regulate exchanges between periphery and CNS, the two most important being blood-brain barrier (BBB) and blood-cerebrospinal fluid barrier (BCSFB). BBB represents, by many orders of magnitude, the largest barrier between periphery and the brain (Fig. 1). It consists of non-fenestrated microcapillary endothelial cells (MCEC) connected together by tight and adherent junctions that prevent paracellular diffusion of molecules between blood and CNS parenchyma (Engelhardt and Sorokin 2009). Specifically expressed transporters on luminal and abluminal sides of MCEC, together with a very limited pinocytosis, allow the tight regulation of exchanges through BBB. Moreover, these exchanges are also strongly regulated by (i) cells adjacent to the abluminal side of MCEC (namely pericytes which cover about 32% of the microcapillary surface;

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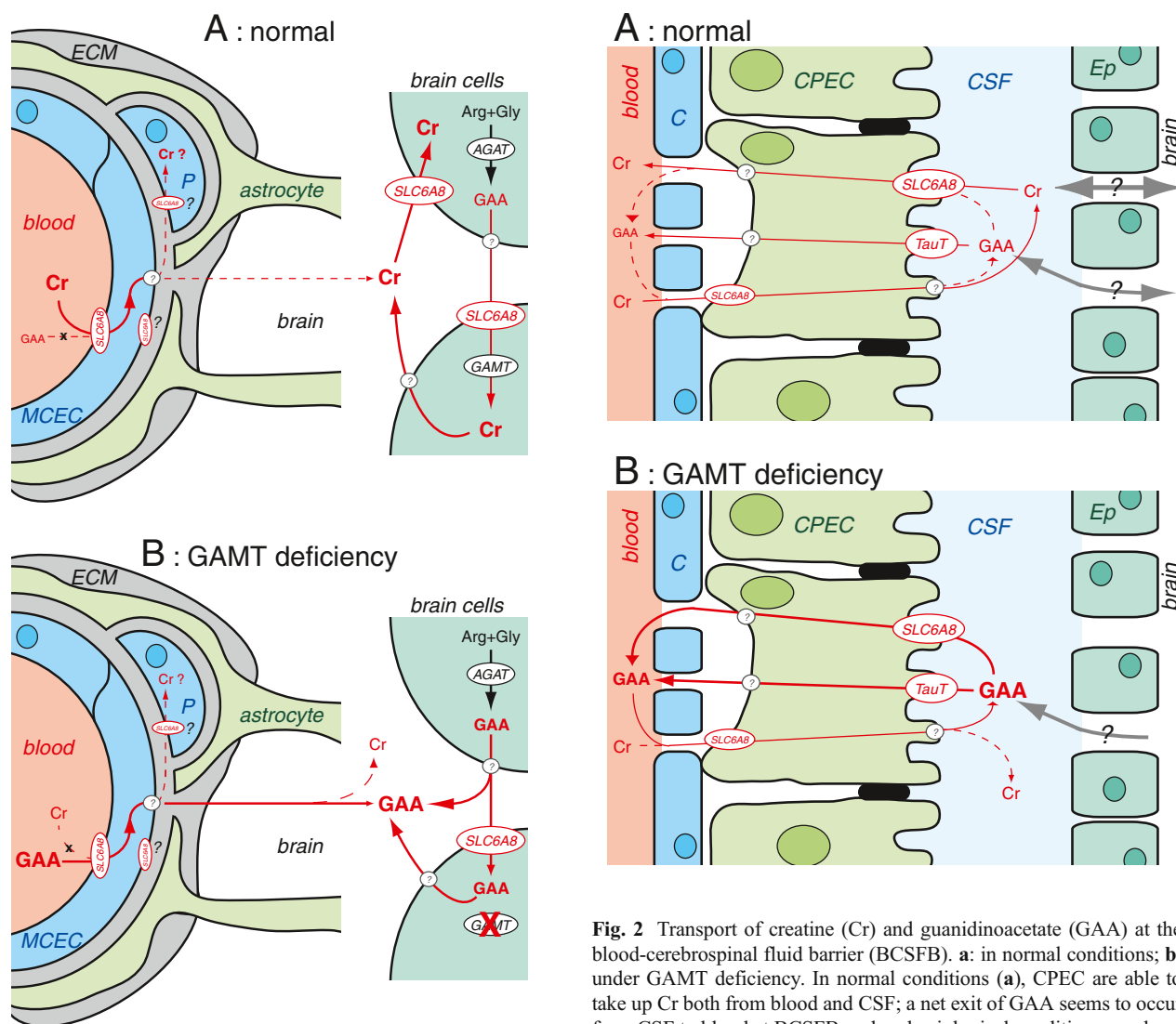


Fig. 1 Transport of creatine (Cr) and guanidinoacetate (GAA) at the blood-brain barrier (BBB). **a:** in normal conditions; **b:** under GAMT deficiency. In normal conditions (**a**), Cr is taken up from periphery through SLC6A8 expressed on MCEC at BBB, but in limited amounts in reason of the absence of SLC6A8 in the surrounding astrocytes and the difficult diffusion of Cr through ECM surrounding BBB. Therefore, the brain needs its own synthetic pathway to ensure sufficient levels of Cr, and do so by expressing AGAT and GAMT. No uptake of GAA from periphery occurs at BBB, except under GAMT deficiency (**b**) and maybe also SLC6A8 deficiency. Arg: arginine; AGAT: arginine:glycine amidinotransferase; Cr: creatine; ECM: extracellular matrix; GAA: guanidinoacetate; GAMT: guanidinoacetate methyltransferase; Gly: glycine; MCEC: microcapillary endothelial cells (BBB); P: pericyte; SLC6A8: Cr transporter

and astrocytic feet which cover more than 98% of the surface made by MCEC and pericytes; Cardoso et al. 2010; Neuwelt et al. 2011) and (ii) a specific extracellular matrix which sheathes MCEC, pericytes and astrocytic feet and limits the diffusion of molecules between MCEC and CNS (Dityatev et al. 2010). BCSFB (Fig. 2), located in lateral, third and fourth

Fig. 2 Transport of creatine (Cr) and guanidinoacetate (GAA) at the blood-cerebrospinal fluid barrier (BCSFB). **a:** in normal conditions; **b:** under GAMT deficiency. In normal conditions (**a**), CPEC are able to take up Cr both from blood and CSF; a net exit of GAA seems to occur from CSF to blood at BCSFB under physiological conditions, predominantly through TauT. This exit of GAA from CSF to blood may be increased under GAMT deficiency (**b**), and uses both TauT and SLC6A8. CPEC: choroid plexus epithelial cells (BCSFB); Cr: creatine; CSF: cerebrospinal fluid; Ep: ependymal epithelium; GAA: guanidinoacetate; SLC6A8: Cr transporter; TauT: taurine transporter

ventricles, is made by unique apical tight junctions between the choroid plexus epithelial cells (CPEC), which are responsible of the secretion of CSF and also express specific transporters allowing regulated exchanges between CSF and blood (Engelhardt and Sorokin 2009; Spector 2010; Wolburg and Paulus 2010).

While it was thought that most, if not all, cerebral Cr is of peripheral origin, AGAT and GAMT are well expressed in CNS where brain cells can synthesize their own Cr (Béard and Braissant 2010; Braissant et al. 2008). In brain parenchyma, SLC6A8 is expressed by neurons and oligodendrocytes. However, while SLC6A8 is also expressed by MCEC at BBB, allowing CNS to import Cr from periphery, it is absent from astrocytes, particularly from their feet lining

BBB (Braissant et al. 2001b; Ohtsuki et al. 2002; Tachikawa et al. 2004). This raised the concept that BBB has a limited permeability for peripheral Cr, and that the brain supplies a part of its Cr by endogenous synthesis (Braissant et al. 2001b; Braissant et al. 2011; Braissant and Henry 2008). However, despite indeed a limited efficiency of brain Cr uptake from periphery, shown *in vivo* both in rodents (Ohtsuki et al. 2002; Perasso et al. 2003) as well as in AGAT- and GAMT-deficient patients (AGAT: OMIM #612718; GAMT: OMIM #612736) treated by Cr (Schulze 2005), Cr (and the intermediate guanidinoacetate, GAA) do cross BBB and BCSFB (Tachikawa and Hosoya 2011).

This review brings together (i) the latest experimental data on the transport of Cr and GAA through BBB and BCSFB with (ii) the clinical evidence shown in the patients deficient for AGAT, GAMT and SLC6A8 (SLC6A8: OMIM #300352), in order to delineate a clearer view of the roles of BBB and BCSFB in the transport of Cr and GAA between periphery and CNS, and on the synthesis and transport of Cr within the brain, both in the mature brain and during development.

Cr and GAA through BBB and BCSFB, and within CNS

It is thought that most of peripheral synthesis of Cr occurs through AGAT in kidney and GAMT in liver (Brosnan and Brosnan 2007), implying the transport of the intermediate GAA through blood circulation. Cr is also synthesized in the mammalian brain (Béard and Braissant 2010; Pisano et al. 1963). The recent knowledge of how AGAT and GAMT, as well as the Cr transporter SLC6A8, are expressed and active at BBB and BCSFB, as well as within CNS, sheds light on the respective importance of Cr synthesis within the brain *versus* Cr and GAA uptake from periphery or release from CNS.

AGAT and GAMT are well expressed in the brain, where they are found, in all CNS structures, in every main cell types (neurons, astrocytes and oligodendrocytes; Braissant et al. 2001b; Nakashima et al. 2005; Schmidt et al. 2004; Tachikawa et al. 2004). We have shown recently that AGAT and GAMT, in most regions of the rat brain, appear expressed in a dissociated way, being rarely co-expressed within the same cell (Braissant et al. 2010). AGAT, but not GAMT, is expressed by MCEC at BBB (Braissant et al. 2001b; Tachikawa et al. 2004), while AGAT and GAMT are expressed by choroid plexus at BCSFB (Braissant et al. 2001b).

SLC6A8 is also expressed throughout the mammalian brain in neurons and oligodendrocytes (Braissant et al. 2001b; Happe and Murrin 1995; Mak et al. 2009; Schloss et al. 1994; Tachikawa et al. 2008). However, in physiological conditions and in contrast to AGAT and GAMT, it cannot be

detected in astrocytes (Acosta et al. 2005; Braissant et al. 2001b; Nakashima et al. 2004), except for very few in cerebellum (Mak et al. 2009). In contrast to its absence from the astrocytic feet lining microcapillaries, SLC6A8 is present in MCEC making BBB, where it is expressed at both luminal and abluminal sides of MCEC (Braissant et al. 2001b; Ohtsuki et al. 2002; Tachikawa et al. 2004) (Fig. 1). At BCSFB, SLC6A8 is mainly expressed at the brush-border membrane of choroid plexus epithelial cells, and to a lesser extent on their basal membrane (Braissant et al. 2001b; Tachikawa et al. 2008) (Fig. 2).

Peripheral Cr can enter the brain. *In vivo* experiments in mouse and rat have demonstrated that CNS can take up Cr from the blood against its concentration gradient, at least to some extent (Ohtsuki et al. 2002; Perasso et al. 2003). However, it has been known for a very long time that this passage of peripheral Cr into the brain is limited. Oral treatment of normal rats with Cr does not increase their CNS Cr content (Chanutin 1927), while oral administration of Cr in healthy human volunteers with about 60 times the daily needs in Cr only leads to a very modest 5% to 10% increase in their brain total Cr (Cr+PCr) after one month of treatment (Dechent et al. 1999). The restricted crossing of peripheral Cr into CNS is also illustrated in Cr orally-treated AGAT- and GAMT-deficient patients who, despite the high doses of Cr used (usually 0.35–2.00 g/kg/day; 20–120 times the normal needs in Cr), replenish their cerebral Cr only slowly, Cr restoration taking months and remaining in most cases only partial (Battini et al. 2002; Ganesan et al. 1997; Item et al. 2001; Schulze 2005; Schulze et al. 1998; Stöckler et al. 1996). GAMT^{-/-} KO mice treated with the high dose of 2 g/kg/day of Cr normalize their brain Cr after 16 to 35 days of treatment, depending on CNS region (Kan et al. 2007).

At BBB (Fig. 1), *in vivo* analysis in mouse has shown that the carrier-mediated uptake of Cr from blood to brain through BBB has an influx transport clearance of 1.6 $\mu\text{L}/(\text{min} \cdot \text{g brain})$ (Ohtsuki et al. 2002), about six times greater than that of sucrose, for which BBB is not permeable, but about 50 times smaller than that of glucose (Hasselbalch et al. 1995). Using TM-BBB4 cells (conditionally immortalized mouse brain capillary endothelial cells) as *in vitro* model for BBB, Ohtsuki et al. showed that Cr is taken up by TM-BBB4 cells by a Na^+/Cl^- -dependent carrier having a K_m of 16 μM and being strongly inhibited by guanidino-propionate and GAA, presenting thus characteristics of SLC6A8 which is expressed at BBB in MCEC (Braissant et al. 2001b; Ohtsuki et al. 2002). SLC6A8 can also mediate the transport of GAA, however with a K_m value ten times greater than for Cr (Tachikawa and Hosoya 2011). Therefore, as GAA levels in blood (1–3.5 μM) are 2–50 times lower than levels of Cr (6–50 μM), it is unlikely that in physiological conditions GAA is taken up from blood by MCEC (Tachikawa et al. 2009). No elimination of GAA

from brain parenchyma through BBB could be observed (Tachikawa et al. 2009).

At BCSFB (Fig. 2), *in vivo* analysis in mouse has also revealed a carrier-mediated uptake of Cr from blood to CSF through CPEC, with an influx transport clearance of 3.6 $\mu\text{L}/(\text{min}\cdot\text{ml CSF})$ (Tachikawa et al. 2008). Tachikawa et al. also showed that GAA can be taken up by TR-CSFB cells (immortalized rat choroid plexus cells) and isolated rat choroid plexus used as *in vitro* model for BCSFB. This uptake of GAA by BCSFB cells appears to be mediated through at least three different mechanisms, two main Na^+/Cl^- -dependent carrier activities having the characteristics of SLC6A8 and of the taurine transporter TauT, and a much smaller transmembrane diffusion (Tachikawa et al. 2008). There appears to be a net efflux of GAA through BCSFB, the elimination clearance of CSF to blood through BCSFB (3.97 $\mu\text{L}/\text{min}$ per rat) being about 33 times higher than the blood-to-CSF influx clearance of GAA through BCSFB (0.12 $\mu\text{L}/\text{min}$ per rat) (Tachikawa and Hosoya 2011). Efflux of GAA from CSF to blood may thus occur through both SLC6A8 and TauT. However, the much higher concentration of Cr (17–90 μM) than GAA (0.015–0.114 μM) in CSF most probably considerably limits the efflux of GAA from CSF through SLC6A8 (Tachikawa et al. 2008).

Cr and GAA at BBB and BCSFB : What can be learned from Cr deficiency syndromes?

CNS is the main organ affected in patients suffering from the Cr deficiency syndromes, inborn errors of Cr biosynthesis and transport caused by mutations in the *GATM*, *GAMT* or *SLC6A8* genes (Item et al. 2001; Salomons et al. 2001; Stöckler et al. 1994). Cr-deficient patients present neurological symptoms in infancy (Battini et al. 2002; DeGrauw et al. 2002; Schulze et al. 1997). In particular, intellectual disability and delays in speech acquisition can be observed (AGAT, GAMT and SLC6A8 deficiencies), as well as intractable epilepsy (GAMT and SLC6A8 deficiencies), autism, automutilating behavior, extrapyramidal syndrome and hypotonia (GAMT deficiency) (Stöckler et al. 2007). With the exception of SLC6A8-deficient heterozygous females where brain Cr deficiency is often partial (Cecil et al. 2003; Valayannopoulos et al. 2011; van de Kamp et al. 2011a), the common phenotype of these three primary Cr deficiencies is the virtual absence of the Cr peak measured by magnetic resonance spectroscopy (MRS) in cortex and basal ganglia (Stöckler et al. 2007).

AGAT- and GAMT-deficient patients can be treated by oral supplementation of Cr. While this strongly improves their neurological status and CNS development, high doses of Cr must be used (usually 0.35–2.0 g/kg/day; 20–120 times the normal needs in Cr) while replenishment of cerebral Cr takes months and only results in partial restoration of

cerebral Cr pools (Battini et al. 2002; Ganesan et al. 1997; Item et al. 2001; Schulze et al. 1998; Stöckler et al. 1996). One exception to these high doses of Cr used to treat Cr-deficient patients is a presymptomatic AGAT patient who received 0.1 g/kg/day of Cr since the age of 4 months (Battini et al. 2006). This interesting case is described in more detail in the following chapter on Cr in immature CNS. A very high dose of creatine (2.0 g/kg/day) was also used in *GAMT*^{-/-} KO mice to analyze their CNS replenishment in Cr (Kan et al. 2007). All these data suggest that BBB has a limited permeability for Cr. Despite improvement of clinical outcome by Cr supplementation, most AGAT- and GAMT-deficient patients remain with CNS developmental problems. Oral supplementation of Cr is inefficient in replenishing brain Cr in SLC6A8-deficient patients, who remain with intellectual disability, severe speech impairment and progressive brain atrophy (Bizzi et al. 2002; Cecil et al. 2001; DeGrauw et al. 2002; Póo-Argüelles et al. 2006). This suggests that Cr cannot enter CNS from periphery by other means than SLC6A8. Attempts to treat SLC6A8-deficient patients with arginine and glycine as precursors of Cr gave some modest encouraging results in two SLC6A8-deficient patients (Chilosi et al. 2008; Wilcken et al. 2008), while it failed to improve the neurological status of many others (Fons et al. 2008; Valayannopoulos et al. 2011; van de Kamp et al. 2011b). BBB and its surrounding astrocytes efficiently express cationic amino acid transporters (CATs 1 and 2B) allowing the import of arginine from periphery to the blood (Braissant et al. 2001a; Braissant et al. 1999). Altogether, data on the treatment of Cr-deficient patients indicate that Cr is taken up in limited amounts by CNS from periphery, and that this modest transport of Cr through BBB most probably exclusively occurs through SLC6A8 (Béard and Braissant 2010; Braissant and Henry 2008) (Fig. 1).

Despite the almost complete lack of Cr observed by MRS, Cr remains present within the brain of Cr-deficient patients (Braissant and Henry 2008). In SLC6A8 deficiency, Cr CSF levels do not differ from age-matched controls (Cecil et al. 2001; DeGrauw et al. 2002; Salomons et al. 2001), while total Cr estimated by MRS has been measured to 1.6 mM in a male patient (21% to 35% of controls; normal range 5.5–7.5 mM; Dezortova et al. 2008). In AGAT deficiency, total Cr levels in cortical gray matter are decreased to 12% of age-matched controls (Battini et al. 2002). In GAMT deficiency, CSF Cr levels are strongly decreased (<2 μM ; normal range: 17–90 μM) (Ensenauer et al. 2004; Schulze et al. 1997; Schulze et al. 2003), while in cortical gray matter total Cr was measured in the 0.2–1.5 mM range (normal range: 5.5–7.5 mM) (Mancini et al. 2005; Stöckler et al. 1994; Dezortova et al. 2008). As the entry of Cr from periphery to CNS appears to exclusively occur through SLC6A8 at BBB (Tachikawa et al. 2009), the presence of normal levels of Cr in CSF of SLC6A8-deficient patients and Cr levels in brain parenchyma between 21%

and 29% of controls must be due to an active CNS endogenous synthesis pathway for Cr (Dezortova et al. 2008; Béard and Braissant 2010; Braissant et al. 2010). As this endogenous brain Cr synthesis pathway is invalidated in AGAT and GAMT deficiencies, the low levels of Cr found in the brain of their respective patients must enter the brain at BBB, through SLC6A8 expressed in MCEC, with the limited efficacy described above (Braissant and Henry 2008; Tachikawa and Hosoya 2011).

GAA accumulation in body fluids is characteristic of GAMT deficiency, where GAA toxicity is responsible for its more complex and specific phenotype. GAA CSF levels in GAMT-deficient patients are 60–1000× higher than in age-matched controls (6.6–15.3 μM versus 0.015–0.114 μM), while GAA was estimated at 3.6 mM within cortical gray matter (normal range: 1.6 mM). GAA can also accumulate in the brain of SLC6A8-deficient patients (Sijens et al. 2005). It was shown that both BBB endothelial cells (Tachikawa et al. 2009) and CNS parenchymal cells (Braissant et al. 2010; Tachikawa et al. 2008) are able to take up GAA by SLC6A8. As the K_m value of SLC6A8 for GAA (269–412 μM) appears ten times higher than that for Cr (29–46 μM) (Tachikawa et al. 2008; Tachikawa and Hosoya 2011), the entry of GAA into CNS in normal conditions must be inhibited by blood Cr levels (Fig. 1a). Under GAMT deficiency however (Fig. 1b), this entry might be facilitated, blood GAA levels becoming higher than Cr levels (Almeida et al. 2004), therefore contributing to GAA accumulation into the GAMT-deficient brain. Endogenous brain AGAT activity must also contribute to GAA accumulation in the CNS of GAMT-deficient patients (Braissant and Henry 2008). Similarly, GAA accumulating in the brain of some SLC6A8-deficient patients (Sijens et al. 2005) is most probably essentially due to brain AGAT activity and the impossibility for GAA to enter GAMT-expressing cells as functional SLC6A8 is lacking (Béard and Braissant 2010; Braissant et al. 2010), and to the lack of SLC6A8-mediated GAA efflux at BCSFB (Tachikawa and Hosoya 2011). Normal levels of GAA in the plasma of SLC6A8-deficient patients most probably makes negligible the contribution of GAA passive diffusion at BBB (Tachikawa et al. 2009) for CNS GAA accumulation in SLC6A8-deficiency. SLC6A8 as well as TauT expressed in the brush-border membrane of choroid plexus at BCSFB may contribute, in GAMT and maybe SLC6A8 deficiencies, to some elimination of GAA from CSF (Tachikawa et al. 2008; Tachikawa et al. 2009; Tachikawa and Hosoya 2011) (Fig. 2b).

Models of transport of Cr and GAA at BBB and BCSFB

Taken together, experimental data on (i) SLC6A8 expression, (ii) functional analysis of Cr and GAA transport and (iii) activity and characterization of SLC6A8 and TauT at

BBB and BCSFB, as well as (iv) the brain characteristics of Cr-deficient patients and the way they respond to Cr treatment, suggest the following models for Cr and GAA transport through BBB and BCSFB (Figs. 1 and 2).

At BBB, SLC6A8 is expressed on both luminal and abluminal sides of MCEC (Fig. 1). In physiological conditions (Fig. 1a), a net entry of Cr from periphery to CNS occurs, most probably through SLC6A8 only. This uptake of peripheral Cr by the brain is limited, due to the absence of SLC6A8 in the astrocytic feet covering most of BBB and the weak diffusion of Cr through the surrounding ECM. CNS can ensure its own Cr synthesis thanks to endogenous expression of AGAT and GAMT, which appear dissociated in most regions of the brain, thus necessitating the transfer of GAA from AGAT- to GAMT-expressing cells through SLC6A8 for completing Cr synthetic pathway. No transfer of GAA occurs through BBB in normal conditions, due to the favorable competition of Cr over GAA for SLC6A8. Under GAMT deficiency however, the competition for SLC6A8 is reversed in favor of GAA, which may cross BBB from periphery and contributes, together with the endogenous brain AGAT activity, to GAA accumulation in the brain of GAMT-deficient patients (Fig. 1b). So far, nothing is known on the use of Cr by pericytes. Equally, the role of SLC6A8 on the abluminal side of MCEC, as well as the way Cr and GAA can leave the cells, are not understood (Fig. 1a).

At BCSFB, SLC6A8 is predominantly expressed on the brush-border apical side of CPEC, as well as weaker on their basal side (Fig. 2). In physiological conditions (Fig. 2a), it is not clear whether a net entry of Cr from periphery, or exit of Cr from CSF, occur at BCSFB. There appears to be a net exit of GAA from CSF to blood, predominantly through TauT. This net exit of GAA from CSF to blood through BCSFB may be stronger under GAMT deficiency (Fig. 2b), and may occur both through TauT and SLC6A8 (favorable competition of GAA over Cr). The exchanges of Cr and GAA between CSF and brain parenchyma, and particularly its extracellular fluid, are not clearly understood so far.

Brain Cr : Different roles for BCSFB and BBB between immature and mature CNS

As proposed above, adult (or mature) CNS might privilege Cr endogenous synthesis versus uptake from periphery. The fetal and perinatal (or immature) brain probably behaves differently. In mammals, the fetal needs in Cr are partly supported by transport of Cr from mother to fetus (Davis et al. 1978; Ireland et al. 2008; Ireland et al. 2011). On the other side, AGAT, GAMT and SLC6A8 are well expressed during embryogenesis, including in CNS (Braissant et al. 2005; Ireland et al. 2009; Wang et al. 2007). However, the

low level of GAMT in early stages suggests that, in contrast to adult brain, fetal and perinatal CNS may predominantly depend on Cr supply from periphery (Braissant et al. 2007). BCSFB develops much earlier than BBB, and is thus the first structure responsible for exchanges between periphery and CNS during development (Fig. 3). The Cr supply to CNS from periphery is probably made possible by the much higher expression of SLC6A8 in CPEC (BCSFB) during development (Braissant et al. 2005) than in the mature brain (Braissant et al. 2001b; Tachikawa et al. 2008), as well as by the very high expression of SLC6A8 in fetal ependymal cells and periventricular zone, as shown in rat (Braissant et al. 2005) and human embryos (O.Braissant and H.R. Widmer, unpublished results) (Fig. 3). In contrast at BBB, SLC6A8 is not detectable in the first stages of MCEC development (Braissant et al. 2005), while it is expressed in mature MCEC (Braissant et al. 2001b; Ohtsuki et al. 2002) (Fig. 3).

The better capacity of immature CNS to take up Cr from periphery is illustrated in two recent studies that showed that the pre-symptomatic treatment of AGAT and GAMT deficiencies appears to completely prevent the phenotypic expression of these diseases (Schulze and Battini 2007). An AGAT-deficient boy, brother of two already affected AGAT-deficient sisters, was diagnosed at birth with the same homozygous mutation as his sisters, and treated orally since

the age of 4 months with Cr monohydrate (0.1 g/kg/day; Battini et al. 2006). Similarly, a GAMT-deficient girl, sister of an already affected GAMT-deficient brother, was diagnosed at birth with the same mutations as her brother, and treated orally since the age of 22 days with Cr monohydrate (0.4 g/kg/day; Schulze et al. 2006). Both patients, over a follow-up of more than 5 years, did not develop the characteristic CNS phenotypic expression of AGAT and GAMT deficiencies (Schulze and Battini 2007). These two cases suggest that Cr plays an essential role in the development of CNS higher cognitive functions, like speech acquisition, during the first months and years of life, and that treatment with Cr before irreversible damage occurs may prevent clinical symptoms of AGAT and GAMT deficiencies permanently. As described above, the pre-symptomatic treatment with Cr in post-natal stages and during the first years of life may also facilitate the entry of Cr into the brain, at stages where BBB is not as tightly regulated as in more mature stages, and where SLC6A8 expression on BBB and BCSFB may still largely facilitate entry of peripheral Cr into the brain (Braissant et al. 2005; Ireland et al. 2009), in contrast to adulthood (Braissant et al. 2001b) (Fig. 3). This facilitated entry of Cr into CNS during the first months and years of life may explain the success of the lower dose of Cr used (0.1 g/kg/day) to treat the above-described pre-symptomatic AGAT-deficient patient (Battini et al. 2006).

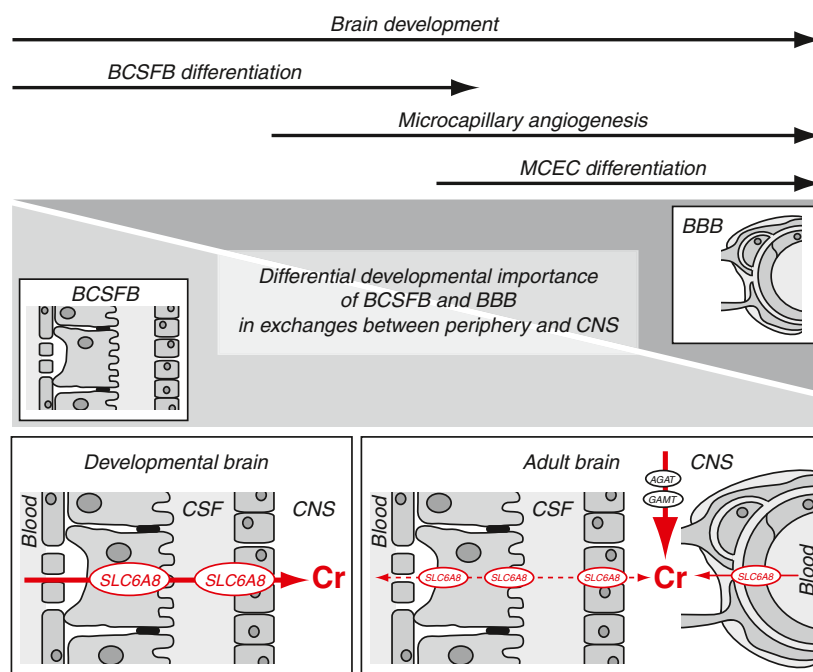


Fig. 3 Transport of creatine (Cr) at blood-cerebrospinal fluid barrier (BCSFB) and blood-brain barrier (BBB) in the developmental *versus* the adult brain. Fetal and perinatal CNS may predominantly depend on Cr supply from periphery. This is probably made possible during development by the high expression of SLC6A8 in CPEC at BCSFB as well as in fetal ependymal cells and periventricular zone. In contrast,

Cr is supplied to the mature brain by its limited uptake from periphery, predominantly at BBB, as well as by endogenous synthesis. AGAT: arginine:glycine amidinotransferase; CNS: central nervous system; Cr: creatine; CSF: cerebrospinal fluid; GAMT: guanidinoacetate methyltransferase; MCEC: microcapillary endothelial cells (BBB); SLC6A8: Cr transporter

Change of BBB properties for Cr in brain pathology?

Numerous pathological states of the brain lead to changes or disturbances of BBB, that can range from adaptation of transporter expression and activity to a frank rupture of BBB (Engelhardt and Sorokin 2009; Neuwelt et al. 2011). This is also true for astrocytes, which are known to become reactive in many CNS pathological conditions (Zhang and Barres 2010). In recent years, we have shown that hyperammonemic conditions lead to a secondary Cr deficiency in brain cells (Bachmann et al. 2004; Braissant et al. 2008; Braissant 2010b), and that Cr co-treatment exerts neuroprotective effects against NH_4^+ toxicity (Braissant et al. 2002; Braissant 2010a; Cagnon and Braissant 2007). While SLC6A8 appears absent from astrocytes in normal, physiological conditions, we could show interestingly that NH_4^+ exposure induces SLC6A8 expression in these same astrocytes (Braissant et al. 2008). Moreover, NH_4^+ exposure also increases both SLC6A8 expression and Cr uptake in MCEC (Bélanger et al. 2007). Thus, BBB in the hyperammonemic CNS might be more permeable to Cr than under physiological conditions, and supplying oral Cr to hyperammonemic neonates or infants might likely contribute to protect their brain development (Braissant 2010a; Braissant 2010b). This has to be put in perspective of the treatment of other brain pathologies with Cr. Indeed, Cr administration can exert protective effects in various neurodegenerative processes, including Huntington's and Parkinson's diseases (Béard and Braissant 2010; Bender et al. 2006; Hersch et al. 2006; Klein and Ferrante 2007). Creatine was also demonstrated to exert neuroprotective effects, both in vitro and in vivo, against anoxic and ischemic damage (Balestrino et al. 2002; Adcock et al. 2002; Lensman et al. 2006) as well as against traumatic brain injury (Sullivan et al. 2000) and spinal cord injury (Hausmann et al. 2002).

Creatine derivatives for a better crossing of BBB?

To counteract the limited permeability of BBB for Cr, and in particular to allow a treatment for SLC6A8-deficient patients, a lot of work has been performed in order to develop more lipophilic derivatives of Cr, that would not need SLC6A8 to be taken up by cells, and that in consequence would cross BBB more easily, possibly by simple trans-membrane diffusion. While some derivatives (e.g., N-amidino-piperidine) were harmful, others (e.g., Cr-Mg-complex acetate, PCr-Mg-complex acetate and Cr ethyl ester) showed interesting neuroprotective effects in hippocampal organotypic cultures (a model with disruption of BBB) (Perasso et al. 2008; Adriano et al. 2011). PCr-Mg-complex acetate and another derivative, Cr benzyl ester, were also demonstrated to cross cell membranes without

the help of the Cr transporter (Lunardi et al. 2006). More interestingly, PCr-Mg-complex acetate and the very recently described creatinyl amino acids showed neuroprotective effects in vivo in rodents, thus suggesting a crossing of BBB (Burov et al. 2011; Perasso et al. 2009). However so far, only the Cr derivative Cr ethyl ester has been used to treat SLC6A8-deficient patients (Fons et al. 2010), but this compound failed to replenish their brain Cr concentration, as well as to improve their neurological status. This is probably due to the non-enzymatic conversion of Cr ethyl ester directly to creatinine, before it reaches CNS (Giese and Lecher 2009). For the treatment of SLC6A8-deficient patients, no data are available so far for the use of other Cr derivatives, in particular those that seem to cross BBB (PCr-Mg-complex acetate and creatinyl amino acids).

Models and conclusions

Taken together, (i) the expression patterns of SLC6A8, AGAT and GAMT at BBB, BCSFB and in the brain, (ii) the functional experiments on Cr and GAA trafficking through BBB and BCSFB both in vivo and in in vitro models, and (iii) the clinical characteristics of Cr-deficient patients, lead to the following models for Cr and GAA transport between periphery and CNS, and for Cr synthesis in CNS (Figs. 1, 2 and 3) (Béard and Braissant 2010; Braissant et al. 2011; Braissant and Henry 2008):

In normal conditions, Cr is taken up by CNS from periphery through SLC6A8 expressed on MCEC at BBB, but in limited amounts because of the absence of SLC6A8 in the surrounding astrocytes and the difficult diffusion of Cr through ECM surrounding BBB. In consequence, the brain needs its own synthesis to ensure sufficient levels of Cr, and does so by expressing AGAT and GAMT. No uptake of GAA from periphery occurs at BBB, except under GAMT deficiency (and maybe also SLC6A8 deficiency). A net exit of GAA seems to occur from CSF to blood at BCSFB under physiological conditions, predominantly through TauT. This exit of GAA from CSF to blood may be increased under GAMT deficiency, and uses both TauT and SLC6A8.

A lot of work is still needed to better understand the transport of Cr, GAA and potentially active Cr derivatives at BBB and BCSFB, and to allow better treatment for Cr-deficient patients, in particular those with SLC6A8 deficiency. In particular, a better knowledge of how Cr and GAA can leave the cells, which so far is very poorly understood (Figs. 1 and 2), may contribute to more focused and rationale treatments.

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