Chronic kidney disease and neurological disorders: are uraemic toxins the missing piece of the puzzle?

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GRAPHICAL ABSTRACT

Narrative review

Chronic kidney disease and neurological disorders: are uraemic toxins the missing piece of the puzzle?

Background

Along with traditional cardiovascular risk factors (such as diabetes, inflammation, hypertension and dyslipidaemia), non-traditional risk factors related to kidney damage (such as uraemic toxins) may predispose patients with CKD to neurological disorders.

Aim

Potential clinical impact of CKD on cerebrovascular and neurological complications

Mechanisms underlying the uraemic toxins’ putative direct actions (based on preclinical and clinical research)

Potential impact of these findings on patient care, and unmet medical needs

Results

Uraemic toxins that might influence brain function:

Small water-soluble compounds:

Asymmetric dimethylarginine (ADMA)

Symmetric dimethylarginine (SDMA)

Dimethylamino-N-oxide (DMAO)

Triethylamine-N-oxide (TEA)

Uric acid, Urea, Methyloxyindole guanidine

Protein-bound compounds:

Indoles (Indolealanine (IA), Indoxyl glucuronide, Indoxyl acid (IAA), Kynurenine)

Creatine (p-cresyl creatine (pCC), p-cresyl glucuronide)

Hippurate (Hippuric acid (HA), 3-Carboxy-4-methyl-5-propyl-2-ureidopropionate (CMP))

Middle molecules:

82 microglobulin (82M), Inteleukin-6, Parathoriod hormone (PTH)

Conclusion

The uraemic toxins that accumulate in the blood in ESRD might explain the association between CKD and cerebrovascular/neurologic complications (at least in part in more advanced CKD stages) and constitute potentially valuable therapeutic targets.

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ABSTRACT

Chronic kidney disease (CKD) perturbs the crosstalk with others organs, with the interaction between the kidneys and the heart having been studied most intensively. However, a growing body of data indicates that there is an association between kidney dysfunction and disorders of the central nervous system. In epidemiological studies, CKD is associated with a high prevalence of neurological complications, such as cerebrovascular disorders, movement disorders, cognitive impairment and depression. Along with traditional cardiovascular risk factors (such as diabetes, inflammation, hypertension and dyslipidaemia), non-traditional risk factors related to kidney damage (such as uraemic toxins) may predispose patients with CKD to neurological disorders. There is increasing evidence to show that uraemic toxins, for example indoxyl sulphate, have a neurotoxic effect. A better understanding of factors responsible for the elevated prevalence of neurological disorders among patients with CKD might facilitate the development of novel treatments. Here, we review (i) the potential clinical impact of CKD on cerebrovascular and neurological complications, (ii) the mechanisms underlying the uraemic toxins’ putative direct action (based on pre-clinical and clinical research) and (iii) the potential impact of these findings on patient care.

Keywords: cardiovascular, CKD, indoxyl sulphate, stroke, uraemic toxins

INTRODUCTION

In 2017, the estimated worldwide prevalence of chronic kidney disease (CKD) was 9%, which corresponds to almost 850 million individuals [1]. Kidney disease has a major effect on overall health, both as a direct cause of morbidity and mortality, and as an important risk factor for cardiovascular disease [1]. CKD perturbs the crosstalk with other organs, with the interaction between the kidneys and the heart having been studied most intensively. However, a growing body of data indicates that diseases of the kidney are associated with diseases of the central nervous system. Mild CKD is highly prevalent in the general population and is already known to be associated with cognitive dysfunction [2]. Indeed, the epidemiologic data suggest that individuals at all stages of CKD have a higher risk of developing cognitive disorders and dementia. Furthermore, patients with CKD are more likely to present with clinically evident stroke and subclinical cerebrovascular disease (e.g. white matter hyperintensities), relative to the general population [3]. Given the high prevalence of kidney disease in the general population, the identification of mechanisms associated with cerebrovascular and neurological complications is critically important in the CKD population.

Along with traditional cardiovascular risk factors (such as diabetes mellitus, inflammation, hypertension and dyslipidaemia), non-traditional risk factors (such as uraemic toxins) related to kidney damage may predispose patients with CKD to neurological disorders (Figure 1). According to the European Uraemic Toxins Work Group (EUTox), uraemic toxins are harmful compounds that accumulate in the body during periods of renal function decline [4]. They are classified into three groups on the basis of their protein-binding properties and molecular weight: protein-bound solutes, middle molecules and small water-soluble solutes [4] (Table 1). Protein-bound toxins and certain large middle molecules are cleared poorly by dialysis. Uraemic toxins may have direct neurotoxic actions (such as astrocyte activation and neuronal death) and/or indirect actions via vascular effects (such as cerebral endothelial dysfunction, calcification and inflammation) [5]. Indeed, various uraemic toxins such as indoxyl sulphate (IS) have been implicated in the pathogenesis of neurological disorders in patients with CKD [6–8].

Here, we review (i) the potential clinical impact of CKD on cerebrovascular and neurological complications, (ii) the mechanisms underlying the uraemic toxins’ putative direct actions (based on pre-clinical and clinical research), (iii) the potential impact of these findings on patient care and (iv) unmet medical needs.

THE CLINICAL IMPACT OF CKD ON CENTRAL NERVOUS SYSTEM COMPLICATIONS

Cerebrovascular disease

A growing body of epidemiologic data suggests that impaired renal function is strongly associated with an increased risk of cerebrovascular disease. Indeed, atrial fibrillation and its major complication—ischaemic stroke—are very common in CKD patients [9]. Individuals in the general population with a slight decrease in their estimated glomerular filtration rate (eGFR) (≤60 mL/min/1.73 m²) have a greater risk of fatal or non-fatal ischaemic stroke, and a low eGFR (≤30 mL/min/1.73 m²) increases the risk of haemorrhagic stroke [3, 10]. In a meta-analysis of 63 cohort studies (encompassing 2 085 225 participants), 20 randomized controlled trials (encompassing 168 516 participants) and 30 392 reported strokes, the stroke risk increased linearly and additively with declining eGFR and increasing albuminuria. The risk of stroke increased by 7% [risk ratio (RR) (95% confidence interval, CI): 1.07 (1.04–1.09)] for every 10 mL/min/1.73 m² decrease in the eGFR. Furthermore, a 25 mg/mmol increment in the albumin–creatinine ratio was associated with a 10% increase in the risk of stroke [RR (95% CI): 1.10 (1.01–1.20)] [11]. Moreover, stroke was the third most common cardiovascular cause of death in patients with CKD [11]. Importantly, CKD and its components (proteinuria and low eGFR) also affect the clinical outcome after ischaemic stroke. Indeed, patients with CKD had significantly higher risks of neurologic deterioration, in-hospital mortality and a poor functional outcome [12]. End-stage renal disease (ESRD) is also associated with other types of cerebrovascular damage, such as cerebral microbleeds and silent cerebral infarcts. In particular, patients on haemodialysis (HD) have an exceptionally high incidence of cerebral microbleeds [13].

Cognitive disorders

Various research results suggest that patients at all CKD stages may have a higher risk of cognitive disorders than
patients without CKD. The eGFR is inversely correlated with overall cognitive function in patients with CKD [14]. The estimated prevalence of cognitive impairment in this population ranges from 30% to 60%, depending on the definition of cognitive impairment that varies between studies [15].

As in the general population, quantifying neurological impairment is particularly difficult to evaluate, indeed the method used to assess cognitive impairment in patients with CKD has a major influence on the outcome. The various screening tests do not have the same degree of accuracy—especially in patients on HD [16]. Neuropsychological assessments must be standardized and comprehensive, since impairments can develop early in CKD and the various skills do not decline at the same rate [17]. However, comprehensive neurological assessments can be long and tiring, and cognitive performance fluctuates in patients on HD over the course of a dialysis session [15,16]. In a study of a small number of patients on HD (n = 28), cognitive performance was worst during the dialysis session itself (probably due to haemodynamic effects) and best shortly before the session or on the day after dialysis [18]. Some data observed an improvement of cognitive performance after an HD session compared with before [19]. Neuropsychological and electrophysiological performance seems to be more stable (and close to normal values) in patients on continuous ambulatory peritoneal dialysis (PD) than in patients on HD [20]. However, Drew et al.’s [21] study of 40 patients did not find a difference in cognitive performance between the hour before the HD session and the first hour of the session, suggesting that cognitive assessment during an HD session is still relevant to screen cognitive disorders in those patients. Executive functions are most frequently impaired in patients with CKD [16]. Sleep disorders and depression frequently affect patients with CKD and are able to interfere with cognitive assessments [22]. The interpretation of cognitive

Table 1. Uraemic toxins that might influence brain function

| Small water-soluble compounds | Protein-bound compounds | Middle molecules |
|-------------------------------|-------------------------|-----------------|
| ADMA                          | Indoles                | B2M             |
| SDMA                          | • IS                   | IL-6            |
| TMAO                          | • Indoxyl glucuronide  | PTH             |
| Uric acid                     | • IAA                  |                 |
| Urea                          | • Kynurenine           |                 |
| Methylguanidine guanidine     | Cresols                |                 |
|                               | • pCS                  |                 |
|                               | • p-cresyl glucuronide |                 |
|                               | Hippurates             |                 |
|                               | • HA                   |                 |
|                               | • CMPF                 |                 |

ADMA, asymmetric dimethylarginine; SDMA, symmetric dimethylarginine; TMAO, trimethylamine N-oxide; CMPF, 3-carboxy-4-methyl-5-propyl-2-furanpropanoic acid.

FIGURE 1: The complicated puzzle of risk factors associated with neurological disorders in patients with CKD. Along with traditional cardiovascular risk factors (such as diabetes, hypertension and dyslipidaemia), non-traditional risk factors related to kidney damage (such as uraemic toxicities) may predispose patients with CKD to neurological disorders.
The use of certain drugs may have an (indirect) adverse effect on cognitive function of patients with CKD. Some drugs frequently used in patients with CKD such as opiates, benzodiazepine, antidepressants and drugs with anticholinergic properties are known to aggravate cognitive disorders or induce delirium [24, 25]. In patients with CKD, poor drug clearance and greater drug penetration through the blood–brain barrier (BBB) might lower the threshold for neurotoxicity and contribute to cognitive dysfunction. Indeed, most patients with CKD take several medications on a daily basis [26–28].

**Movement disorders: Parkinson’s disease**

Movement disorders are among the central nervous system complications in uraemic patients. Asterixis, multifocal myoclonus and restless leg syndrome are well documented in this population and so will not be reviewed here [29, 30].

A few large epidemiologic studies have found an increased risk of Parkinson’s disease risk in patients with CKD [31–34]. All the studies were performed in Asia (in Taiwan and South Korea). In 3,580,435 individuals aged ≥65 years who had received health check-ups from the South Korea National Health Insurance Service, 30,813 developed Parkinson’s disease; renal dysfunction and proteinuria were independent risk factors for the development of Parkinson’s disease, with a hazard ratio (95% CI) of 1.36 (1.31–1.42) for an eGFR of 30–60 mL/min/1.73 m² and 1.74 (1.32–1.63) for an eGFR <30 mL/min/1.73 m² [31]. The progressive association between decreasing of eGFR and increased risk of Parkinson’s disease incidence was confirmed in another study of data from the South Korea National Health Insurance Service [32]. In a Taiwanese study of patients with newly diagnosed ESRD versus controls, the incidence of Parkinson’s disease was 1.55-fold higher in the ESRD cohort than in the control cohort (48.8 versus 31.7/10,000 person-years, respectively), giving an adjusted hazard ratio (95% CI) of 1.73 (1.39–2.15) [33]. Hence, patients with CKD are more prone to develop Parkinsonism, independent of their diabetic status [34]. CKD and Parkinson’s disease share some pathophysiologic mechanisms, such as oxidative stress, hypertension, vitamin D deficiency and anaemia (for a review, see [35]). Nonetheless, the mechanisms that underlie this relationship have not been fully elucidated.

**Anxiety and depression**

Anxiety is associated with poor health outcomes among patients with CKD. A recent meta-analysis reported that the pooled prevalence of anxiety disorders (nine studies, n = 11,071) among patients with CKD was 19%, while that of elevated anxiety symptoms (52 studies, n = 110,739) was 43%, with a higher prevalence in dialysis patients than in pre-dialysis patients [36]. Depression is highly prevalent among patients with CKD and healthcare professionals must be aware of this important issue since it has a major impact on quality of life and is often not recognized. A systematic review and meta-analysis of 55,982 patients with CKD found a prevalence of 26.5% for depressive symptoms and 21.4% for clinically significant depression [37]. Accordingly, the antidepressant prescription rate in patients with CKD is nearly 1.5 times higher than in general population [38]. Anxiety and depression can further reduce a CKD patient’s quality of life, which is already lower than in general population [39].

**Dialysis and transplantation affect central nervous system complications**

Cerebrovascular disease is a common cause of death in patients on long-term dialysis [40]. Indeed, the risk of hospitalization for ischaemic and haemorrhagic stroke is 4- to 10-fold greater in patients on HD than in the general population [11]. It has been suggested that patients on HD have a higher prevalence of silent cerebral infarction (relative to control) [9] and an even higher prevalence of subclinical cerebral lacunar infarctions, microbleeds and loss of white matter integrity (relative to patients with CKD) [3, 41, 42]. In elderly patients, the incidence of stroke rose within a month of starting HD and remained high thereafter, when compared with the period before HD [43]. Although HD is a life-saving renal replacement therapy, a growing body of data shows that the side effects of the HD procedure contribute to cerebrovascular damage. For example, Eldehni et al. [44] showed that patients starting HD treatment had an impressive progression of white matter lesions during the first year of HD. Interestingly, lowering the dialysate temperature was associated with greater intrasession haemodynamic stability and slower progression of white matter lesions [44]. Using intradialytic positron emission tomography [45, 46], near-infrared spectroscopy [46, 47] or ultrasound measurement of the mean flow velocity in the cerebral arteries [48], it has been shown that HD induces an acute reduction in brain perfusion [46–49]. These reductions in brain perfusion were accompanied by an intrasession decline in cognitive functions, including global function, executive function and verbal fluency [49]. Repeated acute reductions in brain perfusion may have harmful long-term consequences. The percentage decline in mean blood flow velocity in the cerebral arteries was significantly correlated with the progression of the white matter hyperintensity burden over a 12-month period [49]. These findings strongly suggest that HD-induced haemodynamic changes have a pathophysiological role in the accelerated progression of brain damage in patients on HD. Accordingly, Polinder-Bos et al. [45] found a significant association between a higher ultrafiltration volume and lower intradialytic cerebral blood flow. However, the researchers did not find a significant association between changes in intradialytic blood pressure course and brain perfusion or brain tissue oxygenation [45, 46]. Likewise, a study of a larger cohort of patients by MacEwen et al. [47] (n = 58) showed that changes in blood pressure were poor predictors of downstream brain ischaemia. It is therefore possible that HD-related factors other than haemodynamic changes are involved, for example an interaction between blood and the extracorporeal circuit, which might lead to HD-induced systemic inflammation and thus endothelial dysfunction. Furthermore, an intradialytic rise in the plasma bicarbonate concentration and blood pH can affect brain perfusion. Indeed, a higher blood pH was associated with a lower cerebral blood flow in most brain regions shortly after the start of HD, relative to the pre-dialysis values [45]. It was also recently
### Table 2. Summary of pre-clinical and clinical studies of direct associations between uraemic toxins and cerebrovascular/neurologic complications

| First author | Model | Uraemic toxin(s) studied | Main findings |
|--------------|-------|--------------------------|---------------|
| **In vitro studies** | | | |
| Oshima et al. [55] | Bulbospinal neurons in the RVLM | Uric acid, IS, Methylguanidine | Uric acid, IS and methylguanidine directly stimulate bulbospinal RVLM neurons via specific transporters, such as OAT1, OAT3, OCT3 and URAT1 |
| Adesso et al. [8] | Glioma cell line (C6) Sera from 18 participants: 4 healthy people, 8 patients with CKD and 6 patients on HD | IS | The sera of patients with CKD induced significant inflammation in astrocyte cells, in proportion to the serum IS concentration. The IS adsorbent AST-120 reduced this inflammatory response. |
| Lin et al. [56] | Human primary astrocytes | IS | IS stimulated the release of reactive oxygen species, increased levels of NRF-2 and reduced the mitochondrial membrane potential. IS also triggered astrocyte apoptosis by inhibiting the mitogen-activated protein kinase pathway |
| Watanabe et al. [57] | Mouse hippocampal neuronal HT-22 cells | Indole, IS, pCS, IAA, Hippurate | Indole, IS, pCS and IAA significantly decreased the viability of HT-22 cells, which was associated with a significant decrease in glutathione levels. |
| **Animal studies** | | | |
| Ohtsuki et al. [58] | Adult male Wistar rats, mature female Xenopus laevis frogs Determination of brain efflux index and OAT3 expression | IS | OAT3 mediates the brain-to-blood transport of IS and is also involved in the efflux of neurotransmitter metabolites and drugs |
| Sato et al. [59] | 4 groups of mice: (i) control (n = 16), (ii) AST-120 (n = 16), (iii) RF (n = 17) and (iv) RF + AST-120 (n = 16); evaluation of uraemic toxin accumulation in different organs, including the brain | IS, pCS | IS and pCS accumulated in the brain of mice with RF. The oral adsorbent AST-120 prevented (to some extent) the tissue accumulation of IS and pCS |
| Mair et al. [60] | CSF and plasma ultrafiltrate were obtained from rats 48 h after a sham operation (control; n = 10) or bilateral nephrectomy (n = 10). The samples were analysed using an established metabolomic protocol | 248 solutes including IS, TMAO, hippurate and urea | The CSF levels of the great majority of uraemic solutes were elevated in rats with RF but typically increased less than in the plasma ultrafiltrate |
| Karbowska et al. [7] | 3 groups of rats: (i) control group (tap water without IS), (ii) experimental group with 100 mg/kg of body weight of IS/day and (iii) experimental group with 200 mg/kg of body weight of IS/day | IS | IS accumulation was greatest in the brainstem IS led to behavioural alterations involving apathetic behaviour, increased stress sensitivity and reduced locomotor and exploratory activity. IS might contribute to the impairment of spatial memory and motor coordination. These results could not be explained completely by changes in cerebral monoamine concentrations and turnovers |
| Bobot et al. [61] | 3 rat models of CKD: (i) an adenine-rich diet, (ii) 5/6 nephrectomy and (iii) AhR-/- knockout mice overloaded with IS in the drinking water. Evaluations of BBB disruption using SPECT/CT, BBB permeability using imaging markers and neurologic impairments using neurobehavioural tests | IS | IS led to BBB disruption. Cognitive impairment in the three models was correlated with serum levels of IS and with BBB disruption. Non-CKD AhR-/- knockout mice were protected against IS-induced BBB disruption and cognitive impairment |
| Sun et al. [62] | 2 groups of mice: (i) kidney-intact controls (n = 20) and (ii) unilaterally nephrectomized (n = 120) mice divided into four groups (n = 15 per group) receiving various doses of pCS (0, 1, 10 and 100 mg/kg/day) intraperitoneally | pCS | Apparent deposition of pCS in the prefrontal cortical tissues was associated with several abnormal behaviours, such as depression, anxiety and cognitive impairment. However, pCS accumulation and behavioural changes were not |

*Continued*
| First author | Model | Uraemic toxin(s) studied | Main findings |
|--------------|-------|--------------------------|---------------|
| For the intervention, two groups of unilaterally nephrectomized mice were treated daily with pCS (100 mg/kg): (i) control (normal saline administration) (n = 16) and (ii) AST-120 (400 mg/kg) via oral gavage | Behavioural evaluation | observed at lower doses of 1 and 10 mg/kg/day in the unilateral nephrectomy group or at doses of 1, 10 and 100 mg/kg/day in the kidney-intact controls. These changes were alleviated by the uraemic toxin adsorbent AST-120 |
| Watanabe et al. [57] | 2 groups of rats: (i) controls (n = 14) and (ii) CKD (n = 15) induced by an adenine-rich diet | No uraemic toxin measurements | The CKD group had larger numbers of pyknotic neuronal cells. The two groups did not differ in spatial learning and memory abilities |

### Clinical studies: observational studies

| Bossola [63] | 80 patients on HD | IL-6 | In a multivariate analysis, there was a direct, inverse correlation between BDI and IL-6 and creatinine. The HARS score was significantly correlated with the PTH levels in a univariate analysis only |
|--------------|------------------|------|------------------|
| Hsu et al. [64] | A cross-sectional study of 209 patients with CKD and a history of depression | IS, pCS, Urea, B2M | Depressive patients had lower IS levels. The levels of urea, B2M and pCS were not significantly associated with depression |
| Yeh et al. [65] | 199 patients with CKD and 84 matched non-CKD participants | IS, pCS | In early-stage CKD, IS was associated with poorer executive function but pCS was not significantly associated with cognitive function |
| Ye et al. [66] | 271 healthy subjects, 596 patients with mild cognitive impairment and 197 patients with Alzheimer’s disease | Uric acid | Higher levels of uric acid were associated with slower cognitive decline |
| Sleeman et al. [67] | 154 patients with newly diagnosed Parkinson’s disease and 99 age-matched controls | Urate, Homocysteine | A lower serum urate concentration was associated with worsening motor function, while a higher homocysteine concentration was associated with cognitive decline and worse motor function |
| Efstathioudou et al. [68] | Mendelian randomization approaches | 28 genetic variants related to the serum uric acid | A Mendelian randomization study did not evidence a clinically relevant causal effect of genetically determined serum urate on a range of cardiovascular and neurovascular outcomes |
| Lin et al. [69] | 260 patients on HD | IS, pCS | Both free IS and free pCS were negatively associated with the CASI and MMSE scores. After controlling for confounders, circulating free IS levels were still negatively associated with MMSE and CASI scores but there was no correlation between free pCS and the total MMSE score or the total CASI score |
**IMPACTS OF URAEMIC TOXINS ON CEREBROVASCULAR DISEASES AND COGNITION**

The progressive loss of kidney function typically observed in CKD is accompanied by the retention of a range of metabolites; this is due to decreased renal clearance and (in some cases) an increase in generation. Many of these solutes (uraemic toxins) have been shown to exert biological activity, hence affecting the functioning of cells and organs, resulting in the uraemic syndrome [4, 54]. As mentioned in the Introduction section, EUTox classifies uraemic toxins as small water-soluble compounds, protein-bound compounds or middle molecules [4, 54].

We systematically searched the literature (the MEDLINE database, up until 31 March 2021) for publications on the relationship between uraemic toxins and neurological complications by combining the following key words (‘uraemic toxins and brain’ + ‘uraemic toxins and stroke’ + ‘uraemic toxins and cognitive’ + ‘uraemic toxins and depression’ + ‘uraemic toxins and anxiety’ + ‘uraemic toxins and neurological’ + ‘uraemic toxins and neurodegeneration’ + ‘uraemic toxins and Parkinson’). A total of 164 articles were analysed; after the exclusion of 56 reviews, 15 redundant articles and 71 irrelevant articles, a total of 22 original articles were selected. The uraemic toxins cited in these publications are listed in Table 1. Most of these studies focused on the direct impact of protein-bound compounds on the brain. Below, we describe the origin of the inculminated uraemic toxins and their putative direct effects on the brain.

**THE ORIGIN OF URAEMIC TOXINS**

The origin of uraemic toxins is presented in the Supplementary data.

**DIRECT EFFECTS OF URAEMIC TOXINS ON THE BRAIN**

Protein-bound uraemic toxins can have direct and/or indirect effects on the brain. Most of the recent literature has focused on direct effects (Table 2).

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**Table 2. Continued**

| First author | Model | Uraemic toxin(s) studied | Main findings |
|--------------|-------|-------------------------|---------------|
| Lin et al. [70] | 230 patients on HD Cognitive functions evaluated with the MMSE and the CASI | IAA, HA | Serum IAA was associated with cognitive impairment, based on the MMSE and CASI scores. There was no correlation between the serum HA level and cognitive status. |
| Li et al. [71] | 222 patients on PD Global cognition (the modified MMSE) and executive function tests | Urea, B2M | Higher middle molecule clearance was independently associated with better performance in general cognition and executive function tests in patients with PD. |
| Linde et al. [72] | 10 kidney transplants recipients and 18 controls (9 patients on HD, and 9 patients with CKD Stages 4 or 5 (eGFR < 30 mL/min/1.73 m²) who were not on dialysis) Extensive neuropsychological assessment | IS, pCS, IAA, TMAO, indoxyl glucuronide, p-creuly glucuronide, phenyl sulphate, kynurenine, tryptophan, kynuraenic acid, tyrosine, phenylalanine and phenylacetylglutamine | The serum concentration of most uraemic toxins decreased significantly within 1 week of kidney transplantation. There were no significant improvements in cognitive function that could be specifically related to kidney transplantation in the first 3 months after the procedure. |
| Sankowski et al. [73] | (i) patients with Parkinson’s disease (n = 18) and (ii) controls (n = 9) Plasma and CSF samples | IS, pCS, TMAO, ADMA, SDMA | In patients with Parkinson’s disease, the CSF: plasma ratios for IS and pCS were higher than in controls. Patients with motor fluctuations had higher CSF levels of pCS (P = 0.0043), IS (P = 0.0361), ADMA (P = 0.0017), SDMA (P = 0.02614) and TMAO (P = 0.0179) than other Parkinson’s patients did. |

WAIS, Wechsler Adult Intelligence Scale; OCT, organic cation transporter; TMAO, trimethylamine N-oxide; ADMA, asymmetric dimethylarginine; CMPF, 3-carboxy-4-methyl-5-propyl-2-furanpropanoic acid; SDMA, symmetric dimethylarginine.
In vitro studies

Some recent data suggest that uraemic toxins can have a direct impact on neurons. For example, Oshima et al. [55] reported that uric acid, IS and methylguanidine directly stimulate neurons in the bulboospinal rostral ventrolateral medulla (RVLM) by increasing the oxidative stress. Watanabe et al. [57] recently examined the effects of uraemic toxins on the mouse hippocampal neuronal cell line HT-22. Indole, IS and indole acetic acid (IAA) significantly decreased the viability of the HT-22 cells, with a parallel decrease in glutathione levels [57]. Similarly, sera from patients with CKD induced significant inflammation in astrocytes; the inflammation was proportional to the serum IS concentration, and treatment with the activated charcoal adsorbent AST-120 reduced this inflammatory response. IS further increased inflammation and oxidative stress in primary central nervous system cells (via nuclear factor-κB and AhR activation) and induced neuron death [6, 8]. In an in vitro study of human astrocytes, IS activated the production of reactive oxygen species and downregulated the production of cell-protective factors like nuclear factor (erythroid-derived 2)-like 2. Moreover, IS triggered astrocyte apoptosis by inhibiting the mitogen-activated protein kinase pathway [56].

Animal studies

Hénaut et al. [74] investigated the cellular and molecular mechanisms by which uraemia worsened the severity of ischaemic stroke after transient middle cerebral artery occlusion (tMCAO) in mice. In the acute phase of stroke recovery, CKD aggravated tMCAO-induced ischaemic brain damage by increasing neuronal loss and apoptosis both in the ischaemic penumbra and the ischaemic core. The increased ischaemic damage observed in CKD mice was associated with impaired post-stroke recovery (relative to non-CKD control mice), as evidenced by greater weight loss, decreased muscle strength, impaired motor coordination and reduced physical resistance to tiredness. An early increase in local inflammation and an impairment in tissue repair in CKD mice were associated with greater ischaemic damage during the acute phase of stroke recovery [74].

It has been suggested that the negative impact of uraemia on cerebrovascular disease and cognition might be due (at least in part) to uraemic toxins in general and their accumulation in the cerebrospinal fluid (CSF) in particular. Indeed, pre-clinical studies have suggested that levels of many uraemic solutes are normally kept in low concentration in the CSF (relative to the plasma ultrafiltrate) by the action of the BBB and blood–CSF barriers. These barriers remain functional when the kidney function fails but cannot prevent the accumulation of uraemic solutes in the CSF [60]. Under normal conditions, the concentration of IS in the brain is reportedly 3.4 times lower than in the serum [60]. Although the source of IS in the brain has not been identified, this limited distribution might be due to the brain-to-blood transfer of IS by the organic anion transporter (OAT) at the BBB [58]. Ohtsuki et al. [58] used the brain efflux index method to examine this transport at the BBB and the involvement of OAT3 in IS transport. These results suggested that OAT3 mediates the brain-to-blood transport of IS and is also involved in the efflux of neurotransmitter metabolites and drugs like acyclovir, cefazolin, baclofen, 6-mercaptopurine, benzoic acid and ketoprofen [58]. Therefore, inhibition of OAT3-mediated brain-to-blood transport might occur in CKD; this would lead to the accumulation of neurotransmitter metabolites and drugs in the brain.

In a mouse model of renal failure (RF), Sato et al. [59] used liquid chromatography–tandem mass spectrometry to determine the levels of IS and p-cresyl sulphate (pCS) in various organs, including the brain. Brain levels of IS and pCS were significantly higher in mice with RF than in control mice. Furthermore, treatment with AST-120—an orally administered polymer that adsorbs low-molecular-weight compounds like indole and p-cresol (precursors of IS and pCS)—was associated with a decrease in brain IS and pCS levels in mice with RF [59].

In a single-photon emission computed tomography (SPECT) imaging study of three rodent models of CKD, Bobot et al. [61] showed that AhR activation by IS leads to disruption of the BBB. Cognitive impairment and BBB disruption were correlated with the serum IS levels [61].

A recent study in the rat confirmed that IS accumulates in the brain (in the brainstem, cerebellum, striatum and hippocampus) and influences the animals’ behavioural profile and cerebral monoamine levels, and thus provided evidence of IS’s neurotoxic activity [7]. The researchers also showed that chronic exposure to IS leads to behavioural alterations, with apathetic behaviour, increased stress sensitivity and reduced locomotor and exploratory activity. Furthermore, IS can contribute to impairments in spatial memory and motor coordination [7]. Watanabe et al. [57] recently reported that rats with adenine-induced CKD presented elevated levels of oxidative stress markers and (in some cases) had pyknotic neuronal cells in the hippocampus.

In unilaterally nephrectomized mice, the serum pCS concentration increased progressively during the administration of pCS at a dose of 100 mg/kg/day. Furthermore, pCS deposition in the prefrontal cortical tissues appeared to be associated with several abnormal behaviours, such as depression, anxiety and cognitive impairment. However, pCS accumulation and behavioural changes were not observed at doses of 1 and 10 mg/kg/day in unilaterally nephrectomized mice or at doses of 1, 10 and 100 mg/kg/day in healthy mice. These changes were alleviated by administration of the adsorbent AST-120 [62].

Clinical studies

Clinical studies have confirmed the above-described preclinical findings in which uraemic toxins are present in the CSF. A recent study measured the concentrations of uraemic toxins (including IS and pCS) in plasma and CSF from control patients and patients with Parkinson’s disease; the CSF–plasma ratio was higher in the latter group. Patients with motor fluctuations had higher levels of uraemic toxins in the CSF but not in the plasma [73].

Only a few clinical studies have evaluated the correlation between neurologic symptoms and uraemic toxin levels. In a cross-sectional study of 204 patients with CKD, individuals
with symptoms of depression had surprisingly lower plasma IS levels [64]. The levels of urea, β2 microglobulin (B2M) and pCS were not significantly associated with depression [64]. In 80 patients on HD, there was a direct, inverse correlation between the Beck Depression Inventory (BDI) score and the levels of interleukin-6 (IL-6) [P = 0.042, odds ratio (95% CI): 1.31 (1.01–1.71)] and creatinine [P = 0.050, odds ratio (95% CI): 0.73 (0.54–1.00)], and the Hamilton Anxiety Rating Scale (HARS) correlated significantly with parathyroid hormone (PTH) levels only in univariate analysis [63]. Indeed, IL-6 has been consistently found to be elevated in stress reactions and depression patients, highlighting the important role of inflammatory processes in depression pathogenesis [75, 76].

Similarly, clinical data on the impact of uraemic toxins on cognitive functions are scarce. A cohort of 199 patients with CKD had poorer cognitive function and higher serum pCS and IS levels than 84 patients with normal renal function. In the patients with CKD, a higher serum IS level was independently associated with poor executive function but the serum pCS level was not associated with cognitive function [65]. A recent study that included patients on HD sought to investigate the association between circulating levels of free IS, pCS, IAA and hippuric acid (HA) on cognitive function, as assessed with the Mini-Mental State Examination (MMSE) and the Cognitive Abilities Screening Instrument (CASI). Both free IS and free pCS levels were negatively associated with the MMSE and CASI scores. After controlling for confounders, the MMSE and CASI scores were still associated with the circulating free IS level but not with the free pCS level [69]. Serum IAA was associated with cognitive impairment, according to the MMSE and CASI scores. There was no correlation between serum HA levels and cognitive function [70]. Cognitive function improves after kidney transplantation [50, 51] suggests that uraemic toxins have a role in this impairment. A recent exploratory study of 10 kidney transplant recipients and 18 non-transplanted patients with CKD sought to evaluate five major cognitive domains (memory, attention and concentration, information processing speed, abstract reasoning and executive function) via an extensive neuropsychological assessment before kidney transplantation and then 1 week and 3 months after transplantation. The researchers did not find any evidence of cognitive changes after kidney transplantation, relative to the control groups. The small number of patients might explain the lack of a significant change. However, the researchers observed clear changes in the serum levels of uraemic toxins after transplantation—even within 3 days [72], when previously reported a marked reduction after 1 month after kidney transplantation [52, 53].

The findings on the relationship between uric acid and cognitive functions are contradictory. Low serum levels of uric acid are associated with dementia and Parkinson’s disease [67]. Therefore, one would expect high levels to be neuroprotective—perhaps because uric acid is a major antioxidant in the plasma [66, 77]. In a clinical trial in non-CKD patients with acute ischaemic stroke, the addition of uric acid to thrombolytic therapy did not increase the proportion of patients who achieved excellent outcome after stroke compared with placebo [RR (95% CI): 1.23 (0.96–1.56)] [78]. A reanalysis of the trial concluded that in women, who usually have lower serum urate levels, the administration of uric acid reduced infarct growth and was better than placebo to reach excellent outcome [RR (95% CI): 2.09 (1.05–4.15)] [79]. However, a high serum uric acid level is a hallmark of vascular disease. Hence, any neuroprotective benefit of hyperuricaemia is probably counterbalanced by the effects of neurovascular disease [68]. Accordingly, gout (in which hyperuricaemia is usually unrelated to CKD) has never been linked to cognitive deficits. Therefore, an association between CKD-related hyperuricaemia and cognitive dysfunction is surprising [80] and should be interpreted with caution, given the large number of other toxins that inevitably accompany CKD.

**INDIRECT EFFECTS OF URAEMIC TOXINS ON THE BRAIN**

Uraemic toxins might also have harmful indirect effects on the brain via oxidative stress [6, 81], vascular dysfunction [82, 83], vascular calcification [84–86], coagulation disorders [87], and cardiovascular diseases like atrial fibrillation and hypertension [88, 89]. These indirect effects have been reviewed in detail elsewhere and will not be described here [5, 83].

**POTENTIAL IMPACTS ON PATIENT CARE**

Uraemic toxins appear to be factors that (in addition to others) specifically influence the increased prevalence of cerebrovascular and neurological signs and symptoms in patients with CKD. Drug treatments that modulate uraemic toxin concentrations (e.g. the administration of intestinal chelators) might prevent the development of harmful effects in patients with CKD. However, there are few therapeutic options. Although phosphate binders can effectively reduce phosphate levels, repositioned products from this drug class do not appear to greatly decrease circulating levels of other uraemic toxins [90]. The orally administered activated charcoal adsorbent AST-120 is widely used in Asian countries to specifically decrease uraemic toxin levels. However, its putative effect on neurological outcomes has yet to be assessed in clinical trials.

It is unlikely that uric acid has a role in brain dysfunction, and so the value of widely used urate-lowering drugs in the treatment of the cerebrovascular and neurological signs and symptoms of CKD can be questioned. Indeed, allopurinol does not modify cognitive impairment in a rare genetic form of hyperuricaemia (Lesch–Nyhan syndrome) [91]. However, high doses of allopurinol and febuxostat reportedly reduce the risk of dementia [92]. An ongoing trial expected to be completed in 2021 is seeking to determine allopurinol’s potential value in ischaemic stroke in patients with eGFR ≥30 mL/min/1.73 m² [93]. Future clinical studies of CKD and cognitive impairment should also consider the use of urate-lowering drugs.

Patients with CKD are frequently affected by gut dysbiosis, which might increase the production of gut-derived uraemic toxins [94]. Probiotics and other nutritional therapies might reduce gut dysbiosis and decrease circulating uraemic toxin levels,
oxidative stress and inflammation [95]. Since protein-bound uraemic toxins are difficult to remove by HD, the gut microbiota might be an alternative target for reducing circulating levels of these compounds and thus their neurologic and other toxic effects in patients with CKD.

New dialysis techniques or methods for better removing protein-bound uraemic toxins might help to reduce the neurologic signs and symptoms that frequently affect patients on HD [96, 97]. Lastly, and as mentioned above, kidney transplantation is followed by a rapid decrease in uraemic toxin levels [56, 74], an improvement in cognitive function [55, 57] and greater white matter integrity [55].

CONCLUSION
It is important to identify factors that explain the association between CKD and cerebrovascular/neurologic complications. The uraemic toxins that accumulate in the blood in ESRD might explain this association (at least in part in more advanced CKD stages) and constitute potentially valuable therapeutic targets.

SUPPLEMENTARY DATA
Supplementary data are available at ndt online.

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CONFLICT OF INTEREST STATEMENT
The authors do not have any conflict of interest relating with this manuscript. The results presented in this article have not been published previously in whole or part, except in abstract format.

APPENDIX
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