Cardiac and Pulmonary Histopathology in Baboons Following Genetically-Engineered Pig Orthotopic Heart Transplantation

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Background:
Although improving, survival after pig orthotopic heart transplantation (OHTx) in baboons has been mixed and largely poor. The causes for the high incidence of early failure remain uncertain.

Material/Methods:
We have carried out pig OHTx in 4 baboons. Two died or were euthanized within hours, and 2 survived for 3 and 8 months, respectively. There was evidence of a significant ‘cytokine storm’ in the immediate post-OHTx period with the elevations in IL-6 correlating closely with the final outcome.

Results:
All 4 baboons demonstrated features suggestive of respiratory dysfunction, including increased airway resistance, hypoxia, and tachypnea. Histopathological observations of pulmonary infiltration by neutrophils, and notably, eosinophils within vessels and in the perivascular and peribronchiolar space, with minimal cardiac pathology, suggested a role for early lung acute inflammation. In one, features suggestive of transfusion-related acute lung injury were present. The 2 longer-term survivors died of (i) a cardiac dysrhythmia with cellular infiltration around the conducting tissue (at 3 months), and (ii) mixed cellular and antibody-mediated rejection (at 8 months).

Conclusions:
These initial findings indicate a potential role of acute lung injury early after OHTx. If this response can be prevented, increased survival may result, providing an opportunity to evaluate the factors affecting long-term survival.

Keywords:
Baboon • Heart Transplantation • Lungs • Pig • Xenotransplantation

Abbreviations:
CPB – cardiopulmonary bypass; NHP – nonhuman primate; OHTx – orthotopic heart transplantation; TRALI – transfusion-related acute lung injury

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**Background**

Complex congenital heart disease in neonates and infants, particularly single ventricle pathology, continues to be associated with significant mortality and morbidity despite advances in surgical and medical techniques [1]. Orthotopic heart transplantation (OHTx) might offer the best prospect of long-term survival of the patient, but the availability of hearts of an appropriate size from deceased human donors remains very limited. For infants with inoperable congenital heart disease or failed single ventricle palliation, bridging by mechanical cardiac assist devices is associated with poor results [2].

We suggest that, with recent advances in the xenotransplantation of genetically-engineered pig hearts [3,4] and kidneys [5-8], bridging with a pig heart may be more successful. However, heart function after pig OHTx in nonhuman primates (NHPs) has been mixed and largely poor [3,9-11]. Early graft failure does not appear to be related to early antibody-mediated complement activation, e.g., hyperacute rejection. Although relatively long-term graft survival has been achieved in a handful of cases [3,4], the cause for the high incidence of early graft failure remains uncertain.

To date, we have a very limited experience of pig OHTx in baboons (n=4), but we have made some observations that may throw light on the factors leading to early and late graft failure. We have carried out 4 such transplants using 2 types of genetically-engineered pigs (Table 1). The anesthetic management, surgical technique, postoperative care, and immunosuppressive and adjunctive regimens have been described previously [12]. Cardiopulmonary bypass (CPB) involved priming with blood, but no further blood transfusions were given. There were no technical complications during OHTx. The immunosuppressive therapeutic regimen of T and B cell depletion, followed by an anti-CD40mAb-based maintenance regimen, was identical to that used successfully in baboons receiving pig kidney grafts [6]. Two baboons died within a few hours, and 2 survived for several months (Table 1).

Although some aspects of this experience have been discussed elsewhere [12,13], we believe the histopathological findings are worthy of comment as they suggest a major role for lung inflammation as a factor in the early deaths, and draw attention to the possible role of eosinophils and transfusion-related acute lung injury (TRALI) which have not been reported previously.

**Material and Methods**

All animal care was in accordance with the “Principles of Laboratory Animal Care” formulated by the National Society for Medical Research and the “Guide for the Care and Use of Laboratory Animals” prepared by the Institute of Laboratory Animal Resources and published by the National Institutes of Health (NIH publication No. 86-23, revised 1985). All procedures

![Table 1. Experimental details of pig orthotopic heart transplants in baboons (n=4).](table)

| Baboon (weight) | Pig genotype (weight) | Cold ischemic time (mins) | Survival | Clinical course | Cause of death |
|-----------------|-----------------------|---------------------------|----------|----------------|----------------|
| B217 (8.5 kg)   | GTKO.hCD55 (13.0 kg)  | 130(a)                    | 3 hours  | Ventricular arrhythmias | Cardiac arrest |
|                 |                       |                           |          | Ventricular fibrillation |                |
| B15416 (9.5 kg) | GTKO.hCD55 (12.1 kg)  | 63(b)                     | 3 months | Sinus tachycardia (days 0-30) | Sudden death – dysrhythmia suspected |
|                 |                       |                           |          | Hypoxia (days 0-7) |                |
|                 |                       |                           |          | Hypertension (days 0-30) |                |
|                 |                       |                           |          | Central line infection (P. aeruginosa – day 14) |                |
|                 |                       |                           |          | Left ventricular hypertrophy (day 30) |                |
| B 1917 (8.3 kg) | GTKO.hCD46. hTBM (9.9 kg) | 77(b)                     | 8 months | Hypoxia (days 0-7) | AMR and CMR |
|                 |                       |                           |          | Hypertension (days 0-90) |                |
|                 |                       |                           |          | Tachycardia (days 0-90) |                |
|                 |                       |                           |          | Increased troponin I (day 180) |                |
|                 |                       |                           |          | Sudden death (day 241) |                |
| B37367 (6.0 kg) | GTKO.hCD46. hTBM (6.0 kg) | 71(b)                     | 4 hours  | Hypoxia, Tachypnea, Tachycardia, Hypertension followed by hypotension, Ventricular fibrillation with resuscitation | Idioventricular rhythm, cardiac arrest |

(a) Stored in cold University of Wisconsin (UW) solution in ice for 60 minutes before OHTx. (b) Perfused with cold del Nido solution, and immediately transplanted. No ice storage. GTKO – α1,3-galactosyltransferase gene-knockout; hCD46 – human complement-regulatory protein CD46; hCD55 – human complement-regulatory protein CD55; hTBM – human thrombomodulin; AMR – antibody-mediated rejection; CMR – cell-mediated rejection.
were approved by the Institutional Animal Care and Use Committee of the University of Alabama at Birmingham.

Results

Clinical Courses

Once CPB was discontinued, all baboons demonstrated tachycardia and hypertension which, in the longer-term survivors, persisted for several weeks (Table 1). In the hours following weaning from the ventilator, all 4 baboons demonstrated features of respiratory dysfunction (e.g., tachypnea, reduced \( P_{O_2} \) and \( O_2 \) saturation (Table 1). Cytokine and histopathology observations suggest a critical role for acute pulmonary dysfunction in the early deaths of 2 baboons, but not in the 2 that survived for 3 and 8 months, respectively.

Cardioplegic arrest was with University of Wisconsin (UW) solution in one case (B217), and the pig heart did not function well following discontinuation of CPB. In the other 3 experiments, in which Del Nido cardioplegia was used, immediate cardiac function (as judged by hemodynamics, echocardiography, and need for inotropic support, was good) [12]. In one case (B37367), the baboon was probably extubated too quickly, and this was followed by hypoxia leading to dysrhythmias.

Acute Inflammatory Cytokine Levels During the Peri-Operative Period and Post Transplantation

In the current report, we summarize previously reported details of the acute inflammatory cytokine response after transplantation (Figure 1) [13]. The most important observation was the much greater increase in IL-6 that occurred in the 2 baboons that died within 4 hours from cardiopulmonary failure (Figure 1). In the 2 longer-surviving baboons, there was a delayed and temporary increase in IL-8 after approximately 2 weeks, but this was associated with features suggesting an infectious complication, and had normalized by 4 weeks.

Histopathology of Pig Hearts and Baboon Lungs

After death of the baboon, the pig heart and baboon lungs were removed and fixed in 10% neutral buffered formalin.
saline for at least 24 h. After gross examination, several blocks of both ventricles, interventricular septum, and all lobes of the lungs were cut, dehydrated, embedded in paraffin, sectioned (5µM), and stained with H&E (and Masson’s trichrome). Histopathological features were independently evaluated by 2 board-certified pathologists (SL and JF) prior to confirmation of the findings reported below.

Short-Term Survivors

**B217 (survived 3 hours):** The heart revealed multiple foci of hemorrhage and myocardial necrosis, most prominently within the left ventricle (Figure 2A, 2B). There was no evidence of fibrin thrombi or neutrophilic infiltrate, suggesting that the histopathologic lesions were secondary to ischemia-reperfusion injury. The lungs did not exhibit significant cellular infiltration in the alveoli or bronchi, but significant numbers of intravascular neutrophils were present within the medium-to-small caliber branches of the pulmonary arteries (Figure 2C, 2D), as well as pulmonary edema (not shown).

**B37367 (survival 4 hours):** The pig heart showed subendocardial hemorrhage and contraction band necrosis, typical of ischemia-reperfusion injury (not shown). The baboon lungs (Figure 3A-3D) showed inflammation affecting both the airways and the vasculature. The airways showed prominent inflammation that extended to the adjacent tissue (Figure 3A), and involved the respiratory epithelium (Figure 3B). The pleomorphic infiltrate included a significant number of eosinophils (Figure 3C). The vasculature was heavily inflamed (Figure 3C), and the inflammation extended to the perivascular and vascular tissue (Figure 3D). As with the airways, the inflammatory response included a significant number of eosinophils that were also seen in the lumen. Of interest, a complete blood count obtained in the immediate postoperative period showed leukopenia (2.7K) with neutropenia (1.89K), and lymphocytopenia.
(0.24K), but normal eosinophil count (0.24K), with a relative eosinophilia (9%).

**Longer-Term Survivors**

**B15416 (survival 3 months):** The baboon remained clinically well, and died suddenly. At necropsy, the pig heart was grossly enlarged compared to the day of transplantation (increase in weight from 30 g to 150 g). Histopathological features of inflammation and fibrosis were present within the myocardial interstitium and around the Purkinje fibers (Figure 4A), suggesting a dysrhythmia as the cause of death, similar to that reported after cardiac allotransplantation [14]. The inflammatory infiltrate included a significant number of eosinophils (Figure 4B). There were chronic and acute/subacute changes in the coronary arteries, including intimal hyperplasia (not shown) and mature fibrin thrombi (Figure 4C). A number of arteries showed endothelial activation (Figure 4D), suggesting the possibility of antibody-mediated rejection. The lungs exhibited mild-to-moderate fibrosis within the alveolar septa, around the pulmonary arteries, and at the pleural surfaces (not shown). Within the alveolar interstitium and surrounding branches of the pulmonary artery, there was a mild lymphocytic infiltrate (not shown). Small foci of calcification in medium-caliber arteries were also seen (not shown).

**B1917 (survival 8 months):** The baboon did well for 7 months, at which time there was an increase in troponin I (not shown). At necropsy, the pig heart was grossly enlarged (increase in weight from 40 g to 264 g). Histologic examination of the coronary arteries showed acute and subacute thrombi (not shown), often in a background of layers of intimal hyperplasia (Figure 5A). The veins, perivascularure, and myocardial interstitium exhibited inflammation that included a high number of eosinophils (Figure 5B and 5C respectively). Within the ventricular septum and at the base of the heart there were areas of remote (dark blue on trichrome) and more recent fibrosis (light blue on trichrome, Figure 5D). Immunohistochemistry confirmed...
antibody-mediated rejection, based on IgM (Figure 6A) and C4d (Figure 6B) localization to the endothelium capillaries. Within the interventricular septum there was extensive fibrosis and lymphoid aggregates that by immunohistochemistry corresponded to significant numbers of CD3+ T cells (Figure 6C) and IgM+ CD20low B cells (Figure 6D, and data not shown). The baboon lungs were unremarkable (not shown). Of interest, a complete blood count obtained prior to euthanasia showed a normal white blood cell count (8.3K), with 71% neutrophils, 12% lymphocytes, and 12% eosinophils, suggesting the vascular eosinophilic infiltrate was accompanied by systemic eosinophilia.

Discussion

For this study, we purposely selected relatively small baboons in an effort to mimic clinical OHTx in infants. Our limited experience to date suggests that respiratory dysfunction, secondary to an acute inflammatory state, may be a key factor in the failure of pig hearts early after OHTx in NHPs. Both CPB and organ xenotransplantation result in an inflammatory response, associated with a major release of cytokines (reviewed in 13). Both pig kidney and heterotopic heart transplantation are associated with a fall in plasma free triiodothyronine (fT3) [6,15], another marker of inflammation, which also occurs after CPB [16-18]. We have measured increases in several cytokines/chemokines, e.g., IL-6, TNF, and MCP-1, which increase post-xenotransplantation [13,19].

In the baboon that received a pig heart protected by University of Wisconsin (UW) cardioplegic solution, we believe that primary graft dysfunction contributed to early death. In the other 3 baboons, in which cardioplegic arrest was with Del Nido solution, early graft function was good. However, when our results of early function are compared with those of others in which a system of continuous graft perfusion was utilized to minimize ischemia [3,4], they suggest that continuous perfusion is
preferable unless the myocardial ischemic period is very short. However, graft function can be preserved using a crystalloid cardioplegic solution if the ischemic period is short.

In the 2 baboons that died within hours we noted marked elevations in IL-6, suggesting an initial cytokine storm. IL-6 is induced by CPB [20], specifically associated with post-CPB left ventricular wall motion abnormalities and myocardial ischemic episodes. Both IL-6 and IL-8 can act as potent chemoattrac-
tants capable of eosinophil and neutrophil recruitment [21-23]. IL-6 plays a major role in the inflammatory responses associat-
ed with both CPB [24-26] and organ xenotransplantation [15].

In a murine model, Kumar et al [27] showed that C-reactive pro-
tein enhances antibody- mediated transfusion-related acute lung injury, and it is well established that IL-6 induces the hepatic synthesis of C-reactive protein. Although we have included to-
cilizumab therapy (to reduce the effect of the inflammatory cy-
tokine, IL-6) in our pig kidney xenotransplantation model for several years [6,28], we did not include it in the present study because our more recent research indicates that it prevented IL-6 binding to baboon tissues but not to pig tissues [29]. When binding to baboon tissues is prevented, the level of IL-6 in the blood increases significantly, and we believe this may be detri-
mental to the pig graft. However, after pig OHTx, if the inflam-
matory response is a major factor resulting in pulmonary dys-
function, preventing binding of IL-6 to the baboon’s lungs with tocilizumab might reduce the injury to the lungs, and thus re-
duce respiratory dysfunction and its effect on the pig cardiac graft. In the Munich experience of pig-to-baboon OHTx, tocili-
zumab was administered [3], and this may have been a factor in reducing post-OHTx inflammatory lung injury.

Figure 5. Histopathology of the pig heart (A-D) in B1917 at necropsy. (A) Coronary artery with layers of intimal hyperplasia, suggesting prior episodes of luminal thrombosis followed by organization. (B) Veins showed subendothelial (and perivascular) inflammation that contained a mixed inflammatory infiltrate that included eosinophils (arrow). (C) The myocardial interstitium showed areas of mixed inflammation including eosinophils (arrows). (D) The myocardial interstitium showed evidence of remote scarring (dark blue; white arrow) and more recent scarring (light blue; black arrow) [trichrome stain].

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Figure 6. Immunohistochemistry of heart in B1917 was consistent with acute antibody-mediated and chronic cell-mediated rejection. Left ventricle with significant IgM (A) and C4d (B) staining of interstitial capillary endothelium is consistent with antibody-mediated rejection. Interventricular septum displays significant numbers of CD3\(^+\) T cells (C) and IgM\(^+\) B cells (D) that is consistent with cell-mediated rejection.

Pulmonary histologic evaluation of these 2 cases revealed striking pulmonary findings. B37367 showed prominent inflammation in the airways, including epithelium and adjacent areas, and in the pulmonary medium- and small-sized arteries (including in the media), and perivascular areas. The pleomorphic infiltrate showed a significant eosinophilic component. To our knowledge, this has not been described in solid organ xenografts. Pancreatic islet xenografts in mice showed strong eosinophilic infiltration (4+), while eosinophils were ‘rare’ in allografts [30]. Peripheral eosinophilia is associated with poor outcome post-lung transplantation [31].

B37367 showed a relative eosinophilia during the immediate postoperative period when the pulmonary and hemodynamic status deteriorated, leading to its demise. Conditions like early-phase SARS-CoV-2 are typically accompanied by eosinophilia [32], and the relative eosinophilia in this baboon points toward a mechanism that promotes eosinophil recruitment to several organs and tissues.

In contrast, B217 displayed an accumulation of neutrophils in the pulmonary microvasculature, and pulmonary edema, mimicking the histopathology findings of transfusion-related acute lung injury (TRALI), a condition in which a blood transfusion triggers the activation of neutrophils in the lung microvasculature [33,34]. (All of the baboons received blood in the CPB machine.) In B217, it seems likely that the cause of death was primarily associated with ischemia-reperfusion injury, possibly resulting from inadequate myocardial protection, but there may have been a contribution from compromised gas exchange and the high inotropic therapy that was necessary.

In our 2 longer-surviving baboons, the evidence for an inflammatory response in the immediate post-transplant period was more modest, correlating with the fact that these baboons survived this period. Nevertheless, B15416 still showed some features of inflammatory lung injury when it suddenly died 3 months’ later.
Both baboons showed evidence of antibody-mediated rejection with endothelial activation, intimal hyperplasia, and thrombi in the coronary arteries. B1917 (the longest survivor) had prominent eosinophilic infiltration around both arteries and veins, suggesting that eosinophils played a role in the thrombotic diathesis. The eosinophilic infiltrate was accompanied by eosinophilia. In B15416 the inflammation centered on the Purkinje fibers, without evidence of myocardial damage from coronary thrombosis, supporting the conclusion that death resulted from a sudden dysrhythmia. In contrast, B1917 showed foci of remote and recent myocardial infarction, associated with a rise in troponin, suggesting that graft arteriopathy had been developing.

In summary, the involvement of eosinophils and the similarities of histologic findings and TRALI require investigation.

In both cases, the increase in size and weight of the hearts had been extensive. In B1917, this was in part associated with the presence of severe rejection, and in both baboons it was probably also associated with the rapid growth of the pig organ, as reported by us and others in relation to both kidney and heart xenotransplantation [3,5,6,35]. The transplantation of hearts from pigs genetically-engineered by growth hormone receptor gene-knockout, as proposed by Hinrichs and his colleagues [36,37], may resolve this problem [38]. Rapamycin has been included in our immunosuppressive regimen for many years, and may have reduced growth of the pig hearts [37].

Conclusions

If the problem of the initial inflammatory response to pig OHTx could be overcome, this might facilitate more consistent long-term survival of the pig heart graft.

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Declaration of Figures' Authenticity

All figures submitted have been created by the authors who confirm that the images are original with no duplication and have not been previously published in whole or in part.

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