Fatal gadolinium-induced encephalopathy following accidental intrathecal administration: a case report and a comprehensive evidence-based review

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ABSTRACT

Gadolinium-based contrast agents (GBCAs) have been suggested as off-label alternatives to iodinebased contrast agents for fluoroscopic imaging during interventional pain procedures. We report a case of accidental intrathecal administration of a GBCA during a neuraxial interventional pain procedure leading to acute gadolinium neurotoxicity, which resulted in encephalopathy and ultimately death. To our knowledge, it is the first published case of fatal intrathecal gadolinium-induced encephalopathy and the first published case of intrathecal gadoteridol causing serious neurologic complications. In addition, the case presented here is placed in context with an associated comprehensive, evidence-based review of the use of gadolinium in interventional pain procedures, addressing gadolinium chemistry and pharmacologic properties, neurotoxicity and radiology. Physicians must be aware that gadolinium poses a significant risk of acute neurotoxicity even in small doses. Until further safety research is performed, GBCAs should not be considered a safe alternative for use in neuraxial interventional spine procedures when there is a risk of inadvertent intrathecal administration.

INTRODUCTION

Gadolinium is a metallic element from the lanthanide series prescribed for use as a contrast agent with MRI.^{1 2} Since the 1980s, gadolinium-based contrast agents (GBCAs) have been administered via the intravenous route prior to MRIs and currently are used in approximately 35% of all MRI examinations.³ Although considered relatively safe in individuals with normal renal function, the administration of intravenous GBCAs has been associated with adverse events.^{4 5} Recently, more attention has been placed on gadolinium toxicity, including nephrogenic systemic fibrosis (NSF) and tissue and brain deposition following intravenous use.^{3 6 7}

GBCAs are also used off label as alternatives to iodine-based contrast agents to assist with visualization in interventional pain procedures in individuals with known hypersensitivity reactions to iodine. Although the practice of injecting GBCAs into the neuraxial space (ie, epidural and intrathecal areas) is not Food and Drug Administration (FDA)-approved, published articles and leading pain management textbooks have suggested GBCAs as alternatives to iodine-based contrast agents. The utilization of GBCAs for interventional pain procedures has not been extensively studied, and

the associated risks have received less attention than other intravenous GBCA-related complications. Lately, safety concerns have been raised regarding GBCA utilization for interventional pain management procedures, especially when there is a risk of intrathecal administration. ¹² ¹³

Based on animal models and human case studies, the off-label use of a GBCA in the intrathecal space either intentionally or unintentionally may pose a significant risk of acute neurotoxicity. 12 14 15 We report a case of inadvertent GBCA intrathecal administration following a neuraxial interventional pain procedure leading to neurotoxicity, encephalopathy, and death. To our knowledge, this is the first published case of fatal intrathecal gadolinium-induced encephalopathy and the first published case of intrathecal gadoteridol (ProHance) causing serious complications secondary to acute neurotoxicity. In addition to reporting this case, we provide a comprehensive review of the use of gadolinium in interventional pain procedures, addressing its radiologic and pharmacologic properties and neurotoxicity, with a focus on neuraxial use for interventional pain procedures.

CASE REPORT

The deceased patient's family consented to the publication of this report. This procedure was not performed by the authors and occurred at an outside outpatient surgery center. A 67-year-old woman with a history of lumbar spinal stenosis secondary to ligamentum flavum hypertrophy presented for a minimally invasive lumbar decompression (MILD). Her medical history was significant for hypothyroidism, hypertension and sleep apnea. It was her second MILD procedure in 4 months. Because of the patient's reported iodine allergy, a GBCA (ie, gadoteridol) was used for fluoroscopic guidance in both MILD procedures for enhanced visualization. Her first MILD procedure was completed without complications. Gadoteridol was also used for her second MILD procedure in order to create the epidurogram. Following the introduction of an 18-gauge Tuohy needle into the epidural space at L5-S1, 5 mL of gadoteridol were injected with reported confirmation of epidural flow. The pain physician's operative note indicated the MILD procedure was completed without any observed complications.

In the recovery room immediately after the completion of the procedure, the patient experienced a severe headache, mental status changes,



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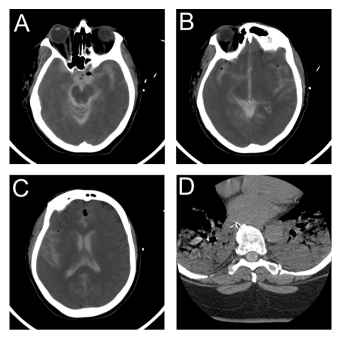


Figure 1 CT head images from postoperative day 0. Cranial CT images demonstrate gadolinium (hyperattenuation) in the basal cisterns (A), third ventricle (B), lateral ventricles, and right sylvian fissure (C). Spine CT imaging taken of the lower thoracic spinal canal displays gadolinium in the intrathecal space (D).

apnea, agitation and increased muscle tone in her arms and legs. Due to her declining condition, she was urgently sent to a hospital for further treatment.

On arrival at the hospital, the patient demonstrated signs of distress including twitching in her eyes and tongue, thrashing and crying out. She also displayed decreased respirations and myoclonic activity. About 20 min after arrival, the patient experienced wide-complex pulseless tachycardia. The patient was intubated and underwent successful cardiopulmonary resuscitation. After 4 min, normal vital signs were restored. The patient then experienced a fever of 103 °F (39.4°C) that did not resolve with acetaminophen and ice packs. The patient also began experiencing seizures that occurred approximately every 20 min. A CT scan of the head was obtained, and the images displayed hyperattenuation in the subarachnoid space that was originally misinterpreted as subarachnoid hemorrhage (figure 1A, B and C). Because of the potential necessity for neurosurgical services, the patient was flown via emergency helicopter to another hospital.

Additional CT and MRI images were taken at this hospital (figures 1D and 2). Reviewing these images and the images taken at the previous hospital, the medical team determined that the images did not indicate a subarachnoid hemorrhage but rather showed an accumulation of GBCA within the intrathecal cerebral spinal fluid (CSF). A diagnosis of encephalopathy secondary to intrathecal injection of a GBCA was made. Brain imaging also revealed infarctions that suggested anoxic brain injury (figure 2). At this point, the neurosurgical team determined it was too late to drain the patient's CSF. The patient's condition continued to deteriorate, culminating in multisystem organ failure and coma. Two weeks after her MILD procedure, the patient was placed into hospice care where she died 4 days later (18 days after the procedure).

The autopsy found no evidence of subarachnoid hemorrhage. Large cortical infarcts were found in the right occipital and parietal lobes. The right insular cortex displayed infarction

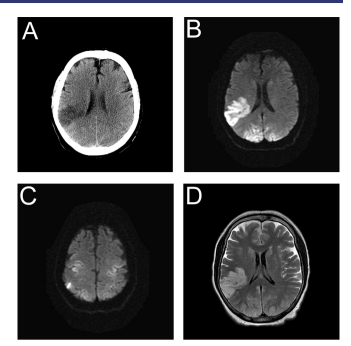


Figure 2 Imaging taken postoperative day 4. Cranial CT imaging (A) displays normal lateral ventricles and evidence of gadolinium-induced cytotoxic edema in the right parietal lobe (low attenuation). MRI DWI show gadolinium-induced cytotoxic edema in the right parietal lobe (B) corresponding to figure 2A. There is also evidence of gadolinium-induced cytotoxic edema in the bilateral occipital region (B) and the bilateral parietal lobes (C). MRI FLAIR reveals cytotoxic edema in the right parietal lobe (D) corresponding to figure 2B. The bright signal indicates that gadolinium is still present in the sulci. DWI, diffusion-weighted imaging; FLAIR, fluid-attenuated inversion recovery.

with borderline cavitation. On microscopic examination, the cerebellum displayed ischemic changes, and the hippocampus and lower pons displayed mild gliosis. The midbrain appeared primarily intact. Cause of death was determined to be encephalopathy caused by the combined factors of gadolinium neurotoxicity and hypoxic–ischemic encephalopathy following cardiopulmonary arrest with successful restoration of circulation.

DISCUSSION AND REVIEW OF THE LITERATURE

To our knowledge, we report the first case report documenting a death following acute GBCA neurotoxicity. 16 In this case, the presence of intrathecal gadolinium confirmed by radiographic imaging suggests that the intrathecal space was inadvertently entered when performing the epidurogram for the MILD. The intrathecal administration of a GBCA resulted in acute neurotoxicity and encephalopathy, including a cascade of cardiovascular and neurologic consequences. The signs and symptoms presented in this case, including alterations in mental status, seizures, respiratory distress and tachycardia, correspond to previous animal research and clinical reports of gadolinium neurotoxicity (table 1). 12 16-21 However, to date, there has not been a reported case of death following the utilization of gadolinium for neuraxial interventional pain procedures. Below we discuss the safety concerns regarding the use of GBCAs for neuraxial interventional pain procedures.

Gadolinium chemistry and relevant pharmacology

Prior to considering the use of GBCAs as an alternative contrast agent for neuraxial interventional pain procedures, it is of critical

Author	Procedure	GBCA* dose used	Signs and symptoms	Outcome
Arlt et al ¹⁷	CT myelogram.	Gadopentetate dimeglumine 20 mL 7 µmol/g brain.†‡	Confusion, nausea, vomiting, dysarthria, somnolence, blurred vision, delirium, limb ataxia, gaze-evoked nystagmus, aggressive behavior, visual and auditory hallucination and incomplete anterograde amnesia.	Day 4: regained alertness Day 7: resolved behavioral disturbances and psychotic symptoms Day 10: discharged Day 56: persistent concentration difficulties
Li et al ¹⁸	MRI myelography.	Gadopentetate dimeglumine 15 mL 5.35 µmol/g brain.†‡	Headache, nausea, vomiting, coma and systemic seizures.	Day 8: regained consciousness Day 22: discharged Day 60: no reported side effects
Kapoor <i>et al</i> ¹⁹	Epidural steroid injection.	Gadodiamide 4 mL 1.43 µmol/g brain.†	Mental status changes, grand-mal seizure, respiratory distress, agitation, hyperglycemia, sinus tachycardia, respiratory acidosis, metabolic alkalosis and amnesia.	Day 8: responded to verbal commands and improved pulmonary function Day 10: discharged Day 182 (6 months): complained of partial seizure-like activity
Park et al ²⁰	CT myelogram.	Gadopentetate dimeglumine 6 mL 2.14 µmol/g brain.†‡	Confusion, global aphasia, vomiting, stupor, severe rigidity, intermittent seizures, fever and high blood pressure.	Day 4: improved cognition Day 15: discharged 1 month: complained of visual disturbances
Nayak <i>et al</i> 16	Administered through side port of an intraventricular catheter.	Gadopentetate dimeglumine 10 mL 3.57 µmol/g brain.†	Agitation, labile blood pressure, aphasia, dysarthria, depressed mentation, right facial droop and increased urine output.	Day 5: performed lumbar CSF drain Day 26: required a tracheostomy for ventilatory support 2 months: still in intensive care. Resolution of the intraventricular GBCA
Reeves et al ²¹	Intrathecal catheter contrast study.	Gadobutrol 2 mL 1.43 µmol/g brain.	Severe spastic pain and spasms in lower extremities.	Day 28: no sequelae
Popescu et al ¹²	L4-L5 interlaminar epidural steroid injection.	Gadobutrol 1.5 mL 1.07 μmol/g brain.†	Vomiting, seizure activity, impaired consciousness and respiratory compromise requiring intubation.	Day 1: extubated and made full recovery

^{*}In this column, volume of GBCA administered is provided alongside the estimated concentration of gadolinium per gram of brain. Calculated using 1400 g as the average weight of the human brain in order to be consistent with calculations from Arlt *et al.*¹⁷

GBCA, gadolinium-based contrast agent.

importance that interventional pain physicians understand relevant gadolinium chemistry, basic mechanisms of action, pharmacology and toxicity. Gadolinium is a trivalent lanthanide ion (Gd³⁺). Free gadolinium ions are toxic. In GBCAs, the gadolinium ions are bound tightly to chelating agents to form a stable complex that reduces the toxicity of the parent compound. The chelating ligand limits the dissociation of the toxic gadolinium ion (Gd³⁺) from the ligand during both the shelf-life and the in vivo time periods. Gadolinium has nine coordination sites; eight sites consist of the bond between gadolinium and

the chelating ligand and the ninth consists of a coordinate bond between gadolinium and the oxygen atom of a water molecule.²³

GBCAs (table 2) are structurally classified as macrocyclic or linear (open chain) based on the chelating ligand and further subdivided into ionic and non-ionic compounds.²³ Macrocyclic GBCAs are considered more stable (ie, the gadolinium ion is more tightly bound to the chelating agent) and are associated with a lower dissociation constant (ie, slower dissociation) and a lower likelihood of dechelation (ie, release of the free gadolinium ion) in comparison to linear GBCAs.²³ ²⁴ Ionic GBCAs dissociate

Table 2	Prescribing informat	tion of different	hrands of	andalinium ²³
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Trade name	Generic name	Structure	Documents the risk of serious nervous system-related complications?	Presents information regarding intrathecal injection?	Molar concentration of gadolinium (mol/L)
Ablavar ⁵⁸	Gadofosveset trisodium	Linear ionic	No	No	0.25
Dotarem ⁵⁹	Gadoterate meglumine	Macrocyclic ionic	Yes	No	0.5
Eovist ⁶⁰	Gadoxetate disodium	Linear ionic	Yes	No	0.25
Gadavist ⁶¹	Gadobutrol	Macrocyclic nonionic	Yes	No	1
Magnevist ⁶²	Gadopentetate dimeglumine	Linear ionic	Yes	No	0.5
MultiHance ⁶³	Gadobenate dimeglumine	Linear ionic	Yes	No*	0.5
Omniscan ⁶⁴	Gadodiamide	Linear non-ionic	Yes†	Yes‡	0.5
Optimark ⁶⁵	Gadoversetamide	Linear non-ionic	Yes	No*	0.5
ProHance ⁶⁶	Gadoteridol	Macrocyclic non-ionic	Yes	No	0.5

^{*}States that it is only for intravenous use.

[†]Intrathecal administration was unintentional.

[‡]Unintentional intrathecal administration was due to drug error.

[†]Specifically indicates the risk of neurotoxicity due to intrathecal administration.

^{\$}States: 'Not for intrathecal use. Inadvertent intrathecal use of gadodiamide has occurred and caused convulsions, coma, sensory and motor neurologic deficits.'

into charged particles when entering solutions and have a higher viscosity and osmolality in comparison to non-ionic GBCAs.² In theory, non-ionic GBCAs may have lower levels of extravasation into soft tissues during intravenous administration. However, linear ionic GBCAs are more thermodynamically stable than linear non-ionic GBCAs and are less likely to undergo dechelation via transmetalation (ie, the exchange of organic groups between metals).^{24 25} Because GBCAs have different structural classifications, disassociation constants and thermodynamic and kinetic stability constants, the risk of gadolinium toxicity differs among available formulations.²³

For neurotoxicity, dechelation resulting in the release of the gadolinium ion is a major concern. At physiologic pH, unassisted dechelation of GBCAs is limited.²³ In vivo, the major method of dechelation is through transmetalation where endogenous cations (eg, Cu²⁺, Ca²⁺, Fe³⁺ and Zn²⁺) compete and displace the gadolinium ion from the chelating agent. ^{16 23 26} When gadolinium is released from a GBCA, it is termed as unchelated, dechelated or disassociated. 27 28 In vivo, when gadolinium is released from a GBCA, it is typically not free, but rather it is bound to a molecule or chemical. These host molecules include precipitated oxides, hydroxides, carbonates or natural biologic chelating agents such as citrate, amino acids, peptides or proteins.²⁸ A significant amount of the research examing gadolinium transmetalation examined the process involving intravenous use, not direct administration of GBCA into the CSF.²² The specific impact of transmetalation on gadolinium toxicity with intrathecal use is unclear.

GBCAs use in interventional pain medicine

In individuals with hypersensitivity reactions to iodine based radiographic contrast agents, GBCAs have been suggested as an alternative agent. ⁹ ¹⁰ ¹² ¹³ ²⁹ Currently, nine GBCAs (table 2) are approved for intravenous use as a contrast agent in MRI protocols.²³ When used off label in the neuraxial space as an alternative to iodinated radiographic contrast media, GBCAs have an intrinsic ability to attenuate X-rays. Maus et al²⁹ examined the relative conspicuity of eight GBCAs (gadofosveset trisodium, gadoterate meglumine, gadoxetate disodium, gadobutrol, gadopentetate dimeglumine, gadobenate dimeglumine, gadodiamide and gadoteridol). Based on the in vitro testing, radiographic contrast levels of each tested GBCA were proportional to the gadolinium molar concentration. Overall, the radiopacity of gadolinium is less than that of iodinated contrast agents. Of the GBCAs tested, gadobutrol had the highest molar concentration of gadolinium (table 2) and had the highest radiodensity. When compared with iodinated radiographic contrast media, gadobutrol's radiographic contrast level existed between iohexol 240 and 140 mg/mL in the kVps ranges tested (70–125 kVp).²⁹

Both peer-reviewed published articles and pain textbooks have suggested that gadolinium may be used as an alternative to iodinated contrast agents for neuraxial pain procedures, often with minimal attention given to the risk of neurotoxicity. 9-12 In addition, despite the limited experience of intrathecal administration (inadvertent and advertent) of gadolinium, it has been stated that it is safe when GBCAs are directly administered into the CSF. 10 11 Regarding interventional pain procedures, Safriel et al published the largest case series to date, examining the use of gadolinium in 92 patients with iodinated radiographic contrast allergies. Their group performed 127 outpatient procedures including discograms, epidural steroid (cervical, thoracic and lumbar) injections via both the interlaminar and transforaminal routes, and facet and nerve root blocks (cervical and

lumbar) injections. The gadolinium doses ranged from 0.2 mL to 15.83 mL (ie, three-level discogram). For epidural steroid injections, the gadolinium dose ranged from 1 mL to 5 mL and for nerve root blocks the dose ranged from 0.2 mL to 1 mL per level. The type of gadolinium used was gadodiamide (linear non-ionic; table 2). For the neuraxial procedures, the authors performed a gadolinium test injection to confirm location of the needle and to exclude intrathecal injection. The volume of the gadolinium test injection was not provided. The authors do not report any occurrences of intrathecal administration. Safriel *et al*⁸ concluded that gadolinium seems to be a safe alternative in spine management procedures in individuals at high risk for a reaction to radiographic iodinated contrast.

Gadolinium toxicity

Gadolinium toxicity can be aligned into four general categories: (1) acute toxicity, including allergic reactions, (2) subacute/ chronic response via cell mediated reactions (eg, NSF), (3) gadolinium storage condition (ie, gadolinium lies primarily inert within the body) and (4) gadolinium deposition disease that involves both acute and subacute components of the immune response.²⁷ For all four categories, genetic and immune roles have been implicated.²⁷ Immediate hypersensitivity reactions to GBCAs are rare and range from 1.5, 8.3 or 16 immediate allergic like adverse events per 10 000 administrations for linear non-ionic, linear ionic and macrocyclic non-ionic GBCAs, respectively. ⁴ Although rare, the immediate hypersensitivity reactions can be fatal.³⁰ The major toxicity concern with intravenous administration of GBCAs is NSF (fibroproliferative disorder) in patients with pre-existing renal disease. Fibrotic lesions in individuals with NSF develop a few days to months after exposure to GBCAs.³¹ Typically, individuals with NSF develop abnormalities in the skin, including swelling and tightness, but the condition can also affect myofascial tissues and body organs including the lungs, diaphragm, myocardium and striated muscle.³² The role of GBCAs and the exact immune mechanism by which the fibrotic lesions are triggered is currently unknown. 31 32 Linear GBCAs (eg, gadodiamide and gadopentetate dimeglumine) have been mainly implicated in the pathogenesis of NSF.²⁶

Recently, much attention has been placed on gadolinium deposition. In 2015, the FDA released the safety communication on the risk of brain deposits with repeated use of intravenous administration of GBCAs for MRI.³³ This safety communication was updated in 2017.34 Gadolinium deposition typically occurs in select areas of the brain including the dentate nuclei, thalamus, caudate nucleus and globus pallidus. 35 36 Gadolinium depositions have been seen with both linear and macrocyclic contrast agents although more often with the former.^{35 36} Iron has been suggested as playing a central role in transmetalation and gadolinium toxicity for both NSF and gadolinium deposition in the brain.²⁶ The areas of the brain typically targeted with gadolinium deposition are intrinsically iron rich.²⁶ In a vast majority of the literature, more attention has been paid to the potential risks following gadolinium deposition in the central nervous system (CNS) after repeated use rather than acute neurotoxicity following a single intrathecal use. Currently, the clinical consequences of gadolinium brain deposition after repeated intravenous gadolinium exposure is unknown. 13 34

Acute neurotoxicity

Although gadolinium deposition and NSF are often the main concerns with the intravenous use of GBCAs, when GBCAs are used in relatively small volumes for neuraxial interventional pain

procedures, the major concern is acute CNS neurotoxicity. When given intrathecally, the blood–brain barrier does not play a role, and no barrier exists at the pial surfaces to the passage of gado-linium.^{37 38} Entry into the brain parenchyma occurs by simple diffusion. When a GBCA is given intrathecally, the amount of brain penetration is variable and often difficult to estimate.³⁹

The exact mechanism by which gadolinium induces neuron disorders and neurotoxicity is unknown. 15 To date, multiple mechanisms for acute neurotoxicity and cell death have been suggested including: (1) impairment of mitochondrial function leading to induced oxidative stress that results in cell apoptosis, ¹⁵ (2) stress-related endoplasmic reticulum alterations in signal transduction, 40 (3) alteration in cell signal processes including protein synthesis and the cell redox state 41 42 and (4) inhibition of cellular calcium hemostasis and modulation of unfolded protein responses. 43 Gadolinium competes with calcium and blocks voltage gated calcium channels at low concentrations.⁵ Gadolinium ions and calcium ions have similar atomic radii, 107.8 pm and 114 pm, respectively. The blocking of these voltage gated calcium channels results in multiple negative consequences including alterations in coagulation, inhibition of smooth, skeletal and cardiac muscle contraction and transmission of nerve impulses.^{5 6 44} Furthermore, the risk of CNS neurotoxicity is increased with gadolinium, because it remains in specific areas of the brain including the cortex, hypothalamus and hippocampus for an extended period of time. 45 Below we will discuss both animal and human studies examining neurotoxicity with GBCAs.

Animal studies examining neurotoxicity of GBCA intrathecal administration

Multiple animal studies have examined intrathecal administration of GBCAs including gadodiamide (linear non-ionic), gadopentetate dimeglumine (linear ionic) and gadobenate dimeglumine (linear ionic). 14 38 46-49 In two studies, Ray et al 14 47 demonstrated dose-dependent GBCA-associated neurotoxic effects in mice. In the first study (table 3), intraventricular administration of gadobenate dimeglumine resulted in acute neurotoxicity over several hours (stereotyped movements and myoclonus) and medium-term neurotoxicity over several days (ataxia and tremor). 14 The neurotoxicity was dose dependent with changes in behavior and twitching beginning at a concentration of 5 µmol/g brain, ataxia and fine muscular tremor beginning at 10 µmol/g brain and an increase in severity of these symptoms at 15 µmol/g brain. No adverse reactions were seen at concentrations of 2.5 µmol/g and 3.3 µmol/g brain. The myoclonic twitching was suggestive of focal seizure activity. In addition, neuropathologic changes developed over 24 hours with the reactive changes persisting for 42 days. Morphologic changes also developed at doses of 5 µmol/g brain and higher. The early changes consisted of loss of oligodendroglia and astroglia and disintegration of the myelin sheath. Axons also showed necrotic changes. Morphologic changes most commonly occurred in the thalamus, hypothalamus, superior olivary nuclei and vestibular nuclei. Lesions were also found in the lumbar and cervical spinal cord at the gray and white matter junctions. The histopathologic analysis of the CNS demonstrated changes similar to those observed in human central pontine myelinolysis.

In a second animal study, Ray *et al*⁴⁷ examined the neurotoxic potential of both gadodiamide and gadopentetate dimeglumine, and both agents were found to be neurotoxic. Gadodiamide also produced dose-dependent acute neurotoxicity and morphologic changes. The gadodiamide produced morphologic changes that were identified in the thalamus, caudate nucleus of the basal

ganglia and cerebellum. The cerebellum was most sensitive to the effects of gadodiamide. Unlike gadopentetate dimeglumine, spinal cord lesions were not identified with the administration of gadodiamide. Both the acute neurotoxicity and morphologic changes were dose dependent with the behavioral changes occurring at 2.5-µmol/g brain and above. The morphologic changes began in the cerebellum at 1.25-µmol/g brain with the thalamic and caudate lesions occurring at higher doses, 10-µmol/g brain.

In a larger animal model (swine), neurotoxicity following intrathecal administration of either gadodiamide or gadopentetate dimeglumine was examined with concentrations ranging from 0.625 mmol/L to 500 mmol/L.⁴⁹ In the animals that were given GBCA concentrations of 500 mmol/L with more cephalad locations of administration, somatomotor irritation and paresis occurred after 24 hours. No effects were seen at the lower concentrations.

Additional animal studies have displayed no signs of neurotoxicity. Toney $et~al^{46}$ examined neurotoxic potential of intrathecally administered gadopentetate dimeglumine in mice at a dose of 2.5 µmol/g brain and demonstrated no significant acute and subacute physiologic and histologic changes. Although, in some of the animal's neural tissue, abnormalities (gliosis and inflammation) were observed. Jinkins $et~al^{38}$ examined intrathecal gadopentetate dimeglumine in rabbits and did not demonstrate seizures or alterations in behavior. The intrathecal toxicity of gadobenate dimeglumine (linear non-ionic) at dosages of 50 and 80 µmol were also studied in another rabbit model. An Neurologic deficits and behavioral changes were not identified. In addition, the histologic examination of the cerebrum, brainstem and spinal cord did not show any signs of toxicity.

Human studies examining neurotoxicity of GBCA intrathecal administration

Although the use of GBCAs in the intrathecal space is not approved by the FDA and some manufacturers' prescribing information (table 1) warns of the risks of intrathecal administration, its utilization and safety have been suggested based on the reports of administration for intrathecal gadolinium-enhanced magnetic resonance myelography and cisternography. ^{11 50-53} When used in these applications, GBCAs are used in small volumes, often in the range of 0.2–0.5 mL, but not exceeding 1 mL. In addition, the GBCA is diluted either with CSF, iodinated-based contrast agents, or normal saline, and injected slowly (eg, 0.2 mL per second). ^{11 50} The procedure is performed to detect abnormalities in the CNS including CSF fistulas.

Even though some have suggested that the use of small volumes of intrathecal GBCAs is safe, acute gadolinium neurotoxicity from both deliberate and inadvertent intrathecal administration has been documented in humans, with signs symptoms that were similar to this case in clinical presentation and timing. 12 16-21 The details of these cases are summarized in table 1. Specifically, three cases have been reported where GBCAs were used for neuraxial interventional pain procedures. 12 19 21 Two of the cases involved inadvertent intrathecal administration when used for interlaminar epidural placement and one case involved the injection of a GBCA for an intrathecal pump catheter contrast study. Acute neurotoxicity signs and symptoms occurred in these cases with both linear and macrocyclic GBCAs and with approximate intrathecal GBCA concentrations ranging from 1.07 µmol/g brain to 7 µmol/g brain (tables 1 and 3). It should be noted that the 1.07 µmol/g brain concentration is less than half of the concentration (2.5 µmol/g brain) found to be safe in animal models (table 3).

Table 3 Comparison of gadolinium concentrations in previous cases of intrathecal use

Study	GBCA used	Subject	Volume used (mL)	Concentration (µmol/g brain)	Induced neurologic complications (Y/N)
This report	Gadoteridol	Human	5	2.3	Yes*
Arlt et al ¹⁷	Gadopentetate dimeglumine	Human	20	7†	Yes
Li <i>et al</i> ¹⁸	Gadopentetate dimeglumine	Human	15	5.35	Yes
Kapoor <i>et al</i> ¹⁹	Gadodiamide	Human	4 and 4	2.86†	Yes
Park <i>et al</i> ²⁰	Gadopentetate dimeglumine	Human	6	2.14†	Yes
Nayak <i>et al</i> 16	Gadopentetate dimeglumine	Human	10	3.57†	Yes
Reeves et al ²¹	Gadobutrol	Human	2	1.43†	Yes
Popescu <i>et al</i> ¹²	Gadobutrol	Human	1.5	1.07†	Yes
Zeng <i>et al</i> ⁵⁰	Gadopentetate dimeglumine	Human	0.2	0.07†	No
	(A)		0.5	0.17†	No
			1	0.36†	No
Tali <i>et al</i> ⁵¹	Gadopentetate dimeglumine	Human	0.5	0.18†	No
	(A)		0.7	0.25†	No
			0.8	0.29†	No
			1	0.36†	No
Tali <i>et al⁵²</i> Albayram <i>et al⁶⁷</i> Dillon ¹¹ Akbar <i>et al⁶⁸</i> Algin and Turkbey ⁵³	Gadopentetate dimeglumine (A, ^{52 53} B ^{67 68} and C ¹¹)	Human	0.5	0.18†	No
Toney <i>et al</i> ⁴⁶	Gadopentetate dimeglumine	Mice	0.01; NA	2.5	No
Ray <i>et al</i> ¹⁴	Gadopentetate dimeglumine	Mice	0.01; NA	2.5	No
			0.015; NA	3.3	No
			0.02; NA	5	Yes
			0.04; NA	10	Yes
			0.06; NA	15	Yes
Ray <i>et al</i> ⁴⁷	Gadodiamide (D)	Mice	NA	1.25	No
			NA	2.5	Yes
			NA	3.75	Yes
			NA	5	Yes
			NA	10	Yes
	Gadodiamide (D)	Mice	NA	2.5‡	No
			NA	10§	No
	Gadodiamide	Mice	NA	10	Yes
	Gadopentetate dimeglumine	Mice	NA	10	Yes

NA indicates not applicable because the volume relates to animal administration not human.

Note: Only included publications that provided sufficient information to calculate concentration in µmol gadolinium per gram of brain.

GBCA, gadolinium-based contrast agent.

In many cases, GBCA acute neurotoxicity signs and symptoms were present within 1 hour of GBCA administration and included mental status changes, cognitive decline, involuntary movements, hallucinations and seizure activity.17; 18; 21 The rapid onset of neurotoxicity signs and symptoms also correlates with animal studies in which symptoms began within 28–220 min after injection. ¹⁴ In addition, autonomic nervous system and cardiovascular symptoms were seen including hypertension, respiratory distress, tachycardia and fever. Signs and symptoms often resolved within days or weeks although extended sequelae were demonstrated in cases. ¹⁶ ¹⁷ ¹⁹ No cases of death from acute neurotoxicity were published prior to this report.

For the case reported here and in previous case reports where imaging was available, both CT and MRI demonstrated GBCA in the subarachnoid space (table 4). 12 16-21 The expected appearances

of subarachnoid gadolinium in both CT and MRI imaging are shown in table 5, and the examples of this contrast deposition in this case are demonstrated in figures 1 and 2. In both the case reported here and the Kapoor *et al*¹⁹ case, the gadolinium in the CSF was initially diagnosed as a subarachnoid hemorrhage. As demonstrated in table 5, gadolinium and subacute bleeding appear similar in both CT and MRI. Furthermore, gadolinium and acute bleeding appear similar in CT and T2 MRI sequences. If there is concern for inadvertent GBCA administration in the intrathecal space, it is essential that all members of the medical team, including the radiologist, are made aware to assist in early and appropriate detection.

Additionally, human studies demonstrate that intrathecal gadolinium directly deposits in the brain in both white and gray matter and that, with this form of delivery, the reduction in gadolinium

A: Mixed with CSF. B: Mixed with saline. C: Mixed with iohexol. D: Mixed with caldiamide.

^{*}Complications were lethal.

[†]Calculated using 1400 g as the average weight of the human brain in order to be consistent with calculations from Arlt et al.¹⁷

[‡]Delivered over 28 hours.

[§]Delivered over 112 hours.

Table 4 Image findings from previous cases of intrathecal gadolinium neurotoxicity				
Author	Image findings			
Arlt et al ¹⁷	Gadolinium enhancement in the subarachnoid space 1 hour after injection, still present on day 4. None detected on day 56.			
Li et al ¹⁸	After 6 days: FLAIR: enhancement in the CSF cisterns and cerebral sulci. T1: enhancement in the CSF cisterns, cerebral sulci, cerebral cortex and deep parieto-occipital white matter. T2: slight enhancement in the corona radiata and centrum semiovale. DWI: hyperintensity in the right insular cortex, bilateral thalamus, basal ganglia and corona radiata. ADC: hypointense signals in the cerebellum, brainstem, basal ganglia and cerebral cortex. MRA: vasospasm in branches of the middle, anterior, and posterior cerebral arteries. After 8 months: MRI still showed some enhancement; ADC and MRA were normal.			
Kapoor <i>et al</i> ¹⁹	FLAIR/T1-weighted images: hyperintensity in subarachnoid space and ventricles.			
Park <i>et al</i> ²⁰	CT showed diffuse high density in subarachnoid space 6 hours after injection. CT normal after 4 days.			
Nayak et al ¹⁶	MR showed enhancement in the basal cisterns and leptomeninges.			
Reeves et al ²¹	Enhancement visualized in intrathecal space during fluoroscopy.			
Popescu <i>et al</i> ¹²	CT and MRI showed enhancement visible in subarachnoid space same day as injection.			

T1: T1-weighted MRI.

T2: T2-weighted MRI.

ADC, apparent diffusion coefficient map; CSF, cerebral spinal fluid; DWI, Diffusion weighted imaging; FLAIR, fluid-attenuated inversion recovery; MRA, MR angiography.

delivery to CNS tissue by the blood–brain barrier does not exist.³⁹ ⁵⁴ ⁵⁵ In six patients who received intrathecal gadolinium (gadopentetate dimeglumine) without a prior history of administration of intravenous GBCA, Öner *et al*⁵⁵ demonstrated the deposition of gadolinium in the deep nuclei of the brain. The glymphatic system (ie, the waste clearance system for the CNS) may assist in distributing the gadolinium throughout the brain.⁵⁵ ⁵⁶

Besides anecdotal reports of providing steroids and performing CSF drainage, treatment is limited for the acute neurotoxicity associated with GBCAs. The treatments are predominantly supportive. Treatments also include the administration of anticonvulsants when seizures occur.

Our case, in addition to the animal studies and previous human reports, including the three neuraxial interventional pain medicine cases, raises significant concerns regarding the utilization of GBCAs as a radiographic contrast alternative in interventional pain medicine. First, acute GBCA neurotoxicity is significant and can result in long-term sequelae including death. Second, the safe dose limits (2.5-µmol/g brain) suggested by animal studies for intrathecal GBCAs do not appear to directly correlate with humans. ^{14 46} Toxicity in humans has occurred at doses of approximately 1-µmol/g brain and in the fatal case reported here, the patient only received approximately 2.3 µmol/g brain. ¹² Li *et al* ¹⁸ suggested that humans may be less tolerant to intrathecal gadolinium than animals; we are in agreement with this suggestion.

Recently, it was suggested that if a GBCA is used for an epidural procedure that the volume should not exceed 1 mL. When performing epidural steroid injections under fluoroscopic guidance, there is the risk of inadvertent dural puncture (approximately 0.5% with the interlaminar approach). Physicians need to be aware that each GBCA has different pharmacologic properties and molar concentrations that do not allow for simplified volume

limit across all GBCAs. In addition, if physicians want to stay below 1 μ mol/g brain (ie, theoretical number based on case reports suggesting neurologic risk) depending on the molar concentration of gadolinium within the GBCA, this would allow volumes of injection between 1.4 mL (1 mol/L of gadolinium) to 5.6 mL (0.25 mol/L of gadolinium). Furthermore, it must be remembered that the radiographic conspicuity increases with the molar concentration of gadolinium in the GBCA and that in order to make a GBCA provide adequate visualization, either a high molar concentration agent needs to be used at a low volume or a lower molar concentration needs to be used at a higher volume. The small volumes needed to stay at low GBCA concentrations in brain tissues may not be adequate for appropriate visualization.

CONCLUSION

In conclusion, gadolinium neurotoxicity is a serious risk and is often not considered when using a GBCA to assist with visualization in neuraxial interventional pain procedures. The use of GBCAs in the intrathecal and epidural spaces is not FDA approved, and as such, there is no clearly defined dose limit. Physicians must be aware that gadolinium poses a significant risk of acute neurotoxicity, even in very small doses. More research is required regarding the intrathecal use of each brand of gadolinium to determine a toxicity profile prior to consideration for use as radiographic contrast agent for neuraxial interventional pain procedures. We hope that any physician who chooses to use gadolinium off label will consider these risks when making that decision and also appropriately inform patients of the inherent risks of gadolinium use for neuraxial interventional pain procedures. Until further research is performed, we believe the use of gadolinium for neuraxial interventional pain procedures should

Table 5			
	Radiographic image characteristics		

	СТ	MRI T1	MRI T2	MRI FLAIR	MRI DWI
Cerebral spinal fluid	Dark	Dark	Bright	Dark	Dark
Gadolinium	Bright	Bright*	Dark	Bright	Bright
Acute blood/hemorrhage	Bright	Dark	Dark	Dark	N/A
Subacute blood/hemorrhage	Bright	Bright	Dark	Bright	N/A

^{*}The appearance of gadolinium on T1 images is concentration dependent. At lower concentrations, gadolinium is hyperintense (bright), and it becomes isointense then hypointense (dark) as the gadolinium concentration increases. T2 imaging is not concentration dependent.

DWI, diffusion-weighted imaging; FLAIR, fluid-attenuated inversion recovery.

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not be considered a viable and safe alternative when iodinated contrast agents are contraindicated.

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Correction: Fatal gadolinium-induced encephalopathy following accidental intrathecal administration: a case report and a comprehensive evidence-based review

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Under the 'Acute neurotoxicity' paragraph on page 725, the last paragraph, point four, states cellular calcium hemostasis, and should read cellular calcium homeostasis.

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