

59 power in the left upper and lower limbs, increased tone,
 60 hyperreflexia, and a positive Babinski sign on the left. No
 61 cranial nerve deficits were documented. Track marks were
 62 present on both forearms, supporting a history of intrave-
 63 nous drug use. No other cutaneous signs of infection were
 64 identified.

65 Magnetic resonance imaging of the brain demonstrated
 66 a 4.4 × 4.3 × 5.1 cm ring-enhancing mass centred in the
 67 right basal ganglia, with extension into the right thalamus
 68 and temporal lobe and mild leftward midline shift
 69 (Figure 1A). There was no radiologic evidence of sinus
 70 disease or extracranial extension. The patient underwent
 71 stereotactic-guided mini-craniotomy and corticectomy for
 72 tissue diagnosis and conservative debulking. Frozen section
 73 examination showed necrotic brain tissue with fungal
 74 hyphae, raising concern for a fungal abscess. Permanent
 75 histopathology demonstrated glial tissue with suppurative
 76 necrosis and numerous fungal spores and hyphae. Periodic
 77 acid-Schiff and Gomori methenamine silver stains highlighted
 78 abundant fungal elements. The formal report described a
 79 necroinflammatory process consistent with a fungal abscess
 80 and found no evidence of malignancy. Species-level
 81 identification was not available. However, the Gomori
 82 methenamine silver (GMS)-stained sections were retrospectively
 83 reviewed by the treating clinical team, in conjunction with
 84 the consultant pathologist, and demonstrated broad pauci-septate
 85 hyphae with

irregular right-angle branching, supporting a diagnosis of
 mucormycosis in the appropriate clinico-radiologic context
 (Figure 2).

Fungal cultures were performed on biopsy material;
 however, growth was negative, and polymerase chain
 reaction-based molecular identification was not pursued
 at our institution at the time of diagnosis. The diagnosis
 therefore rested on the combination of characteristic
 histomorphology on GMS stain and the supporting clinico-
 radiologic context. Serum beta-D-glucan was negative
 (<31 pg/ml), a finding that does not exclude mucormy-
 cosis because Mucorales are typically not detected by
 this assay. Additional workup, including human immu-
 nodeficiency virus serology, fasting glucose, and glycated
 hemoglobin, were unremarkable. Computed tomography
 of the sinuses, chest, abdomen, and pelvis did not reveal
 an extracranial source of infection.

Following biopsy and limited debulking, antifungal
 therapy was started with liposomal amphotericin B at 5
 mg/kg intravenously once daily and isavuconazole at a
 loading dose of 200 mg intravenously every 8 hours for 6
 doses, followed by 200 mg daily. Based on the pathology
 findings and the clinico-radiologic picture, the lesion was
 managed as presumed proven invasive cerebral mucormy-
 cosis according to EORTC/MSGERC Guidelines [6].
 Because of the deep basal ganglia location and predicted
 surgical morbidity, complete excision was not considered

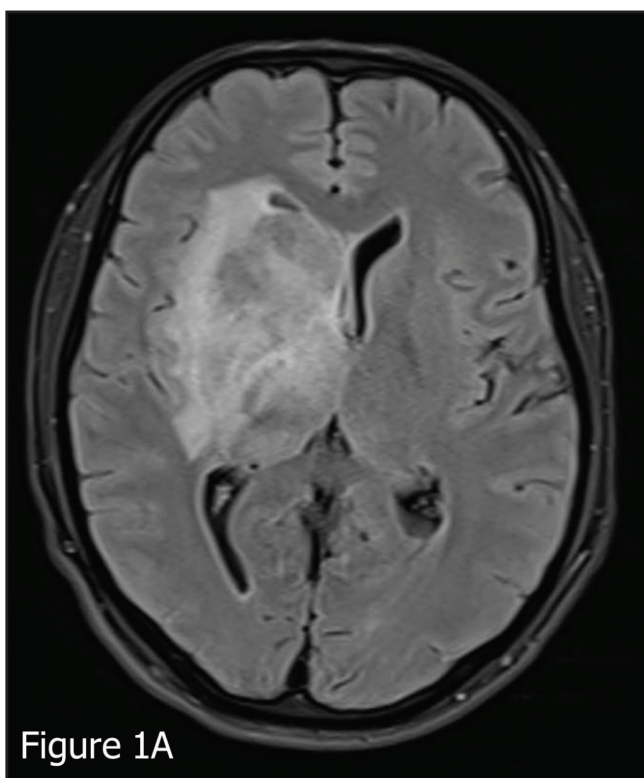


Figure 1A

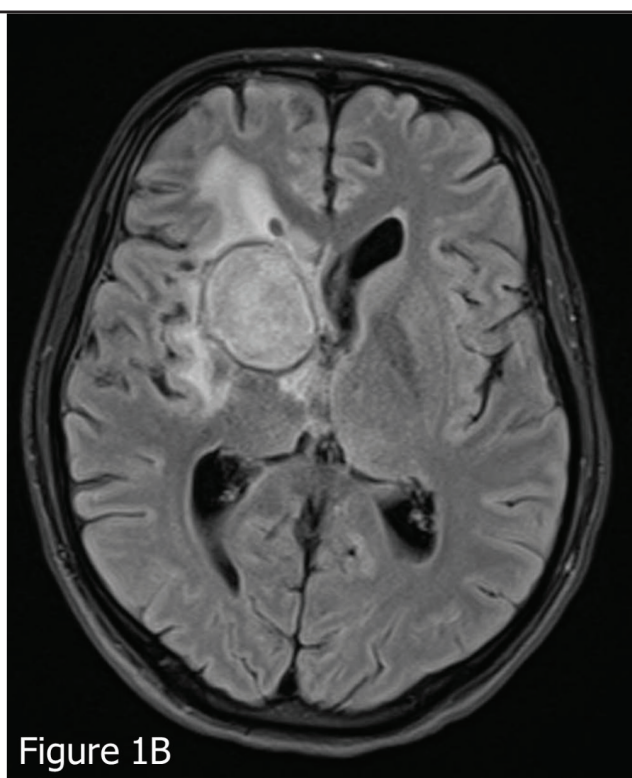
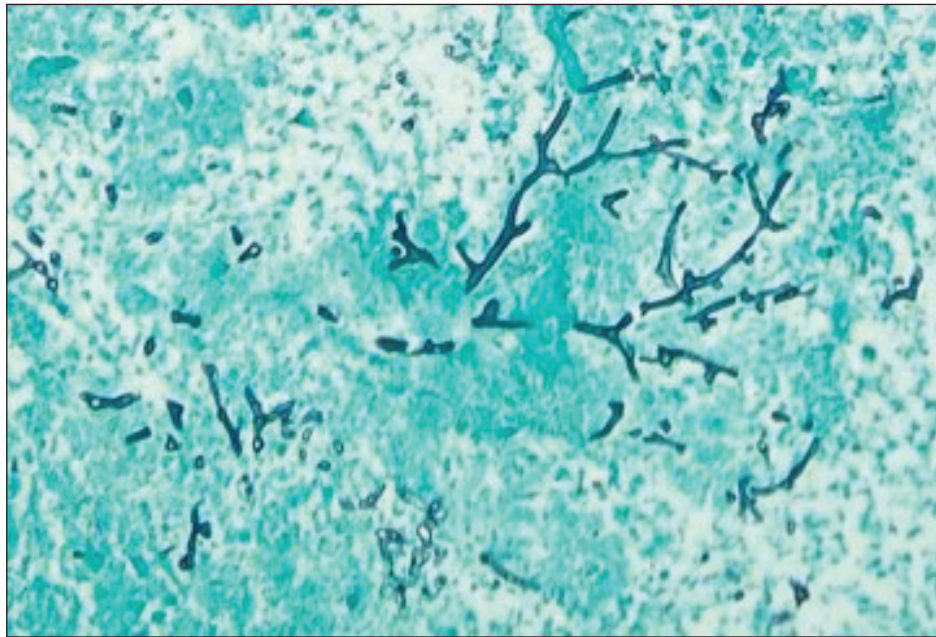


Figure 1B

Figure 1. A: T1-weighted axial view of MRI demonstrating an enhancing right basal ganglia mass measuring 4.4 × 4.3 × 5.1 cm, extending into the right thalamus and right temporal lobe, with mild leftward midline shift. B: T1-weighted axial view of MRI at 16 weeks post-treatment showing reduction in the right basal ganglia mass to (4.0 × 2.9 × 3.0 cm), with decreased peri-lesional edema and improved midline shift.



118
 119 **Figure 2.** Gomori methenamine silver (GMS) stain showing broad, aseptate hyphae with irregular, right-angle branching invading brain
 120 parenchyma (as shown in black arrows)—features consistent with Mucorales species.

121 feasible, and further immediate aggressive resection was
 122 deferred. The patient was later transitioned to prolonged
 123 oral isavuconazole, with infectious diseases follow-up
 124 documenting a planned total treatment duration of 18–24
 125 months because of residual deep-seated disease and
 126 incomplete radiologic resolution.

127 Serial imaging showed no significant interval change
 128 at 6 weeks. At 16 weeks, follow-up magnetic resonance
 129 imaging demonstrated a reduction of the lesion to 4.0
 130 × 2.9 × 3.0 cm, with decreased perilesional edema and
 131 resolution of the midline shift (Figure 1B). Clinically,
 132 the patient showed gradual neurologic improvement but
 133 had persistent left-sided weakness, more pronounced in
 134 the upper limb, and later developed a seizure disorder
 135 requiring levetiracetam. During long-term follow-up, his
 136 course was complicated by interruptions in clinic attend-
 137 ance and medication access, with documented periods off
 138 isavuconazole. On a subsequent visit following a docu-
 139 mented period of isavuconazole, the patient presented
 140 with recurrent headache and vomiting. Clinical assess-
 141 ment raised concern for disease relapse, and reinitiation
 142 of dual antifungal therapy was recommended. Repeat
 143 magnetic resonance imaging was planned at that visit, but
 144 could not be obtained before the patient was lost to fol-
 145 low-up. The absence of imaging confirmation means that
 146 relapse remains clinically suspected rather than radiologi-
 147 cally confirmed, which is acknowledged as a limitation.
 148 Amphotericin-related hypokalemia and hypomagnesemia
 149 were documented and managed with monitoring, hydra-
 150 tion, and electrolyte replacement. Subsequent long-term
 151 follow-up was incomplete because the patient stopped
 152 attending the infectious diseases clinic, and the final

outcome after the planned antifungal course could not be
 153 determined. 154

155 Discussion 155

156 Isolated cerebral mucormycosis is an uncommon but 156
 157 well-recognized presentation of invasive Mucorales infec- 157
 158 tion. Unlike the more typical rhino-orbito-cerebral form, 158
 159 isolated brain involvement has been reported mainly in 159
 160 young, otherwise immunocompetent men with a his- 160
 161 tory of intravenous drug use [3-5,7]. In the patient-level 161
 162 meta-analysis by Kerezoudis et al., intravenous drug use 162
 163 was the dominant risk factor, and basal ganglia involve- 163
 164 ment was a recurring pattern [3]. Similar observations have 164
 165 been reported in earlier case series and reviews [4,5,7,8]. 165
 166 Our case closely fits this clinic-radiologic profile. 166

167 The predilection for the basal ganglia is thought to 167
 168 reflect hematogenous dissemination of fungal elements 168
 169 introduced during injection, followed by seeding of highly 169
 170 vascular deep gray matter structures [3-5,7,8]. Although 170
 171 our patient had no diabetes, human immunodeficiency 171
 172 virus infection, or other conventional immunosuppressive 172
 173 risk factors, intravenous drug use likely represented the 173
 174 major predisposing factor. The absence of sinus, pulmo- 174
 175 nary, or other extracranial disease further supported the 175
 176 diagnosis of isolated cerebral infection. 176

177 Diagnosis remains challenging because clinical presen- 177
 178 tation and neuroimaging may mimic a tumor, pyogenic 178
 179 abscess, or other fungal infections. Definitive diagno- 179
 180 sis depends on tissue examination. In this case, biopsy 180
 181 with histopathology was decisive, demonstrating a fun- 181
 182 gal abscess with broad pauci-septate hyphae and irregu- 182
 183 lar right-angle branching on Gomori methenamine silver 183

184	stain, a morphology that strongly supported mucormycosis. Culture and molecular confirmation were unavailable,	239
185	which is a recognized limitation in many reported cases	240
186	[3,9]. When available, polymerase chain reaction and	241
187	other molecular methods may help support the diagnosis	242
188	in culture-negative disease [9].	243
189		244
190	The initial use of liposomal amphotericin B at 5 mg/kg	245
191	per day alongside concurrent isavuconazole represented	246
192	an individualized clinical decision rather than a guide-	247
193	line-mandated combination. The ECMM/MSGERC 2019	248
194	global guideline strongly recommends liposomal amphi-	249
195	tericin B as first-line therapy for cerebral mucormycosis,	250
196	with doses of up to 10 mg/kg per day considered for CNS	251
197	involvement, while isavuconazole is moderately recom-	252
198	ended either as monotherapy or as a step-down agent	253
199	following initial disease control [10]. Upfront combina-	254
200	tion antifungal therapy is only marginally supported in the	255
201	same guideline, with no specific clinical data available for	256
202	the liposomal amphotericin B plus isavuconazole pairing	257
203	at the time of publication [10]. However, a subsequent	258
204	murine model by Gebremariam et al. demonstrated syner-	259
205	gistic activity of this combination against <i>Rhizopus dele-</i>	260
206	<i>mar</i> and <i>Mucor circinelloides</i> , with an overall survival	261
207	of 80% in the combination arm compared with 50% for	262
208	either agent alone [11].	263
209	In the specific context of CNS mold infections where	264
210	the causative pathogen remains uncertain at the time of	265
211	treatment initiation, the rationale for upfront dual ther-	266
212	apy becomes more defensible, though still not guide-	267
213	line recommended. CNS mold infections may be caused	
214	by <i>Aspergillus</i> species, for which voriconazole or isavu-	268
215	conazole constitutes first-line therapy, or by Mucorales,	269
216	for which liposomal amphotericin B or isavuconazole are	270
217	preferred [12,13]. The combination of both agents, there-	271
218	fore, provided broad empiric coverage for both diagnostic	272
219	possibilities while awaiting further microbiologic results	273
220	[13]. Furthermore, the ECMM/MSGERC mucormycosis	274
221	guideline acknowledges that in cases of extensive disease,	275
222	rapid progression, or poor general condition, the addition	276
223	of isavuconazole or posaconazole to liposomal amphi-	277
224	tericin B can be considered [10]. From a pharmacokinetic	278
225	standpoint, liposomal amphotericin B achieves therapeutic	279
226	concentrations in brain parenchyma despite limited	280
227	cerebrospinal fluid distribution and is recommended at	281
228	higher doses for CNS mucormycosis [10,14], while isavu-	
229	conazole has intermediate CNS penetration and is FDA-	
230	approved for both aspergillosis and mucormycosis [13].	
231	The combination therefore ensured parenchymal antifun-	
232	gal coverage via liposomal amphotericin B alongside an	
233	agent with broader CNS distribution.	
234	CNS aspergillosis and mucormycosis carry extremely	
235	high mortality, and the 2016 IDSA aspergillosis guidelines	
236	acknowledge that combination therapy for CNS disease is	
237	initiated by some practitioners, given the mortality asso-	
238	ciated with this form of dissemination, while recognizing	
	the absence of controlled data supporting improved out-	239
	comes [15]. Given the severity of the disease, the deep	240
	lesion location, and the inability to achieve complete sur-	241
	gical clearance, the clinical team chose upfront dual ther-	242
	apy on the basis of this collective evidence and the need	243
	to maximize antifungal coverage in a high-risk, surgically	244
	inaccessible lesion. Early transition to oral isavuconazole	245
	monotherapy for long-term suppression is consistent with	246
	guideline recommendations for step-down therapy fol-	247
	lowing initial disease stabilization [10]. The deep basal	248
	ganglia location, involvement of eloquent structures,	249
	and expected surgical morbidity favored limited debulk-	250
	ing rather than aggressive excision. Although surgical	251
	debridement is generally recommended when feasible,	252
	selected patients with surgically high-risk lesions may still	253
	achieve short-term disease control with biopsy or limited	254
	debulking combined with antifungal therapy [3,16].	255
	The patient demonstrated interval radiologic improve-	256
	ment at 16 weeks and partial neurologic recovery, although	257
	persistent hemiparesis remained. This residual deficit	258
	likely reflects irreversible injury to deep motor pathways.	259
	His later interruptions in therapy and follow-up compli-	260
	cated long-term assessment and prevented confirmation of	261
	the final treatment outcome. Nevertheless, the early clinical	262
	and radiologic response observed supports the value	263
	of early tissue diagnosis, prompt initiation of antifungal	264
	therapy, and close multidisciplinary follow-up, though the	265
	inability to confirm the final outcome due to loss to fol-	266
	low-up remains an important limitation of this report.	267
	Conclusion	268
	Isolated basal ganglia mucormycosis is a rare but serious	269
	diagnostic consideration in patients with basal ganglia	270
	lesions and a history of intravenous drug use, even in the	271
	absence of traditional immunosuppressive risk factors. It	272
	must be acknowledged that the final treatment outcome in	273
	this case could not be established because the patient was	274
	lost to long-term follow-up, which is a recognized limita-	275
	tion of this report. Nevertheless, the early clinical and	276
	radiologic response observed supports the key educational	277
	messages: tissue diagnosis is essential, and early antifun-	278
	gal therapy with multidisciplinary surgical decision mak-	279
	ing is critical when lesions are deep and surgically high	280
	risk.	281
	What is new?	282
	This case adds to the small published literature on isolated	283
	cerebral mucormycosis in people who inject drugs by doc-	284
	umenting a rare combination of features: an immunocom-	285
	petent host, isolated basal ganglia involvement without any	286
	extracranial source, negative serum beta-D-glucan (highlight-	287
	ing the known insensitivity of this marker for Mucorales), and	288
	partial radiologic and neurologic recovery despite incom-	289
	plete surgical resection. It further illustrates that prolonged	290
	oral isavuconazole represents a clinically viable step-down	291
	and maintenance strategy when complete resection is not	292

293 feasible, and that treatment interruption carries a real risk of
294 relapse even after initial radiologic improvement. The case
295 also demonstrates the diagnostic value of careful morpho-
296 logic review of Gomori methenamine silver-stained sections
297 when culture and molecular confirmation are unavailable.

298 **List of abbreviations**

299	BDG	Beta-D-glucan
300	CNS	Central nervous system
301	CT	Computed tomography
302	GMS	Gomori methenamine silver
303	HbA1c	Glycated hemoglobin
304	HIV	Human immunodeficiency virus
305	IVDU	Intravenous drug use
306	MRI	Magnetic resonance imaging
307	PAS	Periodic acid-Schiff
308	PCR	Polymerase chain reaction

309 **Conflict of Interests**

310 The authors declare that there is no conflict of interest regard-
311 ing the publication of this article.

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314 **Consent for publication**

315 Written informed consent was obtained from the patient for
316 publication of this case report and the accompanying images.

317 **Ethical Approval**

318 Ethical approval is not required at our institution to publish an
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412 <https://doi.org/10.1086/590004> **413**

Summary of the case

NO.	ITEM	DETAILS
1	Patient (gender, age)	30-year-old male
2	Final diagnosis	Isolated basal ganglia cerebral mucormycosis
3	Symptoms	New-onset generalized seizures, altered mental status, and left-sided hemiplegia
4	Medications	Liposomal amphotericin B, isavuconazole, levetiracetam
5	Clinical procedure	Stereotactic-guided mini-craniotomy, biopsy, and conservative debulking