Infections in Transplanted Patients

Margarita Gospodinova¹, Iliyan Todorov¹ and Ilko Bakardzhiev*²

¹Department of Infectious Diseases, Medical University of Varna, Bulgaria
²Medical College, Medical University of Varna, Bulgaria

Received: December 21, 2017; Accepted: December 29, 2017; Published: January 05, 2018

*Corresponding author: Ilko Bakardzhiev, Associate Professor, Medical College, Medical University of Varna, Tsar Osvoboditel 84, Bulgaria, Tel: +359 888 768413; E-mail: varna2008@gmail.com

Abstract

Solid-organ transplantation is a therapeutic option for many end-stage diseases. One of the most common complications in the post-transplant period are different kind of infections. They are caused by some bacterial pathogens such as Legionella spp., Nocardia spp., Salmonella spp. and Listeria monocytogenes, many viral agents – Cytomegalovirus /CMV/, Herpes simplex virus /HSV/, Varicella-zoster virus /VZV/, Epstein-Barr virus /EBV/, and more rarely – fungi and parasites. The onset of the signs and symptoms is essential for establishing the diagnosis – wound infections occur during the first 30 days after transplantation, opportunistic infections – 2 to 6 month later, and beyond 6 months, recipients suffer from the same infections distributed in the general community. Infections in solid-organ transplant recipients are global socio-medical problem, leading to prolonged hospital stay, significant financial losses and disability /and high mortality rate/ of the patients.

Keywords

Infections; Transplantation; Bacteria; Virus; Fungi; Recipient, Donor; CMV;

Solid-organ transplantation is a therapeutic option for many end-stage diseases. According to World Health Organization /WHO/, 115,000 organ transplantations are performed worldwide annually –70% of kidneys, 20% of liver and 10% of heart, lungs and pancreas [15,21]. The term ‘donor’ is used about the patients who donate their organs to the ‘recipient’ – an individual who receive the donor’s organ.

The early period after transplantation is essential for the success of the operation. At this time, severe immunosuppression is induced and recipients are threatened by infections with different microorganisms, which are distributed in the environment [16]. This interaction /microorganisms + macroorganisms + environment/ results in development of different infectious diseases, which take their clinical course in organism with impaired host defense. In such circumstances common infections may lead to severe and generalized clinical forms that are behind the rejection of the transplanted organ and high mortality rate in the recipient.

The risk factors for infectious diseases development in transplanted patients are shown in a table 1. Impaired mechanical barriers or stasis of secretions/excretions in the region of transplanted organ mainly are one of the most common risk factors leading to direct microbial invasion or bacterial overgrowth and subsequent inflammation. So a direct relationship between transplanted organ and the region of presented inflammatory process exists – infections of kidney and urinary tract in kidneys transplanted patients, infections of respiratory system after lungs transplantations etc [6].


<table>
<thead>
<tr>
<th>Host Defense</th>
<th>Examples of Impairment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anatomical barriers to infection</td>
<td>Surgical wound, implanted devices, anastomosis problems, ischemia</td>
</tr>
<tr>
<td>Abnormal flow of body fluids (urine, bile, respiratory secretion)</td>
<td>Post-transplant ureteral, biliary or bronchial obstruction, urinary reflux, neurogenic bladder, aspiration</td>
</tr>
<tr>
<td>Innate immunity (neutrophils, macrophages, monocytes)</td>
<td>Corticosteroids; medication-induced myelosuppression</td>
</tr>
<tr>
<td>Adaptive immunity (T cells and B cells)</td>
<td>Corticosteroids, calcineurin inhibitors, mycophenolate, rituximab, belatacept</td>
</tr>
<tr>
<td>Complement function</td>
<td>Eculizumab</td>
</tr>
<tr>
<td>Splenic function</td>
<td>Transplant-associated splenectomy</td>
</tr>
</tbody>
</table>

Patients underlying conditions also have a potential to damage organ systems. Neurogenic bladder in patient with diabetes mellitus or vesicoureteral reflux in children with congenital anomalies of kidney and the urinary tract are one of the most prevalent, while the bile duct strictures in the course of sclerosing cholangitis or bronchial obstruction during cystic fibrosis are rarely seen [7].

Immunosuppressive treatment, used in the post-transplant period, is associated with immune tolerance establishing which prevents organ rejection, but on the other side, it results in increased risk for infections. Commonly used regimen is a combination among steroids, calcineurin inhibitors – tacrolimus, cyclosporine and mycophenolate/azathioprine causing severe
Infections in Transplanted Patients

leukocytopenia, particularly lymphocytopenia. Significant neutropenia is associated with the use of also mycophenolate, together with trimetoprim-sulfametoxazol /TMP-SMX/ and valgancyclovir. Monoclonal antibodies affect the cellular and humoral immune response. For example, ‘rituximab’ destroys B-cells and decreases the immunoglobulin production, and ‘eculizumab’, which is anti-thymocyte globulin leads to quantitative changes in the complement system [20].

Two groups of infections in transplanted patients – endogenous and exogenous are known, on the base of the route by which microorganisms enter into the human’s body. Endogenous one appears as a result of activated viruses /which stay latent in the human’s body after primary infections/ or bacterial overgrowth in non-sterile anatomic regions /gastrointestinal tract, skin, etc./. Therefore, it is not surprising that commonly isolated agent, responsible for inflammation after transplantation are normal flora for the relevant anatomic region [7,8]. For example, lungs infections in transplanted patients are caused by *Pseudomonas aeruginosa, Aspergillus spp., Pneumocystis jiroveci*; infections after kidneys, liver and pancreas transplantations are caused by *Enterococcus, Escherichia coli, Bacteroides spp.*, *Clostridium spp.*. It is important to know that skin microbiome and many viruses – *cytomegalovirus /CMV/, herpes-simplex virus /HSV/, varicella-zoster virus /VZV/, Epstein-Barr virus /EBV/, human herpesvirus 6 /HHV-6/, hepatitis B virus /HBV/, hepatitis C virus /HCV/, BK virus, John Cunningham virus /JC virus/*, do not cause infections in specific part of the body but may appear in all transplanted patients [1,5,11,12,18].

Exogenous infections result from environment or they are donor-derived. Respiratory viruses such as *influenza, parainfluenza, respiratory syncytial virus /RSV/, adenoviruses, rhinoviruses, human metapneumoviruses /hMPV/* as well as fungi and Nocardia may infect all transplanted patients, but most common lung transplanted. Fecal – oral mode of transmission, which is typical for *Salmonella spp., Listeria spp, Cryptosporidium, Microsporidium* lead to gastrointestinal tract infections, bacteremia or infections of the central nervous system, observed in all transplanted patients without underlying dependence on transplanted organ [3,5,11].

Donor-derived infections /DDI/ are special group of diseases, which also may be seen in all transplanted patients. They are caused by agents, which had been infected donor’s body before transplantation and which transmission is expected to be in the recipient /it is known that donor is infected/. The etiological spectrum is wide, but *CMV, EBV, HBV, Toxoplasma gondii* are the most important one. The DDI may be unexpected, caused by human immunodeficiency virus /HIV/, hepatitis C virus /HCV/, lymphocytic chooriomeningitis virus, *Mycobacterium tuberculosis, rabies, West Nile virus /WNV/* [9,10].

Most cases of infections in transplanted patients are caused by different bacteria strains. After liver transplantations, bacteria are responsible for 33% to 68% of all infections observed in recipients, in heart transplanted – 21-30% of cases, 35% in pancreas transplantations, 47% and 54% of infections in kidney and lungs transplantations respectively. The risk for bacterial infections depends on: the technical implementation of the transplantation, the duration of the hospital stay, immunosuppressive drugs regimen. It is known that bacterial-caused infections are presented in specific period after transplantation, which correlates with the appropriate diagnosis /see table 2/ [1,21].

**Table 2:** Bacterial-caused infections in transplanted patients

<table>
<thead>
<tr>
<th>Type of Transplantation</th>
<th>Causative Agents</th>
<th>Symptoms Onset</th>
<th>Clinical Presentation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Liver</td>
<td>Enterococcus spp, VRE*, anaerobes, Enterobacteriaceae</td>
<td>2 months after transplantation</td>
<td>Hepatitis, liver abscesses, peritonitis</td>
</tr>
<tr>
<td>Lungs</td>
<td>Klebsiella pneumoniae, Pseudomonas aeruginosa, Escherichia coli, Staphylococcus aureus, Enterobacter cloacae</td>
<td>2 weeks after transplantation</td>
<td>Pneumonia, bronchitis, lung abscesses</td>
</tr>
<tr>
<td>Kidneys</td>
<td>Enterococcus, Staphylococcus, Pseudomonas aeruginosa, Gram negative enteric bacteria</td>
<td>After discharging from the hospital</td>
<td>Infections of the operative wound /the risk correlates with high serum creatinine concentration/, urinary tract infections /the risk correlates with prolonged hemodialysis before operation/</td>
</tr>
<tr>
<td>Pancreas</td>
<td>Enterococcus, Staphylococcus, Pseudomonas aeruginosa, Gram negative enteric bacteria</td>
<td>After discharging from the hospital</td>
<td>Infections of the operative wound, intraperitoneal abscesses, pelvic abscesses, peritonitis, perirectal abscesses</td>
</tr>
</tbody>
</table>

Infections in Transplanted Patients

Bacterial agents mentioned below, are one of the most frequently isolated strains as causative agents of infections in transplanted patients:

- **Legionella spp.** - two kind of legionella infection exist: nosocomial and community acquired. Respiratory mode of transmission is realized by inhaling aerosole particles produced by air-conditioners, aerosol filling machine, showers, physiotherapy departments where water procedures are implemented, etc. The kind of infection is not associated with the severity of disease. The first clinical signs and symptoms are presented weeks after transplantation, together with graft rejection manifestation. Consequently, a *Legionella infection in transplanted patients is a specific marker for forthcoming/presented graft rejection*. One of the most important serotypes is *Legionella pneumophila*, *L. micdadei*, *L. bozemanii* and *L. dumoffii*. Clinical course of the disease includes severe pneumonia occurrence, and more rarely some extra-pulmonary manifestations such as hepatitis and peritonitis or pericarditis, which are typically seen in hearts and heart transplanted patients, respectively [7,8].

- **Salmonella spp.** - because of their clinical polymorphism, salmonella infections after transplants are associated with many diagnostic and differential diagnostic difficulties. The risk for infection is strongly associated with antirejection therapy. The most common symptom is fever and blood cultures are positive in 99% [2]. Except of bacteremia, different organ involvements are possible: pyelonephritis, orchitis, perinephric abscesses, lung infections /pneumonia, lung abscess, pleural effusion/, vascular infections /infections of saccular aneurysms, arteriovenous fistula/, sinusitis, meningitis, cholecytitis etc.

- **Listeria monocytogenes** - is a specific causative agent of infections in transplanted patients. The risk period for infection is about 2 months after transplantation. Seasonal variations are typical – high frequency after transplantations during summer and autumn. In 2/3 of the infected persons, central nervous system alone is involved – meningitis, meningoencephalitis, encephalitis. In the rest 1/3 of the patients, just bacteremia with/ without secondary infections of lungs, eyes, heart, and rectum is seen [1,7,8].

- Other bacterial infections – *Mycoplasma hominis* has been reported in liver transplant recipients, *Ureaplasma urealyticum* causes mediastinitis following heart-lung transplantations, and *Rhodococcus equi* - pulmonary infections after heart transplantations [8,13].

With regards to viral agents which cause infections in transplanted patients, microbial spectrum is wider together with significant variations in clinical presentation /in some cases just as a fever of unknown origin/ leading to many difficulties in establishing the right diagnosis such as late diagnosed and misdiagnosed patients. It is behind the high mortality rate – a basic feature of this group of infections.

- **CMV** – is highly associated with severe immunosuppression and is typically seen in the first 3 months after transplantation. CMV infection occurs in 44 to 85% of kidney, heart, and liver transplant recipients [15]. The infection is donor derived / after transplantation of previously infected donors’ organs or cellular blood products/ or endogenous /after reactivation of latent CMV in the post-transplant period/. CMV infections include wide spectrum of clinical signs and symptoms [21]. Most cases are mild, presented just by fever, malaise, leukopenia and thrombocytopenia. Focal infections may occur, commonly involving the parenchyma of transplanted organ: hepatitis in liver transplanted, pancreatitis in pancreas transplanted, pneumonia after heart-lung transplantations, etc [7,8].

- **HSV** – infection is presented as two clinical forms: primary infection in seronegative recipients or reactivation in previously infected. The symptoms occur in the first month after transplantation and may be presented as herpes labialis, pneumonia /with 75% risk for death no matter of acyclovir treatment/, tracheobronchitis, esophagitis, hepatitis, skin involvement [22].

- **VZV** – primary infection with VZV in seronegative recipient is associated with severe forms of chickenpox, complicated with hemorrhagic pneumonia, encephalitis, hepatitis, pancreatitis, purpura fulminans, while reactivation leads to shingles occurrence in the next 6 months, with dissemination tendency, affecting two or more dermatomes [18].

- **EBV** - three forms of EBV infections are typical for transplant recipients – fever of unknown origin, mononucleosis or post-transplantation lymphoproliferative disease /PTLP/. Risk factors for EBV infection are seronegative patients or those treated after transplantation by OKT3 monoclonal antibodies or tacrolimus. Highest frequency of EBV infections is seen in small bowels transplantations – 14%, while EBV is responsible just for 1% of infections in kidney transplant recipients [5,16,17].

- **HBV** – causes fulminant hepatitis in 1% of acutely infected, resulting in life-threatening liver failure. Also, an approved association between chronic infection and liver cirrhosis exists. In both cases, liver transplantation is a choice of treatment, but there is 80-90% possibility for HBV reactivation /the percent is lower in patients with concomitant hepatitis D virus infection/ with 50-60% mortality rate [6,7]. In contrast, mild form of hepatitis appears when recipient is infected after blood transfusion or transplantation of liver from infected donors. The risk period for clinically presented reactivated HBV is 2-6 month after transplantation, ranging from mild to severe, life-threatening hepatitis [12].

- **HCV** - after primary infection with HCV, symptomatic hepatitis occurs just in 25-30%. In 70-80% of acutely infected, the disease takes chronic course with liver cirrhosis in the next 20 years, which is seen in 20-30% of patients, following by hepatic cell carcinoma occurrence in 1-5% of them. Serotype 1b is associated with high morbidity rate and infection in transplanted patients results in high incidence of HCV recurrence [12].

- **HIV** – transplanted HIV positive recipients need of lower doses of immunosuppressive drugs and they develop AIDS approximately 17 months later. However, the mean period from

---

transplantation to AIDS development after primary infection is 32 months. Soon after transplantation, HIV-infected patients present to the hospital with many complaints such as prolonged fever, liver, spleen and lymph nodes enlargement, malaise, loss of appetite and other non-specific symptoms. The risk for opportunistic infections depends on the CD4 count which is similar to “general” population [7,17].

- **Papovaviruses** - Include two main kinds of viruses, so called JC and BK virus. They are associated with progressive multifocal leukoencephalopathy /PMLE/ or rejection of transplanted kidney, respectively and both are reported in renal transplant recipients [19].

Various fungi have specific place as causative agents of many infections in transplanted patients. The risk for fungal infection is estimated to be 5 to 17% in heart, 14 to 22% in heart-lung, 2 to 42% of liver, and 2 to 14% of kidney transplanted recipients. Precipitating factors include using of high dose steroids, previous episode of graft rejection, elderly person, leukopenia or chronic hyperglycemia. Commonly isolated agents are Candida spp., Aspergillus spp., Cryptococcus neoformans, Pneumocystis carinii, Hystomycetes capsulatum, Blastomyces dermatitidis, Paracoccidioides brasiliensis. There are no specific signs in clinical presentation of those infections and symptoms overlap another, common infectious diseases, leading to misdiagnosis and high mortality rate varying between 27-77% [1,11,15,21].

**Mycobacterium tuberculosis** /causing tuberculosis, TBC/ occurs in about 1% of solid-organ transplant recipients with 30% mortality rate [14]. The clinical presentation includes fever, but night sweating and weight loss are not common unlike those who are not transplanted. Nontuberculous mycobacteria, such as M. kansasii, M. avium-intracellulare, M. fortuitum, M. xenopi, M. haemophilum, M. marinum, M. cheloneae, M. abscessus, M. gastri, M. scrofulaceum, and M. thermoresistibile often mimic TBC, but when lungs are involved pyrexia is not a common sign. Clinical course includes chronic infections of skin and joints of the digits, wrists, elbows, ankles, and knees. The risk period for these manifestations is 10 days to 269 months after transplantation [6,7].

From the parasitic infections, Toxoplasma gondii and Strongyloides stercoralis are the most important agents because of their distinctive presentation – infections of the central nervous system 7 years after transplantation, or gastrointestinal tract involvement in the first 6 months in the post-transplant period [4].

Different infections occur in specific patterns after transplantation. A strategic approach to fever (and associated organ system–related manifestations) in the solid organ transplantation recipient includes the following steps /see figure1/:

The problem about infections in transplanted patients is actual in nowadays because of increasing frequency of transplantations worldwide. The recipients are specific risk group for various infectious diseases because they may be infected with different agents, leading to atypical clinical presentation. The specific prophylaxis, early diagnosis of the infection and appropriative treatment are the “three mains” for saving patients’ life.

---

**Figure 1:** Changing timeline of infection after organ transplantation (modified from Fishman J. NEJM. 2007;357(25):2601-14).
References


