Neuroimaging of Propofol Infusion Syndrome: A Case Report and Review of Literature

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Abstract

A school-age boy with a complex medical history underwent a minor elective surgical procedure. Propofol was used for sedation during the procedure. The patient could not be awakened post-operatively. Laboratory findings demonstrated metabolic lactic acidosis, leukocytosis with bandemia, and transaminitis. Neuroimaging demonstrated findings that were consistent with hypoxic-ischemic or toxic-metabolic brain injury involving the bilateral basal ganglia, hippocampi, and cerebellum. The patient’s condition progressively worsened over the course of the following few weeks, and brain death was confirmed by scintigraphy seven weeks later. Prompt neuroimaging in unresponsive patients with suspected propofol infusion syndrome (PRIS) is of critical importance in detecting neurologic injuries, excluding alternative diagnoses, and determining prognostication.

Categories: Pediatrics, Radiology
Keywords: neuroimaging, propofol infusion syndrome, anoxic injury, hypoxic injury, encephalopathy, pediatrics

Introduction

Propofol infusion syndrome (PRIS) is a rare but serious complication of propofol infusion typically reported in seriously ill patients administered high doses of propofol for prolonged duration [1]. Classically, PRIS presents as acute refractory bradycardia, metabolic acidosis, and rhabdomyolysis, which may result in renal failure, cardiac failure, and death in the setting of propofol infusion [1]. Brain injuries or encephalopathy are uncommon manifestations of this syndrome but have been rarely reported [2,3]. Although the mechanism of PRIS is not yet well characterized, recent studies have demonstrated that propofol may inhibit key factors in the electron transport chain and in the transport of long-chain fatty acids, thus impairing beta oxidation of fatty acids and energy production [4-7]. In this case report, we detail neuroimaging findings in a pediatric patient who developed symptoms of PRIS.

Case Presentation

A school-age boy with history of prematurity, cerebral palsy, periventricular leukomalacia, and epilepsy underwent elective surgery for correction of strabismus. The patient’s preexisting neurological conditions were well controlled at the time of surgery; he had normal developmental milestones and was fully alert and oriented with no neurological deficits before the procedure. He was premedicated with midazolam and underwent elective surgery for correction of strabismus. The patient could not be awakened post-operatively. Laboratory analyses following the procedure and, other than an elevated white blood cell count noted immediately after the initial procedure, were well controlled at the time of surgery; he had normal developmental milestones and was fully alert and oriented with no neurological deficits before the procedure. He was premedicated with midazolam and was briefly noted to be tachycardic with a heart rate of 148 beats per minute when first transported to the recovery area, but this finding resolved quickly and was unaccompanied by changes in other vital signs. No seizure activity was noted throughout this period. Unfortunately, failure of the patient to awaken was noted immediately afterwards, while still in the recovery room. Physical examination revealed extremity withdrawal to pain, but the patient was otherwise unresponsive. There were no vital sign abnormalities such as hypoxia, bradycardia, or hypotension during the patient’s surgery or during the post-operative period that were detected to explain the patient’s unresponsiveness. Likewise, the patient’s history did not reveal any exposure to known toxins. He did not have a history of mitochondrial disorder or other metabolic dysfunction. He was afebrile throughout the procedure and, other than an elevated white blood cell count noted immediately after the initial procedure, had no signs or symptoms of any sort of infection. Laboratory analyses following the procedure demonstrated metabolic acidosis, mildly elevated alanine aminotransferase (ALT), and an elevated white blood cell count with a left shift.

An emergently acquired unenhanced CT scan of the patient’s brain demonstrated bilaterally symmetrical areas of hypoattenuation in the basal ganglia, mesial temporal lobes, midbrain, and cerebellum (Figure 1). No definite area of acute hemorrhage was identified. There was no midline shift. MR imaging of the brain without contrast was subsequently obtained and demonstrated extensive restricted diffusion with...
corresponding T2 and fluid-attenuated inversion recovery (FLAIR) hyperintense signal changes in bilateral basal ganglia, hippocampi, cerebral peduncles, and most of the cerebellum (Figure 1). There was significant mass effect on the fourth ventricle without any significant upstream dilatation of the ventricular system. No evidence of periventricular cerebrospinal fluid (CSF) seepage was noted. Following these findings, the patient was admitted to the pediatric intensive care unit (PICU) for further treatment and care.

Unfortunately, the patient’s condition continued to deteriorate with complete loss of response to all stimuli, and he required ventilatory support as he stopped making spontaneous respiratory efforts. A follow up MRI obtained three days after the initial surgical procedure demonstrated interval worsening of the cytotoxic edema involving bilateral basal ganglia, hippocampi, cerebral peduncles, and the cerebellum (Figure 2). In addition, interval development of cerebellar tonsillar herniation and ischemic changes in the upper cervical spinal cord were revealed. Time-of-flight MR angiogram (MRA) and MR venogram (MRV) of the brain were normal.
FIGURE 2: Follow-Up MRI Findings Obtained Three Days After Initial Imaging

Images 2A, 2B, 2C: T2 fluid-attenuated inversion recovery (FLAIR), isotropic diffusion-weighted, and apparent diffusion coefficient (ADC) map images demonstrated increased areas of diffusion restriction in the basal ganglia (blue arrows). Images 2D, 2E, 2F: T2 FLAIR, isotropic diffusion-weighted, and ADC map images demonstrated increased areas of diffusion restriction in the cerebellum (red arrowheads).

Although at this point the patient’s clinical presentation indicated possible brain death, a reliable clinical brain death exam was not feasible on account of cervical cord ischemia. Supportive care of the patient was continued in the PICU. Subsequent MR imaging obtained four weeks later demonstrated persistent diffuse cytotoxic edema involving both cerebral hemispheres and the entire cerebellum. Further worsening of cerebellar herniation and persistent cervical cord ischemia were also evident. Time-of-flight MRA revealed absence of flow-related signal in the circle of Willis or any of its branches, with blooming identified in proximal bilateral middle cerebral arteries, proximal bilateral posterior cerebral arteries, and the basilar artery concerning for widespread arterial thrombosis (Figure 3). MRV showed absence of flow-related enhancement in the superior sagittal sinus, the straight sinus, bilateral transverse sinuses, and bilateral sigmoid sinuses, with corresponding hyperintense signal on T2 weighted imaging, concerning for widespread venous thrombosis (Figure 4).
FIGURE 3: Follow-Up MRI Obtained One Month After Initial Imaging

Image 3A, 3B: Isotropic diffusion-weighted image and apparent diffusion coefficient (ADC) maps demonstrated new, widespread diffusion restriction throughout the cerebral white matter (blue arrows). Images 3C, 3D: Isotropic diffusion-weighted image and ADC map demonstrated demonstrate new, widespread diffusion restriction in the cerebellum (red arrowheads).
FIGURE 4: Time-of-flight MRA Obtained One Month After Initial Imaging

Time-of-flight MR angiogram (MRA) one month after the patient was initially noted to be unresponsive showed absence of flow-related signal in the circle of Willis and branch arteries bilaterally. Red arrowheads indicate the distal internal carotid arteries with a lack of intracranial arterial flow.

A nuclear brain perfusion study performed one week later demonstrated diffuse absence of parenchymal radiotracer uptake, consistent with absence of brain perfusion (Figure 5). Given his failure to awaken after the procedure, the history of recent propofol infusion, immediate post-operative lab findings showing metabolic lactic acidosis, mild elevation of ALT, and the absence of other signs or symptoms pointing to another cause of these ailments, the patient was diagnosed with PRIS.

FIGURE 5: Nuclear Flow Study Obtained at Seven Weeks After Initial Procedure

Anterior (A) and left lateral (B) static images acquired 15 minutes after intravenous injection of 99mTc
function involvement that spared the basal ganglia reportedly made an excellent recovery and regained previous.

Our patient and the case described by Li et al., one other pediatric patient with reported brain hemorrhage of the left caudate nucleus that was observed on follow-up imaging. In contrast, extensive diffusion restriction in the basal ganglia and cerebellum similar to our patient, except that Li et al. diffusion on MRI, the prognosis was very poor.

It may also predict prognosis. In our patient and in another patient whose basal ganglia showed restricted. suspected PRIS patients may allow faster detection and prompt initiation of neuroprotective measures, and maintenance a normoglycemic status in these patients may help to prevent development of PRIS.

Brain injuries or encephalopathy are rarely observed in this syndrome but have been reported [2,3]. Risk factors for PRIS include young age, severe concurrent illness, high fat or low carbohydrate intake, hypoglycemia, coadministration of catecholamine or steroid use, fatty acid oxidation disorders, or high doses of propofol (>4-5 mg/kg/hr) for a prolonged period (>48 hr) [1,5,9,10]. It must be noted that our patient was not genetically tested for inborn errors of metabolism that may have predisposed him to developing PRIS. While there are no specific protocols for management of PRIS, treatment typically includes immediate discontinuation of propofol, correction of pH and electrolyte imbalances, supportive care including extracorporeal membrane oxygenation when necessary, and hemodialysis in select cases where electrolyte or oxygen imbalances cannot be corrected via other means [11,12]. At this time, given the risks associated with prolonged propofol use in young patients, propofol is not currently recommended for long term sedation in pediatric patients [1].

The mechanism of PRIS is proposed to be propofol induced disruption of intracellular adenosine triphosphate (ATP) production, by interference in the mitochondrial electron transport chain or by interference with oxidative metabolism of free fatty acids within mitochondria. Most reported pediatric cases have been observed after continuous infusion of the drug [5,8]. In a small percentage of pediatric patients, severe PRIS-like syndrome after low dose propofol is suspected to be due to enhanced sensitivity from underlying mitochondrial defects. The clinical syndrome in these patients may resemble an acute presentation of patients with genetic mitochondrial diseases. Some authors suggest that close attention to maintaining a normoglycemic status in these patients may help to prevent development of PRIS [8].

Propofol is hypothesized to block the transport proteins in the mitochondrial membrane responsible for influx of long-chain fatty acids, thereby impairing beta oxidation of fatty acids [1,3,4]. Histopathological examination of skeletal and cardiac muscle in PRIS has demonstrated extensive myocyte necrosis and fat accumulation in some cases [5]. Previous studies have also proposed that propofol may block the electron transport chain through inhibition of Complex 1 or Coenzyme Q, thereby inhibiting electron transfer on Cytochrome C [1,5,6]. This pathogenic mechanism helps to explain why, in our patient and in another patient with documented brain injury due to PRIS, regions of the brain with relatively high metabolic activity—the basal ganglia, hippocampi, and cerebellum—demonstrated injury first.

Overall, similar acute findings have been reported on brain imaging after hypoxic brain injury in the setting of carbon monoxide toxicity, including spatial distribution [13-15]. This is not unexpected since both carbon monoxide and propofol action are centered on the mitochondrial electron transport chain, with carbon monoxide targeting complex IV (unlike complex I inhibition by propofol) [16]. Prompt neuroimaging in suspected PRIS patients may allow faster detection and prompt initiation of neuroprotective measures, and it may also predict prognosis. In our patient and in another patient whose basal ganglia showed restricted diffusion on MRI, the prognosis was very poor [2]. In the case reported by Li et al., imaging demonstrated extensive diffusion restriction in the basal ganglia and cerebellum similar to our patient, except that Li et al. also described hemorrhage of the left caudate nucleus that was observed on follow-up imaging. In contrast to our patient and the case described by Li et al., one other pediatric patient with reported brain involvement that spared the basal ganglia reportedly made an excellent recovery and regained previous function [3]. The pertinent details of these cases are compared in Table 1.
| Poretti et al. [2] | Li et al. [3] | Current case report |
|-------------------|--------------|---------------------|
| Age of Patient    | 3 year-old female | 30 year-old female | 6 year-old male |
| Risk factors      | Young age    | None                | Young age    |
| Genetic mitochondrial condition | Not mentioned | No abnormalities in mitochondrial DNA | Not tested |
| Indication for propofol use | Sclerotherapy of large venous malformation | Laparoscopic surgery for colon polyp/ cancer | Surgical repair of bilateral optic esotropia |
| Propofol rate and dose | Not available | Mean rate of 3.3 mg/kg/hr totaling 120 mg in 6 hours | Mean infusion rate of 1.32 mg/kg/hr for 100 min, plus 3 bolus doses totaling 100 mg in same time period |
| Imaging Findings (CT and MRI) | T2 prolongation and diffusion restriction was noted in supra- and infratentorial white matter; these findings were completely resolved on follow up. | Low attenuation was shown in bilateral basal ganglia, on CT. Follow up CT demonstrated hemorrhage in caudate nucleus. Diffusion signal abnormality of the basal ganglia, temporal lobe, and cerebellum bilaterally and a corresponding T1 hypointensity was also noted. | Low attenuation was evident in bilateral basal ganglia and cerebellum on CT. Diffusion restriction of bilateral basal ganglia, cerebellum, and bilateral hippocampi with hyperintensities was noted on T2, FLAIR, and DW MRI sequences. Four weeks later, diffuse cytotoxic edema on DWI was noted, with widespread hyperintensity on T2 and FLAIR sequence, involving the entire cerebral hemispheres and the cerebellum. |
| Laboratory Changes | Increased levels of acetyl and hydroxy-butyryl ketones with generalized elevation of fatty acylcarnitine intermediates. | N/A | Lactic acidosis; also, elevated leukocyte count with a left shift; mildly elevated ALT |
| Clinical outcome | Complete recovery | Death | Death |

**TABLE 1: Review of PRIS Literature Where Relevant Neuroimaging Was Available, Compared to Current Report**

PRIS: propofol infusion syndrome; DNA: deoxyribonucleic acid; FLAIR: fluid-attenuated inversion recovery; DWI: diffusion weighted imaging; ALT: alanine aminotransferase.

Although MR imaging was performed in cases reported by Savard et al. and Mtaweh et al., this was done for other indications well before the development of PRIS [6,8]. Similar to our case, Mtaweh et al. reported PRIS-like condition in a child caused by a single dose of propofol (1.25 mg/kg). In contrast, the case described by Savard et al. involved the development of PRIS in an adult at much higher doses administered for a longer period of time. Both case reports were associated with mitochondrial disorders or gene defects. Our patient had not previously experienced symptoms of any sort of mitochondrial disorder and was therefore never genetically tested for it.

**Conclusions**

PRIS is a rare but potentially fatal complication of propofol infusion. Very little is known about possible neurologic sequelae resulting from PRIS. Knowledge of the imaging findings of PRIS is helpful for establishing more standardized diagnostic and management guidelines in patients with suspected PRIS so that more common injury patterns associated with this syndrome may be identified. Furthermore, awareness about this lethal complication is important to enable urgent neuroimaging for obtunded or unresponsive patients after administration of propofol.

**Additional Information**
Disclosures

Human subjects: Consent was obtained by all participants in this study. Conflicts of interest: In compliance with the ICMJE uniform disclosure form, all authors declare the following: Payment/services info: All authors have declared that no financial support was received from any organization for the submitted work. Financial relationships: All authors have declared that they have no financial relationships at present or within the previous three years with any organizations that might have an interest in the submitted work. Other relationships: All authors have declared that there are no other relationships or activities that could appear to have influenced the submitted work.

References

1. Hemphill S, McMenamin L, Bellamy MC, Hopkins PM: Propofol infusion syndrome: a structured literature review and analysis of published case reports. Br J Anaesth. 2019, 122:448-59. 10.1016/j.bja.2018.12.025
2. Poretti A, Bosemani T, Huisman TAGM: Neuroimaging findings in pediatric propofol infusion syndrome. Pediatr Neurol. 2014, 50:451-2. 10.1016/j.pediatrneurol.2013.12.016
3. Li X, Zhao Z, Liu X, Ma G, Zhu MJ: Encephalopathy associated with propofol infusion syndrome: a case report. Medicine. 2018, 97:e9521. 10.1097/MD.0000000000009521
4. Finsterer J, Frank M: Propofol is mitochondrion-toxic and may unmask a mitochondrial disorder. J Child Neurol. 2016, 31:1489-94. 10.1177/088307381661458
5. Vanlander AV, Jorens PG, Smet J, et al.: Inborn oxidative phosphorylation defect as risk factor for propofol infusion syndrome. Acta Anaesthesiol Scand. 2012, 56:520-5. 10.1111/j.1399-6576.2011.02628.x
6. Savard M, Dupré N, Turgeon AF, Deshiens R, Langevin S, Brunet D: Propofol-related infusion syndrome heralding a mitochondrial disease: case report. Neurology. 2015, 81:770-71. 10.1212/WNL.0000000000001978
7. Trapani GM, Almonte C, Sanna E, Biggio G, Liso G: Propofol in anesthesia. Mechanism of action, structure-activity relationships, and drug delivery. Curr Med Chem. 2000, 7:249-71. 10.2174/0929867003375335
8. Mtsaweh H, Bayr H, Kochanek PM, Bell MJ: Effect of a single dose of propofol and lack of dextrose administration in a child with mitochondrial disease: a case report. J Child Neurol. 2014, 29:NP40-NP46. 10.1177/0883073813509640
9. Krajičová A, Waldau P, Anděl M, Duška F: Propofol infusion syndrome: a structured review of experimental studies and 153 published case reports. Crit Care. 2015, 19:398. 10.1186/s13054-015-1112-5
10. Michel-Macías C, Morales-Barquet DA, Reyes-Palomino AM, Machuca-Vaca JA, Orozco-Guillén A: Single dose of propofol causing propofol infusion syndrome in a newborn. Oxf Med Case Reports. 2018, omy023. 10.1093/omcr/omy023
11. Mayette M, Gonda J, Hsu IL, Mihm FG: Propofol infusion syndrome resuscitation with extracorporeal life support: a case report and review of the literature. Ann Intensive Care. 2013, 3:32. 10.1186/2110-5820-3-32
12. De-Silva SS, Wong R, Coquilhon P, Gavrilița C, Asuncion A: Partial-exchange blood transfusion: an effective method for preventing mortality in a child with propofol infusion syndrome. Pediatrics. 2010, 125:e1493-e1499. 10.1542/peds.2009-1823
13. de Oliveira AM, Paulino MV, Vieira AP, et al.: Imaging patterns of toxic and metabolic brain disorders. Radiographics. 2019, 39:1672-95. 10.1148/rg.2019190016
14. Lo CP, Chen SY, Lee KW, Chen WL, Chen CY, Hsuah CJ, Huang GS: Brain injury after acute carbon monoxide poisoning: early and late complications. AIR Am J Roentgenol. 2007, 189:W205-W211. 10.2214/AJR.07.2425
15. Jeon SB, Sohn CH, Seo DW, Oh BJ, Lim KS, Kang DW, Kim WY: Acute brain lesions on magnetic resonance imaging and delayed neurological sequelae in carbon monoxide poisoning. JAMA Neurol. 2018, 75:436-43. 10.1001/jamaneurol.2017.4618
16. Queiroga CSF, Almeida AS, Vieira HLA: Carbon monoxide targeting mitochondria. Biochem Res Int. 2012, 749845. 10.1155/2012/749845