Ketamine initially introduced as an anesthetic agent in the 1960s, is a receptor complex antagonist n-methyl-d-aspartic acid that has been used as an anesthetic and/or analgesic for many years. In the past decade the use of ketamine as an analgesic or “analgesic adjuvant” appears to have increased. Subanesthetic doses of ketamine have been utilized for analgesia and opioid-sparing effects in chronic pain as well as in refractory cancer pain, and palliative medicine (1-11). Additionally, recently its use studied seems to have grown for chronic noncancer pain (12,13). Its increased use studied overall has largely been for neuropathic pain and especially for the treatment of refractory complex regional pain syndrome (14-17). Ketamine has also been studied to reduce allodynia in patients with complex regional pain syndrome (18). Different routes of administration have been reported to produce beneficial results such as low-dose intranasal (S)-ketamine in patients with complex regional pain syndrome (18).

The mechanism of action has been considered to be mainly a noncompetitive antagonism of the N-methyl-D-aspartic acid (NMDA) receptor [19], however, its mechanisms of action may be extremely complex as ketamine may interact with multiple other receptors (19,20). Ketamine has also been shown to inhibit tumor necrosis factor-alpha (TNF-alpha) and interleukin 6 (IL-6) gene expressions in lipopolysaccharide (LPS)-activated macrophages (21). It has even been speculated that the antiproliferative effects may be responsible for antihyperalgesic effects of ketamine (22).

The ketamine molecule contains a chiral carbon atom marking a chiral center. Ketamine can exist in two forms, or enantiomers; S-ketamine and R-ketamine. The physical properties of the enantiomers are identical, but their interactions with complex molecules, underlying PK/PD parameters, might differ (23). The elimination clearance of S-ketamine is more than that of R-ketamine (24).

Although anesthesiologists and pain specialists alike are familiar with the classic and common potential adverse effects of ketamine, they may not be as familiar with an increasingly recognized potential association of ketamine that clinicians should be aware of when prescribing ketamine. The observation that ketamine may lead to significant urological side-effects was first recognized in chronic ketamine recreational use (25-27).

In the South West of England, Cottrell and colleagues have reported increasing numbers of patients referred to urological services with complications of chronic ketamine use (28). Other cases have been reported in mainland Europe, Canada, Malaysia (26, 29) and the UK (30).

Chen and colleagues discovered 4 female patients, ages 18 to 25 years, who presented to emergency department (ED) with urinary urgency, frequency, and pelvic pain, who had negative work-up and no response to antibiotic therapy were diagnosed with presumed interstitial cystitis (31). More detailed history revealed that they were recreational ketamine users. Their symptoms significantly improved after stopping ketamine abuse (31).

Urological side effects to ketamine have been reported in the last 12 months as an emerging prob-
problem amongst the drug user population (25, 29, 32). Storr and Quibell describe three patients where significant urological symptoms have arisen, possibly as a result of taking ketamine as an analgesic (33). Urinary tract symptoms are becoming increasingly recognized by ketamine abusers; 20–30% of frequent users reporting bladder symptoms (25, 34). Case series demonstrate a temporal link between ketamine use (abuse) and urological symptoms, urinary tract damage and renal impairment (25, 29, 34), with some but not all improving on cessation of ketamine.

Chu and colleagues (25) reported lower urinary tract symptoms of ketamine-induced cystitis include: dysuria, frequency, urgency, incontinence, macroscopic hematuria and suprapubic pain. Investigations typically included negative urine culture (97% in Chu study). Forty-two (71%) patients had cystoscopy that showed various degrees of epithelial inflammation similar to that seen in chronic interstitial cystitis (25). All of 12 available bladder biopsies had histological features resembling those of interstitial cystitis showing various degrees of epithelial inflammation, ulceration, pectechial haemorrhages, neovascularisation and contact bleeding. Urodynamically, either detrusor overactivity or decreased bladder compliance with or without vesico-ureteric reflux was detected to some degree in all of the 47 patients who agreed to have a video cystometrygram (VCMG) (25). Six of the 47 patients (13%) showed vesicoureteral reflux (VUR) as a secondary event to the severely contracted bladder with high detrusor pressure. Thirty patients (51%) had unilateral or bilateral hydronephrosis on renal ultrasonography, and four (7%) showed features suggestive of papillary necrosis on radiological imaging. Eight patients had a raised serum creatinine level (25). Histology shows denuded bladder epithelium, often with eosinophilic infiltration within the lamina propria.

Tsai et al. reported eleven patients with urinary tract symptoms and a history of ketamine abuse (35). The most common complaints were lower urinary tract symptoms, including dysuria, frequency, urgency and gross hematuria. Urinalyses showed nonbacterial pyuria and were negative for tuberculosis. All biopsy specimens showed infiltrations of granulocytes (mostly eosinophils) and mast cells within the bladder tissue. Medications produced only slight clinical improvements. Intravesical instillation of hyaluronan solution was performed for some patients and significant improvement of lower urinary tract symptoms was observed (35).

Ketamine inhibits, in a dose-dependent manner, cardiovascular and visceromotor responses to the acute noxious stimulus of distension of the urinary bladder (36). This effect is likely related to NMDA receptor antagonist actions of ketamine, since other drugs with such pharmacologic properties produced a qualitatively similar inhibition (36). The main site of action of the drug appears to be localized to the spinal cord since intrathecal administration of inhibition of the mean arterial pressure changes (ΔMAP) and electromyographic (EMG) to urinary bladder distention (UBD) (36).

Mason and colleagues reported a case series illustrates the harmful effects of ketamine on the urinary tract and the associated radiological findings. Delayed diagnosis can result in irreversible renal tract damage requiring surgical intervention (37).

Ultrasound demonstrated small bladder volume and wall thickening. CT revealed marked, generalized bladder wall thickening, mucosal enhancement, and perivesical inflammation. Ureteric wall thickening and enhancement were also observed. In advanced cases ureteric narrowing and strictures were identified using both CT and IVU (37).

One proposed mechanism for KIUI is that it only occurs in patients with a disorder of the urine–tissue interface (25). Thus, penetration of toxic urinary compounds (e.g. ketamine and/or its metabolites) into the bladder wall, as also seen in acute cystitis, may contribute to urinary urgency, frequency and pain (38).

Contact cystitis (25, 29) is another possible mechanism. Ketamine, norketamine and hydroxynorketamine can be measured in high quantities in the bladder. It is hypothesized that these directly trigger an inflammatory response through IgE, mast cell degranulation, eosinophil recruitment and enzyme release (25). Another theory proposed (25) is ketamine and its metabolites damage the microvasculature of the bladder potentially leading to the ischemia, fibrosis and abnormal neo-vascularisation identified on histology. Other suggested mechanisms include an autoimmune reaction (25).

Treatment of KIUI is to at least decrease the dose of ketamine and optimally to discontinue ketamine. Therapeutic strategies that may be useful may involve approaches aimed at repair of the urothelium. Elmiron (pentosan polysulfate sodium), a low molecular weight heparin-like compound, has been tried to rebuild the glycosaminoglycan (GAG) layer of the damaged urothelium and a few patients experienced symptom relief after treatment (29).

It appears that illicit synthetic ketamine obtainable
illegally is reasonably “pure”, since when a small amount which was secured with the aid of the Metropolitan Toronto Police Department Narcotics Division, was analyzed it contained only ketamine with trace amounts of dimethyl sulfoxide (29). It would seem unlikely that the urological insult associated with recreational ketamine use is due to excipients or contaminants from “street” ketamine. Thus, any potential associated bladder dysfunction/urologic insult is probably largely at least partly due to direct local effects of ketamine and/or its metabolites.

Although, the potential association of ketamine dose and the incidence of urological insult is unknown; it seems that daily ketamine infusions administered at outpatient treatment centers for pain relief may go at least as high as 300 mg/hr (39). These doses should certainly be enough to potentially contribute to urological insult. Persson has postulated that long-term use may be an important factor in potentially contributing to bladder dysfunction/urologic insult (40).

Although the reported cases have mainly concerned recreational drug users, they are relevant for ketamine used for legitimate long-term prescribed analgesic use as well (40). Persson successfully utilized long-term oral ketamine in efforts to achieve improved analgesia and a reduced opioid dose in a middle-aged patient who had a lumbar “disc herniated” and complained of severe post-surgical chronic low back and leg pain. After about a year of treatment, the patient started experiencing increased urinary frequency, however, if he lowered the ketamine dose the bladder symptoms diminished but the pain increased and vice versa (40).

As the use of ketamine in interventional pain medicine grows, interventional pain specialists should be aware of all the potential therapeutic effects and adverse effects of ketamine. Clinicians should assess for the development of symptoms and signs of bladder dysfunction/irritation in patients on ketamine therapy. Significant consideration should be given to decreasing the dose or discontinuing ketamine therapy in a patient who suddenly develops symptoms or signs of bladder dysfunction and/or irritation during treatment with ketamine.

References


