Intrathecal Use of Gadobutrol for Glymphatic MR Imaging: Prospective Safety Study of 100 Patients

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ABSTRACT

BACKGROUND AND PURPOSE: Intrathecal contrast-enhanced glymphatic MR imaging has shown promise in assessing glymphatic function in patients with dementia. The purpose of this study was to determine the safety profile and feasibility of this new MR imaging technique.

MATERIALS AND METHODS: A prospective safety and feasibility study was performed in 100 consecutive patients (58 women and 42 men, 51 ± 19 years of age) undergoing glymphatic MR imaging from September 2015 to August 2018. Short- and long-term serious and nonserious adverse events were registered clinically and by interview after intrathecal administration of 0.5 mL of gadobutrol (1.0 mmol/mL) along with 3 mL of iodixanol (270 mg I/mL). Adverse events are presented as numbers and percentages.

RESULTS: One serious adverse event (anaphylaxis) occurred in a patient with known allergy to iodine-containing contrast agents (1%). The main nonserious adverse events during the first 1–3 days after contrast injection included severe headache (28%) and severe nausea (34%), though the frequency depended heavily on the diagnosis. After 4 weeks, adverse events had resolved.

CONCLUSIONS: Intrathecal administration of gadobutrol in conjunction with iodixanol for glymphatic MR imaging is safe and feasible. We cannot conclude whether short-duration symptoms such as headache and nausea were caused by gadobutrol, iodixanol, the lumbar puncture, or the diagnosis. The safety profile closely resembles that of iodixanol alone.

ABBREVIATIONS: gMRI = glymphatic MR imaging; iNPH = idiopathic normal pressure hydrocephalus

The glia-lymphatic (or glymphatic) system was described in 2012, providing experimental evidence for direct communication between the CSF of the subarachnoid space and the perivascular spaces of the mouse brain.1 The glymphatic pathways were suggested to be dependent on aquaporin-4 water channels at astrocytic end-feet and are fundamental for the transport and clearance of waste solutes such as amyloid-β and γ protein, which may accumulate in the brain and thereby cause neurodegenerative disease.2 Furthermore, it was suggested that intrathecal contrast-enhanced MR imaging might be used for imaging of the glymphatic circulation in man,3,4 and such studies have recently confirmed communication between the subarachnoid space and the extravascular compartment of all brain regions, thereby extending previous animal data (Fig 1).

The human glymphatic system differs from that in rodents in several ways, first by being much slower.5,6 Moreover, human studies have shown delayed clearance of CSF tracer from all brain regions in individuals with idiopathic normal pressure hydrocephalus (iNPH),5 and particularly from the entorhinal cortex, where volume loss precedes hippocampal atrophy in Alzheimer disease.7 After the discovery of meningeal brain lymphatic vessels in 2015,8,9 observations from human glymphatic MR imaging (gMRI) have suggested a functional link between glymphatic and lymphatic pathways because peak CSF tracer enhancement coincided in time in both the brain and extracranial lymph nodes.10

The risk of gadolinium retention within the human brain after repeat intravenous administrations has been extensively debated.11-13 It has recently been acknowledged that gadolinium-containing MR imaging contrast agents leak from blood to the CSF,14 even in patients without blood-brain barrier dysfunction,15 and probably in substantial amounts through the choroid plexus.16 In principle, an intravenous dosage of contrast agent to

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venous blood therefore also represents a dosage of contrast agent to the CSF. With this background, exploring the safety profile of intrathecal MR imaging contrast agent administration is highly warranted. We performed a prospective safety and feasibility study of 100 individuals undergoing gMRI following intrathecal injection of 0.5 mL of gadobutrol (Gadovist, 1.0 mmol/mL; Bayer Schering Pharma, Berlin, Germany), in conjunction with 3 mL of iodixanol (Visipaque, 270 mg I/mL; GE Healthcare, Piscataway, New Jersey), and recorded immediate, short-term, and long-term adverse events.

MATERIALS AND METHODS

Approvals
The institutional review board (2015/1868), regional ethics committee (2015/96), and the National Medicines Agency (15/04952–7) approved the study. Inclusion was by written and oral informed consent.

Experimental Design and Patients
This observational and prospective study included consecutive patients admitted to the Department of Neurosurgery, Oslo University Hospital, Rikshospitalet, for management of tentative CSF circulation disorders during September 2015 to August 2018 (Table 1).

We used the following exclusion criteria: history of hypersensitivity reactions to contrast agents, history of severe allergy reactions in general, evidence of renal dysfunction, pregnant or breastfeeding women, and age younger than 18 years or older than 80 years.

MR Imaging Protocol
Sagittal T1-weighted spoiled gradient-echo volume scans were obtained using a 3T Ingenia MR imaging scanner (Philips Healthcare, Best, the Netherlands), with equal imaging sequence parameters at all time points. The main imaging parameters were the following: TR = shortest (typically 5.1 ms), TE = shortest (typically 2.3 ms), flip angle = 8°, FOV = 256 × 256 cm, matrix = 256 × 256 pixels (reconstructed to 512 × 512). We sampled 184 overcontiguous slices with 1-mm thickness that were automatically reconstructed to 368 slices with a thickness of 0.5 mm. Each image acquisition lasted 6 minutes 29 seconds.

Intrathecal Administration of Gadobutrol
After precontrast MR imaging (approximately 8 AM), the patient was transported on a mobile table to an adjacent neurosurgery operating room, where an interventional neuroradiologist performed x-ray-guided lumbar puncture. The level that provided the best access to the spinal canal was determined by the neuroradiologist on an individual basis, most typically at level L2/L3 or L3/L4, and, in some cases, at level L4/5. Correct placement of a spinal syringe tip in the subarachnoid space at the lower lumbar level was verified in terms of CSF backflow from the puncture needle (22 ga × 3.5 inches). Subsequently, a mixture of 2–3 mL of 270 mg I/mL of iodixanol (Visipaque) and 0.5 mL of 1.0 mmol/mL gadobutrol (Gadovist) was injected guided by fluoroscopy, to confirm unrestricted distribution of radiopaque contrast agent in the lumbar subarachnoid space. After needle removal, the patients were instructed to rotate around the long axis of the body once before transportation back to the MR imaging suite, to keep the patient in the supine position.

Postcontrast MR Imaging Acquisitions and Image Analysis
Assessment of contrast agent entry at the level of the craniocervical junction (spinal transit time) by means of MR imaging was initiated as soon as possible after intrathecal gadobutrol administration (typically with an approximately 5- to 10-minute delay). Consecutive and identical MR imaging acquisitions covering the cranial compartment and upper neck region with the previously outlined MR imaging protocol parameters were initiated and performed approximately every 10 minutes during the first hour after contrast agent injection. The patients were thereafter instructed to remain supine in bed. Repeat, identical MR imaging acquisitions were then performed approximately every 2 hours after intrathecal gadobutrol administration until afternoon (about 4 PM). Transfer of patients between the neurosurgical department and the MR imaging suite and between the bed and the MR imaging table was performed by the hospital staff to help the
patient remain in the supine position. Beginning in the afternoon, the patients could move freely; new MR imaging acquisitions were repeated after 24 and 48 hours.

The first appearance of any MR imaging contrast agent at the level of the foramen magnum was assessed visually by an experienced neuroradiologist (G.R.) on the sagittal T1-weighted volume scans in the hospital PACS.

**Assessment of Serious and Nonserious Adverse Events**

Serious and nonserious adverse events were recorded systematically by study nurses not otherwise involved in management of patients. Serious adverse events were defined as any untoward medical occurrence that, at any dose, results in the following: death, an immediately life-threatening situation, inpatient hospitalization or prolongation of existing hospitalization, persistent or significant disability or incapacity, or an important medical event that may jeopardize the subject or may require medical intervention to prevent one of the outcomes listed here. Adverse events not having these consequences were considered nonserious.

Assessment of adverse events was performed prospectively during days 1–3 after intrathecal contrast agent administration, after 4 weeks, and finally after about 12 months. The patients were specifically queried for the presence of a defined set of symptoms presenting or being aggravated after intrathecal contrast administration and MR imaging and for symptom duration. These symptoms were chosen according to adverse effects most commonly observed after intrathecal iohexanol: headache (mild/moderate/severe), nausea (mild/moderate/severe), dizziness (mild/moderate/severe), itch, warm feeling, paresthesia, visual problems, cognitive difficulties, muscular spasms, discomfort at the injection site, and tremor. Other symptoms than those specifically requested were listed independent of possible causes. All patients who still reported symptoms at 4 weeks were phoned by the study nurses at a later time, ranging from a few months to up to 12 months, for a final assessment. Report of symptoms at any other contact after 4 weeks was also registered and categorized as “late.”

**RESULTS**

**Patients**

Table 1 presents demographic information about the 100 consecutive patients who were included in the study. The cohort included 58 female and 42 male patients, with a mean of 51 ± 19 years of age. Comorbidity was common.

**Verified Intracranial Distribution of Gadobutrol**

Gadobutrol enhancement in cranial CSF spaces was verified in all individuals (Fig 2). Contrast was still visible intracranially after 24 and 48 hours. The gMRI of 1 patient with iNPH is shown in Fig 3.

The mean time from intrathecal administration of gadobutrol until the first visual detection of the contrast agent at the foramen magnum (spinal transit time) was 20 ± 23 minutes (median, 13 minutes; range, 6–150 minutes). As illustrated in Fig 4, there was some interindividual variation concerning spinal transit time. There were no significant differences in spinal transit time across the patient groups (data not shown). In all patients, scans obtained until 4 PM were assessed visually to ensure the expected magnitude of enhancement in the intracranial CSF spaces, which could be confirmed. Prolonged spinal transit time was, therefore, less likely to be attributed to extradural injection of contrast agent at the lumbar level.

**Serious Adverse Effects**

No patients without a history of prior adverse events experienced serious adverse reactions likely related to intrathecal gadobutrol administration. One woman with established allergy to iodinated radiocontrast agents was, by error, included in the study, even after interaction with the referring doctor, study nurse, the neuroradiologist who managed hospital admittance, and an anesthesiologist. This individual experienced an immediate anaphylactic reaction consisting of skin rash, dyspnea, and a fall in blood pressure, symptoms that were comparable with a previous reaction to iodinated radiocontrast agents (Table 2). Fortunately, after receiving intravenous Ringer-acetate (1000 mL) and surveillance within the intensive care unit for a few hours, no further actions were required. After a few hours, the patient was able to undergo MR imaging, though estimation of spinal transit time was not possible in this individual.

One 80-year-old man with iNPH had a pulmonary embolism, which was attributed to a long train journey and reduced mobility a few days before.

**Nonserious Adverse Events**

The nonserious adverse events within 3 days, at 4 weeks, and at 12 months, respectively, are presented in Table 2.

Nonserious adverse events within the first 1–3 days (delayed) after intrathecal gadobutrol were rather frequent and usually occurred after 10–20 minutes or in the afternoon about 3 PM. Adverse events rarely occurred at days 2 or 3. Twenty-two of 100 individuals reported no symptoms, while 78/100 individuals had symptoms of variable degrees (Table 2). About one-third of individuals experienced the combination of severe headache and nausea. These symptoms were managed conservatively. Whether the high frequency of headache and nausea was caused by the under-
lying disease, the gadobutrol or iohexanol per se, or the spinal puncture could not be determined with certainty.

After 4 weeks, 88 individuals reported no symptoms, while 12 had various symptoms (Table 2). One individual with iNPH reported an itch. In 5 patients with either pineal or arachnoid cysts, 3 had variable degrees of headache, 1 had nausea, and 1 reported dizziness. Three patients reported discomfort at the injection site or in the lumbar region and/or paresthesia, which most likely was related to the lumbar puncture itself.

After 12 months, 1 patient with a pineal cyst had mild headache and dizziness, but these symptoms were the same as she experienced before MR imaging (Table 2).

Table 3 presents nonserious adverse events recorded during days 1–3 according to the diagnosis category. The data show that symptoms like headache, nausea, and dizziness particularly depend on the diagnosis. For example, 15/22 (68%) patients with pineal cysts reported severe headache and nausea, while severe headache and nausea were reported by 2/35 (6%) and 10/35 (29%) patients with iNPH, respectively. Moreover, while no symptoms were reported by 12/35 (34%) patients with iNPH, they were reported by 0/22 (0%) patients with pineal cysts. Notably, the high frequency of symptoms such as headache, nausea, and dizziness reported by patients with pineal gland cysts closely resembles the frequency and character of symptoms they had experienced previously and that represented the main reason for their referral to our hospital in the first place.

**DISCUSSION**

The main observation of this prospective study is that administration of 0.5 mL of gadobutrol (1.0 mmol/mL) is safe in patients with no history of allergic reactions to contrast agents. Nonserious adverse events were rather frequent; however, the occurrence depended heavily on the diagnosis. Among individuals with a pineal cyst, 15/22 (68%) reported severe headache and nausea during days 1–3. We previously reported a high frequency of these symptoms in this patient group in general. We cannot with certainty know whether nonserious adverse events such as headache, nausea, and dizziness after 1–3 days or after 4 weeks were caused by any of the contrast agents or related to the diagnosis itself. Inclusion of other patient categories, such as those with iNPH, most likely would provide other frequencies of reported symptoms.

Because gadobutrol was given in conjunction with iohexanol, to what extent these adverse effects can be attributed to gadobutrol alone remains unanswered.

The study reports prospective data from 100 patients to whom intrathecal gadobutrol was given off-label after special permission from the National Medicines Agency of Norway. Intrathecal doses of MR imaging contrast agents in similar amounts have, however, been used off-label on a clinical basis for years to visualize CSF leakage in individuals with spontaneous intracranial hypotension, as previously reported by others. Intrathecal MR imaging contrast agents were also found to benefit assessment

**FIG 2.** Entry of gadobutrol into the CSF within the intracranial compartment at consecutive MR imaging of a patient with a pineal gland cyst (A) and iNPH (B). Intrathecal administration of 0.5 mL (1.0 mmol/mL) of gadobutrol at the lower lumbar level was preceded by unenhanced, T1-weighted MR imaging (time point zero). After the correct needle position had been verified at fluoroscopy by injection of 3 mL of iohexanol (270 mg I/mL), the patient was transported in the supine position to the MR imaging suite next door, and acquisition of identical, consecutive T1 scans was initiated immediately and performed continuously within the first hour. Typically, the contrast agent had reached the cisterna magna at the first postcontrast scan obtained after approximately 10 minutes (A, thick arrow). In some patients, however, enhancement occurred much later, as in B, where slight enhancement was first depicted at 1 hour (thin arrow). At 6-hour scans, gadobutrol is distributed widely in the CSF of both patients, but less prominently in B, where ventricular reflux can also be noted.
of arachnoid cysts and iNPH, and in diagnosing the cause of otorhinorrhea.

We verified that gadobutrol reached the intracranial CSF compartment in all subjects (except the one with an anaphylactic reaction). The mean spinal transit time was 20 ± 23 minutes (median, 13 minutes; range, 6–150 minutes; Fig 4).

In terms of safety with intrathecal gadolinium injections, the linear contrast agents have been more investigated than the macrocyclic ones. Gadopentetic acid in a dose 0.5–1.0 mmol was well-tolerated in animal studies and was found to have a low risk when given in low doses (0.5–1.0 mmol) to patients. Headache has been the most dominating adverse effect, which also might be related to the spinal puncture. Neurotoxic effects on animal brains were seen when gadopentetic acid was given intraventricularly in a dose of 5.0 μmol/g brain but was not observed when the dose was <3.3-μmol/g brain. Previous studies in patients showed that overdose of gadopentetic acid (6–20 times the normal dose) can be neurotoxic. Hence, 1 patient who, by accident, received 20 mL of gadopentetic acid (7.0 μmol/g brain) developed neurologic deficits (speech problems, visual impairment, fatigue, and psychotic symptoms) lasting 2 weeks, but symptoms had disappeared after 2 months. Moreover, following intrathecal administration of 2.0 mL of gadobutrol, 1 individual reported spastic pain of the lower extremities.

In an adult brain weighing 1400 g, gadobutrol in a dose of 0.5 mmol corresponds to 500-μmol/1400-g brain (0.36-μmol/g brain). Thus, the dose of gadobutrol in this study was on the order of one-tenth of previously appreciated toxic levels. Nevertheless, nonserious adverse events within 1–3 days postcontrast were seen rather frequently, in particular headache and nausea. Intrathecal gadopentetic acid was previously reported to cause headache in 10/36 (27%) patients with iNPH and in 6/20 (30%) patients with arachnoid cysts.

Iodixanol was given intrathecally to verify the correct needle position, which is part of the spinal puncture routine. The safety profile of iodixanol closely resembles that of gadobutrol in combination with iodixanol in this current study. Thus, intrathecal iodixanol caused nonserious adverse events in a comparable proportion of patients as reported here.

While MR imaging contrast agents are approved for intravenous administration only, gadolinium has recently been depicted in CSF samples obtained from individuals even with normal renal function and an intact blood-brain barrier. Given that the concentration of MR imaging contrast agent in CSF exceeded that of blood 4.5 hours after intravenous administration, a typical intravenous dose (0.1 mmol/kg in a 80-kg patient = 8 mmol, half-time in blood is approximately 2 hours) may cause significant passage of contrast agent to the CSF. Except from our previous reports using gMRI, time-dependent quantification of gadobutrol in human CSF after a typical intravenous dose, for

FIG 3. gMRI shown as standardized T1 acquisitions before and 24 and 48 hours after intrathecal gadobutrol in a subject with iNPH, including midsagittal, midaxial, and midcoronal images. In iNPH, gadobutrol clearance from the CSF is delayed compared with reference patients at 24 and 48 hours, respectively. Other typical features of iNPH are early ventricular reflux and subsequent periventricular enhancement of contrast agent, typically most prominent at 24 hours.

FIG 4. The time from intrathecal administration of gadobutrol until the first enhancement of the contrast agent within subarachnoid space of the foramen magnum (spinal transit time) for the 100 patients included in the study. The mean ± SD spinal transit time was 20 ± 23 minutes.
example using T1 maps, has been reported in a case study. Four weeks after 1 single intrathecal administration, gadolinium is not detectable by MR imaging in any brain region, including the basal ganglia. The potential risk inherent in the use of any contrast agent should always be weighed against its potential benefits. After intrathecal contrast-enhanced MR imaging of humans was first proposed, gMRI has been demonstrated to be a promising clinical tool in the assessment of CSF circulation disorders. In particular, delayed lymphatic clearance of gadobutrol, which is anticipated to be cleared through the same pathways as amyloid-β and α, has been proved in a dementia cohort. Contrast enhancement of the entire brain extravascular space is expected to have a large potential for better characterization of disease within the brain and spinal cord but remains yet unexplored.

Limitations
While the present study addressed the safety profile of intrathecal gadobutrol, the major limitation is that we cannot determine with certainty which adverse events solely relate to the MR imaging contrast agent. The nonserious effects we observed might, as well, be caused by the accompanied administration of ioxilan, the spinal puncture, a heavy load of repeat MR imaging, and repeat transport to and from the MR imaging lab.

To what extent the nonserious adverse events are related to the dose of intrathecal gadobutrol remains to be determined. Another study using 0.5 mL of Gd-DTPA or Magnevist (Bayer HealthCare Pharmaceuticals, Wayne, New Jersey), which corresponds to half the dose of gadobutrol used in the present study, seemed to give a better profile concerning nonserious adverse events, though having a high sensitivity in diagnosing the cause of otorhinorrhea. Hence, a lower dose of gadobutrol than used in the present study might give fewer adverse effects. However, because gadobutrol distributes to both the CSF and the entire extravascular compartment, it remains to be determined whether such low doses are sufficient to demonstrate contrast enhancement in deep brain regions, where we have experienced it to be quite marginal. In future studies, lower doses of gadobutrol than used in the present study might give fewer adverse effects. However, because gadobutrol distributes to both the CSF and the entire extravascular compartment, it remains to be determined whether such low doses are sufficient to demonstrate contrast enhancement in deep brain regions, where we have experienced it to be quite marginal. In future studies, lower doses of gadobutrol than used in the present study might give fewer adverse effects. However, because gadobutrol distributes to both the CSF and the entire extravascular compartment, it remains to be determined whether such low doses are sufficient to demonstrate contrast enhancement in deep brain regions, where we have experienced it to be quite marginal.

| Occurrence of adverse events following intrathecal MRI contrast agent administration* | Immediate | Delayed (Days 1–3) | 4 Weeks Late |
|---|---|---|---|
| Serious adverse events | | | |
| Allergy reaction | 1 (1%) | 0 | 0 | 0 |
| Pulmonary embolism | 0 | 1 (1%) | 0 | 0 |
| Nonserious adverse events | | | |
| Headache | | | |
| Mild | 3 (3%) | 1 (1%) | 1 (1%) | |
| Moderate | 13 (13%) | 2 (2%) | 0 | |
| Severe | 28 (28%) | 1 (1%) | 0 | |
| Nausea | | | |
| Mild | 17 (17%) | 0 | 0 | |
| Moderate | 5 (5%) | 0 | 0 | |
| Severe | 34 (34%) | 1 (1%) | 0 | |
| Dizziness | | | |
| Mild | 10 (10%) | 0 | 1 (1%) | |
| Moderate | 11 (11%) | 1 (1%) | 0 | |
| Severe | 17 (17%) | 0 | 0 | |
| Itch | | | |
| Mild | 3 (3%) | 1 (1%) | 0 | |
| Warm feeling | 7 (7%) | 1 (1%) | 0 | |
| Paresthesia | 12 (12%) | 2 (2%) | 0 | |
| Vision problems | 0 | 0 | 0 | |
| Cognitive difficulties | 0 | 0 | 0 | |
| Discomfort at injection site and/or lumbar region | 13 (13%) | 2 (2%) | 0 | |
| Tremor | 0 | 0 | 0 | |

* Data are No. of individuals (percentage in parentheses).

Table 3: Occurrence of nonserious adverse events within days 1–3, depending on diagnosis*

| Nonserious Adverse Events | iNPH (n = 35) | SIH (n = 14) | AC (n = 18) | PC (n = 22) | IIH (n = 6) | cHC (n = 5) |
|---|---|---|---|---|---|---|
| Headache | | | | | | |
| Mild | 1 (3%) | 2 (14%) | 0 | 0 | 0 | 0 |
| Moderate | 2 (6%) | 3 (21%) | 2 (11%) | 4 (18%) | 0 | 2 (40%) |
| Severe | 2 (6%) | 2 (14%) | 6 (33%) | 15 (68%) | 1 (17%) | 2 (40%) |
| Nausea | | | | | | |
| Mild | 6 (17%) | 2 (14%) | 4 (22%) | 4 (18%) | 0 | 2 (40%) |
| Moderate | 1 (3%) | 1 (7%) | 1 (6%) | 1 (5%) | 1 (17%) | 0 |
| Severe | 10 (29%) | 2 (14%) | 5 (28%) | 15 (68%) | 1 (17%) | 1 (20%) |
| Dizziness | | | | | | |
| Mild | 4 (11%) | 2 (14%) | 2 (11%) | 2 (9%) | 0 | 0 |
| Moderate | 1 (3%) | 1 (7%) | 3 (17%) | 6 (27%) | 0 | 0 |
| Severe | 0 | 0 | 5 (28%) | 9 (41%) | 2 (33%) | 1 (20%) |
| Itch | | | | | | |
| Mild | 0 | 0 | 1 (6%) | 0 | 0 | 1 (20%) |
| Warm feeling | 3 (9%) | 1 (7%) | 1 (6%) | 2 (9%) | 0 | 0 |
| Paresthesia | 1 (3%) | 2 (14%) | 3 (17%) | 6 (27%) | 0 | 0 |
| Vision problems | 0 | 0 | 0 | 0 | 0 | 0 |
| Cognitive difficulties | 0 | 0 | 0 | 0 | 0 | 0 |
| Discomfort at injection site and/or lumbar region | 0 | 2 (14%) | 2 (11%) | 7 (32%) | 2 (33%) | 1 (20%) |
| Tremor | 0 | 0 | 0 | 0 | 0 | 0 |
| No adverse events | 12 (34%) | 3 (21%) | 4 (22%) | 0 | 2 (33%) | 1 (20%) |

Note: SIH indicates spontaneous intracranial hypotension; AC, arachnoid cyst; PC, pineal cyst; IIH, idiopathic intracranial hypertension; cHC, communicating hydrocephalus.

* Data are presented as No. of individuals (percentage in parentheses).
ture studies, the effects of lower doses of gadobutrol should be explored. Additionally, other gadolinium-based contrast agents might be examined.

Another limitation is that the study included a high proportion of individuals with symptoms similar those reported after intrathecal gadobutrol. The patient selection most likely affected the symptom profile we also registered at time points after intrathecal injections.

CONCLUSIONS

This study showed intrathecal administration of gadobutrol for gMRI to be safe and feasible. We cannot, with certainty, conclude whether nonserious adverse effects such as headache and nausea were caused by gadobutrol, iohexanol, the lumbar puncture, or the diagnosis.

Disclosures: Svein Are Sirirud Vatnehol—UNRELATED: Payment for Lectures Including Service on Speakers Bureaus: Norwegian Medical Association, Comments: MRI safety lecture for radiologists in training.

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