July 2002

Solid-Organ Transplantation in HIV-Infected Patients

Scott D. Halpern
University of Pennsylvania

Peter A. Ubel
Veterans Affairs Ann Arbor Healthcare System

Arthur L. Caplan
University of Pennsylvania, caplan@mail.med.upenn.edu

Follow this and additional works at: http://repository.upenn.edu/bioethics_papers

Recommended Citation

Publisher URL: http://hdl.library.upenn.edu/1017/5430

This paper is posted at ScholarlyCommons. http://repository.upenn.edu/bioethics_papers/5
For more information, please contact libraryrepository@pobox.upenn.edu.
Solid-Organ Transplantation in HIV-Infected Patients

Abstract
Before the introduction of highly active antiretroviral therapy in the mid-1990s, transplantation centers were understandably reluctant to provide scarce solid organs for patients infected with the human immunodeficiency virus (HIV). However, because treated patients can now expect to live substantially longer than before, many will have end-stage organ disease long before they have life-threatening conditions related to HIV infection. It is therefore time for the transplantation community to readdress the safety, efficacy, and propriety of transplanting scarce organs in HIV-positive patients who need them.

In this article, we provide ethical arguments for viewing transplantation in patients with HIV infection as analogous to transplantation in patients with other chronic illnesses. Accordingly, transplantation in HIV-positive patients should be initiated at major centers and should not be considered experimental. In addition, reimbursement for such procedures should be similar to that for transplantation in other patients, unless evidence accumulates that HIV-infected transplant recipients fare poorly.

Comments
Publisher URL: http://hdl.library.upenn.edu/1017/5430

This journal article is available at ScholarlyCommons: http://repository.upenn.edu/bioethics_papers/5
Sounding Board

Solid-Organ Transplantation in HIV-Infected Patients

Before the introduction of highly active antiretroviral therapy in the mid-1990s, transplantation centers were understandably reluctant to provide scarce solid organs for patients infected with the human immunodeficiency virus (HIV). However, because treated patients can now expect to live substantially longer than before,1,2 many will have end-stage organ disease long before they have life-threatening conditions related to HIV infection.3 It is therefore time for the transplantation community to readdress the safety, efficacy, and propriety of transplanting scarce organs in HIV-positive patients who need them.

In this article, we provide ethical arguments for viewing transplantation in patients with HIV infection as analogous to transplantation in patients with other chronic illnesses. Accordingly, transplantation in HIV-positive patients should be initiated at major centers and should not be considered experimental. In addition, reimbursement for such procedures should be similar to that for transplantation in other patients, unless evidence accumulates that HIV-infected transplant recipients fare poorly.

Current Practices

Although the United Network for Organ Sharing states that asymptomatic HIV-positive patients “should not necessarily be excluded from candidacy for organ transplantation,”4 most centers are concerned that transplantation might harm HIV-positive patients and believe that scarce organs should not be allocated to patients with a poor prognosis.5 In a 1997 survey of directors of U.S. renal-transplantation centers, 88 percent of respondents indicated that they would not consider transplanting an organ in a patient with “asymptomatic HIV-infection who is otherwise a good candidate for transplantation.”6 An earlier survey of Canadian transplantation centers revealed similar views.7 Only a small proportion of U.S. transplantation centers have agreed to participate in a proposed multicenter study of transplantation in HIV-positive patients. There is evidence that most patients and nurses also believe that HIV-positive patients should not be considered for heart transplantation.8 Moreover, insurance companies have generally refused to pay for liver transplantation in HIV-infected patients.9-11

Ethical Considerations

Even before the introduction of highly active antiretroviral therapy, a positive test for antibodies against HIV was rarely, if ever, a legitimate criterion for withholding medical interventions other than transplantation. However, transplantation differs from other interventions with regard to both ethical and medical considerations. Because transplantable organs are scarce, determining the most ethical allocation system requires simultaneous considerations of efficacy, urgency, and equity.12

There are two distinct ethical questions about efficacy: Does transplantation benefit the individual patients? Would it benefit other patients more? With regard to the first question — the question of absolute efficacy — transplantation can certainly help HIV-positive patients with end-stage organ disease.13-23 The second question — concerning relative efficacy — is rarely addressed in the distribution of plentiful resources, but there is a strong moral basis for posing this question when scarce resources are being allocated.24,25 We do not ask whether elderly persons should receive antihypertensive therapy, even though the benefits of long-term treatment are greater for younger persons. However, if transplantation provided substantially less benefit, in terms of survival and quality of life, for HIV-positive patients, then a policy of preferential allocation of organs to HIV-negative patients might be tenable.

Considerations of efficacy are factored into the allocation policies of the United Network for Organ Sharing in two ways: priority is given to candidates whose ABO blood group matches that of a donor, and in the case of kidney transplantation, HLA matching is an additional consideration.8 With the availability of highly active antiretroviral therapy, is it reasonable to suppose that HIV status might still influence efficacy as much as genetic compatibility?

Though there has been no systematic comparison of outcomes between HIV-positive and HIV-negative organ recipients, recent experience suggests that large differences are unlikely. All published reports of transplantation in HIV-positive patients who are receiving multidrug antiretroviral regimens have concluded that, in most cases, HIV infection does not affect the outcome of transplantation.20-23 There are several other, unpublished reports of favorable outcomes several years after transplantation in HIV-positive patients receiving antiretroviral therapy (Olthoff K: personal communication; and Stock P: personal communication).

Even if HIV-positive transplant recipients proved to have somewhat worse outcomes than HIV-negative recipients, relative efficacy is not the sole ethical criterion for determining candidacy. Medical urgency is another important criterion in heart, lung, and liver transplantation. This helps explain why patients who require rapid retransplantation routinely receive it,24 even though expected survival is markedly diminished.24,26,27
Equity (equal access to organs among patients with equivalent need) is also recognized as important by both transplantation specialists and the public. Thus, most programs offer organs to patients infected with hepatitis C virus (HCV), patients with diabetes, older patients (up to a point), and black patients, despite clear evidence that post-transplantation survival is diminished in each of these groups. On ethical grounds alone, there is no justification for providing organs to these groups of patients but not to patients infected with HIV.

MEDICAL CONSIDERATIONS

Might the questionable safety of transplantation in HIV-positive patients justify the discriminatory allocation system? There is an intuitive concern that immunosuppressive therapy might hasten the progression of HIV disease. However, the experience to date suggests that the use of standard immunosuppressive agents in patients with well-controlled HIV infection does not increase their susceptibility to opportunistic infections or malignant conditions. In fact, two common immunosuppressive agents, cyclosporine and tacrolimus, may actually improve outcomes in HIV-positive transplant recipients by inhibiting interleukin-2–dependent T-cell replication, by directly inhibiting HIV replication, or both.

There is also concern about potential interactions between antiretroviral and immunosuppressive agents. Protease inhibitors increase blood concentrations of tacrolimus and cyclosporine by inhibiting specific cytochrome P-450 enzymes. However, experienced centers routinely monitor blood levels of these agents and adjust doses accordingly. The pharmacokinetics of these interactions are similar to those in transplant recipients who require other concomitant medications (e.g., phenytoin) that may induce or inhibit the same enzymes.

Finally, physicians have been concerned that surgery itself might accelerate the progression of HIV disease. However, there is evidence that disease progression is unaffected by surgery, even major surgery requiring cardiac bypass.

CONCERN ABOUT INTRAOPERATIVE TRANSMISSION OF HIV

Another concern is that HIV-positive patients might transmit the virus to members of the transplantation team. However, the risk of patient-to-surgeon transmission of HIV is extremely low and is substantially lower than the risk of transmission of many other infectious diseases, including HCV, which are present in many patients who undergo surgery. Even if a member of a transplantation team were exposed to HIV, various postexposure regimens provide effective prophylaxis against infection.

Many argue that physicians are obligated to accept such small risks because of their unique responsibilities and privileges. Indeed, several major medical associations, including the American College of Surgeons, contend that individual physicians have a duty to provide care to HIV-positive patients. There may also be a legal duty to provide treatment because HIV-positive persons are covered under the Americans with Disabilities Act.

OTHER ARGUMENTS AGAINST TRANSPLANTATION IN HIV-POSITIVE PATIENTS

Some may argue that it is wrong to expand the pool of eligible patients when the supply of organs is already inadequate. This argument erroneously assumes that patients with longer-established rights to receive organs should have priority over those with rights that have been established more recently. Equity dictates that no group of patients be penalized simply because medicine has only recently advanced to a point at which they may benefit from transplantation.

A final argument against transplanting organs in HIV-positive patients is that HIV infection is often, though by no means always, associated with intravenous-drug use or high-risk sexual practices. Such associations, however, have never been a legitimate reason to withhold medical interventions from patients. HCV is also commonly acquired through intravenous-drug use, and hepatitis B virus (HBV) is also commonly acquired through the same high-risk sexual practices as those associated with the transmission of HIV. Yet these diseases, along with alcoholic liver disease, remain the most common indications for liver transplantation in the United States. It is not the role of medicine to adjudicate the morality of personal behavior through selective treatment of life-threatening conditions.

THE BURDEN OF PROOF

It has been proposed that organs be offered to HIV-positive patients only in well-controlled studies until transplantation experts have shown that the outcomes are similar to those for uninfected patients. We disagree with this approach. Proof of relative efficacy has never been required for the initiation of new surgical procedures or for the expansion of indications for established procedures. There was certainly reason to suspect that HCV-positive patients would fare less well after liver transplantation than HCV-negative patients, but systematic comparisons were not required before these patients could receive organs in nonexperimental settings. It is therefore unjust to require such proof before offering transplantation to HIV-positive patients.
Still, some documentation of relative efficacy will eventually be needed to gain the acceptance of physicians, the public, and third-party payers. The best approach is to perform a substantial number of transplantations in HIV-positive patients at major centers and to record all pertinent data from their experience in widely accessible data bases. Investigators will then be able to compare the outcomes for HIV-positive transplant recipients with those for HIV-negative recipients who have similar demographic and disease-related characteristics. Such retrospective analyses are precisely how we learned about outcomes in HCV-positive patients.\textsuperscript{34,35}

PROVISION OF “MARGINAL” ORGANS

A counterproposal is that HIV-positive patients should first merit consideration for the transplantation of so-called marginal, or expanded-criteria, organs — those harvested from older donors, donors infected with HBV or HCV, and, as is increasingly common in renal transplantation, donors with diabetes or hypertension. The outcomes for recipients of marginal organs are typically worse than those for recipients of “ideal” organs.\textsuperscript{57,58} Therefore, in the absence of evidence that HIV-positive patients derive less benefit from transplantation than their HIV-negative counterparts, we see no justification for making HIV seropositivity an indication to use less viable organs. If an HIV-positive patient also happens to have a known coexisting condition to consider, the likelihood that a patient will benefit from transplantation is simply another coexisting condition to consider.

A CALL FOR CONSISTENCY

We believe it would be wrong to transplant an organ in a patient with terminal AIDS for the same reason that it would be wrong to transplant an organ in a patient with widely metastatic hepatocellular carcinoma: both patients would probably die from the coexisting illness before receiving any appreciable benefit from the new organ. However, a broad range of patients who are at low risk for progression to AIDS or AIDS-related death can be identified with the use of base-line CD4+ T-lymphocyte counts and HIV RNA levels.\textsuperscript{39} Furthermore, because changes in the viral load after the initiation of therapy also predict progression to AIDS,\textsuperscript{60} even patients who at one time met the diagnostic criteria for AIDS may benefit from transplantation if they subsequently receive treatment that provides good control of the disease.\textsuperscript{20} Transplantation physicians must often make difficult decisions about how various coexisting conditions influence the likelihood that a patient will benefit from transplantation. The extent of HIV disease is simply another coexisting condition to consider.

Now is the time to remove barriers to transplantation in HIV-positive patients for whom it is otherwise indicated. We should not wait for evidence of relative efficacy to emerge. As with any new patient population, accumulating data will inform future analyses of the appropriateness of transplantation in HIV-positive patients.

Current considerations of efficacy in organ allocation can be summarized as follows. We have substantial evidence of diminished survival among HCV-positive patients, patients with diabetes, black patients, and patients requiring retransplantation, yet we do not prevent transplantation in these groups. In contrast, we have no evidence of poorer survival among otherwise healthy HIV-positive patients who are receiving antiretroviral therapy, yet both overt and covert barriers to transplantation remain. This contradiction is not justifiable according to any ethical theory. Instead, it indicates yet another way in which we continue to discriminate against HIV-positive persons.

Scott D. Halpern, M.S.C.E.
University of Pennsylvania School of Medicine
Philadelphia, PA 19104

Peter A. Ubel, M.D.
Veterans Affairs Ann Arbor Healthcare System
Ann Arbor, MI 48109

Arthur L. Caplan, Ph.D.
University of Pennsylvania School of Medicine
Philadelphia, PA 19104

We are indebted to Drs. Emily Blumberg, Kim Olthoff, Jeffrey Punch, and Peter Stock for their candid discussions and insightful comments as we prepared the manuscript.

REFERENCES


