

Brain Biopsy for Neurological Diseases of Unknown Etiology in Critically Ill Patients

Bertrand Mathon, Malory Favreau, Vincent Degos, Aymeric Amelot, Alexandre Le Joncour, Nicolas Weiss, Benjamin Rohaut, Loïc Le Guennec, Anne-Laure Boch, Alexandre Carpentier, et al.

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79 ABSTRACT

Objectives: Brain biopsy is a useful surgical procedure in the management of patients with suspected neoplastic lesions. Its role in neurological diseases of unknown etiology remains controversial, especially in ICU patients. This study was undertaken to determine the feasibility, safety and the diagnostic yield of brain biopsy in critically ill patients with neurological diseases of unknown etiology. We also aimed to compare these endpoints to those of non-ICU patients who underwent a brain biopsy in the same clinical context.

86 **Design:** Monocenter, retrospective, observational cohort study.

87 **Setting:** A French tertiary center.

Patients: All adult patients with neurological diseases of unknown etiology under mechanical
 ventilation undergoing in-ICU brain biopsy between January 2008 and October 2020 were
 compared to a cohort of non-ICU patients.

91 Interventions: None.

92 **Measurements and Main Results:** Among the 2,207 brain-biopsied patients during the study 93 period, 234 biopsies were performed for neurological diseases of unknown etiology, including 94 29 who were mechanically ventilated and 205 who were not ICU patients. Specific histological 95 diagnosis and final diagnosis rates were 62.1% and 75.9%, respectively, leading to therapeutic 96 management modification in 62.1% of cases. Meningitis on prebiopsy CSF analysis was the 97 sole predictor of obtaining a final diagnosis (2.3 [1.4-3.8]; p=0.02). ICU patients who 98 experienced therapeutic management modification after the biopsy had longer survival 99 (p=0.03). The grade 1 to 4 (mild to severe) complication rates were: 24.1%, 3.5%, 0% and 100 6.9%, respectively. Biopsy-related mortality was significantly higher in ICU patients compared 101 to non-ICU patients (6.9% vs. 0%, p=0.02). Hematological malignancy was associated with 102 biopsy-related mortality (1.5 [1.01-2.6]; p=0.04).

103 **Conclusions:** Brain biopsy in critically ill patients with neurological disease of unknown 104 etiology is associated with high diagnostic yield, therapeutic modifications and postbiopsy 105 survival advantage. Safety profile seems acceptable in most patients. The benefit/risk ratio of 106 brain biopsy in this population should be carefully weighted.

107	KEYWORDS
108	Coma
109	Mechanical ventilation
110	Intensive care unit
111	Cryptogenic neurological diseases
112	Diagnostic workup
113	Biopsy
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135 **INTRODUCTION**

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137 Brain biopsy is a surgical procedure used to obtain histopathological diagnosis and 138 guide the management of patients with suspected neoplastic lesions, for which its diagnostic 139 yield exceeds 95%(1, 2). As an invasive procedure associated with potentially severe 140 complications, its role in the diagnosis of nonneoplastic neurological diseases remains 141 controversial. The reported diagnostic yield for this indication was low in the before year 2000, 142 ranging from 20 to 30%(3, 4) and was associated with high frequency of complications(5). 143 Recent evidence in both adults and children reported better results (68%-83%) (6-9), leading 144 to reappraise of the role of brain biopsy in the diagnosis and therapeutic algorithm of patients 145 with neurological diseases of unknown etiology(10).

In critically-ill patients, invasive procedures are associated with a higher risk of
complication, especially in the setting of altered hemostasis(11). To the best of our knowledge,
no study specifically addressed the role of brain biopsy in neurological diseases of unknown
etiology in this population.

We conducted a retrospective monocenter study to investigate brain biopsy feasibility, diagnostic yield, and safety in critically ill adults with neurological diseases of unknown etiology. We also aimed to compare these endpoints to those of non-ICU patients who underwent a brain biopsy in the same clinical context. This study therefore explores the critical care population and updates our previously published cohort of non-ICU patients (6).

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157 MATERIALS AND METHODS

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159 **Patients**

We retrospectively reviewed the medical records and histology reports of all adults
brain-biopsied at our tertiary medical center, between January 2008 and October 2020.
Patients meeting the following criteria were included: 1) patients in ICU under mechanical

ventilation; 2) neurological disease of unknown etiology or atypical cerebral evolution of systemic and/or neurological underlying diseases; 3) negative comprehensive less-invasive diagnostic work-up including physical examination, laboratory tests including cerebrospinal fluid (CSF) examination obtained by lumbar puncture, radiological examinations and extraneurological histological findings; and 4) indication for brain biopsy validated by a multidisciplinary team.

Patients were not included if the brain biopsy had been obtained before ICU admission or after discharge. Patients who underwent brain biopsy for histological confirmation of an obvious primary or secondary cerebral neoplasm, or brain abscess were not included.

We then compared variables regarding diagnostic yield and safety between the ICU patients included in this study and a cohort of non-ICU patients with neurological diseases of unknown etiology who underwent a brain biopsy during the study period at our institution. The latter met the above-mentioned criteria 2,3 and 4.

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177 Study variables and outcomes

178 The primary endpoint was the frequency of obtaining a specific histological diagnosis. 179 Secondary endpoints were frequency of obtaining a final diagnosis, the occurrence of any brain 180 biopsy-related complications, and postbiopsy survival. Histological results of brain biopsies 181 were categorized into 3 groups: specific lesion, nonspecific lesion, normal brain. Obtaining a 182 specific histological diagnosis was defined as brain biopsy findings of a specific lesion sufficient 183 by itself to make a diagnosis. The final diagnosis was reached by combining the brain biopsy 184 findings integrated with the patient's medical history and the results of the less-invasive 185 diagnostic work-up. Brain biopsies containing specific lesion(s) were classified as contributory 186 to the final diagnosis. Brain biopsies with nonspecific lesion(s) could nonetheless be classified 187 as contributing to a final diagnosis. A multidisciplinary discussion determined whether a brain 188 biopsy with nonspecific lesion(s) contributed to a final diagnosis. During these discussions, 189 participants systematically and comprehensively reviewed each patient's medical history, 190 neurological and extra-neurological findings, less-invasive diagnostic work-up, brain biopsy

191 microbiology and histology results. The treating physician's main hypothetical diagnosis and 192 treatment at the time of biopsy and changes made thereafter were noted. Two senior 193 neuroradiologists analyzed all the imaging studies, including available 3.0 Tesla magnetic 194 resonance imaging (MRI) sequences and multiparametric imaging data. Two senior 195 neuropathologists examined all histological slides. During the multidisciplinary discussion, 196 participants had to agree unanimously that the brain biopsy contributed to making the final 197 diagnosis.

198 Complications related to brain biopsy were monitored during the 30 days following the 199 intervention. In view of existing literature on complications in neurosurgery, we used a 200 previously published grading severity scale tailored for diagnostic intracerebral procedures (6, 201 12):

202 - grade 1: complication visible only on postoperative computed-tomography (CT) scan
 203 (asymptomatic hemorrhage) or transient event that did not require treatment;

- grade 2: transient complication that resolved completely but required treatment;

205 - grade 3: persistent neurological deficit >6 months postbiopsy;

206 - grade 4: biopsy-related death.

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208 Surgical methodology and neuropathological protocol

209 The biopsies were taken under general anesthesia. A stereotactic biopsy technique 210 was used for deep-seated lesions with patients positioned in a Leksell stereotactic frame. An 211 enhanced CT scan or 3D gadolinium-enhanced and FLAIR sequences 1.5 Tesla MRI were 212 performed. When a stereotactic CT was performed, the images were merged with those of the 213 reference MRI. Once these images were acquired, the trajectory and depth were planned 214 according to the lesion to be targeted. Stereotactic coordinates were calculated with Framelink 215 (Medtronic, Minneapolis, MN) software. The biopsy procedures were then performed as 216 previously described (6). We collected up to 10 tissue samples, $\sim 1 \times 10 \text{ mm}$ (2).

217 For cortical and/or meningeal lesions, biopsies were obtained via open craniotomy or 218 a burr hole. We considered a gold standard diagnostic open biopsy to be 1 cm3 of 219 leptomeninges and cortex including grey and white matter. For MRI-negative patients, the 220 biopsy was preferentially taken from the right middle frontal lobe gyrus, unless history, 221 examination or imaging asymmetry suggested another location would provide a higher 222 diagnostic yield.

223 Postoperative CT scan was then performed immediately after the end of biopsy to rule 224 out complications, before transfer to the ICU (12, 13).

The tissue samples collected were divided into several parts for neuropathological, bacteriological, parasitological and virological investigations. When the differential diagnosis included infection, tissue was set aside for microbiology studies. The management of samples in the neuropathology lab relied on the previously described protocol (6). Since 2016, in case of negativity of the first and second-line panels, the remaining samples were used for metagenomic next-generation sequencing (NGS) analysis in patients with encephalitis (14, 15).

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233 Statistical analyses

234 Results for categorical variables, expressed as number (%), were compared with χ^2 235 tests; those for continuous variables, expressed as mean ± standard deviation or median 236 [25th-75th percentile interguartile range (IQR)], were compared using Student's t-test or 237 Wilcoxon's rank test. Normality of continuous variable distribution was assessed with the 238 Shapiro-Wilk test and nonnormally distributed continuous variables were compared using Wilcoxon's rank test. Patients' demographic, clinical and biological characteristics were tested 239 240 in univariable analyses for association with the primary and secondary endpoints. We 241 compared variables regarding diagnostic yield and safety between ICU and non-ICU patients 242 using appropriated tests. Survival between groups were analyzed with the log-rank test. 243 P<0.05 defined statistical significance. Analyses were computed with IBM SPSS Statistics 244 v22.0 software (IBM Corp, Armonk, NY).

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247 Standard Protocol Approvals, Registrations, and Patient Consents

In accordance with the ethical standards of our hospital's institutional review board (N°2214386 - CNIL), the Committee for the Protection of Human Subjects, informed consent was not obtained for demographic, physiologic, and hospital-outcome data analyses because this observational study did not modify existing diagnostic or therapeutic strategies. The manuscript was prepared in accordance with the STrengthening the Reporting of Observational studies in Epidemiology (STROBE) statement.

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- 255

256 **RESULTS**

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258 Study population and characteristics

259 During the study period, 2,207 patients underwent a brain biopsy, of which 234 (10.6%) 260 were performed to investigate a neurological disease of unknown etiology. Twenty-nine were 261 critically ill and 205 were non-ICU patients. The study flowchart is reported in Figure 1. The 262 general characteristics of the study patients and their brain biopsies are reported in 263 Supplemental Digital Content 1 (Table). The male-to-female ratio was 2.6 and the mean age 264 on biopsy-day was 49.4±15.4 years. Clinical manifestations included altered consciousness 265 (100.0%), neurological deficit (55.2%), extra-neurological symptoms (37.9%) and seizures 266 (27.6%). Elevated CSF proteins and meningitis were reported in 70.4% and 44.4%, respectively. Most patients had multifocal (69%), bilateral (69%) or gadolinium-enhanced 267 (58.6%) lesions. The biopsy-targeted lesion was exclusively supratentorial. One patient had 268 269 no lesion on MRI. The most frequent biopsy technique was stereotaxic (65.5%), with MRI-270 guidance (57.9%). Patient's clinical characteristics and organ failures on ICU admission-day 271 and brain biopsy-day are reported in **Table 1**. Patients were mainly admitted in ICU for coma 272 (79.3%) or status epilepticus (17.2%). Brain biopsy-day organ failure supports were 273 mechanical ventilation 100%, renal replacement therapy 17.2%, vasopressors 10.3%, while

274 no patient was under extracorporeal membrane oxygenation. The median pre-biopsy SOFA

score was 4 [4-6]. The median ICU-admission-to-biopsy interval was 11 [6-19] days.

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277 Diagnoses and diagnostic yield-associated factors

278 Brain biopsy analysis showed a specific lesion, nonspecific lesion or normal brain, in 279 18 (62.1%), 10 (34.5%) and 1 (3.4%) patients, respectively. A final diagnosis could be made 280 in 22 (75.9%) patients, with most common diagnoses including infection (44.8%), autoimmune 281 or inflammatory disease (13.8%), malignancy (13.8%) and demyelinating disease (6.9%) 282 (Supplemental Digital Content 2 - Table). One patient had multiple diagnoses(16). Of note, 283 diagnostic yield did not differ significantly between ICU patients and non-ICU patients (75.9% 284 vs. 74.1%, respectively, p=0.8). Comparisons between ICU patients according to contribution 285 of the biopsy to the final diagnosis are presented in Table 2. The univariate analysis retained 286 only the meningitis on pre-biopsy cerebrospinal fluid analysis as being a predictor of obtaining 287 a final diagnosis (odds ratio (OR) [95% confidence interval (CI)], 2.3 [1.4-3.8]; p=0.02).

288

289 Complications and factors associated with them

290 During the month following the biopsy, 10 (34.5%) patients developed a complication 291 (Supplemental Digital Content 3 - Figure). Seven (70%) were grade-1 asymptomatic and 292 diagnosed on systematic post biopsy CT scan. Nine complications (90%) were biopsy site 293 hemorrhages, none leading to surgical evacuation, and one was brain edema requiring 294 corticosteroid administration (grade 2). Two biopsy site delayed hemorrhages were fatal (grade 295 4): one in a patient with acute myeloid leukemia and persistent severe thrombopenia (30 G/L) 296 20 day after the biopsy, and another in a patient with multiple myeloma and hemodialysis on 297 day 3 postbiopsy. Rates of overall complications and mortality were significantly higher in ICU 298 patients compared to non-ICU patients: 34.5% vs. 17.6%, p=0.03 and 6.9% vs. 0%, p=0.02, 299 respectively. In the ICU patient group, no variable was associated with the occurrence of 300 postbiopsy complication, while history of hematological malignancies was significantly 301 associated with biopsy-related mortality (OR 1.5 [1.01-2.6]; p=0.04).

302 Postbiopsy outcomes

303 Brain biopsy findings led to a therapeutic modification in 62.1% of the ICU-patients; 304 significantly less than in non-ICU patients (79.1%, p=0.04). Twelve patients (41.4%) died in 305 the ICU and a total of 14 (48.3%) within the first year postbiopsy (Fig. 2A). The univariate 306 analysis retained low prebiopsy hemoglobin rate (p=0.01), high SOFA score on biopsy day 307 (p=0.04) and history of hematological malignancies (p=0.02) as being associated with in-ICU 308 mortality. In-ICU mortality was significantly lower in patients in whom the biopsy had led to 309 therapeutic changes (22.2% vs. 63.6%, 0.2 [0.03-0.9]; p=0.048). Obtaining of a final diagnosis 310 was not significantly associated with overall survival ($p_{Log-Rank} = 0.39$, Fig. 2B). Patients with 311 therapeutic management modification after biopsy had a higher probability of survival (72.2% 312 vs. 27.2% at 1-year postbiopsy, $p_{Log-Rank}$ =0.03, Fig. 2C).

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315 **DISCUSSION**

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Most neurological diseases in ICU patients do not require brain biopsy for their diagnosis and management. Nevertheless, in some patients with neurological disease of unknown etiology, obtaining a pathological brain sample can be decisive. To the best of our knowledge, we report the first series on the safety and diagnostic yield of brain biopsy in critically ill patients.

322 Owing the retrospective nature of this study, to maximize the identification of brain 323 biopsy-related complications, we used a severity grading scale that also took into account 324 silent hemorrhagic complications. Our rate of postbiopsy asymptomatic hemorrhages (24.1%) 325 is consistent with the rates previously reported in non-ICU patients (7-67%)(17). However, the 326 overall complication and mortality rates (34.5% and 6.9%, respectively) were higher in ICU 327 patients. Hematologic malignancies were the only factor associated with biopsy-related 328 mortality in our series despite these patients had normal hemostasis parameters value on 329 biopsy-day. Indeed, pre-biopsy platelet transfusions do not prevent the risk of delayed biopsy330 site hemorrhage. One patient had a very delayed (up to 20 days postbiopsy) hemorrhagic 331 complication while having severe thrombopenia (30 G/L). Under exceptional circumstances, 332 we thereby think that the patient's platelet count should be maintained over >100 G/L for at 333 least 3 weeks after the biopsy. For ICU patients with hematologic malignancies who are 334 candidates for brain biopsy, the benefit/risk ratio must be therefore carefully weighted. We 335 demonstrated that multiple organ dysfunction or failure do not impede the conduction and high 336 diagnostic yield of brain biopsy. However, based on our own experience, patients on ECMO 337 support are not good candidates for brain biopsy as these devices are associated with 338 profound hemostatic disturbances(18, 19), and therefore were excluded from being considered 339 for brain biopsy.

340 The rate of final diagnosis established with brain biopsy in ICU patients was high in our 341 series (75.9%) and comparable to that obtained in our control group of non-ICU patients 342 (74.9%) and even with recent studies published in non-ICU patients(6-8). Furthermore, 343 although 24.1% of the biopsies were non-contributory for a diagnosis, they excluded infectious 344 diseases or malignancies, thereby enabling therapeutic management to be adapted 345 accordingly(20-23). Since the mid-2010's, the progress of metagenomic next generation 346 sequencing on brain samples has enabled diagnoses that could not be achieved with usual 347 microbiological analyses. In our study, metatranscriptomics identified sequences of viral 348 infections in brain tissues from 3 immunocompromised patients with clinical and pathological 349 signs of encephalitis. The 3 identified pathogens were measles(24), rubella and a novel 350 zoonotic virus called umbre orthobunyavirus(25). Nonetheless, we did not significantly improve 351 our rate of positive biopsies since the introduction of metagenomics (76.5% vs. 75%, 352 respectively), because our growing expertise probably led to retain wider indication of brain 353 biopsy in challenging cases. A systematic literature review compiled 22 patients with 354 encephalitis in which a next generation sequencing analysis on brain tissue provided a 355 previously unsuspected diagnosis(14). The authors reported a diagnostic yield of brain tissue 356 analysis of 50% versus only 20% for CSF. The vast majority of the positive results from brain 357 samples was in immunocompromised patients suggesting that metagenomics may be best

358 applied to a targeted population in whom it will be most rewarding. Introduced in the diagnostic 359 algorithm of encephalitis of unknown etiology, including in ICU patients, this new technique 360 opens perspective for comprehensive and unbiased detection of pathogens and paves the way 361 to further improving in the diagnostic yield of brain biopsy(15, 26). The sole factor associated 362 with obtaining a diagnosis on the brain biopsy was the detection of a meningitis on pre-biopsy 363 CSF analysis. Indeed, brain biopsy was contributory to a final diagnosis in all patients who had 364 a meningitis. This major finding should be borne in mind when evaluating the expected brain 365 biopsy diagnostic yield in a critically ill patient potentially eligible for a brain biopsy. In addition, 366 we confirmed that small or non-contrast-enhanced lesions, and even negative-MRI in 367 immunocompromised patients, were not associated with a low diagnosis rate.

368 Comparing the diagnostic yields in ICU patients, brain biopsy appears to be as effective 369 compared to other solid-organ biopsies. In the literature, percutaneous renal biopsy establish 370 a diagnosis in 69-71% of patients (27, 28), while transvenous renal biopsy obtain a diagnosis 371 in 96% of patients (29). On their side, open lung biopsies contained a specific lesion for 44% 372 of patients in the 2006 study of Kao et al (30), while Philipponnet et al, in 2018, reported a 80% 373 diagnostic-yield (31). The 62%-rate of the rapeutic management changes following brain biopsy 374 observed in our study is in the range with those reported for other organs: 73-78% for open 375 lung biopsies (30, 31) and 21-71% for renal biopsies (27-29). Regarding the safety profile of 376 biopsies, although it is difficult to compare the severity of biopsy-related complications between 377 different organs, brain biopsies do not appear to carry any excessive risks for critically ill 378 patients compared to other biopsies. Thus, previous works reported complications following 379 7.5% to 22% of renal biopsies (27–29) and 20% to 35% of lung biopsies (30, 31). As we report 380 for brain biopsies, fatal complications were observed both in percutaneous renal biopsies (28) 381 and in open lung biopsies (31).

Interestingly, we demonstrated that patients with therapeutic management modification linked to the brain biopsy results had higher probability of survival. Altogether, our results suggest that the contribution of brain biopsy to diagnosis and treatment is undeniable but may be at the cost of complications although most of them were asymptomatic. In that sense, the

risk of postbiopsy serious complications should always be weighed against the risks borne by the natural course of an undiagnosed and untreated acute neurological disease. The latter is more often life-threatening than the former, as supported by the 2 biopsy-attributable deaths versus 14 because of disease-attributed deaths.

390 In the end, it appears fundamental that the indication and feasibility of brain biopsy are 391 evaluated and retained after multidisciplinary discussion between intensivists, neurosurgeons, 392 anesthesiologists and external physicians (neurologists, internal medicine physicians, 393 hematologists...) weighting the benefit-risk balance in every patient. Based on these results 394 and our experience, we propose a decision-making algorithm for the indication and 395 management of brain biopsy in ICU patients with neurological disease of unknown etiology 396 (Fig. 3). This underscores that a number of elements must be needed to consider a brain 397 biopsy in this context. When these criteria are met together, we advocate that this intervention 398 be considered as early as possible in the diagnostic management of these patients.

399 Our study has limitations and strengths. First, it is retrospective, single-center design 400 with a small number of ICU patients, but this is the first series to report on this procedure in 401 critically ill patients. Second, while we compared ICU and non-ICU patients, it was not possible 402 to select relevant matching criteria. Third, we could not provide data on the optimal timing of 403 biopsy as many patients were referred to our institution from distant centers after very variable 404 previous management duration. Last, brain biopsy safety and efficacy in this study relies on 405 the experience of our neurosurgeons, intensivists and neuropathologists, and those results 406 may not be immediately reproducible in every center.

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409 **CONCLUSIONS**

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Brain biopsy in critically ill patients with neurological disease of unknown etiology has
high diagnostic yield and is associated with frequent therapeutic modifications. Safety profile
seems acceptable in most patients, but fatal post biopsy cerebral hemorrhage occurred in two

- 414 patients with hematologic malignancies. The benefit/risk ratio of brain biopsy in this indication
- 415 should be carefully weighted.

418 FIGURE LEGENDS

- **Fig. 1.** Flowchart of patient inclusion in this study on brain biopsy contribution to diagnosis.



- 422 **Fig. 2.** Comparisons of postbiopsy survival between ICU patients and non-ICU patients (**A**).
- 423 Overall survival of the 29 ICU critically ill patients according to the brain biopsy findings (B)
- 424 and the biopsy-related therapeutic management changes (**C**).



425 426

- 427 Fig. 3. Decision-making algorithm for the indication and management of brain biopsy in428 critically ill patients with neurological disease of unknown etiology.
- 429 The red color indicates that brain biopsy is not recommended/contra-indicated in the present430 situation.
- 431 APTT, activated partial thromboplastin time; CSF, cerebrospinal fluid; DTI, direct thrombin
- 432 inhibitors; INR, international normalized ratio; LMWH, low-molecular-weight heparin; UFH,





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Table 1. Clinical characteristics of the 29 ICU critically ill patients with brain					
biopsies.					
Characteristics	Value				
Age, years	49.4 ± 15.4				
Reason for ICU admission					
Disorders of consciousness	23 (79.3%)				
Status epilepticus	5 (17.2%)				
Acute kidney injury	1 (3.4%)				
Admission-day SAPS II score	39 [26-48]				
Admission-day SOFA score	4 [3-6]				
Biopsy-day SOFA score	4 [4-6]				
In-ICU organ-failure support or monitoring on biopsy day					
Mechanical ventilation	29 (100%)				
Renal replacement therapy	5 (17.2%)				
Vasopressor use	3 (10.3%)				
External ventricular drain	3 (10.3%)				
Intracranial pressure monitoring	2 (6.9%)				
Extracorporeal membrane oxygenation	0 (0%)				
Pre-biopsy length of ICU stay, d	11 [6-19]				
Post-biopsy length of ICU stay, d	20 [7-34]				
Mortality					
In-ICU	12 (41.4%)				
Day-90	12 (41.4%)				
Day-180	13 (44.8%)				
Day-365	14 (48.3%)				

Abbreviations: D, days; ICU, intensive care unit; SAPS II, Simplified Acute Physiology Score II; SOFA, Sequential Organ Failure Assessment. Continuous variables are expressed as mean \pm SD or median [25–75th percentile interquartile range]; categorical variables are expressed as n (%).

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Table 2. Univariable Analysis of Variables Associated with Brain Biopsy Contributory to

Final Diagnosis in the 29 ICU Critically III Patients.

Variables		Biopsy	Biopsy		
	All patients	Contributory to	Noncontributory	Ρ	OR 95%CI
	n = 29	Final Diagnosis	to Final		
		n = 22	Diagnosis <i>n</i> = 7		
Male	21 (72.4)	16 (72.7)	5 (71.4)	1.0	
Comorbidity					
Immunocompromised	17 (58.6)	12 (54.5)	5 (71.4)	0.67	
Cardiovascular	8 (27.6)	5 22.7)	3 (42.9)	0.36	
Autoimmune diseases	3 (10.3)	3 (13.6)	0 (0.0)	0.56	
HIV infection	6 (20.7)	4 (18.2)	2 (28.6)	0.61	
Hematological malignancies	6 (20.7)	4 (18.2)	2 (28.6)	0.61	
Organ transplantation	5 (17.2)	4 (18.2)	1 (14.3)	1.0	
Solid-organ tumor	2 (6.9)	1 (4.5)	1 (14.3)	0.43	
Clinical findings before biopsy					
Admission-day SOFA score	4.3±1.8	4.3±1.8	4.4±1.8	0.89	
Biopsy-day SOFA score	4 [4-6]	4 [3.8-6.3]	4 [4-6]	1.0	
Admission-day SAPS II score	39 [25-47.5]	39 [24-48]	47 [31-48]	0.67	
Extra-neurological symptoms	11 (37.9)	9 (40.9)	2 (28.6)	0.68	
Laboratory findings before biopsy					
Meningitis	12/27 (44.4)	12/27 (57.1)	0/27 (0)	0.02	2.3 (1.4-3.8)
Elevated CSF proteins	19/27 (70.4)	15/21 (71.4)	4/6 (66.7)	1.0	
White blood cell count, G/L	9 [6.5-11]	9.4 [6.3-11.4]	7.6 [6.5-10.6]	0.39	
C-reactive protein, >10 mg/L	17/28 (60.7)	12/21 (57.1)	5/7 (71.4)	0.67	
MRI findings before biopsy					
Multifocal lesions	20 (69)	15 (68.2)	5 (71.4)	1.0	
Bilateral lesions	20 (69)	15 (68.2)	5 (71.4)	1.0	
Gadolinium enhancement	17 (58.6)	14 (63.6)	3 (42.9)	0.40	
Meningeal involvement	4 (13.8)	2 (9.1)	2 (28.6)	0.24	
Largest lesion diameter, mm	14.8 [7.5-30.5]	14.9 [9.5-28.3]	11.1 [4.5-36.7]	1.0	

Biopsy-targeted lesion

characteristics

Subcortical	10 (34.5)	8 (36.4)	2 (28.6)	1.0	
Deep-brain	12 (41.4)	10 (45.5)	2 (28.6)	0.67	
Cortical	5 (17.2)	2 (9.1)	3 (42.9)	0.08	
Size >1 cm	18 (62.1)	16 (72.7)	2 (28.6)	0.07	
Gadolinium-enhanced	16 (55.2)	13 (59.1)	3 (42.9)	0.67	
Biopsy technique					
Stereotaxic	19 (65.5)	16 (72.7)	3 (42.9)	0.19	
MRI-guided	11/19 (57.9)	9/16 (56.3)	2/3 (66.7)	1.0	
Cortico-meningeal	2 (6.9)	2 (9.1)	0 (0.0)	1.0	

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Abbreviations: CI = confidence interval; CSF = cerebrospinal fluid; GCS = Glasgow coma score; MRI = magnetic resonance imaging; OR = odds ratio; SAPS II, Simplified Acute Physiology Score II; SOFA, Sequential Organ

Failure Assessment. Continuous variables, expressed as mean ± SD or median [interquartile range (IQR)], were

516 517 518 519 520 compared with Student's t-test or Wilcoxon's rank test; categorical variables, expressed as n (%), were compared with Fisher's exact tests.

Supplemental Digital Content 1 - Table. Patient, Biopsy and Outcome Characteristics with Comparison

According to Patients' Clinical Status

Characteristic	All patients	ICU patients	Non-ICU	р-
	<i>n</i> = 234	<i>n</i> = 29	patients	value
			n = 205	
Male	149 (63.7)	21 (72.4)	128 (62.4)	0.30
Age on biopsy day, years	48 [36-60]	53 [34.5-59.5]	48 [36.5-60]	0.35
Comorbidity				
Immunocompromised	101 (43.2)	17 (58.6)	84 (41.0)	0.07
Cardiovascular	53 (22.6)	8 (27.6)	45 (22.0)	0.5
Autoimmune diseases	41 (17.5)	3 (10.3)	38 (18.5)	0.28
HIV infection	32 (13.7)	6 (20.7)	26 (12.7)	0.24
Hematological malignancies	28 (12.0)	6 (20.7)	22 (10.7)	0.12
Organ transplantation	23 (9.8)	5 (17.2)	18 (8.8)	0.15
Solid-organ tumor	22 (9.4)	2 (6.9)	20 (9.8)	0.62
Clinical findings before biopsy				
Neurological defect	163/231 (70.6)	16 (55.2)	147/202 (72.8)	0.052
Altered consciousness (GCS <15)	76/231 (32.9)	29 (100.0)	47/202 (23.3)	<0.001
Seizure	64 (27.4)	8 (27.6)	56 (27.3)	0.98
Extra-neurological symptoms	76/230 (33.0)	11 (37.9)	65/201 (32.3)	0.55
Fever	38/231 (16.5)	14 (48.3)	24/202 (11.9)	<0.001
Laboratory findings before biopsy				
Meningitis	67/198 (33.8)	12/27 (44.4)	55/171 (32.2)	0.21
Elevated CSF proteins	101/197 (51.3)	19/27 (70.4)	82/170 (48.2)	0.03
White blood cell count, G/L	6.8 [4.8-10.3]	9 [6.5-11]	6.6 [4.7-9.6]	0.02
Hemoglobin, g/dL	12.2 [10.2-13.8]	9.3 [8.2-11.9]	12.4 [10.6-13.9]	0.001
Platelet count, G/L	249 [193-309]	267 [192-358]	246 [193-301]	0.23
<150 G/L	36/225 (16.0)	5 (17.2)	31/196 (15.8)	0.85
<100 G/L	12/225 (5.3)	2 (6.9)	10/196 (5.1)	0.69
C-Reactive protein >10 mg/L	64/225 (28.4)	17/28 (60.7)	47/197 (23.9)	<0.001
MRI findings before biopsy				
Multifocal lesions	146/233 (62.7)	20 (69.0)	126/204 (61.8)	0.45

Bilateral lesions	124/233 (53.2)	20 (69.0)	104/204 (51.0)	0.07
Hydrocephalus	18/232 (7.8)	4 (13.8)	14/203 (6.9)	0.19
Gadolinium enhancement	143/232 (61.6)	17 (58.6)	126/203 (62.1)	0.72
Meningeal involvement	33/232 (14.2)	4 (13.8)	29/203 (14.3)	0.94
Largest lesion diameter, mm	18.4 [11-29.9]	14.8 [7.5-30.5]	19 [11.2-29.9]	0.31
<10 mm	53/226 (23.5)	8/28 (28.6)	45/198 (22.7)	0.50
>10 mm	173/226 (76.5)	20/28 (71.4)	153/198 (77.3)	0.50
>20 mm	107/226 (47.3)	11/28 (39.3)	96/198 (48.5)	0.36
>50 mm	18/226 (8.0)	5/28 (17.9)	13/198 (6.6)	0.04
Biopsy-targeted lesion				
characteristics				
Subcortical	89/232 (38.4)	10 (34.5)	79/203 (38.9)	0.65
Deep-brain	90/232 (38.8)	12 (41.4)	78/203 (38.4)	0.76
Cortical	38/232 (16.4)	5 (17.2)	33/203 (16.3)	0.89
Supratentorial	212 (90.6)	29 (100)	183 (89.3)	0.06
Cerebellum	10 (4.3)	0 (0.0)	10 (4.9)	0.22
Brainstem	10 (4.3)	0 (0.0)	10 (4.9)	0.22
Size >1 cm	166/232 (71.6)	18 (62.1)	148/203 (72.9)	0.23
Gadolinium-enhanced	143/232 (61.6)	16 (55.2)	127/203 (62.6)	0.44
Biopsy technique				
Stereotaxic	172/233 (73.8)	19 (65.5)	153/204 (75.0)	0.28
MRI-guided	136/172 (79.1)	11/19 (57.9)	125/153 (81.7)	0.02
Cortico-meningeal	15/232 (6.5)	2 (6.9)	13/203 (6.4)	0.92
Biopsy-related histology				
Specific lesion	161 (68.8)	18 (62.1)	143 (69.8)	0.40
Non-specific lesion	67 (28.6)	10 (34.5)	57 (27.8)	0.46
Normal brain	6 (2.6)	1 (3.4)	5 (2.4)	0.75
Biopsy-related diagnosis				
Diagnostic biopsy	174 (74.4)	22 (75.9)	152 (74.1)	0.84
Second biopsy	7 (3.0)	0 (0.0)	7 (3.4)	0.31
Biopsy-related complication				
Complication	46/233 (19.7)	10 (34.5)	36/204 (17.6)	0.03
Symptomatic complication	10/233 (4.3)	3 (10.3)	7/204 (3.4)	0.11

Biopsy-related mortality	2 (0.9)	2 (6.9)	0 (0.0)	0.02
Post-biopsy outcomes				
Therapeutic management change	177/230 (77.0)	18 (62.1)	159/201 (79.1)	0.04
Death during follow-up	57 (24.4)	16 (55.2)	41 (20.0)	<0.001
Follow-up, days	323 [107-703]	201 [28-646]	343 [140-712]	0.23

524 525 526 Abbreviations: CSF = cerebrospinal fluid; GCS = Glasgow coma score; MRI = magnetic resonance imaging. Continuous variables, expressed as median [interquartile range (IQR)], were compared with Wilcoxon's rank test; categorical variables, expressed as n (%), were compared with χ^2 tests.

Supplemental Digital Content 2 - Table. Features, diagnosis and outcome of the 29 ICU patients.

Pati ent (ref.)	S ex	Ag e, yea rs	Medical history	Epile psy	CSF	Tar get lesi on >1c m	Ga do +	Biop sy	Complic ation	Final diagnosis	Outco me
1	F	43	HIV	No	Menin gitis	Yes	Ye s	Ster eo	Grade 1	CD8+ encephalitis	Dead day 400
2	М	64	Lung transplant	No		Yes	Ye s	Ster eo	None	Tuberculosis	Alive
3	М	58	0	No	Menin gitis	Yes	No	Ope n	None	Lymphoma	Alive
4	F	30	HIV	Yes	High CSF protein s	Yes	No	Ope n	Grade 1	No	Alive
5	М	36	0	No	Menin gitis	Yes	Ye s	Ope n	None	Gliomatosis cerebri	Dead day 324
6	М	59	Hodgkin	Yes	Menin gitis	Yes	Ye s	Ope n	None	Cerebral vasculitis	Alive
7	M	64	HSCT	No	High CSF protein s	Yes	Ye s	Ope n	Grade 4	No	Dead day 20
8	М	57	Myeloma	No	High CSF protein s	No	Ye s	Ster eo	Grade 4	Paracoccidioido mycosis	Dead day 1
9	М	53	Myeloma/ HSCT	Yes	High CSF protein s	No	Ye s	Ster eo	None	Lymphoma	Dead day 6
10	М	45	0	No	Norma I	Yes	Ye s	Ster eo	None	HSV-1 encephalitis	Alive
11	M	60	Myeloma	Yes	High CSF protein	No	Ye s	Ope n	None	No	Dead day 30
12	М	28	HIV	No	Norma	Yes	Ye s	Ster eo	None	ADEM	Alive
13	М	55	HIV	No	Norma I	No	No	Ster eo	None	PML*	Dead day 21
14	М	69	0	Yes	Menin gitis	No	Ye s	Ster eo	None	Lymphoma	Alive
15	F	76	Lung cancer	Yes	Norma I	No	Ye s	Ope n	None	No	Dead day 591
16	М	31	0	No	Menin gitis	Yes	Ye s	Ster eo	Grade 1	Tuberculosis	Alive
17	М	21	Heart transplant	No		No	No	Ster eo	None	No	Dead day 50
18	M	55	0	No	High CSF protein s	No	No	Ster eo	None	No	Alive
19	М	54	Kidney transplant	No	Menin gitis	Yes	Ye s	Ster eo	None	Aspergillosis	Dead day 3
20 (24)	F	28	HIV	Yes	Norma I	Yes	No	Ope n	None	Measles encephalitis	Dead day 38

21 (25)	F	58	0	Yes	Menin gitis	No lesi on	No	Ope n	Grade 2	Bunyavirus encephalitis	Dead day 12
22	F	68	Liver Transplant	Yes	High CSF protein s	Yes	No	Ope n	None	Rubella encephalitis	Dead day 104
23	М	42	0	No	Menin gitis	No	No	Ster eo	None	Behçet's disease	Alive
24	М	51	0	No	Menin gitis	Yes	No	Ster eo	None	ADEM	Dead day 25
25 (23)	М	37	Obesity	No	Menin gitis	No	No	Ster eo	Grade 1	Multiple angiopathy of Sars-Cov-2 infection	Dead day 37
26 (23)	М	50	Kidney transplant	No	High CSF protein s	No	Ye s	Ster eo	Grade 1	Multiple angiopathy of Sars-Cov-2 infection	Alive
27 (23)	F	77	Obesity, HBP	No	High CSF protein s	Yes	Ye s	Ster eo	Grade 1	Multiple angiopathy of Sars-Cov-2 infection	Dead day 35
28 (16)	F	31	HIV	No	Norma I	Yes	Ye s	Ster eo	Grade 1	Toxoplasmosis + Lymphoma	Alive
29	Μ	33	0	No	Menin gitis	Yes	No	Ster eo	None	Behcet's disease	Alive



