

## Is it currently possible to evaluate the risk posed by PERVs for clinical xenotransplantation?

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1 *Xenotransplantation*

2 **COMMENTARY**

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4 Is it currently possible to evaluate the risk posed by PERVs for  
5 clinical xenotransplantation?

6

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14

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16 Porcine endogenous retroviruses, virus safety, solid organ xenotransplantation, islet cell  
17 xenotransplantation

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20 The risk of transmission of porcine microorganisms is, in addition to the immunological  
21 rejection and the physiological incompatibilities, a major hurdle to the clinical use of pig  
22 cells, tissues and organs for the treatment of organ failure in humans, to overcome the medical  
23 need caused by the increasing lack of human donors. Whereas most of the porcine  
24 microorganisms may be eliminated by early weaning, colostrum deviation, vaccination,  
25 antiviral drugs, animal isolation, Caesarean delivery of newborns, and embryo transfer,  
26 porcine endogenous retroviruses (PERVs) cannot be eliminated this way because they are  
27 integrated in the genome of all pigs [1]. Only a few years before evidence was published that  
28 PERV is able to infect human cells [2], two other retroviruses, simian immunodeficiency  
29 virus of chimpanzees (SIVcpz) and simian immunodeficiency virus from sooty mangabeys  
30 (SIVsm), now called human immunodeficiency viruses 1 and 2 (HIV-1 and 2), invaded the  
31 human population causing the fatal acquired immunodeficiency syndrome (AIDS) [3, 4].  
32 Although HIV and PERVs are not very closely related, the fact that PERV is a retrovirus  
33 makes it so difficult to evaluate its risk [5]. Also, although most retroviruses are  
34 immunosuppressive in the infected host, the absence of an animal model makes it difficult to  
35 show this for PERV [6].

36 In recent years several strategies have been exploited to evaluate the risk posed by  
37 PERV, such as (i) infection experiments *in vitro*, (ii) infection experiments *in vivo* in small  
38 laboratory animals and in nonhuman primates (NHPs) with and without immunosuppression,  
39 (iii) preclinical trials in NHPs transplanting pig cells and organs with and without  
40 immunosuppression, and (iv) clinical trials mostly using encapsulated pig islet cells without  
41 immunosuppression. Despite these substantial efforts, these studies do not allow to make  
42 unequivocal conclusions whether PERVs pose a risk in the case of treatment of humans with  
43 porcine cells, tissues or solid organs, as will be discussed in the sections below.  
44 Unfortunately, there are no alternative approaches to test this in an experimental setting:  
45 essentially clinical trials are needed to answer this question.

46 1. Infection experiments *in vitro*

47 There are three types of PERV, PERV-A and PERV-B which are present in the genome of all  
48 cells and which infect human cells (human-tropic viruses), and PERV-C which is present in  
49 most, but not all pigs and infects only pig cells (ecotropic virus) (for review see [1]). This  
50 means that PERV-A and PERV-B are able to infect different human cells and cell lines *in*  
51 *vitro*, in cell culture [2]. Recombinant viruses between PERV-A and PERV-C (PERV-A/C),  
52 able to infect human cells and characterized by a high replication rate, have been described  
53 [1]. Some human cell lines such as the 293 pig embryonic kidney cell line are highly  
54 susceptible, and after repeated passages of PERV through these human cells, the virus showed  
55 a higher replication rate and genetic changes in its long-terminal repeat (LTR). These viruses  
56 were called “human cell-adapted PERVs” [7]. The lack of the restriction factor apolipoprotein  
57 B-editing catalytic polypeptide-like subunit (APOBEC) and transformation by DNA viruses  
58 are thought to be the reason for the high susceptibility of 293 cells. However, primary cells  
59 including porcine aortic endothelial cells (PAEC) and peripheral blood mononuclear cells  
60 (PBMCs) have also been infected [8, 9]. PBMCs can more effectively be infected with human  
61 cell-adapted PERVs, however it remains unclear whether the virus infection is productive,  
62 e.g., whether the virus infects cells and produces excess progeny [10]. In the case of PAEC, a  
63 productive infection including mRNA production and particle release have been demonstrated  
64 [8]. Güell et al. [11] described the infection of human umbilical vein endothelial cells  
65 (HUVECs) with PERV, and the presence of proviral DNA in the infected cells. However it  
66 was not clear how the HUVECs had been derived and whether the infection was productive.  
67 Furthermore, all infection experiments, including the experiment with HUVECs, have been  
68 performed using PK15 cells or heavily infected 293 cells as virus source, both releasing high  
69 amounts of virus. In contrast, most primary pig cells, for example pig PBMCs, show only a  
70 low PERV expression at the RNA level and no virus release [12]. Only after mitogenic  
71 stimulation of some pig PBMCs, virus particles were released that were able to infect human

72 293 cells [12, 13]. Therefore, these *in vitro* studies have only a limited relevance for the  
73 evaluation whether PERVs pose a risk for xenotransplantation (Table 1).

74

## 75 2. Infection experiments *in vivo*

76 PERV-A and PERV-B infect not only human cells *in vitro*, but also cells of other species  
77 (polytropic viruses), with some exceptions such as mouse cells [1, 14-18]. Based on these  
78 results numerous attempts have been undertaken to establish a small laboratory animal model  
79 of PERV infection (Table 1). However, injection of PERV preparations into mice, rats, guinea  
80 pigs, and minks, with or without immunosuppression, failed to infect these animals [15-19].  
81 Cells from NHPs could also be infected *in vitro*, however in most cases this did not result in a  
82 productive infection [17, 19]. In some cases, e.g., chimpanzee cells, only human cell-adapted  
83 PERVs were able to show infection [20]. When three NHP species, namely baboons, rhesus  
84 monkeys and pig-tailed monkeys, were inoculated with human cell-adapted PERV-A/C, and  
85 the animals were treated daily with three different immunosuppressive drugs (cyclosporine,  
86 everolimus (RAD), and methylprednisolone), no PERV infection was observed during a  
87 follow-up of more than 300 days [17, 21].

88 Inoculation of rats with PERV or PERV-producing cells [22], or pig islet cells [16], as  
89 well as treatment of minks [18] or guinea pigs [22] with PERV did not result in infection.  
90 Only in guinea pigs a transient infection was observed [23]. Mice could not be infected [15,  
91 24], because mice lack a PERV receptor [25]. Noteworthy, early reports on PERV infection of  
92 SCID mice [26, 27] and athymic mice [28] proved to represent an artifact based on  
93 pseudotyping with endogenous murine retroviruses [29, 30]. Mice transgenic for the human  
94 PERV receptor huPAR-2 have been generated, and it was reported that they could be infected

95 with PERV *in vivo* [31]. Although this is the only known *in vivo* model of PERV infection, no  
96 follow-up studies on pathogenic effects of the virus were published.

97 In rhesus and cynomolgus macaques, and baboons, the main virus receptor PAR-1 was  
98 found to be genetically deficient by a mutation at the same position as reported in mice, which  
99 is one explanation for the inefficient infection [32]. The receptor in African green monkeys  
100 does not have this mutation, but nevertheless the replication is quite low [32].

101 To summarize, all small laboratory animal and NHP model systems are not suitable to  
102 evaluate the risk posed by PERVs or to study PERV pathogenesis (Table 1).

103

### 104 3. Preclinical trials in NHPs

105 In recent years a number of pig-to-NHP preclinical xenotransplantation studies have been  
106 performed regarding hearts, kidneys, islet cells, or studies performing perfusion of pig liver,  
107 under immunosuppression (for review see [1]). In all these studies, and also in more recent  
108 transplantations not listed in [1], i.e., studies on islet cell transplantation in marmosets [33]  
109 and cynomolgus monkeys [34], no PERV transmission was observed. However, since the  
110 PERV receptor in NHPs is not functional, these results cannot be used to evaluate the safety  
111 of xenotransplantation using pig cells and organs (Table 1). Hence, it does not make sense to  
112 include the monitoring for PERV transmission in pivotal nonclinical trials before phase  
113 transition to clinical development. Interestingly, some regulatory agencies require such  
114 studies, which are elaborate and time consuming, and essentially not informative.

115

116

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118 4. Clinical trials

119 In the past, more than 200 humans have received a xenotransplantation product comprising  
120 pig cells or tissues including *ex vivo* perfusion of pig organs or pig-cell based bioreactors ([35,  
121 36], reviewed in [1] and [37]). No evidence for virus transmission was obtained using sensitive  
122 PCR-based methods and immunological assays for the detection of antiviral antibodies.  
123 Neither antibodies against PERV as an indirect sign of infection, nor provirus integration in  
124 PBMCs of the patients was observed.

125 During the last years further clinical trials have been performed, including the first  
126 prospective clinical trials under proper regulatory oversight using encapsulated pig islet cells  
127 to treat type one-diabetes in New Zealand [38] and Argentina [39]. Although the clinical  
128 efficacy in these trials was limited, no PERV transmission has been observed [40, 41].

129 In all of these porcine islet clinical trials no immunosuppression was given and the islet  
130 cells were transplanted encapsulated in biopolymers, a procedure which protects from host's  
131 humoral and cellular immune system (immunoglobulins and immune cells), but also which  
132 prevents release of PERVs (Table 1). After some pioneering explorations more than 40 years  
133 ago, transplantation of a large vascularized organ accompanied by an effective pharmaceutical  
134 immunosuppression has still not been performed.

135

136 5. Perspectives

137 Although human cells can be infected with PERVs under specific and somewhat  
138 artificial conditions, i.e., co-culture of human cells with porcine cell lines that do not resemble  
139 primary pig cells regarding PERV expression and virus production, or co-culture of porcine  
140 cells which human target cell lines that do not resemble primary human cells, no PERV  
141 transmission has been observed in the first clinical trials. Also, upon inoculations of PERV

142 particles or PERV-producing cells into small laboratory animals or NHPs, no PERV  
143 transmission has been observed. In addition, no PERV transmission was observed in  
144 preclinical trials transplanting encapsulated pig islets in diabetic NHPs or transplanting  
145 kidneys or hearts into immunosuppressed NHPs. Noteworthy, the trials and infection  
146 experiments in NHPs are limited by the lack of a functional PERV receptor in NHPs. Trials in  
147 humans used mainly encapsulated pig islet cells. Encapsulation prevents immune rejection,  
148 but could also prevent the release of PERV and other pathogens. *Ex vivo* perfusion of pig liver  
149 and spleen by human blood, pig skin transplantations, and injection of pig neuronal cells into  
150 the immunoprotected human brain, have also been performed [35, 36]; but till now  
151 transplantations of vascularized pig organs under chronic immunosuppression have not been  
152 performed. At present there are no additional experimental approaches available to evaluate  
153 whether PERVs pose a risk.

154         During the last years, first reports have been published that PERVs in the genome can  
155 be inactivated by CRISPR/Cas9-mediated gene editing tools [42], and also that this procedure  
156 allows the generation of live pigs with all PERVs being inactivated [43]. Although the  
157 functionality has been shown in *in vitro* cell culture, with inherent low translation value to the  
158 pig-to-human clinical situation as outlined above, it needs to be shown in an *in vivo* situation  
159 that the inactivation of PERVs in the pig donor makes sense, also in relation to the off-target  
160 effects of the gene editing procedure [44].

161 This aside, the possibility of gene editing resulting in inactivated PERVs raised the question  
162 whether conventional pigs can still be used for xenotransplantation, or whether only  
163 CRISPR/Cas9-inactivated pigs have to be used as source animals for future  
164 xenotransplantations [11, 44-46]. PERV proviruses inactivated by CRISPR/Cas9 cannot be  
165 restored by recombination, since in all proviruses the gene coding for the important reverse  
166 transcriptase is destroyed. Recombination or co-packaging between PERVs and human

167 endogenous retroviruses (HERVs) have not been reported [47]. Furthermore, off-target effects  
168 by CRISPR/Cas9 may happen, but they will be detected when analyzing the health of the  
169 animals, and animals with defects will be eliminated.

170 Therefore, two options for the first solid organ xenotransplantations could be foreseen.  
171 First, the use of organs from conventional, non-CRISPR/Cas9-treated animals in well-  
172 controlled trials, e.g., using pigs with absence of PERV-C, low copy number and low  
173 expression of PERV-A and PERV-B. Monitoring of the xenotransplant recipient would be as  
174 proposed by regulatory agencies [48] using highly sensitive PCR-based and immunological  
175 methods. Alternatively, pigs with CRISPR/Cas9-inactivated PERVs could be used. The  
176 monitoring might in first instance be similar as mentioned above, considering that the sense of  
177 the gene editing can not be demonstrated in *in vivo* animal models [44]. Additional strategies  
178 to prevent PERV transmission have been considered such as a vaccine based on neutralizing  
179 antibodies [49-52] and antiretroviral drugs (for review see [53]), which may be used should a  
180 positive detection of PERV occur. With this in mind, it seems feasible to go ahead with  
181 conventional animals as has been done in many trials before.

182

183

#### 184 CONFLICT OF INTEREST

185

186 H-J S is director at SchuBiomed Consultancy, and provides consultancy in the biomedical  
187 sector worldwide. JD and LS have no conflicts of interest.

188

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Table 1 Evaluation of the results of different PERV transmission experiments

Setting, method	Outcome PERV transmission	Possible reason why negative	Possible reason why positive	Outlook, conclusion
Clinical transplantation of pig islet and other cells to humans, ex vivo perfusion	No transmission*	Encapsulated cells, low number, no immunosuppression		No relevance for solid organ transplantation into humans
Preclinical transplantation of different pig organs into non-human primates	No transmission	Absence of functional PERV receptor		No relevance for solid organ transplantation into humans
Infection experiments in vivo in small animals and non-human primates (NHP) with and without pharmaceutical immunosuppression	No transmission**	Absence of PERV receptor, or absence of functional PERV receptor, or low PERV receptor density		No relevance for solid organ transplantation into humans
Infection experiments in vitro	Infection of human cells and cells from other species		Use of high virus load for infection, target cells susceptible due to lack of restriction factors, use of human cell adapted virus	Only limited relevance for transplantation into humans, innate and adaptive immune system not present

\* In some patients microchimerism was detected, e.g., the presence of pig cells, but no infection [35].

\*\*Reports showing that SCID mice were infected with PERV [26-28] were the result of an artefact based on pseudotyping between PERV and endogenous murine retroviruses [29, 30].