A devastating neurological complication occurred [1–4] as a result of the hypercoagulability of COVID-19 pneumonia [5]. An ischemic stroke (or perhaps venous sinus thrombosis) involved the posterior inferior cerebellar artery (PICA) territory with hemorrhagic conversion, severe mass effect, and subsequent brain death despite maximal medical therapy. Rarely [6, 7] seen sequential imaging shows the radiographic development of supratentorial destruction from a posterior fossa hemorrhagic infarction.

Description of the Case
A 61-year-old woman with a pertinent history of metabolic syndrome was hospitalized due to severe hypoxemia from COVID-19 pneumonia, diagnosed by positive nasal swab for SARS-CoV-2 and typical ground glass opacities on chest computed tomography (CT) imaging. She was transferred to the ICU on the 3rd day of hospitalization and was found to have a D-Dimer serum level above the upper limit of detection (> 42,000 ng/mL Fibrinogen Equivalent Units) despite having received subcutaneous prophylactic dose heparin since the time of admission. Imaging confirmed the absence of deep venous thrombosis and pulmonary emboli. She was intubated on the 5th hospital day after failing noninvasive oxygen therapy. Despite deep sedation and paralysis, she required prone positioning for refractory hypoxemia by the 6th day of hospitalization. Management of her respiratory and renal failure necessitated nearly continuous deep sedation, neuromuscular blockade, and renal replacement therapy. On the evening of the 11th day of hospitalization, sedation was weaned at which point she resumed some spontaneous movements as she had during prior sedation holidays. However, early in the morning of the 12th day, movement ceased and she became unresponsive.

CT of the head was performed, which showed a large right cerebellar intraparenchymal hemorrhage with extensive mass effect resulting in basilar cistern effacement and upward transtentorial herniation (Fig. 1a). There was marked hydrocephalus and early loss of posterior fossa gray–white differentiation (Fig. 1b–d). The edema surrounding the hemorrhage was confined to the PICA territory, raising suspicion that a PICA or venous sinus thrombosis underwent hemorrhagic conversion. On exam, only cough and gag reflexes as well as the right corneal reflex were preserved, without central or peripheral motor response to pain. A ventriculostomy was placed as the patient was not a surgical candidate (due to neurological condition, coagulopathy, and multiorgan failure). There was brisk drainage of the ventriculostomy and intracranial pressures (ICPs) measured approximately 30 mmHg despite medical therapy. The neurologic examination progressed to loss of all brainstem reflexes by the 13th day, which was followed by ICP measurements less than 10 mmHg. On the 17th day, CT angiography was obtained which showed diffuse cerebral edema and near complete ventricular collapse, but filling only to the M1 segments bilaterally (Fig. 2a–e).

Despite maximal preoxygenation, she rapidly desaturated in less than a minute with a closed circuit held on end inspiration with maintained expiratory pressure, suggesting that even with tracheal insufflation, an apnea test could not be tolerated in fulminant COVID-19 pneumonia and acute respiratory distress syndrome (ARDS) without use of mechanical ventilation. The remainder of the brainstem reflex and motor exam was completed with only negative findings, and confounding variables were systematically ruled out [8]. With a confirmatory nuclear
study showing no uptake, brain death was declared on
the 19th day of hospitalization (Fig. 3a, b).

**Discussion**
Our case provided the opportunity to observe a sequence
of events after presumed primary brainstem death, which
is seldom captured sequentially with modern CT imag-
ing. Neuroimaging showed the progression of brain
destruction caused by an expansive posterior fossa mass
causing upward (through the tentorial opening) and
downward herniation (through the foramen magnum).
Despite maximal cerebrospinal fluid (CSF) drainage, dif-
fuse cerebral edema obliterated the ventricular system
with initially elevated ICP and later low ICP after hernia-
tion. The process of hemispheric infarction from an acute
posterior fossa mass is illustrated by the sequence of radi-
ographic images:

![Axial CT head images from the 12th day of hospitalization](image)
Fig. 2  a–c Axial, d reconstructed sagittal and e coronal CT angiography head images from the 17th day of hospitalization, after the interval loss of all brainstem reflexes which showed diffuse cerebral edema and near complete ventricular collapse, but preserved filling of the Circle of Willis and M1 segments without further evidence of cortical perfusion, which are also demonstrated in f A-P 3-D reconstructions. The sagittal images demonstrate extensive upward herniation and compression on the deep venous structures.
1. Multiple cerebellar and brainstem infarcts and mass effect from hemorrhagic conversion of the right cerebellum rapidly produces progressive and extensive loss of brainstem function (Fig. 1a, b).

2. Secondary hydrocephalus from tentorial compression and aqueductal obstruction from upward herniation, and by implication compression of the vein of Galen, causes venous congestion and thus a multifactorial increase in ICP, which was not improved with ventriculostomy (Fig. 1b–d).

3. Increased ICP impedes intracranial blood flow into the anterior circulation beyond the M1 segments (Fig. 2a–f). The rest of the brain is seriously irrevocably under-perfused. The posterior circulation was already non-filled at this time.

4. Total brain infarction ensues and blood flow stops secondary to diffuse cerebral edema, proven via radionuclide study (Fig. 3a, b).

Most patients who are declared brain dead have a supratentorial mass lesion or diffuse process that results in infratentorial destruction, but our patient showed the opposite. These images depict the full process of brain infarction from a posterior fossa etiology, and this process is more multifaceted than acute hydrocephalus alone. Once compression nearly totally destroyed the brainstem, the swelling rapidly and indirectly destroys the rest of the brain via upward herniation, compression of the vein of Galen and venous congestion. In such a multifaceted brain injury, ventriculostomy will not improve the condition nor will decompression of the cerebellum. A better understanding of the pathophysiology of posterior fossa lesions with loss of all brainstem reflexes is important in evaluating brain death [7, 9]. This case again confirms that when an examination shows complete loss of brainstem reflexes in a massive posterior fossa lesion, a conserved and likely very predictable progression occurs with no posterior, followed by no anterior, cerebral perfusion. In primary brainstem death (from intrinsic lesions or compression), the revealant vital functions have ceased and other functions are on the verge of ceasing.

Fulminant COVID-19 pneumonia poses logistical challenges in the brain death examination due to unsafe apnea testing in patients with dramatic and diffuse exudative epithelial denudation of alveoli. The COVID-19 pandemic has dramatically reduced organ donation [10], but the effect on the frequency of brain death determination is not known in both COVID-19 positive and negative patients.

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CM drafted the manuscript. All authors revised the manuscript, and participated in the medical care of the patient.

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