

*Briefing to the OPTN Board of Directors on*

# **Recognizing Seasonal and Geographically Endemic Infections in Organ Donors: Considerations for Deceased and Living Donation**

*OPTN Ad Hoc Disease Transmission Advisory Committee*

*Prepared by: Taylor Livelli  
UNOS Policy Department*

## **Contents**

Executive Summary	2
Purpose	3
Background	3
Overall Sentiment from Public Comment	6
Public Comment Themes and Considerations	9
Recommendations	11
Compliance Analysis	15
Conclusion	15
Guidance Document	16
Appendix A: Post-Public Comment Changes	45

# Recognizing Seasonal and Geographically Endemic Infections in Organ Donors: Considerations for Deceased and Living Donation

*Sponsoring Committee:* Ad Hoc Disease Transmission Advisory  
*Public Comment Period:* July 27, 2023 – September 19, 2023  
*Board of Directors Meeting:* December 4, 2023

## Executive Summary

The OPTN Ad Hoc Disease Transmission Advisory Committee (the Committee) is refreshing and combining existing guidance documents for endemic diseases for living and deceased donors. OPTN *Policy 14.4: Medical Evaluation Requirements for Living Donors* requires each living donor hospital to develop and follow a written protocol for identifying and testing donors at risk for transmissible seasonal or geographically defined endemic disease as part of its medical evaluation.<sup>1</sup> This requirement aims to minimize disease transmission through organ transplantation and promote living donor and transplant recipient safety. The Committee created four guidance documents to help living donor hospitals comply with this policy. The Committee reviewed these documents after the Centers for Disease Control and Prevention (CDC) representatives and the OPTN Membership and Professional Standards Committee (MPSC) requested the Committee provide clarity on endemic diseases. The Committee updated and condensed these documents while also adding guidance for deceased donors.

The four guidance documents that will be updated and condensed are:

- *Recognizing Seasonal and Geographically Endemic Infections in Organ Donors: Considerations during Living Donor Evaluation*
- *Preventing Chagas disease in transplant recipients: Donor screening and recipient monitoring*
- *Identifying Risk Factors for West Nile Virus (WNV) During Evaluation of Potential Living Donors*
- *Guidance for Identifying Risk Factors for Mycobacterium tuberculosis (MTB) During Evaluation of Potential Living Kidney Donors*

These documents were implemented from 2012 to 2014 and need nomenclature, testing, and potential donor-derived transmission events (PDDTE) data updates. The revised document will incorporate guidance for screening for endemic diseases for deceased and living donors. It also advises transplant hospitals on recipient and living donor management after transplanting organs from donors with endemic diseases. The revision includes updated epidemiology and the addition of sections on testing turnaround time.

---

<sup>1</sup> OPTN Policy 14.4: Medical Evaluation Requirements for Living Donors (Accessed May 24, 2023)  
[https://optn.transplant.hrsa.gov/media/eavh5bf3/optn\\_policies.pdf](https://optn.transplant.hrsa.gov/media/eavh5bf3/optn_policies.pdf).

The revision of these documents aims to maintain transplant recipient and living donor safety through infectious disease testing by minimizing disease transmission through organ transplantation. The Committee will evaluate the success of this guidance document by monitoring donor-derived transmission events.

This guidance document was issued for public comment from July 27, 2023 to September 19, 2023. The Committee reviewed the public comments and made changes to the document to incorporate feedback, discussed below.

## Purpose

This guidance document aims to decrease donor-derived disease transmissions from organ transplantation. Endemic diseases have high potential for morbidity and possible mortality if transmitted to recipients. As organ offer patterns continue to change, increased awareness and communication for potential endemic diseases across regions is necessary to mitigate risks.

## Background

In 2022, the Endemic Diseases Subcommittee of the Committee reviewed the potential gaps in education and policy regarding certain endemic diseases that present significant patient safety risks, and for which identification and treatment strategies exist but are not in common use. Diseases are endemic to a region when they are consistently prevalent in that specific area.<sup>2</sup> The Subcommittee included subject matter experts from the CDC and the Food and Drug Administration (FDA), along with Committee representatives from organ procurement organizations (OPOs). The MPSC wrote a memo to the Committee on opportunities to improve patient safety, noting concerns regarding a lack of standardized testing and reporting practices of OPOs for endemic diseases.<sup>3</sup> The Committee received support from the OPTN OPO and Operations and Safety Committees for this guidance document.

The document provides guidance for six endemic diseases: Histoplasmosis, Coccidioidomycosis, Strongyloidiasis, Chagas disease, Tuberculosis, and West Nile Virus. It advises transplant hospitals and OPOs on how to screen, who to screen, and management of infected living donors and transplant recipients.

## Histoplasmosis

*Histoplasma capsulatum* is a dimorphic fungus found throughout the world. It is endemic in the United States (U.S.) in the Ohio and Mississippi River Valley and into the Midwest and South-Central states.<sup>4</sup> In most cases, clinical signs of infection in immunocompetent patients are limited to the lungs, though many patients are asymptomatic. Immunocompromised patients may develop severe disseminated disease that is fatal without prompt diagnosis and treatment. Donor-derived histoplasmosis has only rarely been reported.<sup>5</sup>

---

<sup>2</sup> Centers for Disease Control and Prevention. "Principles of Epidemiology." Lesson 1: Introduction to Epidemiology, May 18, 2012. <https://www.cdc.gov/csels/dsepd/ss1978/lesson1/section11.html>.

<sup>3</sup> Ad Hoc Disease Transmission Advisory Committee, OPTN, meeting summary for February 14, 2022, accessed April 13, 2023, [https://optn.transplant.hrsa.gov/media/jn4p42r3/2022\\_02\\_14\\_dtac\\_open\\_summary.pdf](https://optn.transplant.hrsa.gov/media/jn4p42r3/2022_02_14_dtac_open_summary.pdf).

<sup>4</sup> Akram SM, Koirala J. Histoplasmosis. [Updated 2023 Feb 25]. In: StatPearls [Internet]. Treasure Island (FL): StatPearls Publishing; 2023 Jan-. Available from: <https://www.ncbi.nlm.nih.gov/books/NBK448185/>

<sup>5</sup> Ibid.

## Coccidioidomycosis

*Coccidioides immitis* and *Coccidioides posadasii* are dimorphic fungi endemic in arid and semi-arid climates in Washington state and the southwestern part of the U.S., Northern Mexico including areas along the U.S. border, and parts of Central and South America.<sup>6</sup> The most common clinical syndrome is pneumonia often accompanied by joint pains, fatigue, and weight loss. Chronic fibrocavitary pulmonary disease may follow primary infection, and some infected persons have persistent symptomatic pulmonary nodules, reticulonodular disease, or cavitary disease. Immunosuppressed individuals may develop disseminated infection.<sup>7</sup> Multiple cases of donor-derived coccidioidomycosis have been reported, often with poor outcomes.

## Strongyloidiasis

*Strongyloides stercoralis* is a nematode (roundworm). Unlike other parasites, *Strongyloides* can replicate for decades within humans via an autoinfection cycle, resulting in chronic symptomatic or asymptomatic infections that can flourish in the setting of immunosuppression. There are an estimated 370 million infected people worldwide.<sup>8</sup>

*Strongyloides* is most common in tropical, subtropical, or warm temperate climates, including the Caribbean, Mexico, South and Central America, Africa, Southeast Asia, Southern and Eastern Europe, and the Southeastern U.S.<sup>9</sup> In the U.S., a prevalence of up to 6.1% has been reported in select populations, with a much higher prevalence of up to 46.1% in immigrant populations.<sup>10</sup> Infection results primarily from exposed skin contact with soil that is contaminated with *Strongyloides* larvae. Activities that increase the risk of becoming infected include direct contact with contaminated soil (i.e., walking barefoot), unprotected skin contact with human waste or sewage, and occupations that increase contact with contaminated soil, such as farming and coal mining. Rural populations and those of lower socioeconomic status are at higher risk.<sup>11</sup>

## Chagas Disease

Chagas disease is caused by infection with the protozoan parasite *Trypanosoma cruzi* (*T. cruzi*). It is estimated that eight million people are infected in the Americas, with over 300,000 infected individuals currently living in the U.S. Most infections were acquired in endemic regions of Mexico, South and Central America, where the infection is transmitted by the bite of an infected triatomine bug. Infection can also be transmitted from an infected mother during the second and third trimester of pregnancy or

<sup>6</sup> Chow NA, Kangiser D, Gade L, McCotter OZ, Hurst S, Salamone A, Wohrle R, Clifford W, Kim S, Salah Z, Oltean HN, Plumlee GS, Litvintseva AP. Factors Influencing Distribution of *Coccidioides immitis* in Soil, Washington State, 2016. mSphere. 2021 Dec 22;6(6):e0059821. doi: 10.1128/mSphere.00598-21. Epub 2021 Nov 3. PMID: 34730378; PMCID: PMC8565518.

<sup>7</sup> Akram SM, Koirala J. Coccidioidomycosis. [Updated 2023 Feb 25]. In: StatPearls [Internet]. Treasure Island (FL): StatPearls Publishing; 2023 Jan-. Available from: <https://www.ncbi.nlm.nih.gov/books/NBK448161/>.

<sup>8</sup> Mora Carpio AL, Meseeha M. *Strongyloides Stercoralis*. [Updated 2023 Feb 7]. In: StatPearls [Internet]. Treasure Island (FL): StatPearls Publishing; 2023 Jan-. Available from: <https://www.ncbi.nlm.nih.gov/books/NBK436024/>.

<sup>9</sup> Schär F, Trostorf U, Giardina F, Khieu V, Muth S, Marti H, Vounatsou P, Odermatt P. *Strongyloides stercoralis*: Global Distribution and Risk Factors. PLoS Negl Trop Dis. 2013 Jul 11;7(7):e2288. doi: 10.1371/journal.pntd.0002288. PMID: 23875033; PMCID: PMC3708837.

<sup>10</sup> Global Health, Division of Parasitic Diseases and Malaria. "CDC - *Strongyloides* - Epidemiology & Risk Factors." Parasites-*Strongyloides*, December 31, 2018. <https://www.cdc.gov/parasites/strongyloides/epi.html>.

<sup>11</sup> Ibid.

rarely during childbirth, as well as through infected food or drink, and through blood transfusion and organ transplantation.<sup>12</sup>

Most infections are acquired in childhood during residence in an endemic area. The acute phase of infection may be associated with a mild febrile illness and is often unrecognized.<sup>13</sup> Parasitemia clears within eight to twelve weeks without treatment, but individuals remain chronically infected. Infection persists for the individual's life, with chronic disease developing in 20-30%, usually manifesting as cardiac or gastrointestinal disease. Chronic infection can be diagnosed through serologic *T. cruzi* tests.<sup>14</sup>

Blood donor screening in the United States beginning in 2007 identified confirmed infections in donors from 37 states and Puerto Rico, with 57% of all positive tests coming from California and Florida, areas with significant Mexican, Central, and South American immigrant populations.<sup>15</sup> Latent infection with *T. cruzi* can persist for decades, therefore organ donor screening of high-risk individuals using serology testing is utilized at some centers. Transmission of Chagas disease has been studied in 32 organ transplant recipients from 14 seropositive donors in the U.S. Transmission was confirmed in nine recipients from six donors, including three of four (75%) heart transplant recipients, two of ten (20%) liver recipients and two of 15 (13%) kidney recipients. Living organ donor transmissions have been reported in Mexico and South America.<sup>16</sup>

## Tuberculosis

Up to one-third of the world's population is infected with mycobacterium tuberculosis (MTB); however, infection in the U.S. is much less common.<sup>17</sup> Reported cases of MTB disease have been declining in the U.S. since 1992 with most cases occurring in foreign born persons.<sup>18</sup> After initial infection with MTB, most people do not develop MTB disease; the infection disseminates throughout the body and remains dormant. This condition is called tuberculosis infection (TBI) (previously referred to as "latent tuberculosis infection"). In 2022, there were 8,300 reported TB cases<sup>19</sup> in the United States (a rate of 2.5

<sup>12</sup> Global Health, Division of Parasitic Diseases and Malaria. "CDC - Chagas Disease - Epidemiology & Risk Factors." Parasites- American Trypanosomiasis (also known as Chagas Disease), April 11, 2022. <https://www.cdc.gov/parasites/chagas/epi.html>.

<sup>12</sup> Bern C, Kjos S, Yabsley MJ, Montgomery SP. Trypanosoma cruzi and Chagas' Disease in the United States. Clin Microbiol Rev. 2011 Oct;24(4):655-81. doi: 10.1128/CMR.00005-11. PMID: 21976603; PMCID: PMC3194829. - American Trypanosomiasis (also known as Chagas Disease), April 11, 2022. <https://www.cdc.gov/parasites/chagas/epi.html>.

<sup>13</sup> Bern C, Kjos S, Yabsley MJ, Montgomery SP. Trypanosoma cruzi and Chagas' Disease in the United States. Clin Microbiol Rev. 2011 Oct;24(4):655-81. doi: 10.1128/CMR.00005-11. PMID: 21976603; PMCID: PMC3194829.

<sup>14</sup> Ibid.

<sup>15</sup> Chin-Hong, P.V., B.S. Schwartz, C. Bern, S.P. Montgomery, S. Kontak, B. Kubak, M.I. Morris, M. Nowicki, C. Wright, and M.G. Ison. "Screening and Treatment of Chagas Disease in Organ Transplant Recipients in the United States: Recommendations from the Chagas in Transplant Working Group." American Journal of Transplantation 11, no. 4 (January 3, 2011): 672-80. <https://doi.org/10.1111/j.1600-6143.2011.03444.x>.

<sup>16</sup> M.E. Levi, D. Kumar, M. Green, M.G. Ison, D. Kaul, M.G. Michaels, M.I. Morris, B.S. Schwartz, I.A. Echenique, E.A. Blumberg, Considerations for Screening Live Kidney Donors for Endemic Infections: A Viewpoint on the UNOS Policy, American Journal of Transplantation, Volume 14, Issue 5, 2014, Pages 1003-1011, ISSN 1600-6135, <https://doi.org/10.1111/ajt.12666>.

<sup>17</sup> World Health Organization. "Tuberculosis." Tuberculosis . Accessed May 24, 2023. <https://www.who.int/news-room/questions-and-answers/item/tuberculosis#:~:text=About%20one%2Dthird%20of%20the,ill%20with%20TB%20of%2010%25>.

<sup>18</sup> Division of Tuberculosis Elimination, National Center for HIV, Viral Hepatitis, STD, and TB Prevention, Centers for Disease Control and Prevention. "Trends 2021." Tuberculosis (TB), November 9, 2022. <https://www.cdc.gov/tb/publications/factsheets/statistics/tbtrends.htm>.

<sup>19</sup> Provisional CDC data

cases per 100,000 persons).<sup>20</sup> In patients with TBI, the highest risk of reactivation and the development of MTB disease occurs in the first two years after infection and then declines.<sup>21</sup> Since initial infection does result in live MTB in many organs, tuberculosis can be transmitted via non-lung organs used for transplantation from donors who have never had clinical signs or symptoms of MTB disease.<sup>22</sup>

## West Nile Virus

West Nile Virus (WNV) is a ribonucleic acid virus that spreads to humans by mosquito bites. Birds serve as a reservoir for WNV, and humans are a dead-end host for the virus. The virus was first detected in North America in 1999, and since then, human infections have been reported in all U.S. states.<sup>23</sup> The number of human infections varies significantly based on the region and year, with most transmission occurring during the warmer months (July to October) when mosquitoes are most active.<sup>24</sup> While the majority of WNV infections are asymptomatic, about 20% of those infected may develop fever and less than one percent develop neurological complications such as encephalitis or myelitis (involving the spinal cord).<sup>25</sup> This means that even donors who do not display symptoms could still transmit the virus to recipients. While several treatments have been explored, there is currently no established treatment for WNV.

The guidance document focuses on these six endemic diseases due to increased risk for endemic disease transmission through organ transplant. This can be attributed to the movement of populations and broader organ distribution.

## Overall Sentiment from Public Comment

Sentiment is collected on public comment proposals and is measured on a 5-point Likert scale from strongly oppose to strongly support (1-5). These reports are helpful to spot high-level trends but they are not meant as public opinion polls or to replace the substantive analysis below. Generally, public comment sentiment has been supportive of this proposal, as indicated by the total sentiment score of 4.0 by member type and region, with some pockets of concern. Below are graphics that illustrate the sentiment received through public comment.

<sup>20</sup> Division of Tuberculosis Elimination, National Center for HIV, Viral Hepatitis, STD, and TB Prevention, Centers for Disease Control and Prevention. "Trends 2021." Tuberculosis (TB), November 9, 2022.

<https://www.cdc.gov/tb/publications/factsheets/statistics/tbtrends.htm>.

<sup>21</sup> Petruccioli E, Petrone L, Chiacchio T, Farroni C, Cuzzi G, Navarra A, Vanini V, Massafra U, Lo Pizzo M, Guggino G, Caccamo N, Cantini F, Palmieri F, Goletti D. Mycobacterium tuberculosis Immune Response in Patients With Immune-Mediated Inflammatory Disease. *Front Immunol.* 2021 Aug 10;12:716857. doi: 10.3389/fimmu.2021.716857. PMID: 34447382; PMCID: PMC8382688.

<sup>22</sup> Abad CLR, Razonable RR. Donor derived Mycobacterium tuberculosis infection after solid-organ transplantation: A comprehensive review. *Transpl Infect Dis.* 2018 Oct;20(5):e12971. doi: 10.1111/tid.12971. Epub 2018 Aug 12. PMID: 30055041.

<sup>23</sup> Colpitts TM, Conway MJ, Montgomery RR, Fikrig E. West Nile Virus: biology, transmission, and human infection. *Clin Microbiol Rev.* 2012 Oct;25(4):635-48. doi: 10.1128/CMR.00045-12. PMID: 23034323; PMCID: PMC3485754.

<sup>24</sup> Shocket MS, Verwillow AB, Numazu MG, Slamani H, Cohen JM, El Moustaid F, Rohr J, Johnson LR, Mordecai EA. Transmission of West Nile and five other temperate mosquito-borne viruses peaks at temperatures between 23°C and 26°C. *Elife.* 2020 Sep 15;9:e58511. doi: 10.7554/eLife.58511. PMID: 32930091; PMCID: PMC7492091.

<sup>25</sup> L.D. Kramer, West Nile Virus, Editor(s): Brian W.J. Mahy, Marc H.V. Van Regenmortel, *Encyclopedia of Virology (Third Edition)*, Academic Press, 2008, Pages 440-450, ISBN 9780123744104, <https://doi.org/10.1016/B978-012374410-4.00633-6>.

**Figure 1** shows sentiment received from all respondents (regional meeting, online, and email) by their stated member type. Again, there was overall support for the concept, demonstrated by a sentiment score by member type of 4.0. Histocompatibility labs and organ procurement organizations (OPOs), however, showed only slight opposition to the proposal, demonstrated by a 3.8 sentiment score.

**Figure 1: Sentiment by Member Type**

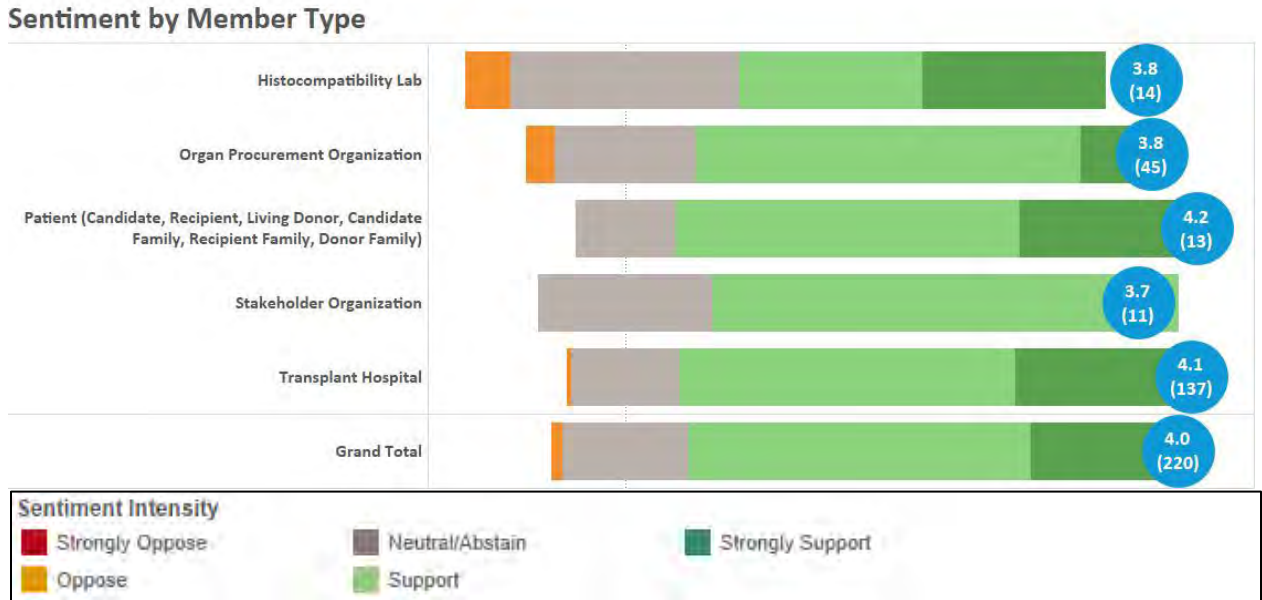
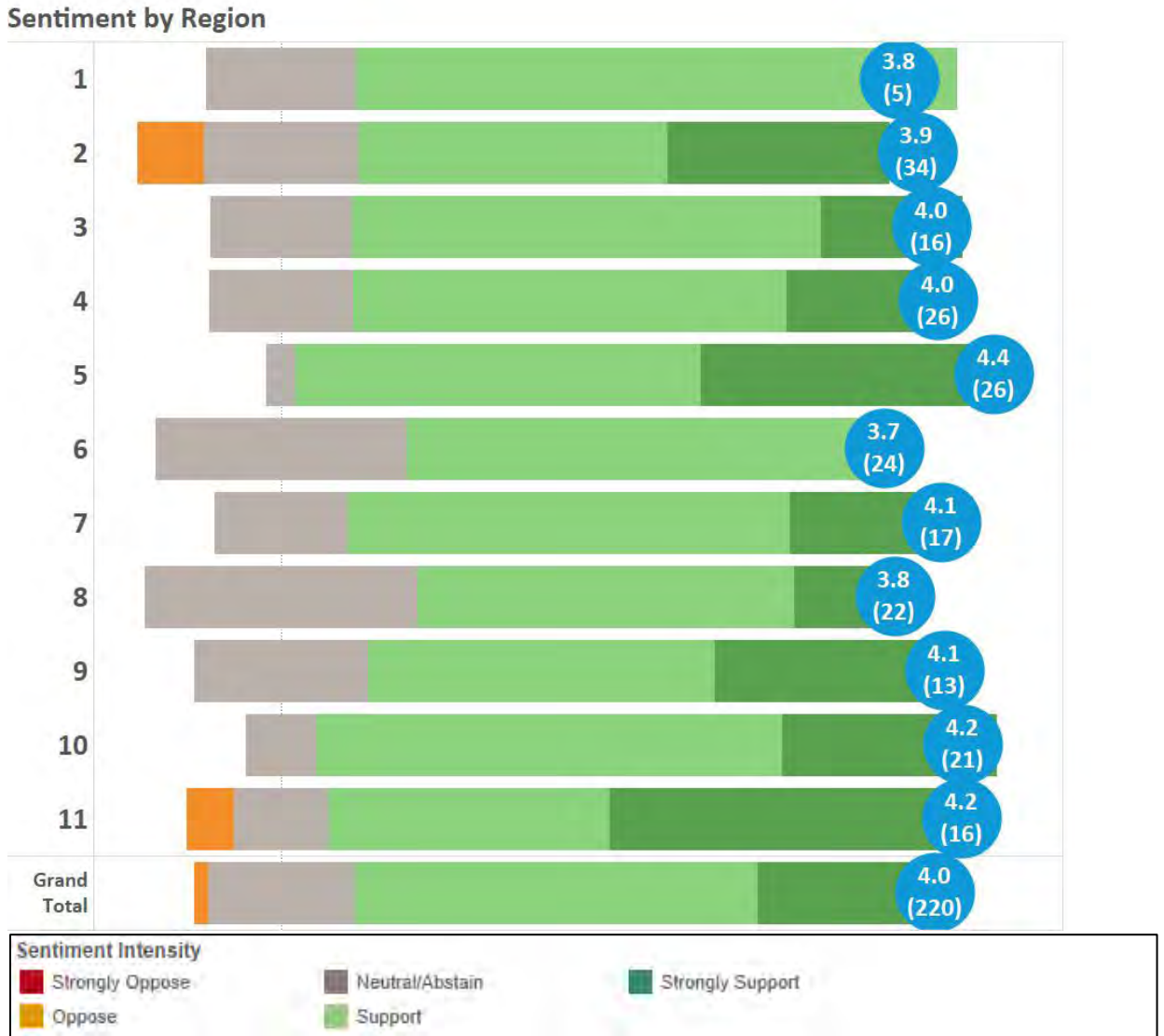




Figure 2 shows sentiment received at regional meetings. Again, overall sentiment was supportive, as indicated by a total sentiment score of 4.0. Opposition was raised in region 2 and region 11.

Figure 2: Sentiment by Region





## Public Comment Themes and Considerations

Comments were generally supportive of this guidance document. In addition to the sentiment score, items out for public comment also provide the opportunity for respondents to submit a substantive written comment. Responses are submitted by members of the public at large, as well as on behalf of regions and committees.<sup>26</sup>

Commenters covered many different topics, including the following themes:

- Support for an updated guidance document
- Cost and availability of recommended testing
- Lack of medical and social history to inform screening
- System enhancements needed for endemic disease screening
- Turnaround time of testing and organ utilization

### Support for an updated guidance document

The majority of public comments supported the guidance document revisions, including comments from all member types. Members agreed that the development of one standard guidance document for screening living and deceased donors for seasonal and geographic endemic infections would help decrease donor-derived infections.

### Cost and availability of recommended testing

Multiple commenters urged the Committee to consider testing availability across OPOs and transplant programs, as well as the additional costs of tests for endemic diseases. Members suggested the Committee investigate test availability, cost acquired by the testing lab to add equipment and reagents, and time it takes to purchase, validate, hire, and train staff in labs to accommodate these additional tests.

The Committee discussed this feedback and will consider this as they explore policy changes for endemic disease screening in the future. Since this guidance document does not mandate any screening tests, the Committee reviewed and discussed the results of this sentiment in public comment and supports sending the guidance to the Board with no changes to testing recommendations due to cost and availability.

### Lack of medical and social history to inform screening

Members voiced concern about the limited information obtained from social and medical history questioning. Members noted that without this information it is challenging to practice targeted screening for endemic diseases. Sentiment echoed that standard medical history questioning may not be able to adequately capture the risk of each donor and is often unreliable.

---

<sup>26</sup> For comments submitted on behalf of the region or committees, the public comment item is discussed at the meeting, OPTN staff draft a summary of the discussion, and the Regional Councillor or Committee leadership review the comment, confirming it is an accurate representation of the discussion that occurred.

The Committee discussed this feedback and does not recommend endemic disease screening when donor history is unknown. This guidance document outlines screening practices and recipient management when medical and social history is known by the OPO and transplant hospital. The Committee decided to move forward with no changes in reference to this sentiment.

## System enhancements needed for endemic disease screening

Members also expressed concern about the challenges of communicating endemic disease testing through the OPTN Computer System. They recommended the Committee explore additional enhancements to facilitate this communication. Recommendations included optional fields in the OPTN Computer System that would allow for timely exchange of information about potential endemic infectious disease risks, updating living donor consent forms to inform potential donors about the disclosure of these test results to the recipient for full transparency, and optional test result fields in the OPTN Donor Data and Matching System for infectious disease testing.

The Committee agrees that enhancements to the OPTN Computer System are needed regarding screening for endemic diseases and recipient management. However, the Committee does not believe these enhancements are in the scope of these guidance document revisions. These enhancements are something the Committee will consider in future project work as they move forward.

## Turnaround time of testing and organ utilization

A couple of comments urged caution over testing turnaround time and its impact on organ utilization. Members voiced concern about their ability to complete testing in a timely manner. They urged the Committee to ensure this testing would not interfere with the ability to utilize organs and delay allocation. The Committee discussed this feedback and agreed they would consider turnaround time and organ utilization if policy changes are pursued by the Committee for mandatory screening for any of these endemic diseases.

## Additional changes

The Committee made additional modifications to the guidance document based on Committee member feedback and singular public comments that did not fall under a specific identified theme. The American Society of Transplantation (AST) recommended revision to the histoplasmosis donor screening section to reflect urine histoplasma antigen as a primary screening method for asymptomatic donors, removing the addition of histoplasma antibodies by complement fixation. The Committee discussed this suggestion and decided to highlight urine histoplasma antigen as the optimal screening method but include histoplasma antibodies by complement fixation as an alternative option. AST also requested the removal of the recommendation to screen donors with a history of pneumonia of unknown type in the past two years because this is overly broad and will likely capture many potential donors who do not have histoplasma. The Committee agreed with this suggestion and limited their screening suggestion to donors with a history of pneumonia in the past 3 to 6 months where the clinical scenario was suspicious of histoplasmosis.

The Committee also made modifications to the guidance document based on Committee member feedback.

This included clarifying:

- ‘History of injection drug use’ to ‘history of non-medical injection drug use’ under the ‘who to screen’ tuberculosis subsection
- ‘Site remote from transplant’ to ‘organ that is not being transplanted’ in the table regarding management of latent or history of active TB in living donors and recipients of these organs
- Risk factors are listed to inform screening under ‘who to screen’ tuberculosis subsection

This also included removing:

- Dose recommendations for treatment since this document does not differentiate between adult and pediatric recipients and donors
- Hunting/taxidermy as a risk factor since it does not apply to these specific endemic diseases
- Race/ethnicity section from ‘who to screen’ tuberculosis subsection

This included the addition of:

- ‘Deceased donor cause of death (if associated with signs/symptoms of infectious illness)’ to risk factors
- ‘Correctional workers’ to risk factors
- Recommendation to consult infectious disease staff in management of recipient and living donor sections

## Recommendations

### Histoplasmosis

The Committee recommends living and deceased donors with signs, symptoms, or radiological findings consistent with active histoplasmosis (cough, fever, sweats, weight loss, non-calcified pulmonary nodules/cavities, or lymphadenopathy)<sup>27</sup> have a thorough medical evaluation including testing for histoplasmosis prior to organ donation. Living and deceased donors at higher risk of transmitting disease include those with recent infection or a prior history of known active histoplasmosis.<sup>28</sup> Screening should be considered for those donors. Potential donors with a history of pneumonia in the past 3 to 6 months where the clinical scenario was suspicious of histoplasmosis should also be considered for screening. Even in endemic areas, universal screening of donors lacking any of the above characteristics is not likely to be productive. Potential donors with the sole finding of a calcified granuloma on chest imaging do not require further testing.<sup>29</sup>

The Committee advises potential living donors with clinical evidence of active histoplasmosis or a serological pattern suggestive of active disease receive infectious disease consultation and treatment prior to organ donation. Recipients of donors with a history of histoplasmosis may not require specific

<sup>27</sup> Kauffman CA. Histoplasmosis: a clinical and laboratory update. *Clin Microbiol Rev.* 2007 Jan;20(1):115-32. doi: 10.1128/CMR.00027-06. PMID: 17223625; PMCID: PMC1797635.

<sup>28</sup> Kovacs CS Jr, Koval CE, van Duin D, de Moraes AG, Gonzalez BE, Avery RK, Mawhorter SD, Brizendine KD, Cober ED, Miranda C, Shrestha RK, Teixeira L, Mossad SB. Selecting suitable solid organ transplant donors: Reducing the risk of donor-transmitted infections. *World J Transplant.* 2014 Jun 24;4(2):43-56. doi: 10.5500/wjt.v4.i2.43. PMID: 25032095; PMCID: PMC4094952.

<sup>29</sup> N. Singh, S. Huprikar, S.D. Burdette, M.I. Morris, J.E. Blair, L.J. Wheat, Donor-Derived Fungal Infections in Organ Transplant Recipients: Guidelines of the American Society of Transplantation, Infectious Diseases Community of Practice†, *American Journal of Transplantation*, Volume 12, Issue 9, 2012, Pages 2414-2428, ISSN 1600-6135, <https://doi.org/10.1111/j.1600-6143.2012.04100>.

prophylaxis. Infectious disease consultation is recommended for these recipients. Clinical and laboratory monitoring for disease is a reasonable approach.<sup>30</sup>

## Coccidioidomycosis

Since many patients with transmissible coccidioidomycosis are asymptomatic and infection is widespread in endemic areas, some experts recommend screening as part of the routine evaluation of all potential donors who reside in endemic areas or who have recently resided or had prolonged stays in such areas.<sup>31</sup> Persons with symptoms consistent with coccidioidomycosis (fevers, weight loss, poorly responding pneumonia) or those with unexplained chest imaging findings (cavities, nodules, lymphadenopathy, reticulonodular infiltrates) require screening as well.<sup>32</sup> Finally, it is reasonable to perform further screening on donors with a known history of coccidioidomycosis, as potential donors with persistently positive serologic studies are more likely to harbor viable organisms. Serological tests (enzyme immunoassay, complement fixation, or immunodiffusion) may be combined with chest imaging.<sup>33</sup>

The Committee advises potential living donors with evidence of active infection receive infectious disease consultation and a thorough evaluation to determine the extent of the infection. Donation from actively infected living donors should be deferred until treatment is complete and evidence of active infection (based on clinical, radiographic, and serological factors) has resolved. Recipients should also be offered infectious disease consultation. Prophylaxis with antifungal medications active against *Coccidioides* species may be effective in preventing disease in recipients of infected living donors.<sup>34</sup>

## Strongyloidiasis

Implementation of the *Improve Deceased Donor Evaluation for Endemic Diseases*<sup>35</sup> proposal will require all deceased donors be screened for *Strongyloides*. Given the high mortality rate of donor-derived *Strongyloides* infection, and the efficacy, safety, simplicity and low cost of prophylactic therapy, universal screening for *Strongyloides* is recommended for all living donors. Serology should be completed using primarily an Immunoglobulin G assay for antibodies to *Strongyloides* pending implementation of *Improve Deceased Donor Evaluation for Endemic Diseases* and is recommended prior to implementation.

Donors infected with *Strongyloides* may still donate. Living donors and recipients who test positive should be offered infectious disease consultation. Living donors can be treated with ivermectin, with no further delay in performing recovery and transplant. Recipients who receive organs from infected and untreated donors should be offered infectious disease consultation and may be treated with ivermectin or albendazole.<sup>36</sup>

<sup>30</sup> Ibid.

<sup>31</sup> Ibid.

<sup>32</sup> Ibid.

<sup>33</sup> Ibid.

<sup>34</sup> Ibid.

<sup>35</sup> "Improve Deceased Donor Evaluation for Endemic Diseases," OPTN, Policy Notice, Accessed October 10, 2023, [https://optn.transplant.hrsa.gov/media/lrxfreqj/dtac\\_endemics\\_policy-notice\\_june23bod.pdf](https://optn.transplant.hrsa.gov/media/lrxfreqj/dtac_endemics_policy-notice_june23bod.pdf).

<sup>36</sup> Fatehi Elzein, Hamad Albahili, Abdelkarim Bahloul, Thamer Alonazi, Adnan Alghamdi, Eid Alsufyani, Abdullatif Musa, Mohammed Alsaheed, Transplant-related strongyloidiasis in solid organ transplant recipients in Saudi Arabia and the Gulf Cooperation Council countries, *International Journal of Infectious Diseases*, Volume 93, 2020, Pages 133-138, ISSN 1201-9712, <https://doi.org/10.1016/j.ijid.2020.01.032>.

## Chagas Disease

Deceased donors who were born in a country currently classified as endemic for Chagas disease by the CDC will require screening according to OPTN *Policy 2.9* pending implementation of *Improve Deceased Donor Evaluation for Endemic Diseases*<sup>37</sup> and is recommended prior to implementation.

Screening should be considered in the following circumstances for living and deceased donors:<sup>38</sup>

- Children of women born in endemic regions if their birth mother’s serology is positive or unknown
- Donors who have resided in an endemic region for more than three months
- Donors who received a blood transfusion in endemic regions and those who have a previous diagnosis of Chagas disease

Serologic testing using an FDA licensed, approved, or cleared assay should be performed on donors with risk factors for Chagas disease.

Living donors who test positive should be informed about the result and offered infectious disease consultation. Confirmatory testing through submission to the CDC or performance of at least two different FDA licensed, approved, or cleared antibody diagnostic tests should be performed.

Potential recipients of *T. cruzi* positive donor organs should be tested for the presence of *T. cruzi* antibody, particularly since they may share unrecognized risk factors with their potential donor. Once transplant has occurred, infectious disease consultation is recommended to coordinate post-transplant testing. In the United States, the CDC can assist with polymerase chain reaction (PCR) and parasitemia testing. Incorporating a PCR testing platform into the post-transplant testing program is recommended, as molecular testing may be positive weeks before parasitemia is detected using standard screening.

## Tuberculosis

The most easily identifiable risk factor for Tuberculosis (TB) is place of birth with estimated risk of TB infection (TBI) of 18.7% among foreign-born persons in the U.S. as compared to 1.8% among U.S. born persons.<sup>39</sup>

The following factors would be expected to increase the risk of TBI and should indicate the need for screening<sup>40</sup>:

- Close contacts of persons with infectious TB disease

<sup>37</sup>“Improve Deceased Donor Evaluation for Endemic Diseases,” OPTN, Policy Notice, Accessed October 10, 2023, [https://optn.transplant.hrsa.gov/media/lrxfreqj/dtac\\_endemics\\_policy-notice\\_june23bod.pdf](https://optn.transplant.hrsa.gov/media/lrxfreqj/dtac_endemics_policy-notice_june23bod.pdf).

<sup>38</sup> Forsyth CJ, Manne-Goehler J, Bern C, Whitman J, Hochberg NS, Edwards M, Marcus R, Beatty NL, Castro-Sesquen YE, Coyle C, Stigler Granados P, Hamer D, Maguire JH, Gilman RH, Meymandi S. Recommendations for Screening and Diagnosis of Chagas Disease in the United States. *J Infect Dis.* 2022 May 4;225(9):1601-1610. doi: 10.1093/infdis/jiab513. PMID: 34623435; PMCID: PMC9071346.

<sup>39</sup> Vernon A. Treatment of latent tuberculosis infection. *Semin Respir Crit Care Med.* 2013 Feb;34(1):67-86. doi: 10.1055/s-0032-1333544. Epub 2013 Mar 4. PMID: 23460007.

<sup>40</sup> Division of Tuberculosis Elimination, National Center for HIV, Viral Hepatitis, STD, and TB Prevention, Centers for Disease Control and Prevention. “Who Should Be Tested for TB Infection.” *Tuberculosis (TB)*, April 14, 2016. <https://www.cdc.gov/tb/topic/testing/whobetested.htm>.

- Those who spend significant time (greater than 3 months) in areas of the world with high rates of TB or those born in these regions including relief work in a country with high TB risk
  - The guidance document contains a map available on the CDC website that outlines these regions
- History of injection of nonmedical injection drug use
- Persons who reside (or ever resided) or worked in institutional settings which resulted in increased risk of exposure to TB (hospitals, nursing homes, correctional facilities, other health care settings, homeless shelters)

Screening strategies for living donors are dependent on history of TBI and whether the donor was treated appropriately. Recommendations include: monitor recipients of living donors, offer infectious disease consultation, consider deferring transplant, and consider additional testing and treatment.

Recipients of an organ from a donor who is tuberculosis skin test (TST) or Interferon Gamma Release Assay (IGRA)-positive, had recent exposure to active TB, or had radiographic evidence of untreated TB should be considered for TBI therapy, following CDC guidance for anti-tuberculosis agents and durations, and offered infectious disease consultation.

## West Nile Virus

Screening strategies for WNV in living and deceased donors include universal year-round testing of all donors versus targeted testing during periods of human WNV activity.

It is advisable to screen deceased donors during months of regional WNV activity. In cases involving deceased donors, the feasibility of conducting WNV testing within a required timeframe is uncertain. As such, the Committee advises against recovering organs from donors who satisfy any of the following criteria: (1) a confirmed WNV infection, (2) a positive WNV nucleic acid testing (NAT) test result, ideally meeting a threshold signal-to-cutoff ratio or verified by repeat testing as recommended by the manufacturer, or (3) clinical manifestations of meningitis, encephalitis, or flaccid paralysis of indeterminate origin, particularly if they originate from regions with documented WNV activity.

Living donors should be screened with WNV NAT within seven to 14 days of donation. Although no studies have examined organ donation after WNV infection, the Committee suggests postponing donation for a minimum of 28 days in living donors, after which NAT and IgM testing should be performed. These donors should be offered infectious disease consultation. If both tests are negative, it is likely that the initial NAT was a false positive, and organ donation can be considered.

Infectious disease consultation is recommended for recipients of these donors. Development of symptoms compatible with WNV infection within the first several weeks of transplant should prompt testing in the recipient. There are currently no specific medications available for the treatment of WNV infection.

## Compliance Analysis

### NOTA and OPTN Final Rule

The Committee submits this guidance document under the authority of the National Organ Transplantation Act (NOTA), which states that the OPTN shall, "adopt and use standards of quality for the acquisition and transportation of donated organs"<sup>41</sup> and under the authority of the OPTN Final Rule, which states, "An OPTN member procuring an organ shall assure that laboratory tests and clinical examinations of potential organ donors are performed to determine any contraindications for donor acceptance, in accordance with policies established by the OPTN."<sup>42</sup> This guidance document will assist living donors in complying with OPTN *Policy 14.4: Medical Evaluation Requirements for Living Donors* and allow OPOs to use standards of quality when allocating organs.

### OPTN Strategic Plan

This guidance document aligns with the strategic plan goal to promote living donor and recipient safety by reducing transmission of endemic infections through organ transplantation.

## Conclusion

This guidance document aims to decrease donor-derived disease transmission from organ transplantation and provide appropriate recommendations for living donor recovery hospitals and OPOs when screening donors for endemic diseases. The document advises transplant hospitals on recipient management after transplanting organs from donors with endemic diseases. Minor public comment changes were made to reflect stakeholder and Committee member suggestions. The Committee will assess the success of this guidance document through decreased endemic disease transmissions in organ transplantation.

---

<sup>41</sup> 42 USC §274(b)(2)(E).

<sup>42</sup> 8 42 CFR §121.6(a).



## Guidance Document

Proposed new language is underlined (example) and language that is proposed for removal is struck through (~~example~~). Heading numbers, table and figure captions, and cross-references affected by the numbering of these policies will be updated as necessary.

1           **Preventing Chagas Disease In Transplant Recipients:**  
2           **Donor Screening And Recipient Monitoring (2014)**

3           Repealed.  
4

5           **Identifying Risk Factors For West Nile Virus (WNV)**  
6           **During Evaluation Of Potential Living Donors (2013)**

7           Repealed.  
8

9           **Guidance For Identifying Risk Factors For**  
10          **Mycobacterium Tuberculosis (MTB) During**  
11          **Evaluation Of Potential Living Kidney Donors (2012)**

12          Repealed.  
13

14          **Recognizing Seasonal And Geographically Endemic**  
15          **Infections In Organ Donors: Considerations During**  
16          **Living Donor Evaluation (2014)**

17          Repealed.  
18

19          **Recognizing Seasonal and Geographically Endemic**  
20          **Infections in Organ Donors:**  
21          **Considerations for Deceased and Living Donation**

## 22 Table of Contents

23	Background	17
24	Summary and Goals	17
25	Developing a Written Protocol for Identifying and Testing Donors	17
26	Recognizing Risk Factors	18
27	FUNGAL INFECTIONS	21
28	PARASITIC INFECTIONS	24
29	BACTERIAL INFECTIONS	31
30	VIRAL INFECTIONS	35

31

## 32 Background

33 Both deceased and living donors are at risk for transmitting endemic diseases that are either  
 34 geographically or seasonally defined. Since 2014, Organ Procurement and Transplantation Network  
 35 (OPTN) policy has required living donor hospitals to develop a written protocol for identification and  
 36 testing potential donors at risk for these endemic infections as part of the medical evaluation process.  
 37 More recently on June 26, 2023, the OPTN Board of Directors approved additional policies<sup>43</sup> to address  
 38 screening of deceased donors for endemic diseases to further reduce the risk for potential donor-  
 39 derived infectious events.

## 40 Summary and Goals

41 The OPTN Ad Hoc Disease Transmission Advisory Committee (DTAC) created this guidance document to  
 42 assist programs in identification of potential living and deceased donors who may carry an increased risk  
 43 of transmitting seasonal or geographically endemic disease to organ recipients. This document will also  
 44 help programs manage recipients who receive organs from donors with endemic diseases. This resource  
 45 is not OPTN policy, so it does not carry the monitoring or enforcement implications of policy. It is not an  
 46 official guideline for clinical practice, nor is it intended to be clinically prescriptive or to define a  
 47 standard of care. This is a resource tool intended to be of educational support for organ procurement  
 48 organizations (OPOs) and transplant centers and is for voluntary use by members.

## 49 Developing a Written Protocol for Identifying and Testing Donors

50 The DTAC reviews potential donor-derived disease transmission events (PDDTE) reported to the OPTN  
 51 for both deceased and living donors. A number of the PDDTE reported are seasonally and geographically  
 52 associated. Some of the reported events resulted in recipient illness or death. Recognition of disease in  
 53 these donors can be especially challenging, as risk factors may not be recognized, and laboratory  
 54 screening is not generally universally performed. This guidance document provides information the  
 55 DTAC believes will help programs and OPOs identify and test donors at risk for transmissible seasonally  
 56 or geographically defined disease. This information is meant to assist the transplant community in

<sup>43</sup> Improve Deceased Donor Evaluation for Endemic Diseases," OPTN, Policy Notice, Accessed October 10, 2023, [https://optn.transplant.hrsa.gov/media/lrxfreqj/dtac\\_endemics\\_policy-notice\\_june23bod.pdf](https://optn.transplant.hrsa.gov/media/lrxfreqj/dtac_endemics_policy-notice_june23bod.pdf).

57 performing organ donor and recipient infectious disease screening procedures as part of the overall  
 58 medical evaluation and recipient management process.

## 59 **Recognizing Risk Factors**

60 There are several factors to consider in determining a donor's risk of transmissible infection. OPOs and  
 61 living donor recovery hospitals should consider the following when screening potential organ donors:

- 62 • Geographic risks (including duration of time spent in a location)
  - 63 ○ Birthplace of the potential donor
  - 64 ○ Home country/region of the U.S., and prolonged residence outside home region, either  
 65 recent or remote
  - 66 ○ Close family members' countries of origin
  - 67 ○ Donor hospital region
  - 68 ○ Occupational or recreational travel to other countries and/or regions
- 69 • Occupational risks
  - 70 ○ Healthcare workers, veterinarian/animal care workers
  - 71 ○ Construction workers, landscapers, park rangers, and other outdoor workers
  - 72 ○ Peace Corps workers, international journalists
  - 73 ○ Current or previous military service, particularly outside the U.S.
  - 74 ○ Medical mission trips
  - 75 ○ Correctional workers
- 76 • Seasonal risks
  - 77 ○ Residence in/travel to warm weather climates with potential insect exposures
- 78 • Hobbies
  - 79 ○ Time living outdoors including camping, swimming in lakes, drinking stream water,  
 80 insect exposures
  - 81 ○ Adventure sports
  - 82 ○ Gardening
- 83 • Significant animal exposure (wild and/or domestic)
  - 84 ○ Large numbers of cats or dogs or any unusual pets
  - 85 ○ Laboratory/research animals
  - 86 ○ Veterinarian/vet assistant
- 87 • Family members and close contacts with potential risk factors
  - 88 ○ Geographic or seasonal infections previously diagnosed in close family members or  
 89 other contacts may predict risk for subclinical infection in the potential donor
- 90 • Personal history of seasonal or geographic infection in the potential donor, even if remote
- 91 • Deceased donor cause of death (if associated with signs/symptoms of infectious illness)

92 The organ donor population has become increasingly geographically diverse, reflecting the enhanced  
 93 mobility and complex migration patterns of the general population. Therefore, it is not practical to list all  
 94 the pathogens that have the potential for transmission through organ transplantation. Parasitic  
 95 infections such as amebiasis, babesiosis, leishmaniasis, schistosomiasis, echinococcosis, and malaria,  
 96 bacterial infections such as brucellosis and melioidosis, fungal infections such as paracoccidioidomycosis  
 97 and talaromycosis, and viral infections such as Eastern Equine Encephalitis Virus have distinct areas of  
 98 endemism and may be transmitted through the organ allograft. In donors with a history of residence in  
 99 developing countries or remote regions, unusual occupational exposure risks, or extensive travel,

100 infectious disease (ID) consultation may be helpful as part of the pre-transplant evaluation for living  
 101 donors or prior to organ acceptance for deceased donors. This guidance document does not replace  
 102 consultation with ID.

103 Table 1 covers several common seasonal and geographically endemic infections that may be transmitted  
 104 from organ donor to recipient.

105 **Table 1: Common Seasonal and Geographically Endemic Infections in Organ Donors**

<u>Disease</u>	<u>Signs/Symptoms in Potential Donor</u>	<u>Known Risk Factors</u>	<u>Potential Testing for Deceased Donors</u>	<u>Potential Testing For Living Donors</u>	<u>Imaging that may be helpful</u>
<u>Histoplasmosis</u>	<u>Fever, night sweats, lymphadenopathy, cough, non-calcified pulmonary nodules or cavities</u>	<u>Residence in Midwestern or South-Central states along the Mississippi or Ohio River Valleys</u>	<u>-Urine and/or serum antigen enzyme immunoassay</u> <u>-Serology: complement fixation and/or immunodiffusion</u>	<u>-Urine and/or serum antigen enzyme immunoassay</u> <u>-Serology: complement fixation and/or immunodiffusion</u>	<u>Chest X-ray (CXR) or CT</u>
<u>Coccidioidomycosis</u>	<u>Fever, joint pains, cough, neck stiffness, headaches, pulmonary nodules or cavities, reticulonodular infiltrates</u>	<u>Residence in endemic areas of Washington state, the Southwestern United States, Northern Mexico, and parts of South &amp; Central America</u>	<u>Serology:</u> <u>-enzyme immunoassay (preferred)</u> <u>-complement fixation</u> <u>-immunodiffusion</u> <u>-Urine or serum antigen testing</u>	<u>Serology:</u> <u>-enzyme immunoassay (preferred)</u> <u>-complement fixation</u> <u>-immunodiffusion</u> <u>-Urine or serum antigen testing</u>	<u>CXR or CT</u>
<u>Strongyloidiasis</u>	<u>Chronic abdominal pain, bloating, heartburn, intermittent diarrhea and constipation, dry cough, skin rashes, and/or eosinophilia, or could be entirely asymptomatic.</u>	<u>Soil exposure in tropical/warm climates. Walking barefoot or unprotected skin contact with human sewage or contaminated soil. Infection may persist for decades.</u>	<u>Serologic testing is required for all<sup>44</sup></u>	<u>Donors could be tested by serology (preferred) and/or stool ova and parasite examination, specifically looking for <i>Strongyloides</i>.</u>	<u>None</u>

<sup>44</sup> Pending policy implementation of *Improve Deceased Donor Evaluation for Endemic Diseases*.

<u>Disease</u>	<u>Signs/Symptoms in Potential Donor</u>	<u>Known Risk Factors</u>	<u>Potential Testing for Deceased Donors</u>	<u>Potential Testing For Living Donors</u>	<u>Imaging that may be helpful</u>
<u>Chagas disease</u>	<u>Most asymptomatic; symptomatic chronic infection may present with cardiomyopathy, cardiac conduction abnormalities, megaesophagus, megacolon</u>	<u>Born or resided in endemic areas of Mexico, South &amp; Central America, child of woman who lived in endemic area, received blood transfusion in endemic area</u>	<u>Required by OPTN policy if donor is born in endemic country<sup>45</sup></u> <u>Serology testing (See Chagas Guideline, Table 3)</u>	<u>Serology testing (See Chagas Guideline, Table 3)</u>	<u>None unless symptomatic with chronic Chagas disease</u>
<u>Tuberculosis</u>	<u>Fever, night sweats, weight loss, cough, recurrent pneumonia, exudative pleural effusion of unknown etiology, lymphadenopathy, non-calcified pulmonary nodules or cavities</u>	<u>Born outside U.S., prolonged residence outside U.S., homeless, alcohol or other substance abuse, jail/prison time, health care worker, known TB exposure</u>	<u>Positive tuberculin skin test (TST) or interferon gamma release assay (IGRA)**;</u> <u>Sputum/BAL AFB smear, culture, nucleic acid amplification, TB PCR; tissue AFB smear, culture, TB PCR</u> <u>**Deceased donors on high-dose steroids may have false-negative IGRA/TST</u>	<u>Positive tuberculin skin test (TST) or interferon gamma release assay (IGRA);</u> <u>sputum/BAL AFB smear, culture, nucleic acid amplification, TB PCR; tissue AFB smear, culture, TB PCR</u> <u>*refer to OPTN Policy 14</u>	<u>CXR, CT thorax, CT abdomen/ pelvis (renal TB)</u>
<u>West Nile Virus</u>	<u>Often asymptomatic; 20% develop acute febrile illness; &lt;1% encephalitis, myelitis</u>	<u>Mosquito exposure, blood transfusion; risk varies by season &amp; location</u>	<u>Nucleic acid test (NAT)</u>	<u>NAT</u>	<u>None</u>

<sup>45</sup> Pending policy implementation of *Improve Deceased Donor Evaluation for Endemic Diseases*.

## 106 **FUNGAL INFECTIONS**

### 107 **Histoplasmosis Guidance**

#### 108 **Background**

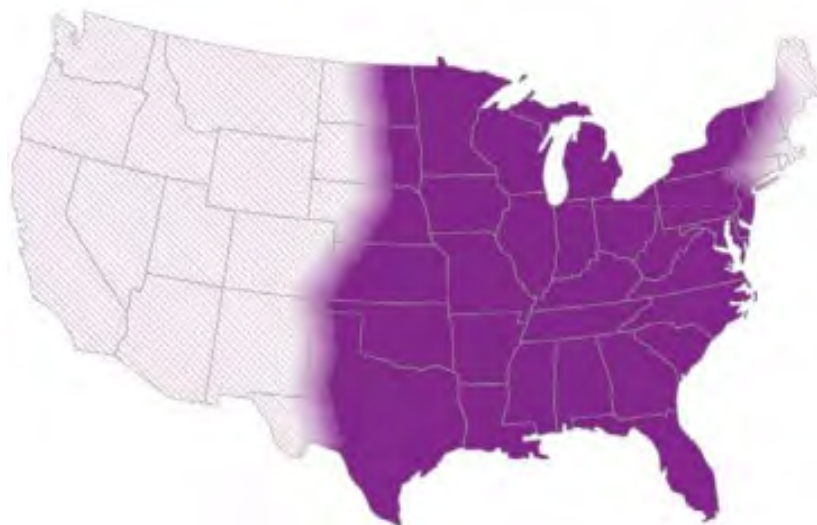
109 *Histoplasma capsulatum* is a dimorphic fungus found throughout the world. It is endemic in the U.S. in  
 110 the Ohio and Mississippi River Valleys and into the Midwest and South-Central states (Figure 1). In most  
 111 cases, clinical signs of infection in immunocompetent patients are limited to the lungs, though many  
 112 patients are asymptomatic. Immunocompromised patients may develop severe disseminated disease  
 113 that is fatal without prompt diagnosis and treatment. Donor-derived histoplasmosis has only rarely been  
 114 reported. In endemic areas, up to 75% of the population is infected with *Histoplasma capsulatum*;  
 115 therefore, no further exposure risk factors need be elicited in residents of these areas.

#### 117 **Who Should be Screened**

##### 118 Living and Deceased Donors

119 Donors at higher risk of transmitting disease include those with recent infection or a prior history of  
 120 known active histoplasmosis and should be screened. Potential donors with a history of pneumonia in  
 121 the past three to six months where the clinical scenario was suspicious of histoplasmosis should also be  
 122 considered for screening. Donors with signs, symptoms, or radiological findings consistent with active  
 123 histoplasmosis (cough, fever, night sweats, weight loss, non-calcified pulmonary nodules/cavities, or  
 124 lymphadenopathy) need a thorough medical evaluation including testing for histoplasmosis prior to  
 125 organ donation. Even in endemic areas, universal screening of donors lacking any of the above  
 126 characteristics is not likely to be productive. Potential donors with the sole finding of a calcified  
 127 granuloma on chest imaging do not require further testing.

128  
 129 **Figure 1: Centers for Disease Control and Prevention’s (CDC) current estimate of**  
 130 ***Histoplasma* endemicity in the United States<sup>46</sup>**



131

<sup>46</sup> CDC. “More Information about the Estimated Areas with Blastomycosis ...” More information about the estimated areas with blastomycosis, coccidioidomycosis (Valley fever), and histoplasmosis in the United States. Accessed June 23, 2023.

132 Darker shading indicates areas where *Histoplasma* is more prevalent. Diagonal shading shows the  
 133 potential geographic range of *Histoplasma*.

134

### 135 *How to Screen*

#### 136 Living and Deceased Donors

137 In asymptomatic potential donors, screening should be conducted through urine and/or serum antigen  
 138 enzyme immunoassays. Alternatively, serological testing can be used to screen those at risk for  
 139 infection. Complement fixation and immunodiffusion testing are the most commonly performed tests.  
 140 Complement fixation titers of 1:8 and 1:16 may just indicate previous infection at low risk for  
 141 transmission. Positive immunodiffusion testing is more concerning for the presence of viable organisms,  
 142 particularly if a H band is present.

143

#### 144 *Management of Infected Living Donors*

145 Potential living donors with clinical evidence of active histoplasmosis (fever, night sweats,  
 146 lymphadenopathy, cough, non-calcified pulmonary nodules or cavities) or a serological pattern  
 147 suggestive of active disease should receive treatment prior to organ donation. Living donors should be  
 148 informed about the result and offered consultation with ID. For patients with histoplasmosis limited to  
 149 the lungs, a reasonable approach would include six to 12 weeks of treatment, with resolution of clinical  
 150 signs and symptoms of histoplasmosis. After treatment and resolution of antigenuria/antigenemia, the  
 151 risk of donor-derived infection is likely low. Given that disseminated histoplasmosis is typically  
 152 associated with other significant medical conditions, it is unlikely that a patient with a history of  
 153 disseminated histoplasmosis would qualify for living organ donation.

#### 154 *Management of Recipients*

155 Recipients of donors with a history of histoplasmosis may not require specific prophylaxis. Once  
 156 transplant has occurred, consultation with ID is recommended. Clinical and laboratory monitoring for  
 157 disease is a reasonable approach. Serial urine or serum antigen testing (every one to three months) for  
 158 one year might identify donor-derived histoplasmosis prior to the development of clinical symptoms.  
 159 Serological tests have low sensitivity after organ transplantation, and most solid organ transplant  
 160 recipients with histoplasmosis have negative serological studies.

161

#### 162 *Infection Avoidance Between Testing and Transplant*

163 In endemic areas, exposure to *H. capsulatum* may occur with many daily activities and is difficult to  
 164 entirely avoid. Nonetheless, in the pre-donation period, certain activities (cave exploration, significant  
 165 time in construction sites, exposure to bird droppings or bat guano, digging soil) should be avoided.

166

## 167 Coccidioidomycosis Guidance

### 168 *Background*

169 *Coccidioides immitis* and *Coccidioides posadasii* are dimorphic fungi endemic in arid and semi-arid  
 170 climates in Washington State and the Southwestern part of the U.S. (Figure 2), Northern Mexico  
 171 including areas along the U.S. border, and parts of Central and South America. The most common clinical

<https://www.cdc.gov/fungal/pdf/more-information-about-fungal-maps-508.pdf>.



172 syndrome is pneumonia often accompanied by joint pains, fatigue, and weight loss. Chronic  
 173 fibrocavitary pulmonary disease may follow primary infection, and some infected persons have  
 174 persistent symptomatic pulmonary nodules, reticulonodular disease, or cavitary disease.  
 175 Immunosuppressed individuals may develop disseminated infection. Multiple cases of donor-derived  
 176 coccidioidomycosis have been reported, often with poor outcomes.

177  
 178

**Figure 2: CDC's current estimate of *Coccidioides* endemicity in the United States**



179 <sup>47</sup>

180  
 181  
 182  
 183  
 184

Darker shading shows areas where *Coccidioides* is more likely to be present. Diagonal shading shows the potential geographic range of *Coccidioides*.

#### Who Should be Screened

##### Living and Deceased Donors

186 One study indicated that 2.1% of persons evaluated for living donation in an endemic region were  
 187 seropositive, suggesting recent or active disease. Since many patients with transmissible  
 188 coccidioidomycosis are asymptomatic and infection is widespread in endemic areas, some experts  
 189 recommend screening as part of the routine evaluation of all potential donors  
 190 who reside in endemic areas or who have recently resided or had prolonged stays in such areas. Persons  
 191 with signs and symptoms consistent with coccidioidomycosis (fever, joint pains, cough, neck stiffness,  
 192 headaches) or those with unexplained chest imaging findings (cavities, nodules, lymphadenopathy,  
 193 reticulonodular infiltrates) warrant screening as well. Finally, it is reasonable to perform further  
 194 screening on donors with a known history of coccidioidomycosis, as potential donors with persistently  
 195 positive serologic studies are more likely to harbor viable organisms.

<sup>47</sup> CDC. "More Information about the Estimated Areas with Blastomycosis ..." More information about the estimated areas with blastomycosis, coccidioidomycosis (Valley fever), and histoplasmosis in the United States. Accessed June 23, 2023. <https://www.cdc.gov/fungal/pdf/more-information-about-fungal-maps-508.pdf>.

196 How to Screen197 Living and Deceased Donors

198 In asymptomatic potential donors, serological tests (enzyme immunoassay, complement fixation, or  
 199 immunodiffusion) may be combined with chest imaging. Patients with suggestive findings on imaging  
 200 (non-calcified nodules, cavities, lymphadenopathy, reticulonodular infiltrates) may require sputum  
 201 cultures or bronchoscopy with culture of lavage fluid, although sensitivity of cultures may be low. For  
 202 prospective donors with normal imaging and negative serologies, the risk of donor-derived infection is  
 203 likely low.

204

205 Management of Infected Living Donors

206 Potential living donors with evidence of active infection should be offered consultation with ID and  
 207 require a thorough evaluation to determine the extent of the infection. Donation from actively infected  
 208 donors should be deferred until treatment is complete and evidence of active infection (based on  
 209 clinical, radiographic, and serological factors) has resolved.

210

211 Management of Recipients

212 Prophylaxis with antifungal medications active against *Coccidioides* species may be effective in  
 213 preventing disease in recipients of infected donors. Fluconazole or itraconazole can be used as  
 214 prophylaxis; fluconazole is more reliably absorbed and interactions with calcineurin inhibitors are less  
 215 significant. Posaconazole and voriconazole are second-line options. Echinocandins or inhaled polyenes  
 216 are not effective. Once transplant has occurred, consultation with ID is recommended. Whether  
 217 additional prophylaxis is needed for recipients of donors who received adequate treatment for  
 218 coccidioidomycosis in the past is unknown. While periodic post-transplant serological monitoring could  
 219 be considered, the sensitivity of serological testing in immunosuppressed patients is likely low.  
 220 Recipients should be closely monitored, when prophylaxis is discontinued, with periodic clinical,  
 221 radiologic, and serologic assessments, especially in the first six to 12 months after cessation of  
 222 prophylaxis.

223

224 Infection Avoidance between Testing and Transplant

225 For people living in endemic areas, completely avoiding exposure to *Coccidioides* is difficult to  
 226 impossible. Highest risk activities include those that result in heavy inhalation of desert soil, particularly  
 227 in the dry periods following the rainy season. Potential donors that develop pulmonary infection or  
 228 illness characterized by fatigue, cough, and/or joint pain after initial screening may require repeat  
 229 evaluation.

230

## **PARASITIC INFECTIONS**

231 Strongyloidiasis Guidance232 Background

233 *Strongyloides stercoralis* is a nematode (roundworm). Unlike other parasites, *Strongyloides* can replicate  
 234 for decades within humans via an autoinfection cycle, resulting in chronic symptomatic or asymptomatic  
 235 infections that can flourish in the setting of immunosuppression. There are an estimated 370 million  
 236 infected people worldwide.

237  
 238 *Strongyloides* is most common in tropical, subtropical, or warm temperate climates, including the  
 239 Caribbean, Mexico, South and Central America, Africa, Southeast Asia, Southern and Eastern Europe,  
 240 and the Southeastern U.S. In the U.S., a prevalence of up to 6.1% has been reported in select  
 241 populations, with a much higher prevalence of up to 46.1% in immigrant populations. Infection results  
 242 primarily from exposed skin contact with soil that is contaminated with *Strongyloides* larvae. Activities  
 243 that increase the risk of becoming infected include direct contact with contaminated soil (i.e., walking  
 244 barefoot), unprotected skin contact with human waste or sewage, and occupations that increase  
 245 contact with contaminated soil, such as farming and coal mining. Rural populations and those of a lower  
 246 socioeconomic status are at higher risk.

247  
 248 Hyperinfection occurs in immunocompromised hosts and can lead to disseminated infection, with  
 249 mortality as high as 85%. From 2009 to 2013, the CDC assisted in investigations of donor-derived  
 250 strongyloidiasis in 11 recipients from seven organ donors and found 86% of the donors to originate from  
 251 *Strongyloides* endemic areas. Prophylaxis with ivermectin was noted to be highly effective in preventing  
 252 disease transmission in the CDC series and in another series reported by an OPO.

### 253 254 *Who Should Be Screened*

#### 255 Living and Deceased Donors

256 Given the high mortality rate of donor-derived *Strongyloides* infection, and the efficacy, safety,  
 257 simplicity and low cost of prophylactic therapy, universal screening for *Strongyloides* is recommended  
 258 for all living donors and is required for deceased donors by OPTN Policy 2.9 pending implementation of  
 259 *Improve Deceased Donor Evaluation for Endemic Diseases*<sup>48</sup>, and is recommended prior to  
 260 implementation.

261  
 262 Donors with a personal history of previously treated *Strongyloides* infection should be at low risk for  
 263 transmission, although repeat (new) infection after treatment can occur, and the autoinfection cycle  
 264 could allow for subclinical persistence. Signs and symptoms include chronic abdominal pain, bloating,  
 265 heartburn, intermittent diarrhea and constipation, dry cough, skin rashes, and/or eosinophilia, or  
 266 persons can be entirely asymptomatic. Although transmission within families is unlikely, a family history  
 267 should trigger screening given the likelihood of similar exposures.

### 268 269 *How to Screen*

#### 270 Living and Deceased Donors

271 Screening is by serology or stool examination. Serology is required for deceased donors pending  
 272 implementation of *Improve Deceased Donor Evaluation for Endemic Diseases*<sup>49</sup> and recommended for  
 273 living donors. Serology is primarily an Immunoglobulin G (IgG) assay for antibodies to *Strongyloides*. The  
 274 enzyme-linked immunosorbent assay (ELISA) is preferable because of its greater sensitivity (90%) and  
 275 specificity (99%), compared with indirect fluorescent antibody (IFA) and indirect hemagglutination (IHA)  
 276 tests. Antibody testing cannot differentiate between current and prior or treated infections; titers do  
 277 not reliably decrease rapidly after treatment. The assays may produce false positive results in patients  
 278 with filariasis and other nematode infections. Serology testing is commercially available, often with  
 279 turnaround times of one to five days.

<sup>48</sup> "Improve Deceased Donor Evaluation for Endemic Diseases," OPTN, Policy Notice, Accessed October 10, 2023, [https://optn.transplant.hrsa.gov/media/lrxfreqj/dtac\\_endemics\\_policy-notice\\_june23bod.pdf](https://optn.transplant.hrsa.gov/media/lrxfreqj/dtac_endemics_policy-notice_june23bod.pdf).

<sup>49</sup> Ibid.

280 Stool testing is less sensitive than serology, as stools are positive only during larval shedding which can  
 281 be intermittent and would be indicated when serological testing is unavailable or when serological  
 282 findings are negative but clinical suspicion of disease is high. A single stool specimen has a sensitivity of  
 283 only 15%–30%, although sensitivity increases to nearly 100% if seven consecutive daily stool specimens  
 284 are examined in an expert laboratory, which may not be practical in the transplant setting. Intestinal  
 285 endoscopy with biopsies may also diagnose *Strongyloides* infection, although it would not be used for  
 286 routine screening.

287  
 288 For living donors, screening should be done with initial testing to allow time for subsequent treatment if  
 289 necessary.

### 291 *Management of Infected Living Donors*

292 Living donors who test positive should be informed about the result and offered consultation with ID.  
 293 Donors infected with *Strongyloides* may still donate. Living donors can be treated with ivermectin, with  
 294 no further delay in organ transplant. Additional serologic or stool testing after treatment is not likely to  
 295 be helpful.

### 297 *Management of Recipients*

298 Once transplant has occurred, consultation with ID is recommended. Recipients who receive organs  
 299 from infected and untreated donors may be treated with ivermectin or albendazole. Ivermectin is well  
 300 tolerated and is preferred (when available) due to superior efficacy. Only oral ivermectin is available in  
 301 the U.S. There may be a drug interaction between ivermectin and the calcineurin inhibitors (tacrolimus  
 302 and cyclosporine), therefore monitoring of drug levels is recommended.

303  
 304 Transplant centers should follow up on such pre-transplant test results and give prophylaxis to  
 305 recipients if their donors were not previously treated. Prophylaxis and treatment doses of anti-parasitic  
 306 medications in this setting are probably the same, although they have not been studied. Most cases of  
 307 donor-derived strongyloidiasis occur several weeks to months after transplant, so centers should have  
 308 adequate time to treat recipients before infection occurs. In cases in which there is insufficient time for  
 309 test results to return and treatment to be given to donors, centers should plan for post-transplant  
 310 treatment of the recipient.

311  
 312 People dually infected with Human T-Cell Lymphotropic Virus-1 (HTLV-1) and *Strongyloides* are more  
 313 likely to develop severe cases of strongyloidiasis and are at higher risk for recurrence. It may be  
 314 worthwhile to screen patients with *Strongyloides* infection for HTLV-1, so the treating clinicians are  
 315 aware of the risk of increased disease severity and recurrence.

316  
 317 Patients from areas of Africa endemic for loiasis (primarily central and western sub-Saharan Africa)  
 318 should be screened for *L. loa* microfilaremia, as administration of ivermectin to those with  
 319 microfilaremia can precipitate life-threatening encephalopathy, known as the Mazzotti reaction.

### 321 *Infection Avoidance between Testing and Transplant*

322 In the absence of exposure, acquisition of acute infection would generally be rare in the several months  
 323 between testing and transplant for most living donors, unless they travel to endemic regions and have  
 324 significant soil exposure. Shoes should be worn in endemic areas to avoid contact with infected soil.

## 325 Chagas Disease Guidance

### 326 Background

327 Chagas disease is caused by infection with the protozoan parasite *Trypanosoma cruzi* (*T. cruzi*). It is  
 328 estimated that eight million people are infected in the Americas, with over 300,000 infected individuals  
 329 currently living in the U.S. Most infections were acquired in endemic regions of Mexico, South and  
 330 Central America, where infection is transmitted by the bite of an infected triatomine bug. Infection can  
 331 also be transmitted from an infected mother during the second and third trimester of pregnancy or  
 332 rarely during childbirth, as well as through infected food or drink, and through blood transfusion and  
 333 organ transplantation.

334  
 335 Most infections are acquired in childhood during residence in an endemic area. The acute phase of  
 336 infection may be associated with a mild febrile illness and is often unrecognized. Parasitemia clears  
 337 within eight to 12 weeks without treatment, but individuals remain chronically infected. Infection  
 338 persists for the individual's life, with chronic disease developing in 20-30%, usually manifesting as  
 339 cardiac or gastrointestinal disease. Chronic infection can be diagnosed through serologic tests for  
 340 antibody to *T. cruzi*.

341  
 342 Blood donor screening in the U.S. beginning in 2007 identified confirmed infections in donors from 37  
 343 states and Puerto Rico, with 57% of all positive tests from California and Florida, areas with significant  
 344 Mexican, Central and South American immigrant populations. Latent infection with *T. cruzi* can persist  
 345 for decades, therefore organ donor screening of high-risk individuals using serology testing is utilized at  
 346 some centers. Transmission of Chagas disease has been studied in 32 organ transplant recipients from  
 347 14 seropositive donors in the United States. Two of 15 (13%) renal transplant recipients had donor-  
 348 derived infection; none of these were living donor transplants. Living organ donor transmissions have  
 349 been reported in Mexico and South America.

### 351 Who Should Be Screened

#### 352 Living and Deceased Donors

353 Deceased donors who were born in a country currently classified as endemic for Chagas disease by the  
 354 CDC must be screened according to OPTN Policy 2.9 pending implementation of *Improve Deceased*  
 355 *Donor Evaluation for Endemic Diseases*<sup>50</sup> and is recommended until then. Screening is recommended for  
 356 living donors born in a country currently classified as endemic for Chagas CDC by the CDC as well.

357  
 358 Screening should also be considered for living and deceased donors in the following circumstances:

- 359 • Children of women born in endemic regions if their birth mother's serology is positive or  
 360 unknown
- 361 • Donors who have resided in an endemic region for more than three months
- 362 • Donors who received a blood transfusion in endemic regions and those who have a previous  
 363 diagnosis of Chagas disease
- 364 • Symptomatic donors who have chronic infection and may present with cardiomyopathy, cardiac  
 365 conduction abnormalities, megaesophagus, megacolon

<sup>50</sup> "Improve Deceased Donor Evaluation for Endemic Diseases," OPTN, Policy Notice, Accessed October 10, 2023, [https://optn.transplant.hrsa.gov/media/lrxfreqj/dtac\\_endemics\\_policy-notice\\_june23bod.pdf](https://optn.transplant.hrsa.gov/media/lrxfreqj/dtac_endemics_policy-notice_june23bod.pdf).

366 Countries currently classified as endemic for Chagas disease by the CDC:

367 Argentina

368 Belize

369 Bolivia

370 Brazil

371 Chile

372 Colombia

373 Costa Rica

374 Ecuador

375 El Salvador

376 French Guiana

377 Guatemala

378 Guyana

379 Honduras

380 Mexico

381 Nicaragua

382 Panama

383 Paraguay

384 Peru

385 Suriname

386 Uruguay

387 Venezuela

388

389 How to Screen

390 Living and Deceased Donors

391 Serology testing using an FDA licensed, approved, or cleared assay (see Table 2) should be performed on  
392 donors with risk factors for Chagas disease. Turnaround time for serology tests ranges from one to 10  
393 days. Tests for parasitemia are not sensitive enough to detect chronic *T. cruzi* infection. Detectable  
394 antibody is usually present within a month of infection and remains present throughout the individual's  
395 life.

**Table 2: Serological Tests Available for *T. cruzi* Infection**

<u>Test name, Manufacturer</u>	<u>Target Antigen</u>	<u>Test Format</u>	<u>Sample Type</u>	<u>FDA-cleared/approved use</u>
<u>Abbott Prism Chagas (T. cruzi [E. coli, recombinant] antigen)<sup>51</sup>, Abbott Laboratories, Abbott Park, IL</u>	<u>T. cruzi recombinant antigens (FP10, FP6, FP3, TcF)</u>	<u>ChLIA<sup>52</sup></u>	<u>Serum/plasma</u>	<u>Donor screening</u>
<u>ORTHO T. cruzi ELISA Test System Ortho-Clinical Diagnostics, Inc. Raritan, NJ</u>	<u>Whole cell lysate</u>	<u>EIA<sup>53</sup></u>	<u>Serum/plasma</u>	<u>Donor screening, individual diagnostics</u>
<u>Chagatest ELISA recombinant v.3.0 <sup>54</sup>Wiener Laboratories S.A.I.C., Rosario, Argentina</u>	<u>Recombinant epimastigote and trypomastigote Proteins</u>	<u>EIA</u>	<u>Serum/plasma</u>	<u>Diagnosis, NOT donor screening test</u>
<u>Hemagen Chagas' Kit, Hemagen Diagnostics, Inc., Columbia, MD</u>	<u>Purified antigens from cultured T. cruzi</u>	<u>EIA</u>	<u>Serum</u>	<u>Diagnosis, NOT donor screening test</u>
<u>Abbott ESA Chagas Assay, Abbott Laboratories, Abbott Park, IL</u>	<u>T. cruzi recombinant antigens (FP10, FP6, FP3, TcF)</u>	<u>Enzyme Strip Assay</u>	<u>Serum/plasma</u>	<u>Supplemental test in donors who test positive with first-line assays, not approved for individual diagnosis</u>

397

398 \*Serologic testing may also be available through the CDC

399

400 Management of Infected Living Donors

401 Living donors who test positive should be informed about the result and offered consultation with ID.

402 Confirmatory testing through a submission to the CDC or performance of at least two different FDA

<sup>51</sup> This test may only be available through local blood bank.

<sup>52</sup> ChLIA, chemiluminescence immunoassay

<sup>53</sup> EIA, enzyme immunoassay

<sup>54</sup> Preferred tests for initial donor screening



403 licensed, approved, or cleared antibody diagnostic tests should be performed. Infected donors may  
404 require further evaluation for chronic Chagas disease and may ultimately require specific treatment.  
405 Evaluation and treatment of these individuals should proceed in accordance with local guidelines, as the  
406 availability of confirmatory diagnostics and anti-parasitic therapy varies.

#### 407 Management of Recipients

409 The risk of transmission of *T. cruzi* infection from an infected donor to an uninfected deceased kidney  
410 donor recipient has been reported to be between 13 and 18%. There are no data regarding living donor  
411 transmissions. It is reasonable to consider accepting donor organs from *T. cruzi* positive donors on an  
412 individual basis with informed consent and close monitoring of the recipient. Potential recipients of a *T.*  
413 *cruzi* positive donor organ should themselves be tested for the presence of *T. cruzi* antibody, particularly  
414 since they may share unrecognized risk factors with their potential living donor. If the recipient tests  
415 negative and the decision is made to proceed with the transplant, recipients must be counseled  
416 specifically about the transmission risk and the need for close monitoring post-transplant, with the  
417 potential need for anti-parasitic therapy that is FDA approved but may lead to side effects that may be  
418 difficult to tolerate.

419  
420 Once transplant has occurred, consultation with ID is recommended to coordinate post-transplant  
421 testing. In the U.S., the CDC can assist with PCR and parasitemia testing. Incorporating a PCR testing  
422 platform into the post-transplant testing program is recommended, as molecular testing may be positive  
423 weeks before parasitemia is detected using standard screening.

424  
425 PCR and parasitemia tests of peripheral blood should be performed using the following schedule:

- 426 • Weekly for the first three months post-transplant
- 427 • Every two weeks for the fourth post-transplant month
- 428 • Monthly thereafter until a minimum of six months post-transplant AND until net state of  
429 immunosuppression is at optimal post-transplant baseline with no evidence of infection or  
430 rejection
- 431 • More frequent monitoring is recommended any time the patient requires antirejection therapy

432  
433 For access to testing of transplant recipients at risk for reactivation of Chagas, contact the CDC  
434 Division of Parasitic Diseases and Malaria at 404-718-4745 (business hours) or 770-488-7100 (nights  
435 and weekends) or via e-mail at [parasites@CDC.gov](mailto:parasites@CDC.gov).

436  
437 Current therapy for Chagas disease is limited to benznidazole and nifurtimox (Lampit<sup>®</sup>, Bayer), both  
438 of which are partially metabolized via cytochrome P450 reductase and may increase tacrolimus and  
439 cyclosporine blood levels. Both drugs have potential adverse effects, but benznidazole is generally  
440 better tolerated and is given twice a day while nifurtimox is given three times a day.

#### 441 **Infection Avoidance between Testing and Transplant**

442  
443 Potential living donors should avoid prolonged travel to endemic regions, particularly rural areas  
444 with primitive housing or significant insect exposure. Consumption of uncooked food or drink from  
445 endemic regions should also be avoided.

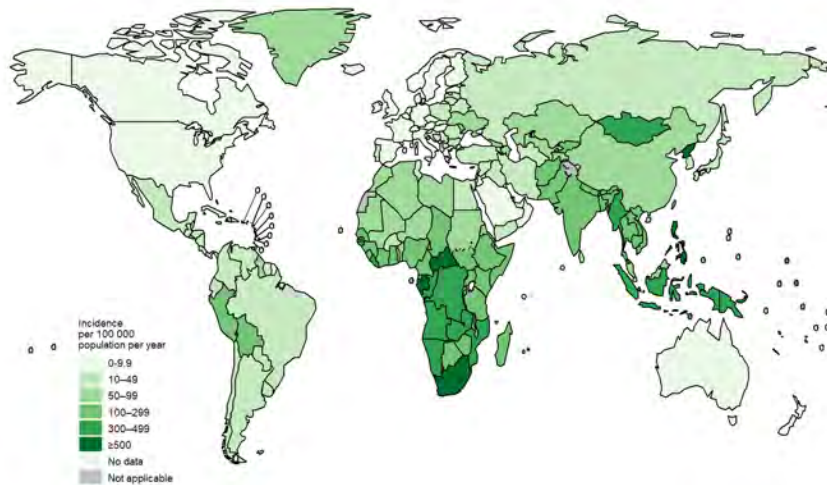
## 446 BACTERIAL INFECTIONS

### 447 Mycobacterium Tuberculosis (MTB) Guidance

#### 448 Background

450 Up to one-third of the world’s population is infected with MTB; however, infection in the United States  
 451 is much less common. Reported cases of MTB disease have been declining in the United States since  
 452 1992 with the majority of cases occurring in foreign-born persons. After initial infection with MTB, most  
 453 people do not develop MTB disease; the infection disseminates throughout the body and remains  
 454 dormant. This condition is called tuberculosis infection (TBI) (previously referred to as “latent  
 455 tuberculosis infection”). In 2022, there were 8,300 reported TB cases<sup>55</sup> in the United States (a rate of 2.5  
 456 cases per 100,000 persons). Since initial infection does result in live MTB in many organs, tuberculosis  
 457 can be transmitted via non-lung organs used for transplantation from donors who have never had  
 458 clinical signs or symptoms of MTB disease.

459 **Figure 3. World Health Organization Estimated Global TB Incidence Rates, 2021<sup>56</sup>**



#### 460

#### 461 Who Should Be Screened

##### 462 Living and Deceased Donors

463 Since TBI precedes the development of MTB disease, similar risk factors would be expected to be  
 464 present. One difference is that since the rate of reactivation decreases with time from infection, donors  
 465 with distant infection would still be at risk for transmission but at a lower risk for MTB disease. The most  
 466 important risk factors for TB infection are:

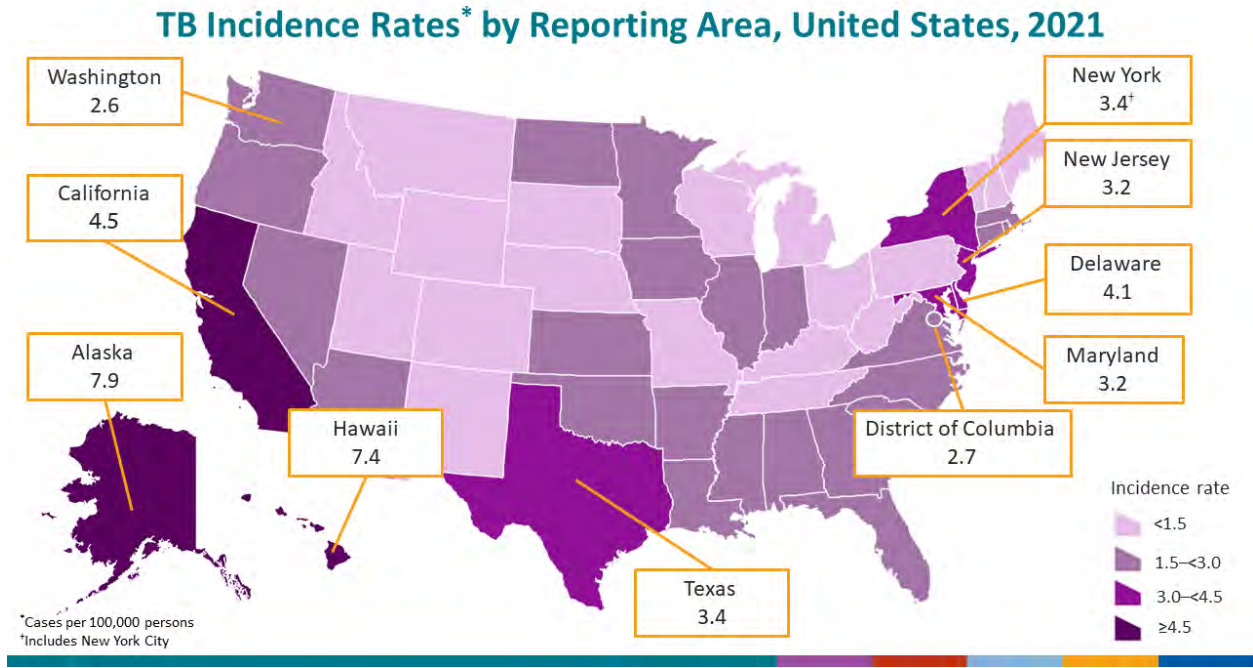
<sup>55</sup> Provisional CDC data

<sup>56</sup> WHO. “2.1 TB Incidence.” World Health Organization, October 27, 2022. <https://www.who.int/teams/global-tuberculosis-programme/tb-reports/global-tuberculosis-report-2022/tb-disease-burden/2-1-tb-incidence>.  
<https://www.who.int/teams/global-tuberculosis-programme/tb-reports/global-tuberculosis-report-2022/tb-disease-burden/2-1-tb-incidence>.

467 powerful easily identifiable risk factor is place of birth with estimated risk of TBI of 18.7% among  
468 foreign-born persons in the U.S. as compared to 1.8% among U.S. born persons. Higher risk countries  
469 are identified by incidence per 100,000 population per year in Figure 3.

470 For individuals from lower risk regions, including the United States, the following factors should help  
471 OPOs and transplant programs make screening decisions:

- 472 • Close contacts of persons with infectious TB disease
- 473 • Those who spend significant time (greater than 3 months) in areas of the world with high rates  
474 of TB or those born in these regions (all but the lightest shaded area of Figure 4) including relief  
475 work in a country with high TB risk.
- 476 • History of non-medical injection drug use
- 477 • Persons who reside (or ever resided) or worked in institutional settings which resulted in  
478 increased risk of exposure to TB (hospitals, nursing homes, correctional facilities, other health  
479 care settings, homeless shelters)
- 480 • Radiographic evidence of prior tuberculosis on chest radiograph
- 481 • Symptoms such as fever, night sweats, weight loss, cough, recurrent pneumonia, exudative  
482 pleural effusion of unknown etiology, lymphadenopathy, non-calcified pulmonary nodules or  
483 cavities

**Figure 4: TB Incidence Rates by Reporting Area, United States, 2021<sup>57</sup>**

485

486 How to Screen487 Living and Deceased Donors

488 While MTB disease can involve almost any organ and cause many syndromes (e.g., meningitis), a careful  
 489 medical history and examination will likely identify most patients with MTB disease. TBI, by definition, is  
 490 asymptomatic, and specific testing is required to identify patients with TBI. The current FDA-approved  
 491 screening methods for TBI in the US include the tuberculin skin test (TST) and the interferon gamma  
 492 release assays (IGRAs): QuantiFERON-TB gold in tube (QFT), and T-SPOT.TB. These tests do not  
 493 differentiate TBI from MTB disease and may be negative during times of MTB disease. The TST test  
 494 requires injection into the skin and a return visit in 48-72 hours to interpret the test. The IGRAs are  
 495 blood tests and may show positive, negative, or indeterminate results.

496 Indeterminate results indicate that the controls did not work in that patient, and this result is more  
 497 common in immunosuppressed patients. One advantage of the IGRAs is that patients who received a  
 498 childhood BCG vaccine (very common outside of North America and Western Europe) are less likely to  
 499 have a false-positive test with IGRAs than with the TST test, due to enhanced specificity of  
 500 the *Mycobacterium tuberculosis* antigens used in the IGRA assays.

<sup>57</sup> CDC. "Health Disparities." Centers for Disease Control and Prevention, November 10, 2022.  
<https://www.cdc.gov/tb/topic/populations/healthdisparities/default.htm>.

501 Management of Infected Living Donors and Recipients

502 Table 3 outlines management of living donors with a history of active TBI and treatment of recipients of  
 503 these organs. In all clinical scenarios, consultation with ID is recommended.

504 **TABLE 3: Management of latent or history of active TB in living donors and recipients of these organs**

<b><u>Clinical Scenario: Living Donor</u></b>	<b><u>Risk for Transmission</u></b>	<b><u>Recommendation</u></b>
<u>History of TBI-treated appropriately</u>	<u>Lower</u>	<u>Monitor recipient clinically</u>
<u>History of TBI-treated insufficiently or not treated or treatment details not clear OR new diagnosis of TBI-positive TST or Interferon gamma release assay found during pre-transplant evaluation; evaluation finds no evidence of active TB</u>	<u>Moderate</u>	<u>Consider deferring transplant if possible until donor has taken some/all of chemoprophylaxis and consider chemoprophylaxis of recipient; monitor clinically.</u>
<u>Unexplained pulmonary apical fibrosis in donor without cavitation and without additional testing</u>	<u>Variable</u>	<u>Defer donation pending further evaluation.</u>
<u>History of MTB disease treated appropriately over two years ago</u>	<u>Lower to moderate</u>	<u>Monitor recipient clinically; consider cultures of previous TB sites if possible. Consider TB prophylaxis of recipient.</u>
<u>History of MTB disease-site remote from transplant (organ not being transplanted) treated appropriately within two years.</u>	<u>Lower to moderate</u>	<u>Monitor recipient clinically; consider cultures of previous TB sites if possible. Suggest chemoprophylaxis of recipient.</u>
<u>History of MTB disease-site remote from transplant (organ not being transplanted) treated insufficiently and/or with other than standard regimen Excluding disseminated or CNS TB.</u>	<u>Higher Increased risk if less than two years since active TB diagnosis.</u>	<u>Defer live donors until adequately treated; recommend cultures of previous TB sites prior to transplant if possible</u>
<u>History of renal MTB disease treated appropriately. (If not treated appropriately donation should be deferred until after appropriate treatment)</u>	<u>Moderate</u>	<u>Verify treatment; monitor clinically; recommend chemoprophylaxis for recipient; recommend cultures of previous TB site(s).</u>

505 Management of Recipients of Deceased Donors

506 For recipients of an organ from a donor who is TST or IGRA-positive, had recent exposure to active TB, or  
 507 had radiographic evidence of untreated TB should be considered for TBI therapy, following CDC  
 508 guidance for anti-tuberculosis agents and durations. Once transplant has occurred, consultation with ID  
 509 is recommended. Specific attention should be given to potential drug-drug interactions of TBI agents  
 510 and immunosuppressive medications with careful monitoring of calcineurin inhibitors.

511  
512 Infection Avoidance between Testing and Transplant

513 Potential living donors should avoid prolonged travel to endemic regions and behavioral exposures that  
 514 increase risk if possible, including institutional settings which may result in increased risk of exposure to  
 515 TB (hospitals, nursing homes, correctional facilities, other health care settings, and homeless shelters).  
 516

**VIRAL INFECTIONS**517 West Nile Virus (WNV) Guidance518 Background519 Epidemiology and pathophysiology

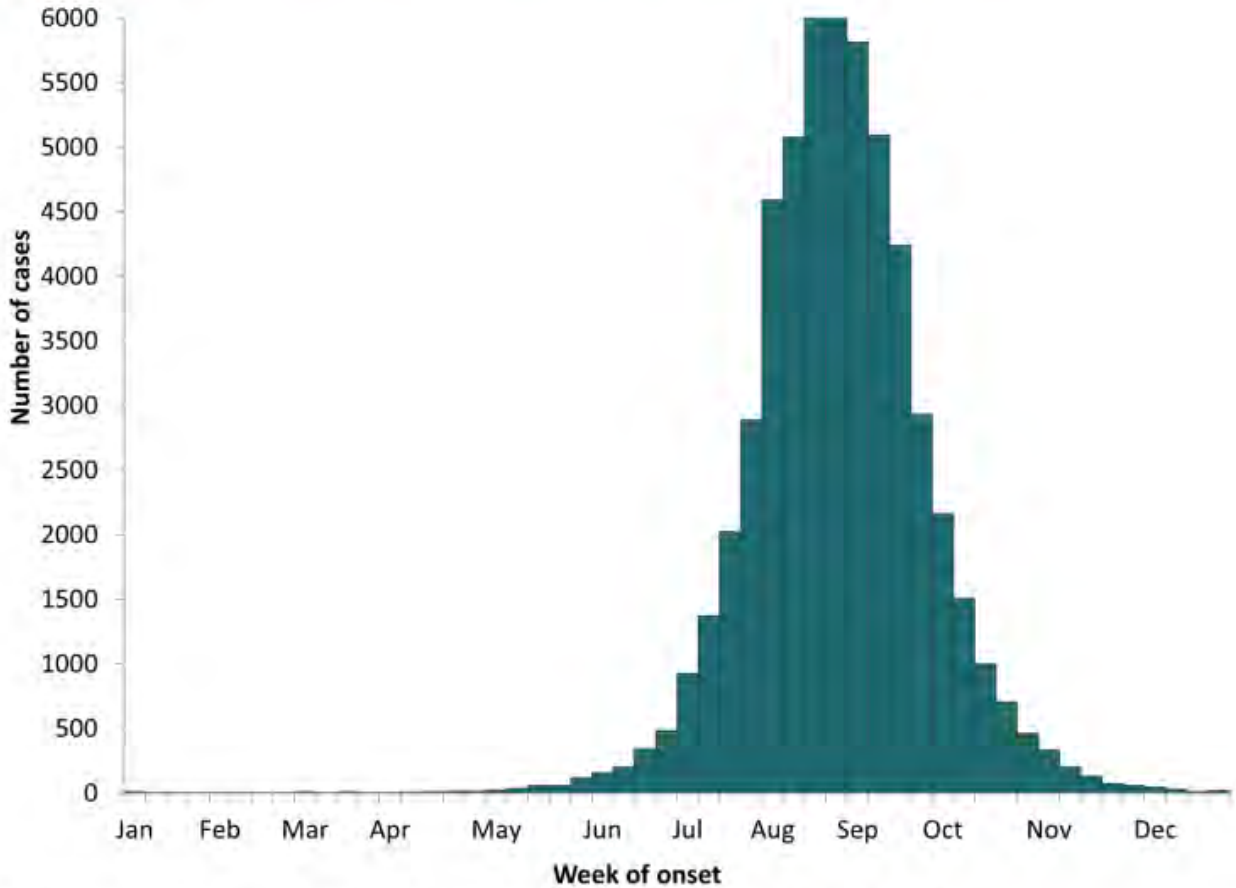
520 WNV is an RNA virus that spreads to humans primarily by the bite of infected mosquitoes, although  
 521 multiple non-vector modes of transmission have been described, including blood transfusion, organ  
 522 transplantation, breastfeeding, intrauterine, and laboratory exposure. Birds serve as a reservoir for  
 523 WNV, and humans are a dead-end host for the virus. The virus was first detected in North America in  
 524 1999, and since then, has spread to all 48 contiguous U.S. states. The number of human infections varies  
 525 significantly based on the region and year, with most transmission (as shown in Figure 5) occurring  
 526 during the warmer months (July to October) when mosquitoes are most active, although warmer  
 527 regions can have a longer season. While the majority of WNV infections are asymptomatic, about 20% of  
 528 those infected may develop fever, and less than one percent develop neurological complications such as  
 529 encephalitis or acute flaccid myelitis (involving the spinal cord). This means that even donors who do not  
 530 display symptoms could still transmit the virus to recipients. While several treatments have been  
 531 explored, there is currently no established treatment for WNV.



532

Figure 5: Months of typical WNV activity in the United States<sup>58</sup>

West Nile virus disease cases reported to CDC by week of illness onset, 1999-2021



Source: ArboNET, Arboviral Diseases Branch, Centers for Disease Control and Prevention

533

534 Who Should Be Screened

535 Living and Deceased Donors

536 Since the transmission of WNV is limited by geography and season, OPOs and transplant programs  
 537 should consider certain factors when determining whether to perform laboratory testing on a potential  
 538 donor. A positive test result obtained during a period of WNV inactivity is more likely to represent a  
 539 false positive than when the test was performed during periods of higher activity. Any false positive  
 540 result could result in unnecessary delays to transplant.

<sup>58</sup> CDC. "West Nile Virus Disease Cases Reported to CDC by Week of Illness Onset ..." West Nile virus cases reported to CDC by week of illness onset, 1999-2021. Accessed June 23, 2023. <https://www.cdc.gov/westnile/resources/pdfs/data/WNV-Week-Onset-1999-2021-P.pdf>.



541 Factors to consider in evaluating donors for possible WNV infection:

- 542 • Has human infection with WNV virus been recognized locally this WNV season?
- 543 • Has the donor travelled to an area with human WNV activity this WNV season?
- 544 • Has the donor ever been diagnosed with WNV fever or WNV neuroinvasive disease?
- 545 • Has the donor had an undifferentiated febrile illness within the current WNV season?
- 546 • Has the donor had significant mosquito exposure this WNV season?

547 *Geographic and seasonal factors to consider.*

548 Screening strategies for WNV in donors include universal year-round testing of all donors versus  
 549 targeted testing during periods of human WNV activity. As the yield of testing during the winter months  
 550 is likely low and given the potential for IgM to remain positive after resolution of the illness, year-round  
 551 testing is discouraged unless programs are unable to manage the complexity of more targeted testing  
 552 strategies. Targeted testing strategies could include testing during a pre-determined timeframe, or  
 553 testing donors only during times of human WNV activity in the area where the donor lives or has  
 554 traveled (i.e., triggered strategy).

555 Triggering strategies can involve communicating with local blood collection organizations and  
 556 determining whether they have shifted from minipool to individual donation (ID)-NAT screening,  
 557 indicating that a WNV-reactive donor has been identified within a 50-mile radius of the triggering zip  
 558 code. Another method involves using WNV human and non-human surveillance data collected by state  
 559 and local health departments. Alternatively, testing could begin each year during the local WNV  
 560 transmission season as determined by historical data (typically July through October, or longer intervals  
 561 in warmer regions).

562 Given that human WNV disease incidence varies greatly from year to year and even county to county,  
 563 targeted testing based on real-time measurement of local WNV activity at local blood banks might be  
 564 the most cost-effective strategy and will reduce the number of false positives. This strategy is, however,  
 565 more complex and time-consuming than a fixed seasonal strategy. Some institutions may already be  
 566 using a local or regional blood bank testing lab to test other material (e.g., peripheral stem cells used in  
 567 stem cell transplantation). In contrast, the strategy of testing during a defined time period regardless of  
 568 local WNV activity would be simpler to implement, but both costs and false positive rates would be  
 569 expected to be higher. **Table 4** describes the advantages and disadvantages of different testing  
 570 strategies.

571

**Table 4: Advantages and Disadvantages of Different Testing Strategies**

	<u>Year-round testing (1)</u>	<u>Seasonal testing (2)</u>	<u>Triggered testing (3)</u>
<u>Ease of implementation</u>	<u>Easy</u>	<u>Intermediate</u>	<u>Difficult</u>
<u>Positive predictive value</u>	<u>Lower during periods of human WNV inactivity</u>	<u>Intermediate depending on level of human WNV activity</u>	<u>High</u>
<u>Cost effectiveness</u>	<u>Least</u>	<u>Intermediate</u>	<u>Most</u>

572

573

574

575

576

577

578

1. This strategy is not recommended.
2. Typically July through October, but time period should be guided by local historical WNV data, in consultation with state and local health departments
3. Testing triggered by switch from minipool to individual blood donation-NAT testing in zip codes of residence of donors. Testing stops when WNV activity no longer noted and blood banks switch back to minipools. Routine communication with local blood banks is required.

579

How to Screen

580

Living and Deceased Donors

581

582

583

584

585

586

587

588

589

To screen for WNV, healthcare professionals use NAT and IgM serologic tests. Currently, there are two FDA-licensed NAT donor screening assays; these are routinely performed at blood banks or some reference labs but are not commonly available in hospital laboratories. Serologic tests which rely on the response of the immune system to infection with WNV, particularly the IgM assay performed on serum and cerebrospinal fluid, are commonly used to diagnose neuroinvasive WNV. IgM is detectable for a median of about five months after infection but may persist longer, even years after infection, and is not used in blood donors to screen for WNV, although it may be performed for use in donor counseling. IgG typically remains positive for life and would not be useful to routinely screen for WNV. Table 5 describes the characteristics of NAT and IgM testing.

590

**Table 5: Tests that could be used to screen for WNV infection**

	<u>Nucleic Acid Tests (NAT)(1)</u>	<u>IgM (serology)(2)</u>
<u>Available Tests</u>	<u>Procleix West Nile Virus Assay COBAS TaqScreen West Nile Virus Test</u>	<u>Various</u>
<u>FDA licensed for organ screening</u>	<u>Yes</u>	<u>No</u>
<u>Availability</u>	<u>Blood bank testing labs Reference labs</u>	<u>Reference labs Commercial labs State public health labs</u>
<u>False positive rate</u>	<u>Low</u>	<u>Likely higher than NAT, but not evaluated for donor screening</u>
<u>Indicates active infection</u>	<u>Yes</u>	<u>Remains positive for median of five months (up to seven years documented); active infection may have cleared</u>
<u>Required for blood donor screening</u>	<u>Yes</u>	<u>No</u>

- 591 1. Should be used as part of any testing strategy
- 592 2. Consider in combination with NAT testing but will increase false positive rate

593 Deceased donors

594 It is advisable to screen deceased donors during months of regional WNV activity. WNV NAT testing is a  
595 viable option, but the results may not be available before transplantation. It is important to note that  
596 WNV transmission has occurred in donors who tested negative on NAT, and that there is potential for  
597 false positive NAT test results if protocols are not in place to improve specificity, as are used by blood  
598 collection organizations, or if testing is performed outside of the WNV transmission season, decreasing  
599 the pretest probability.

600 In cases involving deceased donors, the feasibility of conducting WNV testing within a required  
601 timeframe is uncertain. As such, the Committee advises against proceeding with donors who satisfy any  
602 of the following criteria: (1) a confirmed WNV infection, (2) a positive WNV NAT test result, ideally  
603 meeting a threshold signal-to-cutoff ratio or verified by repeat testing as recommended by the  
604 manufacturer, or (3) clinical manifestations of meningitis, encephalitis, or flaccid paralysis of  
605 indeterminate origin, particularly if they originate from regions with documented WNV activity.

606 Living donors

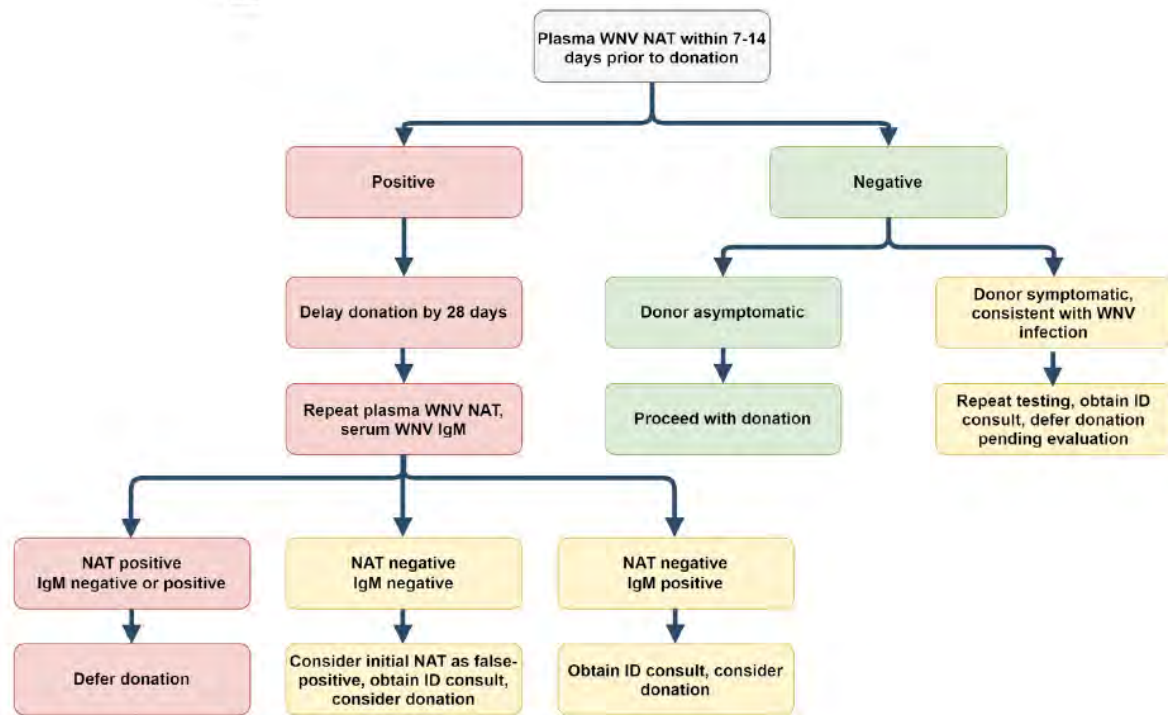
607 Living donors should be screened with WNV NAT within seven to 14 days of donation.

608

609 Management of Infected Living Donors

610 Living donors who test positive should be informed about the result and offered consultation with ID.  
611 Although no studies have examined organ donation after WNV infection, we suggest postponing  
612 donation for a minimum of 28 days in living donors, after which NAT and IgM testing should be  
613 performed. If negative, it is likely that the initial NAT test was a false positive, and organ donation can be  
614 considered. If the NAT is negative but IgM is positive, this likely reflects viral clearance and organ  
615 donation can be considered. However, if the NAT remains positive, organ donation should be  
616 postponed. (Refer to Figure 6 for additional information).

617

**Figure 6: Living donor screening recommendations for WNV<sup>59</sup>**

618

### 619 Management of Recipients

620 Development of symptoms compatible with WNV disease within the first several weeks of transplant  
 621 (and up to six weeks, since transplant recipients can have prolonged incubation periods with WNV  
 622 infection) should prompt testing in the recipient. There are currently no specific medications available  
 623 for the treatment of WNV infection. Once transplant has occurred, consultation with ID is  
 624 recommended.

625

### 626 Infection Avoidance between Testing and Transplant

627 We advise potential living donors to take measures to prevent mosquito exposure during periods of  
 628 mosquito activity, such as wearing loose-fitting, long sleeve shirts and pants, using EPA-registered insect  
 629 repellent, and avoiding outdoor activities during dawn and dusk when mosquitoes are most active.  
 630 Further, a potential living donor should report febrile illnesses to his or her transplant center. WNV  
 631 diagnostic testing should be performed if clinical evaluation suggests the possibility of WNV infection in  
 632 the potential living donor.

<sup>59</sup> Anesi, JA, Silveira, FP; the AST Infectious Diseases Community of Practice. Arenaviruses and West Nile Virus in solid organ transplant recipients: Guidelines from the American Society of Transplantation Infectious Diseases Community of Practice. *Clin Transplant*. 2019; 33:e13576. <https://doi.org/10.1111/ctr.13576>

## **SHARING INFORMATION**

633  
634  
635  
636  
637  
638  
639  
640  
641  
642  
643  
644  
645  
646  
647  
648

### *Recommendations for Sites Participating in Paired Organ Exchanges*

Transplant hospitals participating in organ exchanges should consider sharing their protocols for screening donors for seasonal and geographically endemic infections in addition to related test results. This will provide recipient centers more information about the donor screening process at the donor organ center.

### *When to Report a Potential Donor-Derived Transmission Event*

If the recipient is suspected to be at risk for disease transmission either by the OPO or a transplant center, a potential donor-derived disease transmission event should be reported to the OPTN Improving Patient Safety Portal per OPTN Policies 15.4 Host OPO Requirements for Reporting Post-Procurement Test Results and Discovery of Potential Disease Transmissions and 15.5 Transplant Program Requirements for Communicating Post Transplant Discovery of Disease or Malignancy. This promotes prompt intervention for other recipients of the same donor’s organs.

649 **References**

650

651 Abad, Cybele L., and Raymund R. Razonable. "Donor Derived Mycobacterium Tuberculosis Infection  
 652 after Solid-Organ Transplantation: A Comprehensive Review." *Transplant Infectious Disease* 20, no. 5  
 653 (2018). <https://doi.org/10.1111/tid.12971>.

654 Abanyie FA, Gray EB, Delli Carpini KW, et al. Donor-Derived Strongyloides stercoralis Infection in Solid  
 655 Organ Transplant Recipients in the United States, 2009–2013. *AJT* 2015;15(5):1369-1375

656

657 Anesi JA, Silveira FP. Arenaviruses and West Nile virus in solid organ transplant recipients: guidelines  
 658 from the American Society of Transplantation Infectious Diseases Community of Practice. *Clinical*  
 659 *Transplantation*. 2019;33(9):e13576. <https://doi.org/10.1111/CTR.13576>

660 Assi M, Martin S, Wheat LJ, et al. Histoplasmosis after solid organ transplant. *Clin Infect Dis* 2013 ; 57  
 661 (11) : 1542.

662

663 Bennett DE, Courval, JM. Prevalence of tuberculosis infection in the Unites States population: the  
 664 national health and nutrition examination survey, 1999-2000. *American Journal of Respiratory and*  
 665 *Critical Care Med* 2008; 177 (3):348-355 2008

666 Center for Biologics Evaluation and Research, U.S. Food and Drug Administration. Complete list of DSA  
 667 for infectious agents and HIV diagnostic assays. Accessed May 18, 2023. [https://www.fda.gov/vaccines-](https://www.fda.gov/vaccines-blood-biologics/complete-list-donor-screening-assays-infectious-agents-and-hiv-diagnostic-assays#AntiT.%20Cruzi%20Assays%20(Detect%20antibodies%20to%20Trypanosome%20cruzi))  
 668 [blood-biologics/complete-list-donor-screening-assays-infectious-agents-and-hiv-diagnostic-](https://www.fda.gov/vaccines-blood-biologics/complete-list-donor-screening-assays-infectious-agents-and-hiv-diagnostic-assays#AntiT.%20Cruzi%20Assays%20(Detect%20antibodies%20to%20Trypanosome%20cruzi))  
 669 [assays#AntiT.%20Cruzi%20Assays%20\(Detect%20antibodies%20to%20Trypanosome%20cruzi\)](https://www.fda.gov/vaccines-blood-biologics/complete-list-donor-screening-assays-infectious-agents-and-hiv-diagnostic-assays#AntiT.%20Cruzi%20Assays%20(Detect%20antibodies%20to%20Trypanosome%20cruzi)).

670 Center for Biologics Evaluation and Research, FDA. "Biologics Guidances." U.S. Food and Drug  
 671 Administration. Accessed May 24, 2023. [https://www.fda.gov/vaccines-blood-biologics/guidance-](https://www.fda.gov/vaccines-blood-biologics/guidance-compliance-regulatory-information-biologics/biologics-guidances)  
 672 [compliance-regulatory-information-biologics/biologics-guidances](https://www.fda.gov/vaccines-blood-biologics/guidance-compliance-regulatory-information-biologics/biologics-guidances).

673 Centers for Disease Control and Prevention. "Health Disparities." Health Disparities in TB, November 10,  
 674 2022. <https://www.cdc.gov/tb/topic/populations/healthdisparities/default.htm>.

675 Centers for Disease Control and Prevention. "Histoplasmosis Maps." Histoplasmosis Maps, January 14,  
 676 2021. <https://www.cdc.gov/fungal/diseases/histoplasmosis/maps.html>.

677 Centers for Disease Control and Prevention. "History, Epidemiology, and Risk Factors." Chagas Disease:  
 678 What U.S. Clinicians Need to Know, August 2012.  
 679 [https://www.cdc.gov/parasites/cme/chagas/lesson\\_1/4.html](https://www.cdc.gov/parasites/cme/chagas/lesson_1/4.html).

680 Centers for Disease Control and Prevention. "Incidence." 2020 State and City TB Report, January 13,  
 681 2022. <https://www.cdc.gov/tb/statistics/indicators/2020/incidence.htm>.

682 Center for Disease Control and Prevention. Parasites – Strongyloides: Epidemiology & Risk Factors.  
 683 December 31, 2018. Accessed at <https://www.cdc.gov/parasites/Strongyloides/epi.html> on March 22,  
 684 2023.

- 685 Centers for Disease Control and Prevention. “Valley Fever Maps.” Fungal Diseases, May 21, 2020.  
686 <https://www.cdc.gov/fungal/diseases/coccidioidomycosis/maps.html>.
- 687 Centers for Disease Control, Division of Vector-Borne Diseases, WNV Home  
688 Page <http://www.cdc.gov/ncidod/dvbid/westnile/index.htm>
- 689 Centers for Disease Control and Prevention. “Treatment for Latent TB Infection and TB Disease.” Centers  
690 for Disease Control and Prevention, March 22, 2023.  
691 <https://www.cdc.gov/tb/topic/treatment/default.htm>.
- 692 Chin-Hong PV, Schwartz BS, Bern C, Montgomery SP, Kontak S, Kubak B, Morris MI, Nowicki M, Wright C,  
693 Ison MG. Screening and treatment of Chagas disease in organ transplant recipients in the United States:  
694 recommendations from the Chagas in transplant working group. Am J Transplant 2011 Apr ;11(4) :672-  
695 80.
- 696  
697 “Diagnosis and Management of Tuberculosis in Transplant Donors: A Donor- Derived Infections  
698 Consensus Conference Report”. Morris MI, Daly JS, Blumberg E, Kumar D, Sester M, Schluger N, Kim SH,  
699 Schwartz BS, Ison MG, Humar A, Singh N, Michaels M, Orlowski JP, Delmonico F, Pruett T, John GT,  
700 Kotton CN. Am J Transplant. 2012 Aug 6.
- 701 Galgani JN, Ampel HM, Blair JE, et al. 2016 Infectious Diseases Society of America (IDSA) Clinical Practice  
702 Guideline for the Treatment of Coccidioidomycosis. CID 2016;63:e112-e146.
- 703  
704 Huprikar S, Bosserman E, Patel G, Morre A, et al. Donor-derived Trypanosoma cruzi infection in solid  
705 organ recipients in the United States, 2001-2011. Am J Transplant 2013 Sep;13(9):2418-25.
- 706  
707 Kusne S, Tranto S, Covington S, et al. Coccidioidomycosis transmission through organ transplantation: A  
708 report of the OPTN Ad Hoc Disease Transmission Advisory Committee. AJT 2016;16:3562-3567.iv Blair  
709 JE, Mulligan DC. Coccidioidomycosis in healthy persons evaluated for liver or kidney donation. Transpl  
710 Infect Dis 2007; 9 (1): 78.
- 711  
712 Le M, Ravin K, Hasan A, et al. Single donor-derived strongyloidiasis in three solid organ transplant  
713 recipients: case series and review of the literature. Am J Transplant 2014;14:1199-206.
- 714  
715 World Health Organization. Chagas disease (also known as American trypanosomiasis), April 13, 2022.  
716 Accessed at [https://www.who.int/news-room/fact-sheets/detail/chagas-disease-\(American-](https://www.who.int/news-room/fact-sheets/detail/chagas-disease-(American-trypanosomiasis))  
717 [trypanosomiasis\)](https://www.who.int/news-room/fact-sheets/detail/chagas-disease-(American-trypanosomiasis)) on March 22, 2023.
- 718  
719 Limaye AP, Connolly PA, Sagar M, et al. Transmission of Histoplasma capsulatum by Organ  
720 Transplantation. New England Journal of Medicine 2000; 343 (16): 1163.
- 721  
722 Levi ME, Kumar D, Green M, Ison MG, Kaul D, Michaels MG, Morris MI, Schwartz BS, Echenique IA,  
723 Blumberg E, et al. Considerations for screening live kidney donors for endemic infections: a viewpoint on  
724 the UNOS policy. Am J Transplant 2014 May;14(5):1003-11.
- 725  
726 Martín-Dávila P, Fortún J, López-Vélez R, Norman F, Montes de Oca M, Zamarrón P, González MI,  
727 Moreno A, Pumarola T, Garrido G, Candela A, Moreno S. Transmission of tropical and geographically



728 restricted infections during solid-organ transplantation. Clin Microbiol Rev. 2008 Jan;21(1):60-96. Doi:  
729 10.1128/CMR.00021-07. PMID: 18202437; PMCID: PMC2223841.

730

731 Miller R, Assi M. Endemic fungal infections in solid organ transplant recipients – Guidelines from the  
732 American Society of Transplantation Infectious Disease Community of Practice. Clin Transplantation  
733 2019;33:e13553.

734

735 Neglected Tropical Diseases, Strongyloides. World Health Organization, 2014. (Accessed April 24, 2014,  
736 at [http://www.who.int/neglected\\_diseases/diseases/strongyloidiasis/en/.](http://www.who.int/neglected_diseases/diseases/strongyloidiasis/en/))

737

738 Pierrotti LC, Carvalho NB, Amorin JP, et al. Chagas disease recommendations for solid-organ transplant  
739 recipients and donors. Transplantation 2018;102(2S-2):S1-S7.

740

741 Parasites – Strongyloides. Centers for Disease Control and Prevention, 2014. (Accessed April 24, 2014, at  
742 [http://www.cdc.gov/parasites/strongyloides/.](http://www.cdc.gov/parasites/strongyloides/))

743

744 Roxby AC, Gottlieb GS, Limaye AP. Strongyloidiasis in transplant patients. Clin Infect Dis 2009;49:1411-  
745 23.

746

747 Schwartz BS, Mawhorter SD. Parasitic infections in solid organ transplantation. Am J Transplant 2013  
748 Mar;13 Suppl 4:280-303

749

750 Singh N, Huprikar S, Burdette SD, et al. Donor-derived fungal infections in organ transplant recipients:  
751 guidelines of the American Society of Transplantation, infectious diseases community of practice. Am J  
752 Transplant 2012; 12 (9): 2414.

753

754 Subramanian, AK, Theodoropoulos, NM; on behalf of the Infectious Diseases Community of Practice of  
755 the American Society of Transplantation. Mycobacterium tuberculosis infections in solid organ  
756 transplantation: Guidelines from the infectious diseases community of practice of the American Society  
757 of Transplantation. Clin Transplant. 2019; 33:e13513. <https://doi.org/10.1111/ctr.13513>

758 U.S. Food and Drug Administration. Guidance for Industry: Use of Nucleic Acid Tests to Reduce the Risk  
759 of Transmission of West Nile Virus from Donors of Whole Blood and Blood Components Intended for  
760 WHO. “Global Tuberculosis Report.” World Health Organization. Accessed May 18, 2023.  
761 [https://www.who.int/teams/global-tuberculosis-programme/tb-reports.](https://www.who.int/teams/global-tuberculosis-programme/tb-reports)

#

## Appendix A: Post-Public Comment Changes

New language that was proposed following public comment is underlined and highlighted (example); language that is proposed for removal following public comment is struck through and highlighted (example).

### **Recognizing Seasonal and Geographically Endemic Infection in Organ Donors: Considerations ~~during Deceased and Living Donor Evaluation~~ for Deceased and Living Donation**

#### Table of Contents

<u>Background</u>	17
<u>Summary and Goals</u>	17
<u>Developing a Written Protocol for Identifying and Testing Donors</u>	17
<u>Recognizing Risk Factors</u>	18
<u>FUNGAL INFECTIONS</u>	21
<u>PARASITIC INFECTIONS</u>	24
<u>BACTERIAL INFECTIONS</u>	31
<u>VIRAL INFECTIONS</u>	35

#### Background

Both deceased and living donors are at risk for transmitting endemic diseases that are either geographically or seasonally defined. Since 2014, Organ Procurement and Transplantation Network (OPTN) policy has required living donor hospitals to develop a written protocol for identification and testing potential donors at risk for these endemic infections as part of the medical evaluation process. More recently on June 26, 2023, the OPTN Board of Directors approved additional policies<sup>60</sup> to address screening of deceased donors for endemic diseases to further reduce the risk for potential donor-derived infectious events.

#### Summary and Goals

The OPTN Ad Hoc Disease Transmission Advisory Committee (DTAC) created this guidance document to assist programs in identification of potential living and deceased donors who may carry an increased risk of transmitting seasonal or geographically endemic disease to organ recipients. This document will also

<sup>60</sup> Improve Deceased Donor Evaluation for Endemic Diseases," OPTN, Policy Notice, Accessed October 10, 2023, [https://optn.transplant.hrsa.gov/media/lrxfreqj/dtac\\_endemics\\_policy-notice\\_june23bod.pdf](https://optn.transplant.hrsa.gov/media/lrxfreqj/dtac_endemics_policy-notice_june23bod.pdf).

help programs manage recipients who receive organs from donors with endemic diseases. This resource is not OPTN policy, so it does not carry the monitoring or enforcement implications of policy. It is not an official guideline for clinical practice, nor is it intended to be clinically prescriptive or to define a standard of care. This is a resource tool intended to be of educational support for organ procurement organizations (OPOs) and transplant centers and is for voluntary use by members.

## Developing a Written Protocol for Identifying and Testing Donors

The DTAC reviews potential donor-derived disease transmission events (PDDTE) reported to the OPTN for both deceased and living donors. A number of the PDDTE reported are seasonally and geographically associated. Some of the reported events resulted in recipient illness or death. Recognition of disease in these donors can be especially challenging, as risk factors may not be recognized, and laboratory screening is not generally universally performed. This guidance document provides information the DTAC believes will help programs and OPOs identify and test donors at risk for transmissible seasonally or geographically defined disease. This information is meant to assist the transplant community in performing organ donor and recipient infectious disease screening procedures as part of the overall medical evaluation and recipient management process.

## Recognizing Risk Factors

There are several factors to consider in determining a donor's risk of transmissible infection. OPOs and living donor recovery hospitals should consider the following when screening potential organ donors:

- Geographic risks (including duration of time spent in a location)
  - Birthplace of the potential donor
  - Home country/region of the U.S., and prolonged residence outside home region, either recent or remote
  - Close family members' countries of origin
  - Donor hospital region
  - Occupational or recreational travel to other countries and/or regions
- Occupational risks
  - Healthcare workers, veterinarian/animal care workers
  - Construction workers, landscapers, park rangers, and other outdoor workers
  - Peace Corps workers, international journalists
  - Current or previous military service, particularly outside the U.S.
  - Medical mission trips
  - **Correctional workers**
- Seasonal risks
  - Residence in/travel to warm weather climates with potential insect exposures
- Hobbies
  - **Hunting/dressing game, taxidermy**
  - Time living outdoors including camping, swimming in lakes, drinking stream water, insect exposures
  - Adventure sports
  - Gardening
- Significant animal exposure (wild and/or domestic)
  - Large numbers of cats or dogs or any unusual pets
  - Laboratory/research animals

- Veterinarian/vet assistant
- Family members and close contacts with potential risk factors
  - Geographic or seasonal infections previously diagnosed in close family members or other contacts may predict risk for subclinical infection in the potential donor
- Personal history of seasonal or geographic infection in the potential donor, even if remote
- **Deceased donor cause of death (if associated with signs/symptoms of infectious illness)**

The organ donor population has become increasingly geographically diverse, reflecting the enhanced mobility and complex migration patterns of the general population. Therefore, it is not practical to list all the pathogens that have the potential for transmission through organ transplantation. Parasitic infections such as amebiasis, babesiosis, leishmaniasis, schistosomiasis, echinococcosis, and malaria, bacterial infections such as brucellosis and melioidosis, fungal infections such as paracoccidioidomycosis and talaromycosis, and viral infections such as Eastern Equine Encephalitis Virus have distinct areas of endemicity and may be transmitted through the organ allograft. In donors with a history of residence in developing countries or remote regions, unusual occupational exposure risks, or extensive travel, infectious disease (ID) consultation may be helpful as part of the pre-transplant evaluation for living donors or prior to organ acceptance for deceased donors. This guidance document does not replace **expert ID evaluation consultation with ID.**

Table 1 covers **a number of several** common seasonal and geographically endemic infections that may be transmitted from organ donor to recipient.

**Table 1: Common Seasonal and Geographically Endemic Infections in Organ Donors**

Disease	Signs/Symptoms in Potential Donor	Known Risk Factors	Potential Testing for Deceased Donors	Potential Testing For Living Donors	Imaging that may be helpful
Histoplasmosis	Fever, night sweats, lymphadenopathy, cough, non-calcified pulmonary nodules or cavities	Residence in Midwestern or South-Central states along the Mississippi or Ohio River Valleys	<b>Serology</b> <b>-complement fixation immunodiffusion</b> <b>-EIA-Urine or serum antigen testing</b> <b>-Urine and/or serum antigen enzyme immunoassay</b> <b>-Serology: complement fixation and/or immunodiffusion</b>	<b>Serology</b> <b>-complement fixation immunodiffusion</b> <b>-EIA-Urine or serum antigen testing</b> <b>-Urine and/or serum antigen enzyme immunoassay</b> <b>-Serology: complement fixation and/or immunodiffusion</b>	Chest X-ray (CXR) or CT

Disease	Signs/Symptoms in Potential Donor	Known Risk Factors	Potential Testing for Deceased Donors	Potential Testing For Living Donors	Imaging that may be helpful
Coccidioidomycosis	Fever, joint pains, cough, neck stiffness, headaches, pulmonary nodules or cavities, reticulonodular infiltrates	Residence in endemic areas of Washington state, the Southwestern United States, Northern Mexico, and parts of South & Central America	Serology: -enzyme immunoassay (preferred) -complement fixation -immunodiffusion -Urine or serum antigen testing	Serology: -enzyme immunoassay (preferred) -complement fixation -immunodiffusion -Urine or serum antigen testing	CXR or CT
Strongyloidiasis	Chronic abdominal pain, bloating, heartburn, intermittent diarrhea and constipation, dry cough, skin rashes, and/or eosinophilia, or could be entirely asymptomatic.	Soil exposure in tropical/warm climates. Walking barefoot or unprotected skin contact with human sewage or contaminated soil. Infection may persist for decades.	Serologic testing is required for all <sup>61</sup> *pending policy implementation of <i>Improve Deceased Donor Evaluation of Endemic Diseases</i>	Donors could be tested by serology (preferred) and/or stool ova and parasite examination, specifically looking for <i>Strongyloides</i> .	None
Chagas disease	Most asymptomatic; symptomatic chronic infection may present with cardiomyopathy, cardiac conduction abnormalities, megaesophagus, megacolon	Born or resided in endemic areas of Mexico, South & Central America, child of woman who lived in endemic area, received blood transfusion in endemic area	Required by OPTN policy if donor is born in endemic country <sup>62</sup> Serology testing (See Chagas Guideline, Table 3) *pending policy implementation of <i>Improve Deceased Donor Evaluation of Endemic Diseases</i>	Serology testing (See Chagas Guideline, Table 3)	None unless symptomatic with chronic Chagas disease

<sup>61</sup> Pending policy implementation of *Improve Deceased Donor Evaluation for Endemic Diseases*.

<sup>62</sup> Pending policy implementation of *Improve Deceased Donor Evaluation for Endemic Diseases*.

Disease	Signs/Symptoms in Potential Donor	Known Risk Factors	Potential Testing for Deceased Donors	Potential Testing For Living Donors	Imaging that may be helpful
Tuberculosis	Fever, night sweats, weight loss, cough, recurrent pneumonia, exudative pleural effusion of unknown etiology, lymphadenopathy, non-calcified pulmonary nodules or cavities	Born outside U.S., prolonged residence outside U.S., homeless, alcohol or other substance abuse, jail/prison time, health care worker, known TB exposure	Positive tuberculin skin test (TST) or interferon gamma release assay (IGRA)**; Sputum/BAL AFB smear, culture, nucleic acid amplification, TB PCR; tissue AFB smear, culture, TB PCR **Deceased donors on high-dose steroids may have false-negative IGRA/TST	Positive tuberculin skin test (TST) or interferon gamma release assay (IGRA); sputum/BAL AFB smear, culture, nucleic acid amplification, TB PCR; tissue AFB smear, culture, TB PCR *refer to OPTN Policy 14	CXR, CT thorax, CT abdomen/pelvis (renal TB)
West Nile Virus	Often asymptomatic; 20% develop acute febrile illness; <1% encephalitis, myelitis	Mosquito exposure, blood transfusion; risk varies by season & location	Nucleic acid test (NAT)	NAT	None

## FUNGAL INFECTIONS

### Histoplasmosis Guidance

#### *Background*

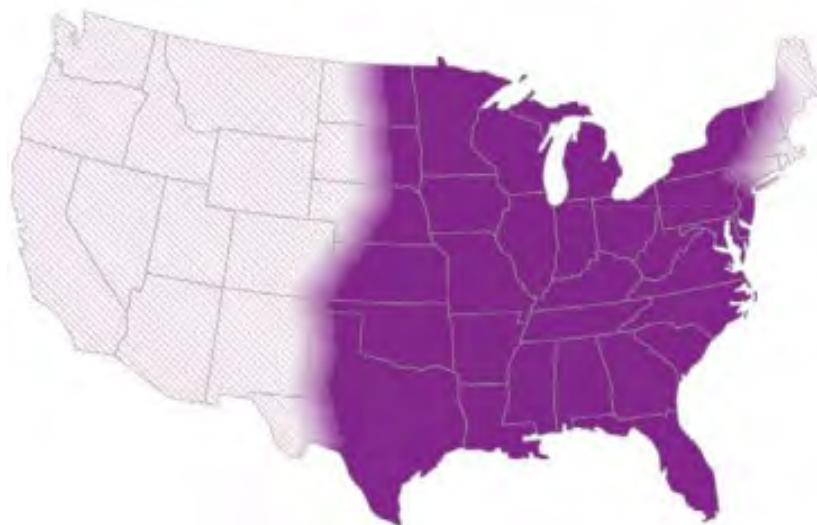
*Histoplasma capsulatum* is a dimorphic fungus found throughout the world. It is endemic in the U.S. in the Ohio and Mississippi River Valleys and into the Midwest and South-Central states (Figure 1). In most cases, clinical signs of infection in immunocompetent patients are limited to the lungs, though many patients are asymptomatic. Immunocompromised patients may develop severe disseminated disease that is fatal without prompt diagnosis and treatment. Donor-derived histoplasmosis has only rarely been reported. In endemic areas, up to 75% of the population is infected with *Histoplasma capsulatum*; therefore, no further exposure risk factors need be elicited in residents of these areas.

#### *Who Should be Screened*

##### Living and Deceased Donors

Donors at higher risk of transmitting disease include those with recent infection or a prior history of known active histoplasmosis and should be screened. Potential donors with a history of pneumonia in the past **two years three to six months** where the clinical scenario was suspicious of histoplasmosis should also be considered for screening. Donors with signs, symptoms, or radiological findings consistent with active histoplasmosis (cough, fever, night sweats, weight loss, non-calcified pulmonary nodules/cavities, or lymphadenopathy) need a thorough medical evaluation including testing for histoplasmosis prior to organ donation. Even in endemic areas, universal screening of donors lacking any of the above characteristics is not likely to be productive. Potential donors with the sole finding of a calcified granuloma on chest imaging do not require further testing.

**Figure 1: Centers for Disease Control and Prevention’s (CDC) current estimate of *Histoplasma* endemicity in the United States<sup>63</sup>**



<sup>63</sup> CDC. “More Information about the Estimated Areas with Blastomycosis ...” More information about the estimated areas with blastomycosis, coccidioidomycosis (Valley fever), and histoplasmosis in the United States. Accessed June 23, 2023.



Darker shading indicates areas where *Histoplasma* is more prevalent. Diagonal shading shows the potential geographic range of *Histoplasma*.

### *How to Screen*

#### Living and Deceased Donors

In asymptomatic potential donors, Serological testing can be used to screen those at risk for the infection. Complement fixation and immunodiffusion testing are the most commonly performed tests. screening should be conducted through urine and/or serum antigen enzyme immunoassays. Alternatively, serological testing can be used to screen those at risk for infection. Complement fixation and immunodiffusion testing are the most commonly performed tests. Complement fixation and immunodiffusion testing are the most commonly performed tests. Complement fixation titers of 1:8 and 1:16 may just indicate previous infection at low risk for transmission. Positive immunodiffusion testing is more concerning for the presence of viable organisms, particularly if a H band is present.

### *Management of Infected Living Donors*

Potential living donors with clinical evidence of active histoplasmosis (fever, night sweats, lymphadenopathy, cough, non-calcified pulmonary nodules or cavities) or a serological pattern suggestive of active disease should receive treatment prior to organ donation. Living donors should be informed about the result and offered consultation with ID. For patients with histoplasmosis limited to the lungs, a reasonable approach would include six to 12 weeks of treatment, with resolution of clinical signs and symptoms of histoplasmosis. After treatment and resolution of antigenuria/antigenemia, the risk of donor-derived infection is likely low. Given that disseminated histoplasmosis is typically associated with other significant medical conditions, it is unlikely that a patient with a history of disseminated histoplasmosis would qualify for living organ donation.

### *Management of Recipients*

Recipients of donors with a history of histoplasmosis may not require specific prophylaxis. Once transplant has occurred, consultation with ID is recommended. Clinical and laboratory monitoring for disease is a reasonable approach. Serial urine or serum antigen testing (every one to three months) for one year might identify donor-derived histoplasmosis prior to the development of clinical symptoms. Serological tests have low sensitivity after organ transplantation, and most solid organ transplant recipients with histoplasmosis have negative serological studies.

### *Infection Avoidance Between Testing and Transplant*

In endemic areas, exposure to *H. capsulatum* may occur with many daily activities and is difficult to entirely avoid. Nonetheless, in the pre-donation period, certain activities (cave exploration, significant time in construction sites, exposure to bird droppings or bat guano, digging soil) should be avoided.

---

<https://www.cdc.gov/fungal/pdf/more-information-about-fungal-maps-508.pdf>.

## Coccidioidomycosis Guidance

### Background

*Coccidioides immitis* and *Coccidioides posadasii* are dimorphic fungi endemic in arid and semi-arid climates in Washington State and the Southwestern part of the U.S. (Figure 2), Northern Mexico including areas along the U.S. border, and parts of Central and South America. The most common clinical syndrome is pneumonia often accompanied by joint pains, fatigue, and weight loss. Chronic fibrocavitary pulmonary disease may follow primary infection, and some infected persons have persistent symptomatic pulmonary nodules, reticulonodular disease, or cavitary disease. Immunosuppressed individuals may develop disseminated infection. Multiple cases of donor-derived coccidioidomycosis have been reported, often with poor outcomes.

**Figure 2: CDC’s current estimate of *Coccidioides* endemicity in the United States**



64

Darker shading shows areas where *Coccidioides* is more likely to be present. Diagonal shading shows the potential geographic range of *Coccidioides*.

### Who Should be Screened

#### Living and Deceased Donors

One study indicated that 2.1% of persons evaluated for living donation in an endemic region were seropositive, suggesting recent or active disease. Since many patients with transmissible coccidioidomycosis are asymptomatic and infection is widespread in endemic areas, some experts recommend screening as part of the routine evaluation of all potential donors who reside in endemic areas or who have recently resided or had prolonged stays in such areas. Persons with signs and symptoms consistent with coccidioidomycosis (fever, joint pains, cough, neck stiffness, headaches) or those with unexplained chest imaging findings (cavities, nodules, lymphadenopathy, reticulonodular infiltrates) warrant screening as well. Finally, it is reasonable to perform further

<sup>64</sup> CDC. “More Information about the Estimated Areas with Blastomycosis ...” More information about the estimated areas with blastomycosis, coccidioidomycosis (Valley fever), and histoplasmosis in the United States. Accessed June 23, 2023. <https://www.cdc.gov/fungal/pdf/more-information-about-fungal-maps-508.pdf>.

screening on donors with a known history of coccidioidomycosis, as potential donors with persistently positive serologic studies are more likely to harbor viable organisms.

### *How to Screen*

#### *Living and Deceased Donors*

In asymptomatic potential donors, serological tests (enzyme immunoassay, complement fixation, or immunodiffusion) may be combined with chest imaging. Patients with suggestive findings on imaging (non-calcified nodules, cavities, lymphadenopathy, reticulonodular infiltrates) may require sputum cultures or bronchoscopy with culture of lavage fluid, although sensitivity of cultures may be low. For prospective donors with normal imaging and negative serologies, the risk of donor-derived infection is likely low.

#### *Management of Infected Living Donors*

Potential living donors with evidence of active infection **should be offered consultation with ID** and require a thorough evaluation to determine the extent of the infection. Donation from actively infected donors should be deferred until treatment is complete and evidence of active infection (based on clinical, radiographic, and serological factors) has resolved.

#### *Management of Recipients*

Prophylaxis with antifungal medications active against *Coccidioides* species may be effective in preventing disease in recipients of infected donors. Fluconazole or itraconazole can be used as prophylaxis; fluconazole is more reliably absorbed and interactions with calcineurin inhibitors are less significant. Posaconazole and voriconazole are second-line options. Echinocandins or inhaled polyenes are not effective. **The recommended duration and dose of prophylactic agents has not been well defined but would typically consist of 400 mg of fluconazole daily for at least three to six months, although in some circumstances lifelong prophylaxis is recommended. Once transplant has occurred, consultation with ID is recommended.** Whether additional prophylaxis is needed for recipients of donors who received adequate treatment for coccidioidomycosis in the past is unknown. While periodic post-transplant serological monitoring could be considered, the sensitivity of serological testing in immunosuppressed patients is likely low. Recipients should be closely monitored, when prophylaxis is discontinued, with periodic clinical, radiologic, and serologic assessments, especially in the first six to 12 months after cessation of prophylaxis.

#### *Infection Avoidance between Testing and Transplant*

For people living in endemic areas, completely avoiding exposure to *Coccidioides* is difficult to impossible. Highest risk activities include those that result in heavy inhalation of desert soil, particularly in the dry periods following the rainy season. Potential donors that develop pulmonary infection or illness characterized by fatigue, cough, and/or joint pain after initial screening may require repeat evaluation.

## PARASITIC INFECTIONS

### Strongyloidiasis Guidance

#### *Background*

*Strongyloides stercoralis* is a nematode (roundworm). Unlike other parasites, *Strongyloides* can replicate for decades within humans via an autoinfection cycle, resulting in chronic symptomatic or asymptomatic infections that can flourish in the setting of immunosuppression. There are an estimated 370 million infected people worldwide.

*Strongyloides* is most common in tropical, subtropical, or warm temperate climates, including the Caribbean, Mexico, South and Central America, Africa, Southeast Asia, Southern and Eastern Europe, and the Southeastern U.S. In the U.S., a prevalence of up to 6.1% has been reported in select populations, with a much higher prevalence of up to 46.1% in immigrant populations. Infection results primarily from exposed skin contact with soil that is contaminated with *Strongyloides* larvae. Activities that increase the risk of becoming infected include direct contact with contaminated soil (i.e., walking barefoot), unprotected skin contact with human waste or sewage, and occupations that increase contact with contaminated soil, such as farming and coal mining. Rural populations and those of a lower socioeconomic status are at higher risk.

Hyperinfection occurs in immunocompromised hosts and can lead to disseminated infection, with mortality as high as 85%. From 2009 to 2013, the CDC assisted in investigations of donor-derived strongyloidiasis in 11 recipients from seven organ donors and found 86% of the donors to originate from *Strongyloides* endemic areas. Prophylaxis with ivermectin was noted to be highly effective in preventing disease transmission in the CDC series and in another series reported by an OPO.

#### *Who Should Be Screened*

##### **Living and Deceased Donors**

Given the high mortality rate of donor-derived *Strongyloides* infection, and the efficacy, safety, simplicity and low cost of prophylactic therapy, universal screening for *Strongyloides* is recommended for all living donors and is required for deceased donors by OPTN Policy 2.9 pending implementation of *Improve Deceased Donor Evaluation for Endemic Diseases*<sup>65</sup>, and is recommended prior to implementation.

Donors with a personal history of previously treated *Strongyloides* infection should be at low risk for transmission, although repeat (new) infection after treatment can occur, and the autoinfection cycle could allow for subclinical persistence. Signs and symptoms include chronic abdominal pain, bloating, heartburn, intermittent diarrhea and constipation, dry cough, skin rashes, and/or eosinophilia, or persons can be entirely asymptomatic. Although transmission within families is unlikely, a family history should trigger screening given the likelihood of similar exposures.

#### *How to Screen*

##### Living and Deceased Donors

---

<sup>65</sup> "Improve Deceased Donor Evaluation for Endemic Diseases," OPTN, Policy Notice, Accessed October 10, 2023, [https://optn.transplant.hrsa.gov/media/lrxfreqj/dtac\\_endemics\\_policy-notice\\_june23bod.pdf](https://optn.transplant.hrsa.gov/media/lrxfreqj/dtac_endemics_policy-notice_june23bod.pdf).

Screening is by serology or stool examination. Serology is required for deceased donors pending implementation of *Improve Deceased Donor Evaluation for Endemic Diseases*<sup>66</sup> and recommended for living donors. Serology is primarily an Immunoglobulin G (IgG) assay for antibodies to *Strongyloides*. The enzyme-linked immunosorbent assay (ELISA) is preferable because of its greater sensitivity (90%) and specificity (99%), compared with indirect fluorescent antibody (IFA) and indirect hemagglutination (IHA) tests. Antibody testing cannot differentiate between current and prior or treated infections; titers do not reliably decrease rapidly after treatment. The assays may produce false positive results in patients with filariasis and other nematode infections. Serology testing is commercially available, often with turnaround times of one to five days.

Stool testing is less sensitive than serology, as stools are positive only during larval shedding which can be intermittent and would be indicated when serological testing is unavailable or when serological findings are negative but clinical suspicion of disease is high. A single stool specimen has a sensitivity of only 15%–30%, although sensitivity increases to nearly 100% if seven consecutive daily stool specimens are examined in an expert laboratory, which may not be practical in the transplant setting. Intestinal endoscopy with biopsies may also diagnose *Strongyloides* infection, although it would not be used for routine screening.

For living donors, screening should be done with initial testing to allow time for subsequent treatment if necessary.

#### *Management of Infected Living Donors*

Living donors who test positive should be informed about the result and offered consultation with ID. Donors infected with *Strongyloides* may still donate. Living donors can be treated with ivermectin, with no further delay in organ transplant. Additional serologic or stool testing after treatment is not likely to be helpful.

#### *Management of Recipients*

Once transplant has occurred, consultation with ID is recommended. Recipients who receive organs from infected and untreated donors may be treated with ivermectin or albendazole. Ivermectin is well tolerated and is preferred (when available) due to superior efficacy. Only oral ivermectin is available in the U.S. There may be a drug interaction between ivermectin and the calcineurin inhibitors (tacrolimus and cyclosporine), therefore monitoring of drug levels is recommended.

Transplant centers should follow up on such pre-transplant test results and give prophylaxis to recipients if their donors were not previously treated. Prophylaxis and treatment doses of anti-parasitic medications in this setting are probably the same, although they have not been studied. Most cases of donor-derived strongyloidiasis occur several weeks to months after transplant, so centers should have adequate time to treat recipients before infection occurs. In cases in which there is insufficient time for test results to return and treatment to be given to donors, centers should plan for post-transplant treatment of the recipient.

People dually infected with Human T-Cell Lymphotropic Virus-1 (HTLV-1) and *Strongyloides* are more likely to develop severe cases of strongyloidiasis and are at higher risk for recurrence. It may be

---

<sup>66</sup> Ibid.

worthwhile to screen patients with *Strongyloides* infection for HTLV-1, so the treating clinicians are aware of the risk of increased disease severity and recurrence.

Patients from areas of Africa endemic for loiasis (primarily central and western sub-Saharan Africa) should be screened for *L. loa* microfilaremia, as administration of ivermectin to those with microfilaremia can precipitate life-threatening encephalopathy, known as the Mazzotti reaction.

### *Infection Avoidance between Testing and Transplant*

In the absence of exposure, acquisition of acute infection would generally be rare in the several months between testing and transplant for most living donors, unless they travel to endemic regions and have significant soil exposure. Shoes should be worn in endemic areas to avoid contact with infected soil.

## **Chagas Disease Guidance**

### *Background*

Chagas disease is caused by infection with the protozoan parasite *Trypanosoma cruzi* (*T. cruzi*). It is estimated that eight million people are infected in the Americas, with over 300,000 infected individuals currently living in the U.S. Most infections were acquired in endemic regions of Mexico, South and Central America, where infection is transmitted by the bite of an infected triatomine bug. Infection can also be transmitted from an infected mother during the second and third trimester of pregnancy or rarely during childbirth, as well as through infected food or drink, and through blood transfusion and organ transplantation.

Most infections are acquired in childhood during residence in an endemic area. The acute phase of infection may be associated with a mild febrile illness and is often unrecognized. Parasitemia clears within eight to 12 weeks without treatment, but individuals remain chronically infected. Infection persists for the individual's life, with chronic disease developing in 20-30%, usually manifesting as cardiac or gastrointestinal disease. Chronic infection can be diagnosed through serologic tests for antibody to *T. cruzi*.

Blood donor screening in the U.S. beginning in 2007 identified confirmed infections in donors from 37 states and Puerto Rico, with 57% of all positive tests from California and Florida, areas with significant Mexican, Central and South American immigrant populations. Latent infection with *T. cruzi* can persist for decades, therefore organ donor screening of high-risk individuals using serology testing is utilized at some centers. Transmission of Chagas disease has been studied in 32 organ transplant recipients from 14 seropositive donors in the United States. Two of 15 (13%) renal transplant recipients had donor-derived infection; none of these were living donor transplants. Living organ donor transmissions have been reported in Mexico and South America.

### *Who Should Be Screened*

#### Living and Deceased Donors

Deceased donors who were born in a country currently classified as endemic for Chagas disease by the CDC must be screened according to OPTN Policy 2.9 pending implementation of *Improve Deceased*

*Donor Evaluation for Endemic Diseases*<sup>67</sup> and is recommended until then. Screening is recommended for living donors born in a country currently classified as endemic for Chagas CDC by the CDC as well.

Screening should also be considered for living and deceased donors in the following circumstances:

- Children of women born in endemic regions if their birth mother’s serology is positive or unknown
- Donors who have resided in an endemic region for more than three months
- Donors who received a blood transfusion in endemic regions and those who have a previous diagnosis of Chagas disease
- Symptomatic donors who have chronic infection and may present with cardiomyopathy, cardiac conduction abnormalities, megaesophagus, megacolon

Countries currently classified as endemic for Chagas disease by the CDC:

Argentina  
 Belize  
 Bolivia  
 Brazil  
 Chile  
 Colombia  
 Costa Rica  
 Ecuador  
 El Salvador  
 French Guiana  
 Guatemala  
 Guyana  
 Honduras  
 Mexico  
 Nicaragua  
 Panama  
 Paraguay  
 Peru  
 Suriname  
 Uruguay  
 Venezuela

### *How to Screen*

#### Living and Deceased Donors

Serology testing using an FDA licensed, approved, or cleared assay (see Table 2) should be performed on donors with risk factors for Chagas disease. Turnaround time for serology tests ranges from one to 10 days. Tests for parasitemia are not sensitive enough to detect chronic *T. cruzi* infection. Detectable antibody is usually present within a month of infection and remains present throughout the individual’s life.

<sup>67</sup> “Improve Deceased Donor Evaluation for Endemic Diseases,” OPTN, Policy Notice, Accessed October 10, 2023, [https://optn.transplant.hrsa.gov/media/lrxfreqj/dtac\\_endemics\\_policy-notice\\_june23bod.pdf](https://optn.transplant.hrsa.gov/media/lrxfreqj/dtac_endemics_policy-notice_june23bod.pdf).



**Table 2: Serological Tests Available for *T. cruzi* Infection**

Test name, Manufacturer	Target Antigen	Test Format	Sample Type	FDA-cleared/approved use
Abbott Prism Chagas (T. cruzi [E. coli, recombinant] antigen) <sup>68</sup> , Abbott Laboratories, Abbott Park, IL	<i>T. cruzi</i> recombinant antigens (FP10, FP6, FP3, TcF)	ChLIA <sup>69</sup>	Serum/plasma	Donor screening
ORTHO T. cruzi ELISA Test System Ortho-Clinical Diagnostics, Inc. Raritan, NJ	Whole cell lysate	EIA <sup>70</sup>	Serum/plasma	Donor screening, individual diagnostics
Chagatest ELISA recombinant v.3.0 <sup>71</sup> Wiener Laboratories S.A.I.C., Rosario, Argentina	Recombinant epimastigote and trypomastigote Proteins	EIA	Serum/plasma	Diagnosis, NOT donor screening test
Hemagen Chagas' Kit, Hemagen Diagnostics, Inc., Columbia, MD	Purified antigens from cultured <i>T. cruzi</i>	EIA	Serum	Diagnosis, NOT donor screening test
Abbott ESA Chagas Assay, Abbott Laboratories, Abbott Park, IL	<i>T. cruzi</i> recombinant antigens (FP10, FP6, FP3, TcF)	Enzyme Strip Assay	Serum/plasma	Supplemental test in donors who test positive with first-line assays, not approved for individual diagnosis

\*Serologic testing may also be available through the CDC

### *Management of Infected Living Donors*

Living donors who test positive should be informed about the result **and offered consultation with ID.** Confirmatory testing through a submission to the CDC or performance of at least two different FDA

<sup>68</sup> This test may only be available through local blood bank.

<sup>69</sup> ChLIA, chemiluminescence immunoassay

<sup>70</sup> EIA, enzyme immunoassay

<sup>71</sup> Preferred tests for initial donor screening

licensed, approved, or cleared antibody diagnostic tests should be performed. Infected donors may require further evaluation for chronic Chagas disease and may ultimately require specific treatment. Evaluation and treatment of these individuals should proceed in accordance with local guidelines, as the availability of confirmatory diagnostics and anti-parasitic therapy varies.

### *Management of Recipients*

The risk of transmission of *T. cruzi* infection from an infected donor to an uninfected deceased kidney donor recipient has been reported to be between 13 and 18%. There are no data regarding living donor transmissions. It is reasonable to consider accepting donor organs from *T. cruzi* positive donors on an individual basis with informed consent and close monitoring of the recipient. Potential recipients of a *T. cruzi* positive donor organ should themselves be tested for the presence of *T. cruzi* antibody, particularly since they may share unrecognized risk factors with their potential living donor. If the recipient tests negative and the decision is made to proceed with the transplant, recipients must be counseled specifically about the transmission risk and the need for close monitoring post-transplant, with the potential need for anti-parasitic therapy that is FDA approved but may lead to side effects that may be difficult to tolerate.

Once transplant has occurred, ~~expert ID consultation~~ consultation with ID is recommended to coordinate post-transplant testing. In the U.S., the CDC can assist with PCR and parasitemia testing. Incorporating a PCR testing platform into the post-transplant testing program is recommended, as molecular testing may be positive weeks before parasitemia is detected using standard screening.

PCR and parasitemia tests of peripheral blood should be performed using the following schedule:

- Weekly for the first three months post-transplant
- Every two weeks for the fourth post-transplant month
- Monthly thereafter until a minimum of six months post-transplant AND until net state of immunosuppression is at optimal post-transplant baseline with no evidence of infection or rejection
- More frequent monitoring is recommended any time the patient requires antirejection therapy

For access to testing of transplant recipients at risk for reactivation of Chagas, contact the CDC Division of Parasitic Diseases and Malaria at 404-718-4745 (business hours) or 770-488-7100 (nights and weekends) or via e-mail at [parasites@CDC.gov](mailto:parasites@CDC.gov).

Current therapy for Chagas disease is limited to benznidazole and nifurtimox (Lampit<sup>®</sup>, Bayer), both of which are partially metabolized via cytochrome P450 reductase and may increase tacrolimus and cyclosporine blood levels. Both drugs have potential adverse effects, but benznidazole is generally better tolerated and is given twice a day while nifurtimox is given three times a day.

### **Infection Avoidance between Testing and Transplant**

Potential living donors should avoid prolonged travel to endemic regions, particularly rural areas with primitive housing or significant insect exposure. Consumption of uncooked food or drink from endemic regions should also be avoided.

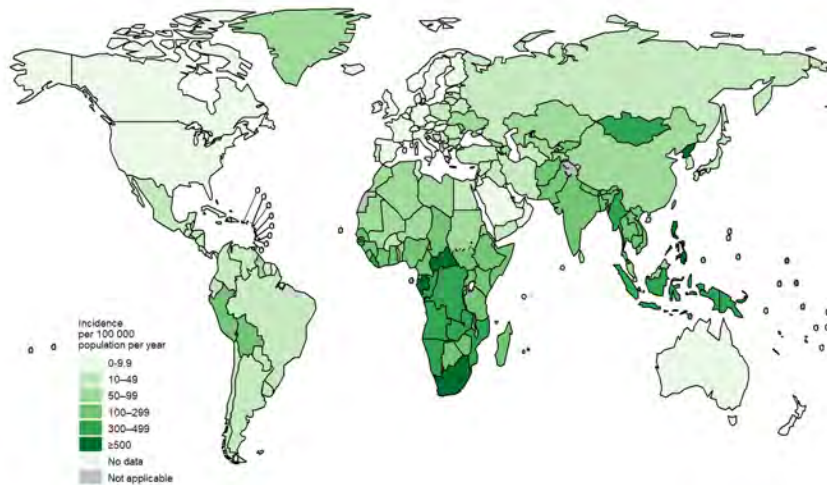
## BACTERIAL INFECTIONS

### Mycobacterium Tuberculosis (MTB) Guidance

#### Background

Up to one-third of the world's population is infected with MTB; however, infection in the United States is much less common. Reported cases of MTB disease have been declining in the United States since 1992 with the majority of cases occurring in foreign-born persons. After initial infection with MTB, most people do not develop MTB disease; the infection disseminates throughout the body and remains dormant. This condition is called tuberculosis infection (TBI) (previously referred to as "latent tuberculosis infection"). In 2022, there were 8,300 reported TB cases<sup>72</sup> in the United States (a rate of 2.5 cases per 100,000 persons). Since initial infection does result in live MTB in many organs, tuberculosis can be transmitted via non-lung organs used for transplantation from donors who have never had clinical signs or symptoms of MTB disease.

**Figure 3. World Health Organization Estimated Global TB Incidence Rates, 2021<sup>73</sup>**



#### Who Should Be Screened

##### Living and Deceased Donors

Since TBI precedes the development of MTB disease, similar risk factors would be expected to be present. One difference is that since the rate of reactivation decreases with time from infection, donors with distant infection would still be at risk for transmission but at a lower risk for MTB disease. The most

<sup>72</sup> Provisional **CDC data**

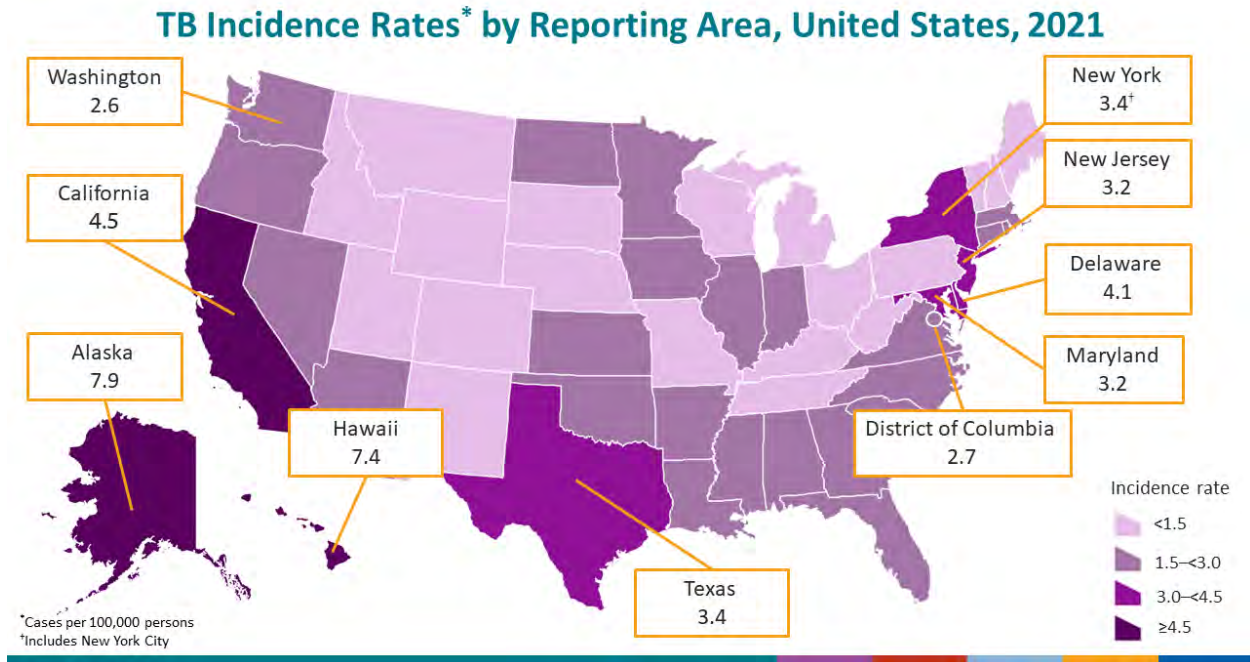
<sup>73</sup> WHO. "2.1 TB Incidence." World Health Organization, October 27, 2022. <https://www.who.int/teams/global-tuberculosis-programme/tb-reports/global-tuberculosis-report-2022/tb-disease-burden/2-1-tb-incidence>.  
<https://www.who.int/teams/global-tuberculosis-programme/tb-reports/global-tuberculosis-report-2022/tb-disease-burden/2-1-tb-incidence>.

powerful easily identifiable risk factor is place of birth with estimated risk of TBI of 18.7% among foreign-born persons in the U.S. as compared to 1.8% among U.S. born persons. Higher risk countries are identified by incidence per 100,000 population per year in Figure 3.

For individuals from lower risk regions, including the United States, ~~would be expected to increase the risk of TB:~~ the following factors should help OPOs and transplant programs make screening decisions:

- Close contacts of persons with infectious TB disease
- Those who spend significant time (greater than 3 months) in areas of the world with high rates of TB or those born in these regions (all but the lightest shaded area of Figure 4) including relief work in a country with high TB risk.
- ~~History of injection drug use~~ History of non-medical injection drug use
- Persons who reside (or ever resided) or worked in institutional settings which resulted in increased risk of exposure to TB (hospitals, nursing homes, correctional facilities, other health care settings, homeless shelters)
- ~~The percentage of TB cases that occur in Hispanic or Latino, Black or African American, and Asian persons is higher than expected based on the percentage of these populations in the U.S. population.~~
- Radiographic evidence of prior tuberculosis on chest radiograph
- Symptoms such as fever, night sweats, weight loss, cough, recurrent pneumonia, exudative pleural effusion of unknown etiology, lymphadenopathy, non-calcified pulmonary nodules or cavities

Figure 4: TB Incidence Rates by Reporting Area, United States, 2021<sup>74</sup>



### How to Screen

#### Living and Deceased Donors

While MTB disease can involve almost any organ and cause many syndromes (e.g., meningitis), a careful medical history and examination will likely identify most patients with MTB disease. TBI, by definition, is asymptomatic, and specific testing is required to identify patients with TBI. The current FDA-approved screening methods for TBI in the US include the tuberculin skin test (TST) and the interferon gamma release assays (IGRAs): QuantiFERON-TB gold in tube (QFT), and T-SPOT.TB. These tests do not differentiate TBI from MTB disease and may be negative during times of MTB disease. The TST test requires injection into the skin and a return visit in 48-72 hours to interpret the test. The IGRAs are blood tests and may show positive, negative, or indeterminate results.

Indeterminate results indicate that the controls did not work in that patient, and this result is more common in immunosuppressed patients. One advantage of the IGRAs is that patients who received a childhood BCG vaccine (very common outside of North America and Western Europe) are less likely to have a false-positive test with IGRAs than with the TST test, due to enhanced specificity of the *Mycobacterium tuberculosis* antigens used in the IGRA assays.

<sup>74</sup> CDC. "Health Disparities." Centers for Disease Control and Prevention, November 10, 2022. <https://www.cdc.gov/tb/topic/populations/healthdisparities/default.htm>.

## Management of Infected Living Donors and Recipients

Table 3 outlines management of living donors with a history of active TBI and treatment of recipients of these organs. **In all clinical scenarios, consultation with ID is recommended.**

**TABLE 3: Management of latent or history of active TB in living donors and recipients of these organs**

Clinical Scenario: Living Donor	Risk for Transmission	Recommendation
History of TBI-treated appropriately	Lower	Monitor recipient clinically
History of TBI-treated insufficiently or not treated or treatment details not clear OR new diagnosis of TBI-positive TST or Interferon gamma release assay found during pre-transplant evaluation; evaluation finds no evidence of active TB	Moderate	Consider deferring transplant if possible until donor has taken some/all of chemoprophylaxis and consider chemoprophylaxis of recipient; monitor clinically.
Unexplained pulmonary apical fibrosis in donor without cavitation and without additional testing	Variable	Defer donation pending further evaluation.
History of MTB disease treated appropriately over two years ago	Lower to moderate	Monitor recipient clinically; consider cultures of previous TB sites if possible. Consider TB prophylaxis of recipient.
History of MTB disease-site remote from transplant ( <b>organ not being transplanted</b> ) treated appropriately within two years.	Lower to moderate	Monitor recipient clinically; consider cultures of previous TB sites if possible. Suggest chemoprophylaxis of recipient.
History of MTB disease-site remote from transplant ( <b>organ not being transplanted</b> ) treated insufficiently and/or with other than standard regimen Excluding disseminated or CNS TB.	Higher Increased risk if less than two years since active TB diagnosis.	Defer live donors until adequately treated; <b>consider consult with infectious diseases specialist</b> ; recommend cultures of previous TB sites prior to transplant if possible
History of renal MTB disease treated appropriately. (If not treated appropriately donation should be deferred until after appropriate treatment)	Moderate	Verify treatment; monitor clinically; recommend chemoprophylaxis for recipient; recommend cultures of previous TB site(s). <b>consider consult with infectious diseases specialist.</b>

### *Management of Recipients of Deceased Donors*

For recipients of an organ from a donor who is TST or IGRA-positive, had recent exposure to active TB, or had radiographic evidence of untreated TB should be considered for TBI therapy, following CDC guidance for anti-tuberculosis agents and durations. Once transplant has occurred, consultation with ID is recommended. Specific attention should be given to potential drug-drug interactions of TBI agents and immunosuppressive medications with careful monitoring of calcineurin inhibitors.

### *Infection Avoidance between Testing and Transplant*

Potential living donors should avoid prolonged travel to endemic regions and behavioral exposures that increase risk if possible, including institutional settings which may result in increased risk of exposure to TB (hospitals, nursing homes, correctional facilities, other health care settings, and homeless shelters).

## **VIRAL INFECTIONS**

### **West Nile Virus (WNV) Guidance**

#### *Background*

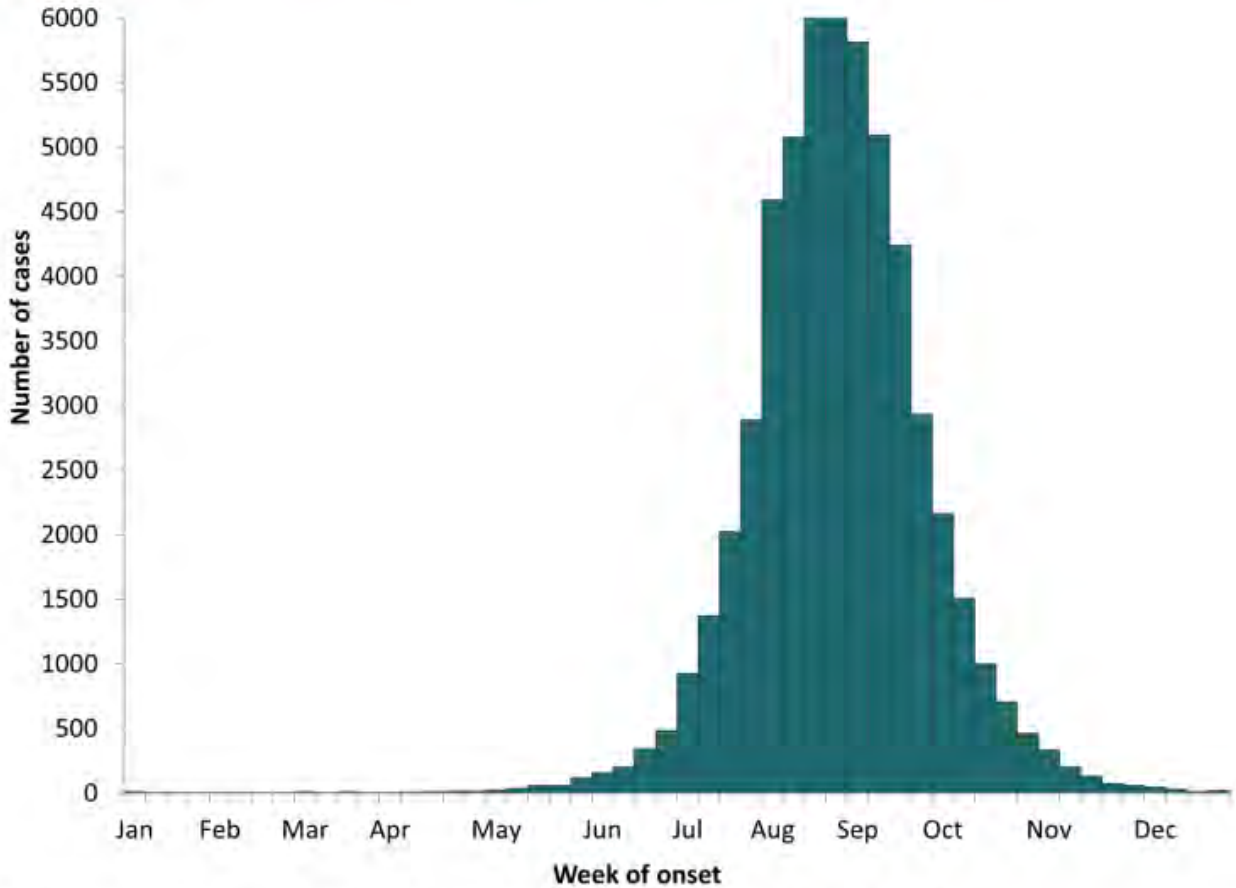
#### *Epidemiology and pathophysiology*

WNV is an RNA virus that spreads to humans primarily by the bite of infected mosquitoes, although multiple non-vector modes of transmission have been described, including blood transfusion, organ transplantation, breastfeeding, intrauterine, and laboratory exposure. Birds serve as a reservoir for WNV, and humans are a dead-end host for the virus. The virus was first detected in North America in 1999, and since then, has spread to all 48 contiguous U.S. states. The number of human infections varies significantly based on the region and year, with most transmission (as shown in Figure 5) occurring during the warmer months (July to October) when mosquitoes are most active, although warmer regions can have a longer season. While the majority of WNV infections are asymptomatic, about 20% of those infected may develop fever, and less than one percent develop neurological complications such as encephalitis or acute flaccid myelitis (involving the spinal cord). This means that even donors who do not display symptoms could still transmit the virus to recipients. While several treatments have been explored, there is currently no established treatment for WNV.



Figure 5: Months of typical WNV activity in the United States<sup>75</sup>

**West Nile virus disease cases reported to CDC by week of illness onset, 1999-2021**



Source: ArboNET, Arboviral Diseases Branch, Centers for Disease Control and Prevention

*Who Should Be Screened*

Living and Deceased Donors

Since the transmission of WNV is limited by geography and season, OPOs and transplant programs should consider certain factors when determining whether to perform laboratory testing on a potential donor. A positive test result obtained during a period of WNV inactivity is more likely to represent a false positive than when the test was performed during periods of higher activity. Any false positive result could result in unnecessary delays to transplant.

<sup>75</sup> CDC. "West Nile Virus Disease Cases Reported to CDC by Week of Illness Onset ..." West Nile virus cases reported to CDC by week of illness onset, 1999-2021. Accessed June 23, 2023. <https://www.cdc.gov/westnile/resources/pdfs/data/WNV-Week-Onset-1999-2021-P.pdf>.

Factors to consider in evaluating donors for possible WNV infection:

- Has human infection with WNV virus been recognized locally this WNV season?
- Has the donor travelled to an area with human WNV activity this WNV season?
- Has the donor ever been diagnosed with WNV fever or WNV neuroinvasive disease?
- Has the donor had an undifferentiated febrile illness within the current WNV season?
- Has the donor had significant mosquito exposure this WNV season?

*Geographic and seasonal factors to consider.*

Screening strategies for WNV in donors include universal year-round testing of all donors versus targeted testing during periods of human WNV activity. As the yield of testing during the winter months is likely low and given the potential for IgM to remain positive after resolution of the illness, year-round testing is discouraged unless programs are unable to manage the complexity of more targeted testing strategies. Targeted testing strategies could include testing during a pre-determined timeframe, or testing donors only during times of human WNV activity in the area where the donor lives or has traveled (i.e., triggered strategy).

Triggering strategies can involve communicating with local blood collection organizations and determining whether they have shifted from minipool to individual donation (ID)-NAT screening, indicating that a WNV-reactive donor has been identified within a 50-mile radius of the triggering zip code. Another method involves using WNV human and non-human surveillance data collected by state and local health departments. Alternatively, testing could begin each year during the local WNV transmission season as determined by historical data (typically July through October, or longer intervals in warmer regions).

Given that human WNV disease incidence varies greatly from year to year and even county to county, targeted testing based on real-time measurement of local WNV activity at local blood banks might be the most cost-effective strategy and will reduce the number of false positives. This strategy is, however, more complex and time-consuming than a fixed seasonal strategy. Some institutions may already be using a local or regional blood bank testing lab to test other material (e.g., peripheral stem cells used in stem cell transplantation). In contrast, the strategy of testing during a defined time period regardless of local WNV activity would be simpler to implement, but both costs and false positive rates would be expected to be higher. **Table 4** describes the advantages and disadvantages of different testing strategies.

**Table 4: Advantages and Disadvantages of Different Testing Strategies**

	<b>Year-round testing (1)</b>	<b>Seasonal testing (2)</b>	<b>Triggered testing (3)</b>
<b>Ease of implementation</b>	Easy	Intermediate	Difficult
<b>Positive predictive value</b>	Lower during periods of human WNV inactivity	Intermediate depending on level of human WNV activity	High
<b>Cost effectiveness</b>	Least	Intermediate	Most

4. This strategy is not recommended.
5. Typically July through October, but time period should be guided by local historical WNV data, in consultation with state and local health departments
6. Testing triggered by switch from minipool to individual blood donation-NAT testing in zip codes of residence of donors. Testing stops when WNV activity no longer noted and blood banks switch back to minipools. Routine communication with local blood banks is required.

### How to Screen

#### Living and Deceased Donors

To screen for WNV, healthcare professionals use NAT and IgM serologic tests. Currently, there are two FDA-licensed NAT donor screening assays; these are routinely performed at blood banks or some reference labs but are not commonly available in hospital laboratories. Serologic tests which rely on the response of the immune system to infection with WNV, particularly the IgM assay performed on serum and cerebrospinal fluid, are commonly used to diagnose neuroinvasive WNV. IgM is detectable for a median of about five months after infection but may persist longer, even years after infection, and is not used in blood donors to screen for WNV, although it may be performed for use in donor counseling. IgG typically remains positive for life and would not be useful to routinely screen for WNV. Table 5 describes the characteristics of NAT and IgM testing.

**Table 5: Tests that could be used to screen for WNV infection**

	<b>Nucleic Acid Tests (NAT)(1)</b>	<b>IgM (serology)(2)</b>
<b>Available Tests</b>	Procleix West Nile Virus Assay COBAS TaqScreen West Nile Virus Test	Various
<b>FDA licensed for organ screening</b>	Yes	No
<b>Availability</b>	Blood bank testing labs Reference labs	Reference labs Commercial labs State public health labs
<b>False positive rate</b>	Low	Likely higher than NAT, but not evaluated for donor screening
<b>Indicates active infection</b>	Yes	Remains positive for median of five months (up to seven years documented); active infection may have cleared
<b>Required for blood donor screening</b>	Yes	No

3. Should be used as part of any testing strategy
4. Consider in combination with NAT testing but will increase false positive rate

### *Deceased donors*

It is advisable to screen deceased donors during months of regional WNV activity. WNV NAT testing is a viable option, but the results may not be available before transplantation. It is important to note that WNV transmission has occurred in donors who tested negative on NAT, and that there is potential for false positive NAT test results if protocols are not in place to improve specificity, as are used by blood collection organizations, or if testing is performed outside of the WNV transmission season, decreasing the pretest probability.

In cases involving deceased donors, the feasibility of conducting WNV testing within a required timeframe is uncertain. As such, the Committee advises against proceeding with donors who satisfy any of the following criteria: (1) a confirmed WNV infection, (2) a positive WNV NAT test result, ideally meeting a threshold signal-to-cutoff ratio or verified by repeat testing as recommended by the manufacturer, or (3) clinical manifestations of meningitis, encephalitis, or flaccid paralysis of indeterminate origin, particularly if they originate from regions with documented WNV activity.

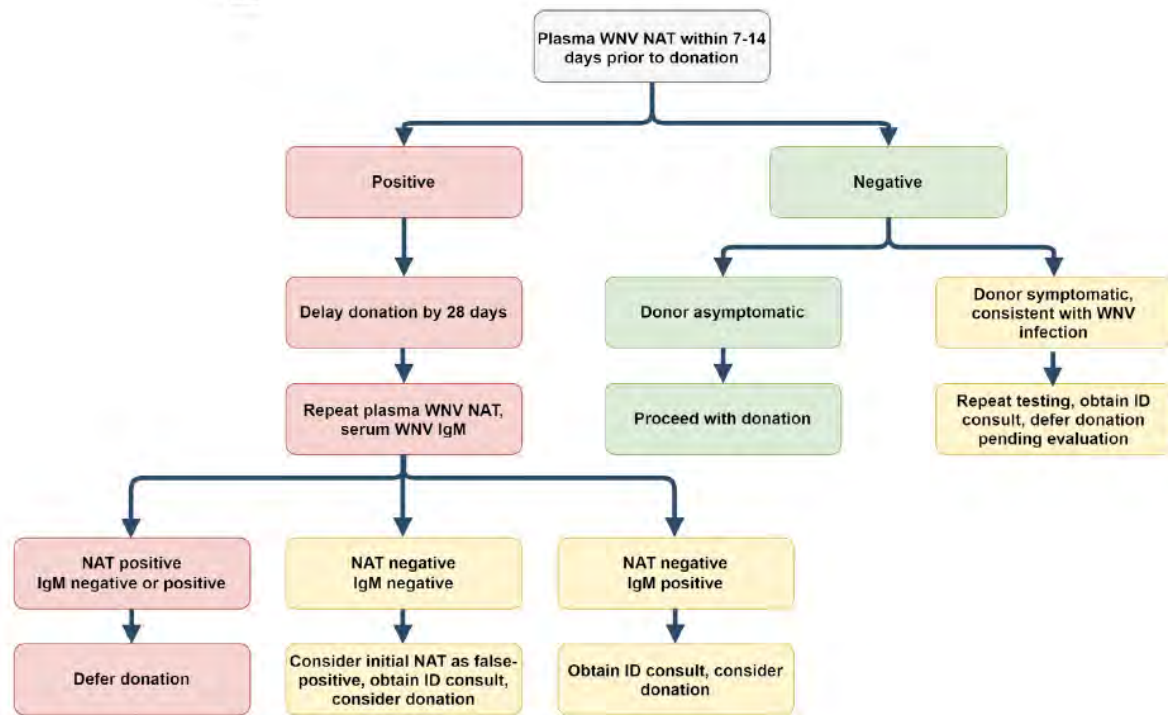
### *Living donors*

Living donors should be screened with WNV NAT within seven to 14 days of donation.

### *Management of Infected Living Donors*

#### **Living donors who test positive should be informed about the result and offered consultation with ID.**

Although no studies have examined organ donation after WNV infection, we suggest postponing donation for a minimum of 28 days in living donors, after which NAT and IgM testing should be performed. If negative, it is likely that the initial NAT test was a false positive, and organ donation can be considered. If the NAT is negative but IgM is positive, this likely reflects viral clearance and organ donation can be considered. However, if the NAT remains positive, organ donation should be postponed. (Refer to Figure 6 for additional information).

Figure 6: Living donor screening recommendations for WNV<sup>76</sup>

### Management of Recipients

Development of symptoms compatible with WNV disease within the first several weeks of transplant (and up to six weeks, since transplant recipients can have prolonged incubation periods with WNV infection) should prompt testing in the recipient. There are currently no specific medications available for the treatment of WNV infection. **Once transplant has occurred, consultation with ID is recommended.**

### Infection Avoidance Between Screening and Transplant

We advise potential living donors to take measures to prevent mosquito exposure during periods of mosquito activity, such as wearing loose-fitting, long sleeve shirts and pants, using EPA-registered insect repellent, and avoiding outdoor activities during dawn and dusk when mosquitoes are most active. Further, a potential living donor should report febrile illnesses to his or her transplant center. WNV diagnostic testing should be performed if clinical evaluation suggests the possibility of WNV infection in the potential living donor.

## SHARING INFORMATION

<sup>76</sup> Anesi, JA, Silveira, FP; the AST Infectious Diseases Community of Practice. Arenaviruses and West Nile Virus in solid organ transplant recipients: Guidelines from the American Society of Transplantation Infectious Diseases Community of Practice. *Clin Transplant*. 2019; 33:e13576. <https://doi.org/10.1111/ctr.13576>

## Recommendations for Sites Participating in Paired Organ Exchanges

Transplant hospitals participating in organ exchanges should consider sharing their protocols for screening donors for seasonal and geographically endemic infections in addition to related test results. This will provide recipient centers more information about the donor screening process at the donor organ center.

## When to Report a Potential Donor-Derived Transmission Event

If the recipient is suspected to be at risk for disease transmission either by the OPO or a transplant center, a potential donor-derived disease transmission event should be reported to the OPTN Improving Patient Safety Portal per OPTN Policies 15.4 *Host OPO Requirements for Reporting Post-Procurement Test Results and Discovery of Potential Disease Transmissions* and 15.5 *Transplant Program Requirements for Communicating Post Transplant Discovery of Disease or Malignancy*. This promotes prompt intervention for other recipients of the same donor’s organs.

### Appendix\*

**Table 6: OPTN Ad Hoc Disease Transmission Advisory Committee Data on PDDTE**

Endemic Disease	Number of Proven or Probable Transmissions since 2008
Histoplasmosis	19
Coccidioidomycosis	10
Chagas disease	7
Strongyloidiasis	39
Tuberculosis	21
West Nile Virus	0

**\*Updated biennially by the DTAC**

## References

No changes following public comment.