Review

The Great Masquerade: Donor-derived Infections with Uncommon Central Nervous System Pathogens

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Abstract

Donor-derived infections (DDI) are an infrequent event in solid organ transplant (SOT) due to advances in screening recommendations, prophylaxis, and surveillance of common infections. However, unexpected pathogen transmission can still occur when a donor is not known to be infected prior to organ procurement, which can lead to significant morbidity and mortality in the organ recipient. Solid organ donors with central nervous system (CNS) pathogens are an uncommon but deadly source of unexpected DDI. Clinically recognizing these CNS infections in a potential deceased donor is enormously challenging as many are clinically silent or overshadowed by other confounding events. Because of this, expert panels caution against transplanting organs from decedents who die with possible or proven encephalitis of unknown etiology. In this review, we discuss the epidemiology, donor characteristics, and outcomes of cases of DDI in SOT recipients with unusual CNS pathogens, and provide a discussion on methods of identifying and reporting possible DDI with these pathogens.
Keywords
Solid organ transplant; meningoencephalitis; central nervous system pathogens; donor-derived infection

1. Introduction

Solid organ transplant (SOT) is considered a definitive treatment for many terminal end-organ diseases; however, demand has far outpaced supply. The number of those waiting for life-saving organ transplantation has doubled during the last two decades, while the number of transplants has only increased by 30% [1, 2]. Organs from donors with known (i.e., expected) treatable infections, or at increased risk of infections, are increasingly being utilized to offset this need [1, 3].

Donor-derived infections (DDI) are infrequent due to advances in screening recommendations, prophylaxis, and surveillance of common infections. However, unexpected pathogen transmission can still occur when a donor is not known to be infected prior to organ procurement, leading to significant morbidity and mortality [2, 4-7]. In order to improve organ transplantation safety in the United States (U.S.), the Organ Procurement & Transplant Network (OPTN)/ United Network for Organ Sharing (UNOS) created the Disease Transmission Advisory Group (DTAG) in 2005, later becoming the ad hoc Disease Transmission Advisory Committee (DTAC). According to data from OPTN/UNOS ad hoc DTAC, unexpected DDI are exceedingly rare, occurring in less than 0.2% of SOT recipients [8]. Such events can occur when the donor has an asymptomatic, subtle, or atypical presentation of a transmissible infection at the time of death [9]. Procedural measures may fail to capture unexpected DDI when screening is not done for geographically endemic pathogens, screening is done too early (e.g., serology fails to detect early infection), or the donor has an infection that is not routinely screened for [2, 3].

Solid organ donors with central nervous system (CNS) pathogens are an uncommon but potentially deadly source of DDI [10]. During the past two decades, donor-derived CNS pathogens, such as West Nile virus (WNV), *Balamuthia mandrillaris*, lymphocytic choriomeningitis virus (LCMV), and rabies virus, among others, have been reported in clusters of SOT recipients [11, 12]. Clinically recognizing these CNS infections in potential deceased donors is enormously challenging as many are clinically silent, overshadowed by other confounding events (e.g., trauma, cerebral disease, or overdose), and donor information is often limited [3, 6, 10]. Overall, the lack of effective therapies, delayed recognition due to the pathogens’ rarity, and the underutilization of reporting systems has resulted in devastating outcomes in SOT recipients [6, 11-13]. Because of these factors, expert panels caution against transplanting organs from decedents who die with possible or proven encephalitis of unknown etiology [8, 12, 13].

We reviewed the published literature focusing on DDI with unusual CNS pathogens. More common forms of CNS infection (e.g., bacterial meningitis, cryptococcosis, and toxoplasmosis) are not included in this review. We discuss the epidemiology, donor characteristics, and outcomes of these challenging cases, and discuss methods of identifying and reporting possible DDI with CNS pathogens. To that end, we queried Medline, Ovid, Embase, and Web of Science using controlled
vocabulary and natural language terms for tissue and organ donor, organ transplantation, transplant recipients, CNS disease, CNS infection, CNS trauma, encephalitis, meningitis, meningoencephalitis, donor-derived, and transplant-derived. Our literature search spanned fifty years and was limited to articles published in English. Abstracts were then manually screened by the authors to ensure that the selected publications featured cases of unexpected donor-derived infections with CNS pathogens after SOT.

2. Viral Pathogens

2.1 The Arboviruses

West Nile Virus (WNV) is a mosquito-borne Flavivirus and one of the most widely distributed arboviruses worldwide, with an area of circulation covering several continents [14, 15]. Since its emergence in North American in 1999, it has become the most common etiological agent of arboviral encephalitis in the Western Hemisphere [12]. The virus is maintained in a bird-mosquito-bird life cycle and is transmitted to humans incidentally through the bite of Culex mosquitoes [12, 14, 15]. While the majority (80%) of infected immunocompetent individuals remain asymptomatic, up to 20% develop symptoms of fever, headache, myalgias, transient rash, and gastrointestinal symptoms [14-16]. Less than 1% develop neuroinvasive disease characterized by acute flaccid paralysis, Parkinsonian cogwheel rigidity, meningitis, encephalitis, meningoencephalitis, and asymmetric muscle weakness [14-16]. However, neuroinvasive disease may be more common in transplant recipients, where the risk from donor-derived WNV may be as high as 50%-75% [15-17]. During the last two decades, there have been approximately eight clusters of donor-derived WNV (Table 1), starting in 2002 when four organ recipients from a common donor developed febrile illnesses [18, 19]. Encephalitis developed in three of the four recipients. On retrospective review of the blood products given to the donor, one unit of plasma was positive for WNV on quantitative polymerase chain reaction (PCR) [18, 19]. Similarly, in 2008, a heart recipient developed WNV encephalitis after transplantation from a donor who was also infected by blood products [20]. In 2005, the Center for Disease Control and Prevention (CDC) reported two SOT recipients (liver and lung) who developed neuroinvasive WNV after receiving organs from a common donor who died from a traumatic brain injury. Of the two kidney recipients, one had an asymptomatic WNV infection and the other was not infected. The donor’s family revealed mosquito exposure and a febrile illness prior to death and the donor’s serology demonstrated prior infection [21]. This suggests that WNV may be transmitted from a previously infected donor who mounted an immune response but likely had virus persisting in their organs. As recently as 2011, a cluster of four SOT recipients was diagnosed with donor-derived WNV from a common donor with developmental cognitive delay who developed fevers, encephalopathy, and muscle weakness prior to death [22, 23]. No workup of possible CNS infections was performed. After WNV infection was detected in one recipient, the donor’s archived clinical samples were tested, revealing WNV infection by serology and PCR [23]. Of the reported 23 SOT recipients from WNV infected donors, 21 became infected after an average of 12.4 days (7 – 20 days) post-transplant, emphasizing the high rate of disease transmission and short duration to onset of symptoms (Table 1). While 13 survived, four died and an additional four had severe neuroinvasive disease without a reported outcome. As there is no treatment or prophylaxis of proven benefit for WNV infection, treatment largely consists of supportive care. However, several management strategies have been outlined.

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in case reports [15]. The U.S. Food and Drug Administrations (FDA) has approved screening and diagnostic tests for WNV, including nucleic acid tests (NAT) and IgM serology [24]. In the U.S., blood banks screen for WNV and, in 2013, OPTN mandated that living donors be screened for WNV in endemic areas [15]. At this time, there are no established WNV screening recommendations in deceased donors. WNV should be included in the differential diagnosis of any SOT recipient with fever and neurological symptoms after transplantation when donor screening is not done.
### Table 1: Solid organ transplant transmitted cases of West Nile virus.

| Cluster Year, location | Donor Age/sex, COD | Donor risk-factor | Donor testing | WNV Procured organs | Symptom onset after SOT (days) | Recipient WNV serum testing | Recipient WNV CSF testing | WNV Treatments | Outcomes |
|------------------------|--------------------|-------------------|---------------|---------------------|-------------------------------|-----------------------------|-----------------------------|----------------|----------|
| 2002, Georgia and Florida, USA [18, 19] | 20/F, trauma | Received infected blood products | Serum IgM (-), PCR (+), viral culture (+) | Heart | 10 | PCR (+) | IgM (+) | None | NI disease, survived |
| | | | | Liver | 7 | IgM (+) | Not tested | None | NI disease, survived |
| | | | | Kidney | 17 | IgM, IgG (-) | | | |
| | | | | Kidney | 14 | IgM (+) | IgM (+) | None | |
| | | | | Liver | 13 | IgM (+) | IgM (+), PCR (+) | Omr-IgG-am | |
| 2005, New York and Pennsylvania, USA [21] | N/A, head trauma | Outdoor exposure, febrile illness | Serum IgG (+), IgM (+), PCR (-) | Lung | 16 | IgM, IgG (+) | IgM and IgG (+) | Omr-IgG-am | |
| | | | | Kidney | — | IgG (+), PCR (+) | Not tested | Omr-IgG-am | Survived |
| | | | | Kidney | — | IgM, IgG, PCR (-) | Not tested | Omr-IgG-am | Not infected |
| 2008, Louisiana, USA [20, 25] 2009, California, USA [26] | 18/M, GSW head | Infected blood donor Probable mosquito bite | Serum IgM (-), IgG (-), PCR (-) | Heart | 8 | IgM (+) | IgM (+) | Supportive care | NI disease, survived |
| | | | | Liver | 15 | IgM (+), IgG (-), PCR (-) | IgM (+) | IVIG | NI Disease, survived |
| Year | Location | Age | Sex | Disease History | Travel to Endemic Area | Test Results | Diagnosis | Outcome |
|------|----------|-----|-----|-----------------|------------------------|--------------|-----------|---------|
| 2009 | Italy    | 78  | F   | ICH             | Yes                    | Liver: PCR (+) | Not tested | Survived |
|      |          |     |     |                 |                        | Liver: IgM (-)| Not tested | None    |
|      |          |     |     |                 |                        | Liver: IgG (+)| Not tested | None    |
| 2010 | USA      | 55  | M   | Head trauma     | Probable mosquito bite | Kidney: IgM (+), PCR (-) | None | Survived |
|      |          |     |     |                 |                        | Kidney: IgM (-), PCR (+) | No | Survived |
|      |          |     |     |                 |                        | Kidney: IgG (+) | None | Survived |
|      |          |     |     |                 |                        | Heart: IgM, IgG, PCR (-) | None | None |
|      |          |     |     |                 |                        | Liver: IgM, IgG (+), PCR (-) | None | Survived |
|      |          |     |     |                 |                        | Liver: IgG (+) | None | Survived |
| 2011 | Italy    | 43  | F   | Head trauma     | Yes                    | Liver: PCR (-) | Not tested | None |
|      |          |     |     |                 |                        | Liver: IgG (+), PCR (-) | None | Survived |
|      |          |     |     |                 |                        | Liver: IgM, IgG, PCR (-) | None | N/A |
|      |          |     |     |                 |                        | Kidney: IgM, IgG, PCR (-) | None | Survived |
|      |          |     |     |                 |                        | Kidney: IgG (+) | None | Survived |
| 2011 | USA      | 56  | M   | Febrile encephalopathy | Yes                    | Liver: IgG (+), PCR (-) | None | Survived |
|      |          |     |     |                 |                        | Liver: IgG (+), PCR (-) | None | Survived |
|      |          |     |     |                 |                        | Liver: IgM, IgG, PCR (-) | None | Survived |

**Note:** PCR = Polymerase Chain Reaction, IgM = Immunoglobulin M, IgG = Immunoglobulin G, FFP = Fresh Frozen Plasma, Omr-IgG-am = Oromi IgG antibody, INF = INF-alfa-2b, IVIG = Intravenous Immunoglobulin, WNV Ab = West Nile Virus Antibody, N/A = Not Applicable, NI = Nonspecific Inflammation.
|                | Kidney | PCR (+), IgM, IgG (-) | PCR (+), IgM (-) | FFP |
|----------------|--------|-----------------------|------------------|-----|
| PCR            | 10     |                       |                  |     |
| IVIG, INF, NI  |        |                       |                  |     |
| disease, died  |        |                       |                  |     |

Abbreviations. COD, cause of death; SOT, solid organ transplant; CSF, cerebrospinal fluid; USA, United States of America; F, female; M, male; WNV, West Nile virus; IgM, immunoglobulin M; IgG, immunoglobulin G; N/A, not available; PCR, polymerase chain reaction; IHC, immunohistochemistry staining; NI, neuroinvasive; Omr-IgG-am, immune globulin with high antibody titers against WNV; GSW, gunshot wound; ICH, intracranial hemorrhage; IVIG, intravenous immunoglobulin; Ab, antibody; FFP, fresh frozen plasma; LN, lymph node; INF, interferon.

a Asymptomatic; b Prophylactic treatment; c Pre-transplant.
Eastern equine encephalitis virus (EEEV) is a mosquito-borne arbovirus of the genus Alphavirus that is endemic to eastern North America [14, 31]. In nature, EEEV cycles between mosquitoes and birds in forest wetlands and it incidentally infects humans through the bite of an Aedes or Culex mosquito [14, 31]. Most cases of EEEV are asymptomatic or they present with a non-specific febrile illness, with less than <5% developing neuroinvasive disease manifesting as meningitis or encephalitis [14]. While rare, EEEV causes one of the most severe arboviral diseases in North America, with approximately 30%-50% case fatality, and many survivors suffer from residual neurological sequelae [14, 31, 32]. During the past two decades, the national arboviral disease surveillance system (ArboNET) has received 121 reports of human disease from 20 U.S. states, predominantly along the Atlantic and Gulf coasts [32]. To date, there has been only one published report of donor-derived EEEV [31]. In 2017, three SOT recipients (lung, heart, and liver) developed encephalitis one week after transplantation. The common donor had succumbed to a gunshot wound and had no history of febrile or neurological illness. Two of the three recipients died and the surviving recipient recovered with a residual tremor. The donor’s serum was positive for EEEV RNA and all organ recipients also showed evidence of EEEV infection [14, 31]. All three SOT recipients developed neuroinvasive EEEV, suggesting a possible increased risk with donor-derived transmission. This cluster illustrates the need to consider EEEV as a cause of encephalitis in SOT recipients residing in endemic areas within the U.S.

2.2 The Arenaviruses

Lymphocytic choriomeningitis virus (LCMV) is an Old World arenavirus found in Europe and the Americas and is transmitted by infected rodents [14, 15]. Infection in humans occurs through the inhalation of aerosolized excreta or saliva in dust, bites, and contact with blood from an infected rodent [14, 15]. In immunocompetent patients, LCMV typically presents as an asymptomatic or mild self-limited febrile illness [12, 15]. More severe cases may present as aseptic meningitis or encephalitis with very low case fatality rates (<1%) [15]. In transplant recipients, LCMV results in severe meningoencephalitis and multisystem organ failure resulting in very high mortality rates [12, 14]. Several clusters of SOT donor-derived LCMV infections have been reported during the last two decades (Table 2). In 2003, four SOT recipients with a common donor developed an LCMV-like infection, resulting in their deaths [33]. Laboratory testing revealed LCMV in all four SOT recipients; however, extensive donor testing revealed no LCMV. Ultimately, no source for the recipients’ infection was identified [33]. Four SOT recipients were diagnosed with LCMV in 2005, three of whom died. Their infections were traced back to their donor, who died from an ischemic stroke but had contact with an asymptomatic infected pet hamster [33]. In 2008, the CDC investigated two SOT recipients with hepatic insufficiency, multiorgan failure, and death, which lead to the diagnosis of LCMV. The donor was homeless and died with an abnormal cerebrospinal fluid (CSF) analysis, emphasizing the risk that decedents with aseptic meningitis or encephalitis of unknown cause pose as donors [34]. The fourth cluster of donor-derived LCMV infections developed after the donor died from cerebral edema due to diabetic ketoacidosis. Interestingly, a corneal procurement was also performed from the donor, but no evidence of infection was ever detected in the recipient [35]. In 2013, the CDC reported a fifth cluster of SOT-derived LCMV from a donor who died from a large intracranial bleed [36]. One SOT recipient (liver) died, while the two kidney recipients survived with neurological sequelae and graft failure. A fourth cornea recipient
remained asymptomatic [36]. Overall, a prominent clinical feature of most patients with donor-derived LCMV infection is hepatitis [15]. Of the 20 SOT recipients reviewed, all become infected an average of 14.6 days (2-23 days) post-transplant (Table 2). All but five SOT recipients died as a result of their DDI. Laboratory diagnosis of LCMV is usually made by detecting IgM and IgG antibodies (CNS or serum) or by PCR or viral cultures (CNS) in a highly specialized viral pathogens laboratory (e.g., the CDC’s Viral Special Pathogens Branch). LCMV as a cause of DDI is under-recognized and likely underdiagnosed because of its rarity and nonspecific clinical characteristics. It is important for transplant providers to be aware that LCMV predisposes transplant recipients to severe, often fatal, disease, while infected donors may be asymptomatic. Low provider awareness, inaccessibility of commercial diagnostic testing, and no proven effective therapies result in delayed diagnoses and subsequent poor outcomes in SOT recipients with donor-derived LCMV.

In 2008, Palacios et al. described a cluster of SOT recipients in Australia who all died of a febrile illness approximately one month post-transplant (Table 2) [37]. All three received organs from a common donor who died of a cerebral haemorrhage ten days after returning from a trip to rural former Yugoslavia. Extensive infectious diseases work-up, including PCR assays for Lyssavirus, herpes viruses 1-8, Flavivirus, Alphavirus, hantavirus, Crimean-Congo hemorrhagic fever virus, and Rift Valley fever virus, among others, were negative [37]. Multiple tissues were extracted from the SOT recipients for unbiased high-throughput sequencing, which yielded a new candidate arenavirus related to LCMV [37].
Table 2 Solid organ transplant transmitted cases of lymphocytic choriomeningitis virus.

| Cluster Year, location | Donor Age/sex, COD | Donor risk-factor | Donor LCMV testing | Organs donated | Symptom onset after SOT (days) | Recipient LCMV serum/tissue testing | Recipient LCMV CSF/CNS testing | LCMV Treatment | Outcomes |
|-----------------------|--------------------|------------------|-------------------|----------------|-------------------------------|-------------------------------------|-------------------------------|----------------|---------|
| 2003, Wisconsin, USA [33] | 51/M, ICH | Unknown | Serum IgG, IgM (-); tissue IHC (-); viral culture (-) | Lung | 4 | IgM, IgG (-); tissue IHC (+) | Not done | None | Died |
|                      |                    |                  | Liver Early^b     | Liver          | 23                            | Tissue IHC (+)                       | Brain IHC (+)                  | None | Died |
|                      |                    |                  | Kidney            | Kidney         | 22                            | IgG, IgM (-); tissue IHC (+)         | Viral culture (+)               | None | Died |
|                      |                    |                  |                   | Liver          | Early^b                       | IgM (+), IgG (-); tissue IHC (+); viral culture (+) | Brain IHC (-) | Cidofovir, IVIG | Died |
|                      |                    |                  |                   |                |                              | Tissue IHC (+); PCR (+); IgG, IgM (+) |                          | None | Died |
| 2005, Rhode Island and Massachusetts, USA [33, 38] | 45/F, stroke and ICH | Infected pet hamster | Serum PCR (-); IgG, IgM (-); tissue IHC (-); viral culture (-) | Lung | 3 | IgG, IgM (-); IHC (+); PCR (+); viral culture (+) | None | Died |
|                      |                    |                  | Kidney            |                | 17                            | Tissue IHC (+); PCR (+); IgM (+), IgG (-); viral culture (+) |                           | None | Died |
| Year, Location          | Age, Sex, Condition                       | Route of Infection | Symptoms                                                                 | Treatment | Outcome |
|-------------------------|------------------------------------------|--------------------|---------------------------------------------------------------------------|-----------|---------|
| 2007, Australia\[37\]   | 57/M, ICH Travel to endemic area         | Serum and tissue   | PCR (-); IgM, IgG (+)                                                    | IV and oral ribavirin | Died    |
|                         |                                          | PCR (+); viral     | culture (+); IgM (+), IgG (-)                                            |            |         |
|                         |                                          | IHC (+); N/A       |                                                                            |            |         |
|                         |                                          | Liver              | PCR (+); IgG, IgM (+)                                                     | None      | Died    |
|                         |                                          |                   | PCR (+); IgG, PCR (-)                                                     | None      | Died    |
|                         |                                          | Kidney             | PCR (+); IGG, IgM (-)                                                     | Not tested| Died    |
|                         |                                          |                   | Not tested                                                                |            |         |
|                         |                                          | Kidney             | PCR (+); viral culture (+); Not tested                                    | None      | Died    |
| 2008, Massachusetts, USA\[34\] | 49/M, febrile encephalitis | Homeless, probable rodent exposure | Serum IgG, IgM (+)                                                        | IVIG, ribavirin | Died    |
|                         |                                          | Kidney             | IgM (+), viral culture (+); Not tested                                    |            |         |
|                         |                                          |                    | tissue IHC (+); Not tested                                                |            |         |
|                         |                                          | Kidney             | PCR (+)                                                                   |            |         |
|                         |                                          |                    |                                                                            |            |         |
| 2011, Arkansas, USA\[35\] | 13/F, Diabetic ketoacidosis and cerebral edema | Probable rodent exposure | LN PCR (+); serum IgG, IgM (-); cornea PCR (-), IHC (-)                 | None      | Survived |
|                         |                                          | Liver              | PCR (+); tissue IHC (+)                                                   |            |         |
|                         |                                          |                    |                                                                            |            |         |
|                         |                                          | Cornea             | IgG, IgM (-)                                                              | Not tested| None    |
|                         |                                          |                    |                                                                            |            |         |
| Organ          | PCR Status | Immunohistochemical Status | Test Status | Treatment | Outcome |
|----------------|------------|----------------------------|-------------|-----------|---------|
| Lung           | (+); tissue IHC (+) | Not tested                 | None        | Died      |
| Kidney         | 7          | PCR (+); IgG, IgM (-)      | Not tested  | None      | Died    |
| Kidney         | 2          | PCR (-); IgG, IgM (-); tissue IHC (-) | CSF PCR (+); IgG, IgM (-) | None | Survived |
| Cornea         | —          | PCR (-); IgG, IgM (-)      | Not tested  | None      | Survived |

| Organ          | PCR Status | Immunohistochemical Status | Test Status | Treatment | Outcome |
|----------------|------------|----------------------------|-------------|-----------|---------|
| Liver          | 20f        | IgM (+); PCR (+)           | Not tested  | IV ribavirin, IVIG | Died    |
| Kidney         | 20f        | IgM (+); PCR (+)           | Not tested  | Oral ribavirin | Survived |
| Kidney         | 20f        | IgM (+)                    | Not tested  | IV ribavirin, IVIG | Survived |

**2013 Iowa, USA [36]**

| Patient | Age/M | Cause of death | Disease Symptoms | Treatment | Outcome |
|---------|-------|----------------|------------------|-----------|---------|
| 49/M    | ICH   | Outdoor exposures | Body tissue excluding CNS tissue | Aortic endothelial cells PCR (+) | Died |

**Abbreviations.** COD, cause of death; LCMV, lymphocytic choriomeningitis virus; SOT, solid organ transplant; CSF, cerebrospinal fluid; CNS, central nervous system; USA, United States of America; F, female; M, male; IgM, immunoglobulin M; IgG, immunoglobulin G; IHC, immunohistochemical staining; ICH, intracranial hemorrhage; IVIG, intravenous immunoglobulin; PCR, polymerase chain reaction; IV, intravenous; N/A, not available; LN, lymph node; a Body tissue excluding CNS tissue; b Early unspecified - symptoms developed early post-transplant; c Test modality not specified; d Arenavirus related to LCMV; e asymptomatic; f Approximately 20 days post-transplant.
2.3 Rabies Virus

Rabies virus (RABV) is a neurotropic virus of the *Lyssavirus* genus with an almost worldwide distribution [12, 14]. Human infections occur after contact with saliva from the bite or scratch of an infected animal, most commonly a bat in the U.S. Thereafter, the virus is taken up by peripheral nerves and transported over a period of weeks to months to the CNS, where it causes uniformly fatal encephalitis in the host [12, 14]. Clinical manifestations of RABV evolve from a nonspecific prodrome of fever, malaise, and headache, followed by encephalopathy and paresthesias, and, finally, to hydrophobia, coma, and death [12]. Human-to-human transmission, which can occur through the utilization of contaminated tissues or organs (Table 3), was first described in the U.S. after a corneal transplant in 1978 and additional cases have since been reported [39-43]. Donor-derived RABV after SOT was not described until 2004 when four recipients (liver, two kidneys, and arterial segment) with a common donor developed encephalitis within 30 days of transplantation [44]. Prior to death, the donor had difficulty swallowing and soon after developed fevers and encephalopathy. He was discovered to be positive for cocaine use and brain imaging demonstrated an intracranial hemorrhage. During contact tracing, it was discovered that the donor had been bitten by a bat before falling ill [44]. Another cluster of donor-derived RABV was reported soon after in Germany [42]. The donor was an otherwise healthy young female who had been bitten by a dog while traveling in India and subsequently developed encephalopathy, which progressed rapidly to respiratory distress and cardiac arrest. Brain imaging showed massive cerebral edema and CSF analysis revealed aseptic meningitis. Corneal tissue and solid organs (liver, lung, kidneys, and pancreas) were transplanted. Three of the SOT recipients developed symptoms like the donor, prompting re-examination of archived tissues and established the diagnosis of RABV. Only the liver recipient, who had been previously vaccinated against RABV 20 years earlier, survived [42]. In 2013, a deceased-donor kidney recipient was admitted for fevers, progressive extremity weakness, paresthesias, encephalopathy, and excessive salivation 17 months post-transplant [45]. He had positive RABV serologies without prior history of vaccination or animal exposure; thus, DDI was considered. Retrospective review of the donor found that he had been bitten by a raccoon months before developing paresthesias, seizures, autonomic dysfunction, and death. Brain imaging was unremarkable at the time, and his CSF analysis revealed aseptic meningitis, but archived tissue analysis was positive for RABV viral RNA. The donor’s other SOT recipients (right kidney, heart, and liver) received post-exposure prophylaxis and the RABV vaccine, developed neutralizing antibody response, and remained asymptomatic [45, 46]. RABV transmission through SOT has also been reported in China recently. Between 2015 and 2017, four SOT recipients were diagnosed with RABV, presumably transmitted from two donors who died from viral encephalitis of unknown etiology and acute disseminated encephalomyelitis (ADEM) [47-49]. A cluster of probable donor-derived RABV was also reported in China in 2016, which led to the death of two kidney recipients. The liver recipient from the same donor, who died of pneumonia and multi-organ failure, did not develop signs or symptoms of RABV encephalitis and was not tested for RABV [50]. Probable donor-derived RABV transmitted through SOT has also been reported recently in Kuwait [51, 52]. Of 24 reported SOT recipients with proven or probable donor-derived RABV, 17 developed symptoms an average of 118 days (20-580 days) post-transplant and all 17 died as a result (Table 3). RABV encephalitis is uniformly fatal and, thus,
should be considered in SOT recipients presenting with encephalitis of unknown etiology, even months after transplantation, in order to initiate post-exposure prophylaxis (i.e., human rabies immune globulin and inactivated rabies vaccine) as soon as possible in related recipients. No single RABV test is sufficient for establishing the diagnosis antemortem [53]. Highly specialized laboratories can use saliva (PCR or viral culture), serum (PCR or viral culture), CSF (antibody testing), or skin biopsies (RABV antigen) containing cutaneous nerves to detect RABV. Several of these reported cases highlight the omission of historical donor data by transplant teams and family members and emphasize the importance of having a centralized reporting system for potential DDI, such as UNOS. Considering the high rates of RABV transmission and, if unrecognized, death, potential donors with progressive neurological symptoms or encephalitis of unknown cause should be screened with extreme prejudice.
Table 3 Solid organ transplant transmitted cases of rabies virus.

| Year, location | Donor Age/sex, COD | Donor risk-factor | Donor testing | RABV Organ donated | Symptom onset after SOT (days) | Recipient serum/tissuea testing | RABV CSF/CNS testing | RABV Treatment | Outcome |
|----------------|--------------------|-------------------|--------------|--------------------|-------------------------------|-------------------------------|------------------------|---------------|---------|
| 2004, Texas, USA [44] | N/A, Encephalopathy, ICH | Bat bite | Serum IgM, IgG (+) | Liver | Earlyb | Tissue IHC IgM, IgG (+); Tissue IHC IgM, IgG (+) | N/A | None | Died |
| | | | | Kidney | Earlyb | Tissue IHC IgM, IgG (-) | EM rhabdovirus particles; IHC (+) | None | Died |
| | | | | Arterial graft | N/A | Tissue IHC (+); IgM (-), IgG (+) | N/A | None | Died |
| | | | | Cornea | — | PCR (-) | Not tested | HRIG + RABV vaccine | Survived |
| 2005, Germany [42] | 26/F, Aseptic meningitis, cerebral edema | Dog bite in India | Brain tissue EM (+) rhabdovirus particles and DFA (+) | Lung | 20 | PCR (+); viral culture (+) | PCR (+) | HRIG + RABV vaccine, INF-a, IV ribavirin | Died |
| | | | | Kidney | 35 | PCR (+) | PCR (+) | HRIG + RABV vaccine, ribavirin, INF-a, amantadine | Died |
| Year | Country | Sex | Age | Cause | Dogs | Rabies Vaccine | Death | Description |
|------|---------|-----|-----|-------|------|----------------|-------|-------------|
| 2013 | USA     | Male | 50 years | Seizure, autonomic dysfunction, misdiagnosed with ciguatera poisoning | Racoon bite | HRIG + RABV vaccine, ribavirin, INF-a, amantadine, INF-a-2a | Died | Kidney + pancreas PCR (+); viral culture (+) PCR (+) | 35 months |
| 2015 | China   | Boy | 6 years | Viral encephalitis of unknown cause | Dog contact | HRIG + RABV vaccine | Survived | Kidney PCR (+) | 44 months |
| 2016 | China   | Boy | 11 years | Viral encephalitis, screened for rabies (ELISA neg) | Unknown | HRIG + RABV vaccine | Died | Kidney PCR (+) | 40 months |
| 2017 | China   | 11/F | ADEM | Unknown | Unknown | HRIG + RABV vaccine | Died | Kidney PCR (+) | 10 months |

Note: HRIG = Human Rabies Immune Globulin
RABV = Rabies Virus
PCR = Polymerase Chain Reaction
IgG = Immunoglobulin G
IgM = Immunoglobulin M
IHC = Immunohistochemistry
INF-a = Interferon alpha
INF-a-2a = Interferon alpha-2a
CNS = Central Nervous System
| Year | Source | Gender | Cause of Death | Dog Bite Location | Cornea<sup>cd</sup> | Explanted corneas PCR (+) | HRIG + RABV Vaccine | Survived |
|------|--------|--------|----------------|-------------------|---------------------|--------------------------|----------------------|----------|
| 2017 | Kwait<sup>f</sup> | Male, cardio-pulmonary arrest | Dog bite in India | N/A | — | Not tested | None | Died |
|      |        |        |                |                   | Kidney 3.5 months | IgM, IgG (-) | N/A | Died |
|      |        |        |                |                   | Kidney 3 months | N/A | N/A | Died |
|      |        |        |                |                   | Heart N/A | N/A | N/A | Died |
|      |        |        |                |                   | Liver N/A | N/A | N/A | Died |

Abbreviations. COD, cause of death; RABV, rabies virus; SOT, solid organ transplant; CSF, cerebrospinal fluid; CNS, central nervous system; USA, United States of America; N/A, not available; ICH, intracranial hemorrhage; EM, electron microscopy; F, female; M, male; IgM, immunoglobulin M; IgG, immunoglobulin G; IHC, immunohistochemical staining; PCR, polymerase chain reaction; DFA, direct fluorescence antibody; Ab, antibody; INF-a, interferon alfa; IV, intravenous; ADEM, acute disseminated encephalomyelitis; ELISA, enzyme-linked immunosorbent assay.

<sup>a</sup> Body tissue excluding CNS tissue; <sup>b</sup> Symptoms developed within 30 days post-transplant; <sup>c</sup> Two corneas transplanted into two recipients and reported results apply to both recipients; <sup>d</sup> Asymptomatic; <sup>e</sup> Archived serum: patient had history of prior RABV vaccination 20-years prior; <sup>f</sup> Probable RABV; <sup>g</sup> Died from pneumonia.
2.4 Tick-Borne Encephalitis Virus

Tick-borne encephalitis virus (TBEV) is a tick-borne Flavivirus found throughout Europe and northern Asia, where it has become a significant health problem [14, 54]. Infection with TBEV ranges from asymptomatic to severe encephalitis and death, and diagnosis is typically confirmed by serum and CSF serology [14, 54]. The virus is maintained in Ixodes ticks and cycles through rodents but can infect humans through either an infected tick bite or consumption of contaminated foods (e.g., raw goat’s milk) in <1% of cases [14]. Few cases of SOT-derived TBEV transmission have been reported [54]. In 2012, a cluster of donor-derived TBEV was reported in Poland, which resulted in the death of three recipients (two kidneys and liver). The same viral strain of TBEV was found in all recipients (brain tissue or CSF PCR) and the donor (brain tissue PCR), who resided in an endemic area but died as a result of trauma after a motor vehicle accident. All three recipients presented to the hospital within one-to-two months after transplantation with meningeal signs and fever [54]. Two patients in this cluster were treated with acyclovir for possible herpes encephalitis; unfortunately, there is no specific proven drug therapy for TBEV. Although rare, transplant providers should consider screening for TBEV in donors who have a history of residing or travel to areas of endemicity during warm seasons.

3. Parasitic Pathogens

3.1 Primary Amebic Meningoencephalitis

Organ procurement from SOT donors dying from a febrile encephalitis without a documented cause is frequently associated with disease transmission and, thus, should be avoided [5]. A notable exception to this is a SOT donor with proven Naegleria fowleri meningoencephalitis [5]. N. fowleri is a small, free-living ameba found worldwide, usually in bodies of warm water [55, 56]. It is the etiologic agent of the devastating and rapidly fatal primary amebic meningoencephalitis (PAM), which presents with fever, headache, meningismus, and rapid neurological deterioration [55]. In 95% of cases, death occurs within a few days of presentation [55]. In 1997, Kramer et al. reported the first case of SOT from a donor who died of undiagnosed PAM, and no subsequent infection occurred in the recipients (two kidneys and liver) despite not receiving prophylactic antiamebic therapy [56]. In 2008, organs (kidneys, pancreas, lung, and liver) from a donor known to have died from PAM were transplanted with no post-transplant infectious complications at six months [55]. Because N. fowleri is exclusive to the CNS and does not have systemic manifestation, transmission through organ transplantation does not occur and should not preclude organ donation [5, 55].

3.2 Granulomatous Amebic Encephalitis

Balamuthia mandrillaris is a small, free-living amebae found ubiquitously in soil worldwide and is the cause of fatal granulomatous amebic encephalitis (GAE) in humans [12, 57]. Infection is characterized by space-occupying brain lesions leading to neurological deficits [12]. Optimal treatment of GAE is not known [57]. Infection occurs through the inhalation of contaminated dust or inoculation through breaks in the skin and infection can present with skin lesions, followed months to years later by GAE [57]. Diagnosis is often made only after death; however, three tests
can establish the diagnosis: serum indirect immunofluorescence assay (IFA), immunohistochemistry (IHC), and indirect immunofluorescence (IIF) staining [58]. Rarely, cases of SOT donor-derived GAE have been reported (Table 4) [57]. In 2009, the CDC reported cases of encephalitis among two kidney recipients from a common donor [59]. The donor was a four-year-old child who often played outdoors and who had succumbed to a subarachnoid hemorrhage following a transient febrile illness, which was diagnosed as post-influenza ADEM. Prior to death, brain imaging showed numerous small enhancing lesions and CSF analysis revealed aseptic meningitis. Post-mortem brain examination revealed *B. mandrillaris* amebae and all SOT recipients were empirically treated. One kidney recipient died despite therapy, while the other survived with significant neurological sequelae. A heart and liver recipient, who was also empirically treated, never developed signs or symptoms of GAE [59]. A second cluster of donor-derived GAE was reported by the CDC in 2010 [60]. The donor was a young male who worked in landscaping, died from a stroke, and had a notably large skin lesion on his back that had been present for six months. Two of four SOT recipients (liver, kidney, and pancreas) from this common donor developed neurological symptoms and GAE was diagnosed on brain biopsies. Despite the initiation of therapy, both recipients died, while the heart and kidney recipients remained asymptomatic after pre-emptive therapy [60]. It is possible that GAE is more common than thought but under-diagnosed in SOT donors with encephalitis of unknown etiology. In 2013, a third cluster was reported, involving five SOT recipients across three U.S. states from a young donor who died from traumatic head injuries [61, 62]. Diagnosis of GAE was made in the liver recipient who experienced a rapid neurologic decline and died within one month of transplantation. Multi-drug pre-emptive therapies (Table 4) were immediately started in the remaining four SOT recipients who remained asymptomatic with positive *B. mandrillaris* antibody titers, suggesting early diagnosis and therapeutic intervention may prevent fatal GAE [62]. Of the 12 reports of SOT recipients from donors who died with GAE, all but three were infected and four subsequently died (Table 4). Symptoms developed an average of 18.8 days (17-26 days) post-transplantation, suggesting an accelerated course in SOT recipients.
### Table 4 Solid organ transplant transmitted cases of *Balamuthia mandrillaris*.

| Clusters Year, location | Donor Age/sex, COD | Donor risk-factor | Donor GAE testing | Organs donated | Symptom onset after SOT (days) | Recipient GAE serum/tissue testing | Recipient GAE CSF/CNS testing | GAE Treatment | Outcomes |
|-------------------------|-------------------|------------------|-------------------|----------------|-------------------------------|---------------------------------|-------------------------------|---------------|----------|
| 2009, Kentucky and Mississippi, USA [57, 59] | 4/M, post-influenza ADEM, ICH | Soil exposure | Brain tissue IHC (+); PCR (+) | Kidney | 20 | Not tested | HP (+) ameba; IHC (+); PCR (+) CSF PCR (+); culture (+) | PEN, SDZ, FLY, FLC, AZT, MLF | Died |
| | | | | Kidney | 20 | Not tested | Serum (-); tissue (-) | CSF (-) | PEN, SDZ, FLY, FLC, AZT, MLF | Survived |
| | | | | Heart\(^b\) | | | Serum (-); tissue (-) | CSF (-) | PEN, AZT, FLC | Not infected |
| | | | | Liver\(^b\) | | | Serum (-); tissue (-) | CSF (-) | PEN, AZT, FLC, SDZ | Not infected |
| | | | | Liver | 17 | IHC (+) | IHC (+); PCR (+) | SDZ, PYR, AMB | Died |
| 2010, Arizona, USA [57, 60] | 27/M | Landscaper | Ab titer (+) | Kidney + pancreas | 19 | IHC (-), Ab titer (+) | HP (+); IHC (+); culture (+); PCR (+) | MLF, AMB, ALB, FLC, AZT | Died |
| | | | | Kidney\(^b\) | | | Tissue HP, IHC, IFA (-); Ab titer (+) | Not tested | PEN, FLC, AZT, MLF, TMP/SMZ\(^c\) | Survived |
| | | | | Heart\(^b\) | | | Tissue HP, IHC, IFA (-); Ab titer (-) | Not tested | PEN, FLC, AZT, SDZ, MLF\(^c\) | Not infected |
| 2013, Alabama, Homeless | 39/M, | Homeless | Ab titer | Liver | 18 | Positive\(^d\) | CSF (-) | None | Died |
| Location       | Seizures | Kidney | Heart | Vascular Graft | Ab titer | CSF | MLF, AZT, ALB, SDZ, FLC | Survived |
|----------------|----------|--------|-------|----------------|----------|-----|-------------------------|----------|
| Florida, Texas, USA [61] | (+)      | —      | —     | —              | Ab titer (+) | CSF (-) | MLF, AZT, ALB, SDZ, FLC | Survived |
|                |          |        |       |                |           |      |                         |          |

Abbreviations. COD, cause of death; GAE, granulomatous amebic encephalitis caused by *B. mandrillaris*; SOT, solid organ transplant; CSF, cerebrospinal fluid; CNS, central nervous system; USA, United States of America; F, female; M, male; ADEM, acute disseminated encephalomyelitis; HP, histopathologic tissue examination, ICH, intracranial hemorrhage; PCR, polymerase chain reaction; IHC, immunohistochemical staining; PEN, pentamidine; SDZ, sulfadiazine; FLY, flucytosine; FLC, fluconazole; AZT, azithromycin; MLF, miltefosine; AMB, amphotericin; PYR, pyrimethamine; ALB, albendazole; TMP/SMZ, trimethoprim/sulfamethoxazole; IFA, indirect immunofluorescence assay; Ab, antibody. a Body tissue excluding CNS tissue; b Asymptomatic; c Preemptive therapy; d Diagnosis of GAE made on autopsy and confirmed by CDC using unspecified specimens and tests.
3.3 Microsporidiosis

Microsporidia are a diverse group of intracellular, spore-forming parasites closely related to fungi and their taxonomic classification has been heavily revised and debated numerous times [63, 64]. Fifteen microsporidian species have been identified as human pathogens, including *Encephalitozoon cuniculi* [64, 65]. Based on seroprevalence data, the most common presentation of microsporidiosis may be asymptomatic disease [65]. Generally, microsporidiosis causes a spectrum of disease ranging from self-limiting gastrointestinal disease (e.g., diarrhea and malabsorption) to life-threatening disseminated infections [65]. This infection has been well characterized in HIV-infected patients; however, recently, SOT donor-derived infections have also been reported. In 2014, a kidney recipient developed headache, encephalopathy, and rapid neurologic decline and died [63]. *E. cuniculi* was detected by PCR in the CNS tissue of the deceased transplant recipient. Two other recipients from the common donor developed neurologic symptoms and encephalitis, but no gastrointestinal symptoms were reported and prompt initiation of treatment with albendazole led to their recovery. The three SOT recipients developed symptoms an average of 72.3 days (56-91 days) post-transplantation. The donor was a middle-aged female who died after multiple endovascular repairs for arteriovenous malformations. A retrospective analysis of the donor’s archived serum indicated an active *E. cuniculi* infection at the time of her death, but no gastrointestinal symptoms were reported [63]. Unlike another reported cluster of microsporidiosis after SOT [65], this cluster described donor-derived microsporidiosis infection in three SOT recipients that led to the rare presentation of disseminated neurological disease without gastrointestinal symptoms [63]. This points to a need for transplant providers to maintain awareness of microsporidiosis as a possible cause of donor-derived CNS infections in SOT recipients with encephalitis and emphasizes that prompt recognition and treatment is essential for optimal SOT-recipient outcomes.

4. Prions

4.1 Transmissible Spongiform Encephalopathy

Transmissible spongiform encephalopathy (TSE) is a general term for prion-related diseases caused by the misfolded proteins, which became well known in the 1980s due to a sudden increase in the incidence of bovine spongiform encephalopathy (BSE) or mad cow disease [66]. Creutzfeldt-Jakob disease (CJD) and variant Creutzfeldt-Jakob disease (vCJD) (i.e., BSE in humans) are rare, uniformly fatal, transmissible, neurodegenerative prion diseases with no known treatment [66, 67]. Clinically, these diseases are characterized by progressive dementia and neuron loss, with an average latency period of 10 years [67]. While the majority of CJD has been acquired through inheritance or sporadically, numerous reports have documented its transmission through contaminated tissues, such as dura mater grafts [68] and corneas [67], among others. Between 1974 and 2006, ten cases of corneal transplant-transmitted CJD have been reported [67]. Rare reports of possible SOT-transmitted CJD have been published, including a liver recipient who died of CJD two years after transplantation [69] and a kidney recipient who died with CJD [70]. The liver recipient’s donor died of a cerebral aneurysm and had no known history of neurological disease, but there was speculation that the donor received TSE-contaminated pooled plasma [69].
The kidney recipient’s donor history was not reported, but another kidney recipient from the same donor died as well after a seizure [70]. In 2014, a retrospective study was conducted in the United Kingdom to look for organ or tissue-associated TSE but found no evidence of transplant-transmitted vCJD [71, 72]. To date, no proven donor-derived CJD or vCJD has been reported in SOT recipients; however, transplant providers assessing transplant recipients with progressive neurologic symptoms or encephalopathy should consider donor-derived TSE as a possibility, even if rare, as the infection control concerns would be enormous. Emphasis should be placed on the use of donor registries and screening tools to detect and prevent potential TSE-containing tissues and organs.

5. Discussion

5.1 Recognizing and Reporting

Unexpected DDI are infrequent and DDI with CNS pathogens are exceedingly rare at <0.05% of reported events [10]. Nonetheless, they account for medically important and devastating causes of DDI and poor outcomes in SOT recipients. We reviewed the published literature on DDI with rare CNS pathogens, including arboviruses (EEEV and WNC), arenaviruses (LCMV), TBEV, RABV, PAM, GAE, microsporidiosis, and prions, and we excluded more common forms of CNS infection (e.g., bacterial meningitis, cryptococcosis, and toxoplasmosis).

The clinical course and preceding events of donors are not always clearly defined and transplant providers should be aware of the risk of unexpected disease transmission in cases exhibiting “warning signs” [10]. These include young deceased donors who died with cerebral vascular accidents, seizures, or CNS imaging abnormalities, potential SOT donors who died with a febrile encephalitis of unknown cause, and potential donors who died from traumatic events with high-risk exposures. Likewise, transplant providers should maintain a high index of clinical suspicion for DDI with CNS pathogens in any SOT recipient presenting with atypical or neurological symptoms of unknown etiology. It is imperative to recognize that these pathogens often do not follow a normal timeline (i.e., incubation) and SOT recipients with DDI such as CNS pathogens can present at any time after transplantation, even years later [73].

As demonstrated by several of the cases discussed in this review, outside of routine screening, individual transplant centers and national regulations can vary in the extent of infectious disease investigation performed on potential SOT donors. Candidates should be evaluated with a thorough investigation of their medical (e.g., prior infections, vaccinations), social (e.g., drug use, sexual practices, incarceration), travel (e.g., itinerary, duration), and exposure (e.g., wild or domestic animals, occupation, areas of residence, hobbies) histories [74]. This historical data should be cross-referenced with information from donors’ friends, family, or legal next-of-kin. Decisions to screen donors for the pathogens covered in this review should be driven by endemicity and their history [10]. In the U.S., OPTN/UNOS has mandated that deceased donors are tested for cytomegalovirus (CMV), Epstein-Barr virus (EBV), human immunodeficiency virus (HIV), hepatitis B (HBV) and C (HCV) viruses, syphilis, and toxoplasmosis, in addition to routine cultures [74]. Testing of living donors includes CMV, EBV, HIV, HBV, *Mycobacterium tuberculosis*, toxoplasmosis, and syphilis. Testing for seasonal or geographical pathogens, such as WNV, *Strongyloides*, *Coccidioides*, and Chagas, should be done on a case-by-case basis based on history [74]. Lastly, reporting a potential DDI to UNOS should be completed with haste and not delayed by testing.
5.2 Clinical Pearls for Assessing and Reporting Donor-Derived CNS Infections

- Any SOT recipient with post-transplant encephalitis or neurological symptoms at any time following transplantation should be worked up for possible donor-derived CNS infection.
- Any suspected donor-derived infection should be reported to the relevant OPO and national transplant authority (UNOS in the U.S.) and should not wait for test results.
- Infectious Diseases consultation and public health involvement is strongly recommended.
- Presumptive donor-derived encephalitis workup should include a detailed timeline of symptoms prior to the donor’s death and history of local exposures, travel history, house-hold pet and wild animal exposures, review of complete vaccination history, hobbies, and occupation.
- Targeted testing of donor archived tissues should be completed with assistance from the CDC and the CDC’s Division of High-Consequence Pathogens and Pathology (DHCPP): [https://www.cdc.gov/ncezid/dhcpp/index.html](https://www.cdc.gov/ncezid/dhcpp/index.html).

6. Conclusion

As demonstrated in this review, recognizing potential donor-derived CNS infections is exceptionally difficult, but it is of the highest clinical importance. Thus, the OPTN’s policy requires Organ Procurement Organizations (OPOs) and organ transplant hospitals in the U.S. to report any concern of a DDI transmission event to the OPTN’s Patient Safety System. To assist transplant providers with the difficult task of identifying donors at high risk of transmitting CNS pathogens, the ad hoc DTAC created guidance to highlight indicators of possible at-risk donors [8, 10, 75]. This guidance also includes questions that OPOs and transplant providers should consider when completing the screening of a potential donor [75]. Utilization of such resources and careful clinical assessment, combined with a high index of suspicion, of potential SOT donors with ambiguous or confounding diagnoses will undoubtedly reduce the incidence of DDI with these often fatal CNS pathogens.

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Author Contributions

All authors contributed to this manuscript equally.

Competing Interests

The authors have declared that no competing interests exist.

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