Transgenic pigs designed to express human α-galactosidase to avoid humoral xenograft rejection

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Abstract The use of animals as a source of organs and tissues for xenotransplantation can overcome the growing shortage of human organ donors. However, the presence of xenoreactive antibodies in humans directed against swine Gal antigen present on the surface of xenograft donor cells leads to the complement activation and immediate xenograft rejection as a consequence of hyperacute reaction. To prevent hyperacute rejection, it is possible to change the swine genome by a human gene modifying the set of donor’s cell surface proteins. The gene construct pGal-GFPBsd containing the human gene encoding α-galactosidase enzyme under the promoter of EF-1α elongation factor ensuring systemic expression was introduced by microinjection into a male pronucleus of the fertilised porcine oocyte. As a result, the founder male pig was obtained with the transgene mapping to chromosome 11p12. The polymerase chain reaction (PCR) analysis revealed and the Southern analysis confirmed transgene integration estimating the approximate number of transgene copies as 16. Flow cytometry analysis revealed a reduction in the level of epitope Gal on the cell surface of cells isolated from F0 and F1 transgenic animals. The complement-mediated cytotoxicity assay showed increased viability of the transgenic cells in comparison with the wild-type, which confirmed the protective influence of α-galactosidase expression.

Keywords Epitopes · Transgenic pigs · Xenotransplantation

Introduction

Decreasing the frequency of chronic graft rejections, decreasing the toxicity of immunosuppressive treatments, inducing tolerance, and ensuring a sufficient number of graft organs are currently the most important challenges of modern transplantology. While the number of people awaiting transplantation is increasing every year, the availability of organs for transplantation is limited. In 2011, according to the Organ Procurement and Transplantation Network (OPTN), the number of registered donors in the United States of America was 11,713 (including living donors, which constituted 42.12 %), and the number of patients awaiting transplantation was 112,728. There was a total demand for 122,438 organs, while the number of transplantations was only 23,745. For example, 16,857 patients awaited liver transplantation, while the available organs could cover only 34.34 % of the demand. In fact, in 2011, 5,288 liver transplantations were conducted in 31 % of awaiting patients (OPTN 2011). The introduction of improvements in the system of organ registration and distribution and better general health of the society obtained through promoting healthy lifestyle are not enough to improve the short supply of potential organs and transplantation tissues. This situation resulted in increased interest in studies on alternative sources of organs. Transplantation...
involving the use of organs from other mammals as human transplants are called xenotransplantation. They have become the most obvious strategy of solving the problem and a chance to save lives.

From both the genetic and physiological points of view, the phylogenetic proximity of apes and humans is the strongest argument standing for xenotransplantations. However, it is also one of the biggest obstacles, involving the risk of infections, particularly with animal viruses. Due to the large phylogenetic distance and similar anatomical and physiological parameters, especially in respiratory and circulatory systems, choosing pigs as a donor of tissues and organs for transplantation seems to be justifiable. The size diversity between different types of pigs allows for choosing organs of size adjusted to different recipients. Pigs, as an easy-breeding species, can be housed under strict barrier conditions and screened for pathogenic xenozoonosis at regular intervals (Cooper 2012). However, it should be expected that the large phylogenetic distance will be the reason for a number of immunological complications after pig-to-human transplantations. Binding the naturally occurring preformed antibodies of recipient to Galα(1,3)Gal epitopes on the endothelial surface makes apes reject grafts from pigs by means of hyperacute rejection (HAR). The Galα(1,3)Gal epitope is formed by connecting a molecule of galactose to N-acetyllactosamine by the 1,3-glycosidic bond in the presence of α1,3-galactosyltransferase (GGTA1, α1,3GT) (Sandrin and McKenzie 1994). Both the enzyme and the sugar residue are not present in humans and apes of the Old World (Galili et al. 1988). It is suggested that, in ancestors of order primates, the α1,3-galactosyltransferase gene-knockout (GTKO). If dom-
utes of D-galactose from the Gal antigen entirely. At the same time, it is suggested that, in the case of isolated enzymes, GLA and α1,2-fucosyltransferase (α1,2FT) coexpression is used. The enzyme α1,2-fucosyltransferase competes with α1,3-galactosyltransferase, which is responsible for the synthesis of Galα(1,3)Gal epitope, for the same substrate, N-acetyllactosamine (N-lac), catalysing the addition of fucose, which results in the formation of the so-called H structure, commonly present in mammals (Larsen et al. 1990; Sandrin et al. 1995). The additive effect of both enzymes resulting in the total elimination of Gal antigen expression on the cell surface was observed in the case of cotransfection of different types of cells with genes coding for GLA and α1,2FT (Osman et al. 1997; Yan et al. 2003; Jia et al. 2004).

The aim of this study was to prepare an expressive pGal-GFPBsd gene construct containing the sequence coding for enzyme α-human galactosidase controlled by the promoter of EF-1α elongation factor ensuring the systemic expression of transgene and introducing it to the genome of pig by microinjection. In this study, we present the results characterising transgenic pigs with human α-galactosidase expression using molecular, cytotagenic and functional approaches. The results are confronted with data obtained for TG1154 transgenic pig with α1,2-fucosyltransferase expression (Lipiński et al. 2010) in order to conduct cumulative transgenesis in the future for cumulating the activity of both enzymes in one organism for the needs of xenotransplantation.

Materials and methods

Gene construct

The pGal-GFPBsd gene construct was obtained by introducing cDNA coding for α-galactosidase (1,290 bp) into a
pTracer-EF/Bsd A (5,987 bp) plasmid containing a strong, constitutive promoter of 1α subunit of human elongation factor EF (hEF-1α) (1,179 bp) and a poly(A) sequence of cattle GH gene (225 bp). Vector pTracer-EF/Bsd A expresses the blasticidin resistance gene as a fusion product with the Cycle 3 GFP, allowing the selection and non-invasive detection of recombinant proteins (Invitrogen). The coding sequence was amplified on single-strand cDNA template synthesised by oligo(dT)20 primers and the SuperScript III kit (Invitrogen). Total RNA was isolated from human white blood cells from peripheral blood with the method based on guanidine isothiocyanate (Chomczynski and Sacchi 1987). Modified primers were used for amplification. Additionally, a sequence recognised by the EcoRI enzyme (5′-GGCGA TTCCATGCGCTGAGGAACCAGA-ACTACA-3′) was placed in the F primer and a sequence recognised by the NotI enzyme (5′-GGCGGCGCCTTAAAGTAAGTGCTTATT AATGACATCTGG-3′) was placed in the R primer.

Polymerase chain reaction (PCR) of 25 μl was conducted in the following conditions: 94 °C, 60 s; 58 °C, 60 s; 72 °C, 90 s; 35 cycles. The reaction mixture contained 1 μl of cDNA, 10 pM of primers, 7.5 nM dNTP, 1× buffer and 1 U of DNA Taq polymerase (Sigma). PCR product was digested by EcoRI (Fermentas) and NotI (Fermentas), purified and ligated with the pTracer-EF/Bsd A vector, which was digested with the same endonucleases. The product was sequenced using automated genetic analysers (Applied Biosystems Prism). The construction prepared for microinjection was purified by the PureLink™ HiPure Plasmid DNA Maxiprep Kit (Invitrogen) and then linearised with the Eaml105I enzyme.

Microinjection

Exogenous DNA as a pGal-GFPBsd gene construct was inserted into the male pronucleus of a fertilised egg cell. Microinjection was performed with the use of two micromanipulators under a reverse microscope with differential interference contrast. The applied magnification was objective 40× X eyepiece = 450×. DNA was injected into the male pronucleus with the use of an injector or a syringe connected to the holder for a manipulator. The volume of inserted DNA was 2–5 pl. DNA was inserted until the moment when the pronucleus increased its volume by 50 %. The DNA concentration was 2–4 ng/μl. After the injection, the damaged zygotes were removed. Selected zygotes with the GFP signal detected were transplanted into the fallopian tubes of synchronised recipients.

Screening of transgene

Analysis included genomic DNA isolation from biopsies of potentially transgenic animals with the use of proteinase K. PCR was conducted on DNA template. Two pairs of primers were used for amplification. Both F primers were complementary to the hEF-1α promoter sequence, and both R primers were complementary to the sequence of α-galactosidase (Fig. 1). A PCR product of 464 bp is formed with the use of the first pair of F1 (5′-GGGGAAGGGTTTTATGCGATG GAG-3′) and R1 (5′-CTGGCTCTTCTCCTGACGATCAG-3′) primers, whereas a second product of 861 bp is formed with the use of the F2 (5′-ACCACTTTGCAGCGAGAAAG ATG-3′) and R2 (5′-GGCGAATCCATGAGGAAAG-3′) primers. PCR was conducted in 20 μl in the following conditions: 94 °C, 40 s; 60 °C, 30 s; 72 °C, 60 s; 35 cycles. The reaction contained 100 ng of genomic DNA, 10 pM of primers, 7.5 nM of dNTP, 1× buffer and 1 U of DNA Taq polymerase (Sigma).

Southern hybridisation analysis

For Southern analysis, total genomic DNA from the wild-type and analysed transgenic pig was extracted from white blood cells from peripheral blood with the method based on guanidine isothiocyanate. 5.6 μg of each DNA was digested with restriction enzyme EcoRI and hybridised with a probe of 464 bp. The probe was a 32P-dATP-labelled PCR product of 464 bp formed on plasmid DNA (pGAL-GFPBsd) with the use of the first pair of primers (F1 and R1) used for integration analysis (described above). The EcoRI enzyme digests the genomic DNA of pig as well as the pGAL-GFPBsd sequence (only one site). Digestion of the pGAL-GFPBsd sequence with EcoRI gives a fragment of 7.3 kb that is identical to the size of the introduced genetic construct. Additionally, for determining the transgene copy number, to the digested DNA of wild-type pig, known amounts of plasmid-containing transgene insert DNA was added. Then, DNA samples were electrophoresed using 1 % agarose gel in 1× TBE, 40 V overnight. Denatured and neutralised DNA immobilised in dried gel was hybridised to a transgene-specific probe overnight (55 °C, 5 × SSPE, 0.1 %SDS).

Determination of the absolute number of transgene copies in transgenic pigs

Since 1 ng of DNA corresponds to the molecular mass of 606,060,600 MDa and the molecular mass of the swine genome amounts to 1,820,000 MDa, therefore, 1 ng of swine genomic DNA contains approximately 333 copies of the genome. 5.6 μg of swine genomic DNA sample per gel line were used, hence, each of the examined samples contained about 1,864,800 copies of the swine genomic DNA. In order to calculate the number of copies of the transgene incorporated into a single genome of transgenic animals, the absolute number of copies of the transgene in titration samples (respectively, 124,122,350, 62,061,175, 31,030,587, 15,515,293,
7,757,646 copies) was divided by the number of copies of the genomic DNA. In the case of heterozygotic transgenic animals, the transgene occurs only on one of the pair of homologue chromosomes; therefore, in order to calculate the number of transgene copies built into one of the chromosomes, the result obtained from the calculations was multiplied by 2.

Skin fibroblasts isolation and cultivation

The experimental material comprised skin fibroblasts obtained from the tissues of wild-type and transgenic pigs. Sterile tissues were rinsed in PBS solution containing antibiotics (50 μg/ml gentamicin sulphate, 100 IU penicillin and 50 μg/ml streptomycin). Skin fragments were cut and digested in 5 ml trypsin-EDTA solution in PBS and then centrifuged. The cell sediment was suspended in enriched DMEM medium (20 % FBS, 1 % antibiotic/antimycotic; Sigma). Culturing was carried out at the temperature of 37 °C at 5 % CO2 content. Transgenic cultures were checked for GFP signal. Cultures of full confluence were passaged to new 25 cm² culture flasks.

Mapping of the pGal-GFPBsd transgene

Transgene mapping was conducted by fluorescent in situ hybridisation (FISH) with the use of in-vitro-cultivated cell lines. Pig karyotype evaluation was performed before conducting FISH. It involved the trypsinisation of chromosome preparations with trypsin and staining them with Giemsa stain solution. The obtained chromosome bands were analysed with the use of a microscope with a camera and appropriate software (Ikaros, MetaSystems). FISH was used for random primer hybridisation of the pGal-GFPBsd gene construct into the metaphase chromosomes obtained earlier. Hybridisation was conducted at 37 °C for 17 h. Signal detection was performed with the use of antibodies against biotin and detected by fluorescein. In order to enable observation under a fluorescent microscope, DAPI/Antifade was used for dying. The image analysis was performed with the use of an Olympus 3X41 fluorescent microscope and the digital image analysis system MetaSystems 4000 ISIS Version 5.0. Transgene location was determined based on the standardised domestic pig karyotype (Gustavsson 1988).

Flow cytometry

Having reached full confluence, the skin fibroblasts cultivated in vitro were washed with Hanks solution and trypsinised (0.25 % trypsin, 0.02 % EDTA). Fibroblasts were collected by centrifugation. The cells were suspended and incubated in PBS containing lectin (10 μg/ml) and 5 % FBS, at 4 °C, for 45 min. Lectin BS-IB4 isolated from Griffonia simplicifolia (Bandeiraea simplicifolia) conjugated with AlexaFluor 647 fluorochrome (Invitrogen) was used for the specific detection of Gal epitope. The flow cytometer (cell sorter) BD FACS Aria™III (Becton Dickinson) with four lasers (375 nm, 405 nm, 488 nm and 633 nm), 11 fluorescence detectors and dispersed laser light detectors – forward scatter (FSC) and side scatter (SSC) were used for the analysis of Gal epitopes on the surface of examined cells. Optical alignment and functional stability tests were conducted with the CST system (Cytometer Setup and Tracking; Becton Dickinson). FACSFlow (Becton Dickinson) was used as the sheath fluid. The following cytometer configuration was applied: 85 μm cell sorting nozzle and sheath fluid pressure of 45 psi. The characteristics of cells were obtained based on two non-fluorescent parameters, FSC and SSC, and one fluorescent parameter, red fluorescence (FL1 detector), from the AlexaFluor 647 fluorochrome, with detection obtained by the use of a 660/20 optical filter. Red laser (633 nm) was used for AlexaFluor 647 fluorochrome excitation. Cytometric analysis was conducted based on specific detector settings, which convert the collected signals into a logarithmical scale. The FSC detector was equipped with an optical filter of neutral density in order to weaken the dispersed light reaching it and enable conversion of the population of analysed cells on a two-dimensional scale to a graph showing the relations between the FSC and SSC. The
threshold was set at the signal of the FSC detector. The data presented on the logarithmical scale were divided into four decades in the form of the signal surface from different detectors (FSC-A, SSC-A and FL1-A) and analysed with the use of FACS DIVA software (Becton Dickinson). The analysis of fluorescence signals preceded the doublets discrimination procedure with the use of height versus width scatter signals measurement, in order to discriminate single events (cells) from conglomerates. The populations were then defined by gating in the dot plots of red fluorescence (FL1) versus side scatter signals (SSC). Each sample was analysed four times. Determining the levels of Gal epitope on the surface of the analysed cells was conducted based on the median values of fluorescence signals from the FL1 detector in populations defined on a two-dimensional graph (FL1 and SSC).

Human serum cytotoxicity test

Having reached full confluence, the skin fibroblasts cultivated in vitro were rinsed with Hanks solution and trypsinised. The fibroblasts were collected by centrifugation and the debris was washed with Hanks solution and suspended in 200 μl of one of three mediums: (A) 50% DMEM and 15% FBS in PBS, basic medium; (B) 50% HS (human serum) in PBS, test medium; (C) 50% thermally inactivated HS in FBS in PBS, basic medium; (B) 50% HS (human serum) in PBS, control medium. The cells were incubated at 37 °C for 30 min. Then, 20 μl of 0.4% trypan blue was added to 20 μl of cell suspension and incubated for 5 min at room temperature. Living and dead cells were counted in Bürker’s chamber with a volume of 0.9 mm³. Cells that were over or touching the top and left line of each square of 1 mm² were counted, while the cells that were over or touching the bottom or right line were ignored. Each analysis was repeated 12 times. The percentage of living cells was calculated for each repetition. Then, the average numbers of living cells, dead cells and all cells from 12 repetitions were calculated.

Results

Obtaining transgenic pigs with α-galactosidase expression

The pGal-GFPBsd gene construct was obtained by introducing cDNA coding for α-galactosidase into a pTracer-EF/Bsd A plasmid. The correctness of the gene construct was confirmed by sequencing. The prepared pGal-GFPBsd gene construct (Fig. 1) was introduced based on the standard method of microinjection into one of the pronuclei of the zygote. 580 fertilised egg cells from 48 donors were subjected to microinjection. 529 zygotes were transplanted into the fallopian tubes of synchronised recipients (average of 33 zygotes/animal). From eight recipients, 37 piglets were obtained in the F0 generation, which were subjected to DNA analysis in order to detect the transgene by PCR. The analysis showed that one male piglet in the F0 generation (TG252) incorporated the transgene (1/37; 2.7 %) (Fig. 2a). When the male TG252 reached puberty, its sperm was used for the artificial insemination (AI) of wild-type sows. As a result of AI, 75 piglets have been obtained so far, among which 12 were transgenic (12/75; 16 %). Changes in the size of the products analysed by PCR were not observed in transgenic pigs from the F1 generation, which means that the transgene is stably integrated with the genome and passed between consecutive generations (Fig. 2b). To confirm transgene integration, we conducted Southern analysis for all transgenic animals, which confirmed the presence of transgene in all of them. We show the results for TG252 (F0) and three representatives of the F1 generation, TG1183, TG1036 and TG1040, in Fig. 3. By using known concentrations of the transgene mixed with wild-type DNA, we were able to estimate the copy number in each of the animals. In the current analysis, the approximate number of transgene copies integrated was estimated as 16.

From screening by Southern analysis, it can be determined that transgene is present, intact, in how many copies and what orientation. Neither of the other commonly used techniques yielded nearly the same amount of information. PCR analysis does not allow to quantify the copy number and, usually, can only confirm the presence of the transgene and cannot provide information as to whether the intactness or orientation of the transgene. Despite its limitations, the technique is useful as a preliminary screening method. The use of PCR allows the identification of potential transgenic animals and reduces the number of samples for Southern analysis (Tinkle and Jay 2002).

Cytogenetic analysis

Karyotype evaluation of the transgenic animals was performed by the Giemsa method. The obtained pattern of bands on chromosomes was analysed and compared with the standard karyotype of domestic pig. The number, structure and architecture of chromosomes were unaffected in both the F0 (TG252) and F1 (TG1183) generations (Fig. 4a, b). In both generations (F0 and F1), the transgene was located only on chromosome 11, short arm, region 1, band 2. It was stable and integrated with genomic DNA (Fig. 4c, d). The transgene was integrated as concatemers of an estimated 16 copies on chromosome 11p12.

Flow cytometry

Flow cytometry was used for checking whether the transgene expression results in the presence of functional α-galactosidase responsible for removing terminal residues of α-D-galactose.
from the Galα(1,3)Gal epitope, which, in consequence, should be connected with the reduction of the Gal antigen level on the surface of cells in pigs. Skin fibroblasts derived from wild-type animals, TG252 (F0) and TG1183 (F1), were used for the analysis. Before the analysis of α-galactosidase functionality on the protein level carried out by measuring the amount of Gal antigen on the cell surface by flow cytometry, the enzyme was labelled by BS-IB4 lectin conjugated with AlexaFluor 647 fluorochrome. BS-IB4 lectin isolated from *Griffonia simplicifolia* (Bandeiraea simplicifolia) specifically binds Gal antigen and the fluorescent intensity of AlexaFluor 647 allows for the determination of the relative amount of labelled antigens on the surface of examined cells compared to non-transgenic cells. The median values of fluorescence signals from FL1 detector (AlexaFluor 647) are proportional to the Gal epitope levels on the surface of the analysed cells. Staining with BS-IB4 lectin showed a significant reduction of fluorescence intensity in the case of fibroblasts from both the founder from the F0 generation (TG252) as well as the transgenic animal from the F1 generation compared to control fibroblasts of wild-type animals. From the obtained results, it can be concluded that the transgene integrated with genomic DNA of pigs is fully functional, was incorporated into an active transcription region and is expressed.

The analysis showed decreased levels of Gal antigen in transgenic animals in F0 and F1 compared to wild-type animals. The reduction of Gal epitope expression was 58.21 %, from the Galα(1,3)Gal epitope. The decrease in Gal antigen levels was confirmed by polymerase chain reaction (PCR) and Southern blot analysis. The PCR was used to amplify DNA fragments of 464 bp and 861 bp. The separation of DNA fragments was conducted in 1.5 % agarose gel. A Southern blot was hybridised with a probe of 464 bp. The probe detects a 7.3-kb band that represents the pGAL-GFPBsd genetic construct (black arrow). The red arrow probably indicates the band of non-specific hybridisation of the probe.
59.67 %, 56.67 % and 65.09 %. The level of Gal epitope expression observed in F1 animals was similar to the F0 animal (TG252). The fluorescence intensity decrease in TG252 was 2.59\times in relation to the wild-type control, while for the pig from F1 (in this study TG1183 and, additionally, TG1036 and TG1040) was 2.86–2.31\times (Fig. 5b). Additionally, fibroblasts from cell lines derived from the TG1154 transgenic pig with the expression of $\alpha_{1,2}$-fucosyltransferase, obtained by the study group in the previous years (Lipiński et al. 2010), were used for comparative analyses. For the transgenic pig TG1154, a relative decrease of more than two and a half (62.8 %) in the amount of Gal antigen on the cell surface was observed in relation to the wild-type control, measured by median values of fluorescence intensity after staining with lectin BS-IB4 (Fig. 5a).

**Human serum cytotoxicity test**

Another test verifying the pGal-GFPbsd transgene functionality was the survival rate analysis of skin fibroblasts derived from TG252 (F0) and TG1183 (F1) in comparison to the survival rate of other wild-type cell lines in the presence of human serum acting as a source of complement system elements. Activation of the complement system results in a number of reactions which effect the formation of a complex attacking the membrane. It is responsible for forming channels...
The analyses conducted for the test medium, where living cells constituted only 22.05 % (+/−3.65 %), showed that the compounds present in the serum affected adversely the survival rate of wild-type pigs compared to basic medium (89.87 %, +/−6.09 %), with no elements of the human complement system present in the human serum. The survival rate analysis showed that there is no statistically significant difference between the survival rate of cells in the basic medium (A) and in the control medium (C), so it is not the source of toxic compounds. The survival rate was determined by the incubation of cell suspension with a solution of trypan blue, a dye which penetrates the structure of dead cells whose membrane is fragmented.

The complexity of processes responsible for initiating immunological reaction does not allow a one-dimensional approach to xenotransplantation. Several strategies need to be combined for obtaining an additive effect in one transgenic organism in order to increase the chance of stopping the immunological aggression against the cells of the xenograft and its consequent survival. Modern genetic engineering is facing a serious problem of producing, in the shortest possible time, polytransgenic animals by interbreeding monotransgenic animals or simultaneous transfection with several transgenes in mono- or polycistronic transgene expression. In the case of xenotransplantation, the opportunity to develop and incorporate multiple genetic modifications simultaneously will be limited until each mechanism is identified. It was proved that, in the case of coinjection of two similar gene constructs, the expression of both transgenes shows
effectiveness similar to that observed in the case of single transgenes introduced into two different animals (Jankowsky et al. 2001; Goldman et al. 2004).

Hyperacute rejection of xenografts is the basic immunological limitation that makes transplanting animal organs in humans unsuccessful. Experimental data and some clinical experience show that there are three main directions that can be followed: depletion of anti-pig antibodies by the development of homozygous pigs for α,3-galactosyltransferase gene-knockout (GTKO), enzymatic removal of Galα1,3-Gal epitope or complement inhibition, or some combination of these approaches (Pierson 2009). As we started our experiments in 2002, there were several concerns about pigs with the GTKO phenotype. First is the possible indispensability of Gal in pigs and its essentiality for cellular function. Second is the presence of another galactosyltransferase which is capable of producing the Gal xenoantigen on glycolipids. And, finally, the deletion of the Galα1,3-Gal epitope would expose new, potentially immunogenic antigens (Gock et al. 2011). In the human organism, even 90 % xenoreactive antibodies recognise the Galα(1,3)Gal epitope, but there is still the remaining 10 % to be challenged. As a consequence, we chose two strategies involving molecular modifications of the animal genomes aiming at preventing the hyperacute rejection from the immunological system. The first strategy described in this study helps to prevent HAR by means of a reduction of the epitope through the introduction and expression of a sequence coding for α-galactosidase, which eliminates the effect of α1,3-galactosyltransferase as a result of removing the terminal α-D-galactose in the Galα(1,3)Gal structure. Microinjection allowed us to obtain a transgenic pig with an integrated, functional and stable pGal-GFP-Bsd transgene. Both the transgenic hog TG252 as well as animals in the F1 generation were characterised by a GLA expression level ensuring the derived lines of fibroblast protection from the destructive human complement system in the cytotoxicity test. The survival rate measured in the basic medium and in test medium B containing 50 % human serum. The survival rate was measured by an average number of living cells compared to the total number of counted cells in 12 repetitions. The percentage cell survival rate is shown on the y-axis.

In previous studies, we focused on attempting to decrease the level of Gal antigen synthesis on the surface of cells by replacing it with a commonly accepted H antigen (0 antigen of the ABO blood group system) by means of the expression of functional human enzyme α1,2FT in transgenic pigs (Jura et al. 2004; Lipiński et al. 2010). Pigs carry an endogenous gene coding for α1,2-fucosyltransferase; however, the H epitope produced by this enzyme is not expressed on the endothelial surface of blood vessels (Oriol et al. 1993; Sharma et al. 1996). A high level of reduction in the synthesis of the Galα(1,3)Gal epitope observed in vivo on the surface of the endothelium is, hence, an effect of artificially generated competition between the endogenous α1,3GT and α1,2FT introduced experimentally. The analysis we conducted showed a significant reduction in Gal antigen on the surface of fibroblasts derived from transgenic animals, with additional copies of the α1,2FT gene introduced to their genomes compared to the number of epitopes occurring on the surface of cells in wild-type animals. The reduction of Gal antigen on the cell surface in transgenic animals is, hence, a result of H transferees, which competes with 1,3-galactosyltransferase for the acceptor molecule. It is precisely correlated with the observed, increased levels of H antigen (Jura et al. 2004; Lipiński et al. 2010).

We plan to interbreed the described transgenic animals and obtain the simultaneous expression of α1,2-fucosyltransferase.

Fig. 6 Survival rate analysis with standard deviations for cells from wild-type animal (blue), from F0 (TG252) with α-galactosidase expression (red), F1 animal (TG1183) with α-galactosidase expression (green) and TG1154 with the expression of α1,2-fucosyltransferase (purple).
and α-galactosidase. Through the combination of two modifications, we will try to obtain the cumulative effect of decreasing the level of Galα(1,3)Gal epitope. The research to date confirmed that GLA expression decreased the level of Galα(1,3)Gal epitope by 78%, and the coexpression of α-galactosidase and 1,2-fucosyltransferase eliminated it entirely from the surface of SV40-immortalised aortic porcine endothelial cells (PEDSV15) (Jia et al. 2004). The enzyme α1,2FT located within the medial compartment of the Golgi apparatus cisterns shows its activity earlier than the trans endogenous α1,3GT present in the trans compartment. The maturing oligosaccharide, when migrating through the Golgi apparatus from cis to trans compartments, cannot accept the terminal galactose in a reaction catalysed by α1,3GT because of the previous fucosylation of N-acetyllactosamine resulting from α1,2FT activity (Hartel-Schenk et al. 1991). The simultaneous transformation of COS cells with α1,2FT and α1,3GT coding genes showed the shift of the expression profile towards the production of H epitope (Sandrin et al. 1995). The expression of α1,2FT does not result in total Galα(1,3)Gal epitope elimination, so introducing additional copies of genes coding for α1,3GT would supplement the activity of 1,2-fucosyltransferase. Introducing gene coding for α-galactosidase removing the terminal α-D-galactose should result in a further reduction of the amount of functional Galα(1,3)Gal epitope formed as a result of competition between α1,3GT and α1,2FT. Given that the effect of GLA activity, when a solution of this enzyme is administered, is transient, the integration of several strategies preventing HAR is even more advisable. Regardless of the initial reduction in the binding of IgM antibodies, after ex vivo perfusion of pig kidneys with endo-β-galactosidase C solution, the resynthesis of Galα(1,3)Gal epitope on the cell surface and increased XNA reactivity are observed. Obtaining a proper therapeutic effect would be linked to the necessity of constant supplementation of the recipient with GLA solution (Ogawa et al. 2000). More recently, Alexion Pharmaceuticals produced a combined GTKO with H-transferase transgenic pig to “compete out” any potential antigen-producing transferrases, including the putative secondary galactosyltransferase (Ramsoondar et al. 2003).

Other strategies of preventing hyperacute rejection of xenotransplants conducted in our team involve enzymatic regulation of the complement system, which concerns blocking the formation of membrane-attacking complex, and inactivation of the α1,3GT gene responsible for the final Galα(1,3)Gal structure (unpublished data). The existence of genetic modifications ensuring the expression of possibly the highest number of regulators of the complement system and altering the distribution of sugars on the surface of pig cells is quite likely to contribute to both quantitatively and qualitatively limiting the use of pharmacological treatment, which weakens the immunological system by blocking its functions. Only animals with numerous genetic modifications obtained through the interbreeding of organisms with single modifications or through cotransfection with several gene constructs will become a reservoir of organs for medical purposes (Cooper 2012).

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