Enzymatic engineering of the porcine genome with transposons and recombinases

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Abstract

Background: Swine is an important agricultural commodity and biomedical model. Manipulation of the pig genome provides opportunity to improve production efficiency, enhance disease resistance, and add value to swine products. Genetic engineering can also expand the utility of pigs for modeling human disease, developing clinical treatment methodologies, or donating tissues for xenotransplantation. Realizing the full potential of pig genetic engineering requires translation of the complete repertoire of genetic tools currently employed in smaller model organisms to practical use in pigs.

Results: Application of transposon and recombinase technologies for manipulation of the swine genome requires characterization of their activity in pig cells. We tested four transposon systems—Sleeping Beauty, Tol2, piggyBac, and Passport in cultured porcine cells. Transposons increased the efficiency of DNA integration up to 28-fold above background and provided for precise delivery of 1 to 15 transgenes per cell. Both Cre and Flp recombinase were functional in pig cells as measured by their ability to remove a positive-negative selection cassette from 16 independent clones and over 20 independent genomic locations. We also demonstrated a Cre-dependent genetic switch capable of eliminating an intervening positive-negative selection cassette and activating GFP expression from episomal and genome-resident transposons.

Conclusion: We have demonstrated for the first time that transposons and recombinases are capable of mobilizing DNA into and out of the porcine genome in a precise and efficient manner. This study provides the basis for developing transposon and recombinase based tools for genetic engineering of the swine genome.

Background

Recent developments in livestock transgenesis, including somatic cell nuclear transfer (SCNT, cloning) [1], and stem cell biology [2,3] have energized plans to engineer the pig genome for both agricultural and emerging biomedical markets. Although pronuclear injection (PNI) and SCNT are proven methods for gene supplementation and gene targeting, respectively, more sophisticated meth-
ods for manipulating the pig genome have been lacking. Tandem gene targeting and SCNT provides a method for the precise introduction of transgenes or alternate alleles, but the inherent inefficiency of homologous recombination and donor-cell senescence limits its efficiency. Transgenesis by random integration of naked DNA has proven much more efficient for gene supplementation, whether using PNI or SCNT. However, random integration of naked DNA is often accompanied by transgene instability [4,5], transgene concatemerization [6,7], loss of transgene expression due to methylation [8-13], and short deletions, inversions and duplications at the site of transgene integration [14-25]. In addition, the lack of precision associated with random integration of naked DNA limits transgene manipulation and control post-integration.

DNA "cut and paste" transposons have been widely used for precise and efficient delivery of DNA expression cassettes into invertebrate and plant genomes. Over the past ten years, several DNA cut and paste transposon systems have been shown to function in vertebrate cells, including Sleeping Beauty (SB) [26,27], Passport (PP) [28,29], Tol2 [30,31], and piggyBac (PB) [32-34]. In addition, transposons have been used for germline transgenesis of fish [35-37], frogs [38-40], and mice [32,41-43] and for transgenesis of mouse somatic and embryonic stem cells [44-46]. It is noteworthy that although transposons function in a wide array of cell types, their efficiency can differ from species to species or even within various cell types of one species. The function and efficacy of vertebrate transposons in pig cells had not previously been examined. Demonstration that one or more transposon systems functions efficiently in porcine cells would provide a rationale for investigating their use in PNI and SCNT. In addition, the precision of transpositional transgenesis (TnT) provides a segue to the development of conditional expression systems for application in pigs and porcine cells.

Many genes have roles in multiple tissues and/or at multiple times during growth and development. Due to a requirement for strict regulation, global ectopic transgene-expression or gene-knockout will be an implausible approach for many targets. To overcome these limitations, binary systems based on transcriptional transactivation or DNA recombination have been developed and applied in model organisms for conditional gene-expression or silencing [47]. Although the tetracycline transcriptional activator system [48] has been demonstrated to function in transgenic pigs [49,50], recombinases have not. Cre and Flp recombinases catalyze a conservative DNA recombination event between two short recombinase recognition sites (RRS), loxP and FRT, respectively [51]. This results in the deletion or inversion of the DNA between two RRS-depending on their orientation. Deletion or inversion of sequences in transgenes can be used as genetic switches to activate or silence gene expression in specific cells, at particular times, or under prescribed conditions. Applications beyond conditional gene expression include the removal/recycling of selectable markers or transgenes [52] or chromosome engineering [53]. The successful application of recombinase technologies to porcine genetics requires the demonstration of Cre and/or Flp activity in porcine cells and the efficient delivery of RRS sites and recombinase-based expression vectors to the porcine genome.

In order to assess the utility of DNA transposons and recombinases for enzymatic engineering of the porcine genome, we tested four transposon systems and two recombinases. The SB, PP, Tol2, and PB transposon systems were able to function in cells derived from pig tissues and significantly improved the rate of transgenesis in vitro. Cre and Flp recombinases were capable of removing antibiotic selection cassettes in porcine cells and conditionally activating transgenes in porcine cells, demonstrating the potential for their applications to "leave no trace" and/or conditional porcine genetic engineering.

Results
Sleeping Beauty activity in porcine cells
To test the ability of the SB transposon systems to mediate transposition into the porcine genome, a transposon vector (pT2-FloxP-PTK) and a transposase expression vector (pKUb-SB11) were constructed (Fig 1A). The transposon vector encodes a puromycin-thymidine kinase (PuroATK, PTK) fusion protein [54] between the inverted repeats of the SB transposon system. The PTK cassette was flanked by both FRT and loxP sites so that it could be used as a substrate for testing both Cre and Flp recombinases (see below). Pig fetal fibroblasts (PFF) or porcine endometrial gland epithelium (PEGE) cells were transfected with the PTK transposon along with the SB expression vector, a vector encoding non-functional SB (pKUb-SBΔDDE), or a β-galactosidase expression vector (pCMV-β). After the transfection period, cells with integrations were rendered resistant to puromycin selection, and formed clonal cell colonies after 9–12 days. Clones were stained with methylene blue and quantified (Fig. 1B). The transposase catalyzed 2.5× (PFF) -10× (PEGE) more colony formation versus transfection with a non-functional transposase (ΔDDE) or β-galactosidase. This difference in the rate of clone formation corresponds to TnT versus the background rate of non-transpositional transgenesis.

Multiple transposon systems function in porcine cells
The success of the SB transposon system prompted investigation of three additional transposon systems. In addition to resteting the SB transposon system in PEGE cells, we also tested PP (an additional member of the Tc1 trans-
**Figure 1**

*Sleeping Beauty function in pig cells.* A) Diagrams of the DNA vectors transfected into pig cells. pT2-FloxP-PTK is the experimental SB transposon. The transposon is flanked by inverted terminal repeats (ITR). The puromycin phosphotransferase-thymidine kinase fusion protein (PTK) is flanked by recombinase recognition sites, FRT and loxP, for Flp and Cre, respectively. pKUb-SB11 is the source of transposase and is expressed from the ubiquitin promoter (Ub). pKUb-SBΔDDE is a non-functional version of transposase because of an internal deletion within the catalytic domain. pCMV-β functions as negative control. B) The colony forming ability of pT2-FloxP-PTK in pig fetal fibroblast (PFF) and porcine endometrial gland epithelium (PEGE) was determined by counting puromycin resistant colonies after plating 60,000 cells on 10 cm dishes when pT2-FloxP-PTK was co-transfected with pKUb-SB11 (+SB), pKUb-SBΔDDE (+SBΔDDE), or pCMV-β (+βgal). The addition of functional transposase (+SB) versus a non-functional transposase (SBΔDDE) or pCMV-β (βgal) was determined to be significant by analysis with an unpaired t-test (p-values < 0.000002).
poson family [55]), Tol2 (a member of the hAT transposon family [56]), and PB, the founding member of the piggyBac transposon family [57]). PEGE cells are one of a few immortalized pig cell lines available, transfected consistently (8–15%), and form tight non-migrating clonal colonies - essential characteristics for the colony forming assays performed. The PTK expression cassette was placed between inverted repeats corresponding to each transposon; pKT2P-PTK, pPTnP-PTK, pGTol2P-PTK, and pPBT-PTK, respectively (Fig. 2A). PEGE cells were co-transfected with each of these transposons along with their corresponding transposase expression construct; pKUb-SB11, pKC-Pts1, pCMV-Tol2, or pKc-PB, respectively. Each transposon vector was also co-transfected with pCMV-β to determine the background rate of non-transpositional integration. Transfected PEGE cells were placed under puromycin selection for 9–12 days, colonies fixed, stained, and enumerated. Again, transfection of PEGE cells with both components of the SB system (Fig. 2B) resulted in over 200 colonies per 60,000 plated cells, or about 3.3% of transfected cells based on an average 10% transfection efficiency. This represented a 13.5-fold increase over transfection without transposase. Similar enhancements to transgenesis were seen for all the transposon systems. PP produced an average of over 100 colonies per 60,000 cells; a 5-fold increase over transfection without transposase (Fig 2C). The inclusion of Tol2 transposase resulted in the generation of puromycin resistant colonies at a rate 21-fold over transfections without transposase (Fig 2C). The inclusion of Tol2 transposase resulted in the generation of puromycin resistant colonies at a rate 21-fold over transfections without transposase (Fig 2D), producing on average over 240 colonies per 60,000 cells. The PB transposon system (Fig. 2E) yielded an average of over 320 colonies per 60,000 cells (about 5% of transfected cells), representing a 28-fold increase over transfection without transposase.

**Molecular characterization of transposition**

Integration of DNA transposons produces target-site duplications upon integration into the genome. Analogous to SB and other Tc1 type transposons, the target site preference for PP is a TA dinucleotide. Target-site preference for the PB transposon is a TTAA tetranucleotide [33]. Integration of Tol2 results in a target-site duplication of eight bases but does not rely on specific primary sequence, instead targeting a characteristic local deformation of DNA [58]. Blocked linker-mediated PCR was used to clone junction fragments after transfection of PEGE cells with each transposon system. Characteristic integration footprints were observed for each transposon system (Fig 3). Junction sequences were compared to sequences in GenBank using BLAST [59]. Despite the small amount of contemporaneous porcine genome sequence available, some flanking DNAs of each transposon system were found to have high identity to the pig genome, in most cases in abundant repetitive elements. This demonstrates bona fide transposition into the porcine genome for each transposon class.

One characteristic advantage of transposase-mediated integration is the precise incorporation of one or more independently transposed gene expression cassettes, without adjacent plasmid vector. In order to observe representative integration events, DNA was isolated from 8 or 9 selected clones from each transposon and analyzed by Southern hybridization (Fig 4). Non-transposase mediated integrations, often head to tail concatamer repeats, have a predictable hybridizing fragment size following restriction enzyme digestion. However, transposon mediated events have unique DNA outside of the ITRs and therefore have unpredictable and varying fragment lengths. The enhancement of transgenesis by transposition (as detected by increased colony formation) was substantiated by the presence of inserts of varying size in cellular clones, in most cases without concatemers. The level of TnT can also be measured by counting the number of independent integrations per cellular clone. The more active transposons Tol2 and PB, display multiple (up to 15) independent integration events. The wild-type PP transposon system mediated a single integration event per cellular clone, reflecting its lower activity in PEGE cells, whereas the engineered SB system displayed an intermediate number of insertions.

**CRE/FLP activity in porcine cells**

To test the ability of Cre and/or Flp recombinase to function in porcine cells pT2-FloxP-PTK (Fig. 1A) was transfected into PEGE cells along with SB. These clones were obtained from preliminary transfections that were selected under very stringent drug conditions that favored high-copy integrations, particularly non-transposition events. DNA from puromycin resistant clones was isolated and analyzed by Southern hybridization. Isolated clones contained multiple copies of the PTK transgene due to non-transpositional integration, as indicated by concatemers and concatamer junction bands (Fig 5). PTK transgenic clones were subsequently transfected with pPGK-nlsCre, pKT2P-nlsFlp, or pKT2C-EGFP. Excision of the PTK cassette was detectable in transiently transfected cells by PCR, and the sequence of the excision product confirmed by sequencing (data not shown). Transfected cells were placed under selection with gancyclovir for 10–14 days and colonies counted (Fig. 5C). Only cells that had excised the PTK gene could withstand gancyclovir selection. As expected for concatemers, we observed a low level of transgene instability as evidenced by the appearance of gancyclovir resistant clones upon transfection with pKT2C-EGFP. A much more pronounced recombinase stimulated elimination of the PTK cassette was demonstrated by elevated resistant colony formation for 7 out of 8 of the clones transfected with either pPGK-nlsCre or
pKT2P-nlsFlp. While Cre and Flp are both active in PEGE cells, in all cases Cre mediated recombination/excision matched or exceeded that observed for Flp. A single clone (6) never showed evidence of PTK elimination. The Southern analysis (Fig. 5B), revealed a fragment of pT2-FloxP-PTK likely resulting from the integration of a shortened PTK expression cassette lacking at least one flanking RRS. This clipped PTK transgene is therefore unable to be removed by recombinase-mediated excision.

**CRE-activated gene expression**

To further demonstrate the functionality of the transposon-based Cre recombinase system for use in porcine genome engineering, a SB transposon containing a Cre-activated gene expression cassette was constructed- pTC-loxPTK-G (Fig. 6A). The PTK gene would be transcribed by the mini-CAGs promoter and efficiently terminated by three complete poly(A) signals (triple stop) in the intact pTC-loxPTK-G [60]. Cre recombination results in deletion
**Figure 3**

**Examples of transposon insertion junctions.** Transposon junctions amplified from PEGE cells are shown in groups of five with expected non-transposed vector sequence (lowercase) highlighted above. From top to bottom, SB (ITR-L), PP (ITR-R), Tol2 (ITR-L), and PB (ITR-R), and. Target site duplications (bold) for each transposon are separated from genomic DNA and corresponding (ITR) by a space.
Figure 4
Southern Blot of PEGE Clones. Individual puromycin resistant PEGE colonies were isolated and expanded for Southern analysis A) SB, B) PP, C) Tol2, and D) PB. Each transposon donor plasmid transfected into PEGE cells is diagramed with restriction endonuclease sites used for DNA digestion and the probe fragment indicated (diagonal lined rectangle). Expected concatemer sizes (vertical lined arrow) / smallest possible transposition event (open arrow) for each transposon are 5159/3335 bp, 5083/3275 bp, 6285/3346 bp, and 5140/3320 bp, respectively. The positions of the marker bands are indicated by black dots on the right of each blot with sizes of 12, 10, 8, 6, 5, 4, and 3 kb are shown.
of the PTK/triple-stop cassette, thereby juxtaposing the mini-CAGS promoter and the downstream gene expression cassette and enabling transcription of the GFP gene. Conditional activation of GFP expression was assessed by microscopy and flow cytometry after transient transfection of PEGE cells with pTC-loxPTK-G in the presence or absence of pPGK-nlsCRE (Fig. 6B). There was no GFP observed in cells transfected with pTC-loxPTK-G alone, whereas about 10–12% of the cells were GFP+ when transfected with pPGK-nlsCre. This corresponds well with the average transfection efficiency of PEGE cells, indicating that the Cre excision reaction is very efficient in transiently transfected cells.

To further examine the efficiency of Cre recombinase in transiently transfected cells, conditional removal of the PTK/triple stop expression cassette was assessed by selection in puromycin following co-transfection of PEGE cells with pTC-loxPTK-G and Cre, β-galactosidase, SB, or Cre + SB. Transfected cells were plated under puromycin selection for 9–12 days, stained with methylene blue, and enumerated to quantify the efficiency of PTK/triple stop elimination prior to or after integration into the genome (Fig. 6C). Addition of pPGK-nlsCRE to the transfection, alone or in combination with pKUb-SB11 reduced puromycin-resistant colony counts to levels significantly lower than that observed for pKUb-SB11 or pCMV-β, which alone result in TnT and non-transpositional transgenesis with an intact PTK gene expression cassette, respectively. Therefore, Cre recombinase excision activity in transiently transfected PEGE cells approaches 100%, especially with regard to plasmids available for transposition by SB transposase.

Although this particular co-transfection with pTC-loxPTK-G and SB suffered from a low transfection efficiency (≈5%) that reduced TnT (compare Fig 6C to 1B), puromycin resistant clones were expanded for characterization by Southern hybridization (Fig 7). Analysis indicated TnT with 1 to 4 transposon integrations per clone. Although,
Figure 6

A CRE-Activated Transgene. A) An illustration of the Cre-activated transgene vector. The full vector, pTC-loxPTK-G, produces PTK from the mini-CAGs promoter. Transcriptional leakage into the downstream gene, GFP, is limited due to the incorporation of three full poly-adenylation signals, a so-called triple-stop. Recombination by Cre eliminates PTK and triple-stop, activating GFP expression from pTC-lox-G. B) pTC-loxPTK-G was transfected into PEGE cells with (+Cre) or without (-Cre) pPGK-nlsCre. Cells were monitored for GFP expression by fluorescent microscopy (image inserts) and flow cytometry. The percentage of cells expressing GFP was dependent on co-transfection with pPGK-nlsCre. C) PEGE cells were transfected with pTC-loxPTK-G along with pPGK-nlsCre (+Cre), pCMV-β (+βgal), pKUb-SB11 (+SB), or pKUb-SB11 and pPGK-nlsCre (+SB +Cre). The cells were plated in puromycin selective media and colonies were counted.
clones 7, 10 and perhaps 11 contained hybridizing species near what would be expected for non-transpositional integration, their molar representation was equal to that of single copy inserts, not multicopy concatemers. Clones 7 and 10 also harbored hybridizing species smaller than was expected for transposition. These fragments likely represent non-transposase mediated DNA recombination events. The proportion of non-transpositional integrations detected by Southern analysis (1 in 4) corresponds well with the observed unfacilitated rate of transgenesis as determined by colony count for this transfection.

pTC-loxPTK-G clones were generated to analyze the efficiency of recombinase-directed selection-cassette recycling and the conditional activation of gene expression from a variety of porcine genomic loci. Puromycin resistant clones were transfected with pPGK-nlsCRE and scored for gancyclovir resistance (Fig 7C). All gancyclovir resistant clones expressed GFP, although variation in the intensity of GFP was observed (data not shown) depending on the parental clone source. This expression variance is expected due to the influence of porcine sequence adjacent to the sites of transposon integration, a phenomenon commonly referred to as "position effect". A significant increase in the efficiency of selection cassette recycling was demonstrated in the presence of single copy inserts when compared to multicopy concatemers (Fig 7C vs 5C). In addition, activation of GFP expression upon recombinase-based excision from integrated transposons demonstrates the efficacy of Cre-dependent conditional gene expression in transgenic porcine cells.

Discussion
Multiple transposons and recombinases are active in porcine cells

This work demonstrates for the first time the capability of four DNA transposon systems, SB, Tol2, PB, and PP, to enzymatically facilitate precise and efficient transpositional transgenesis in porcine cells. We have also established for the first time that Cre and Flp DNA recombinases are active in porcine cells. The combination of these DNA directed enzyme systems provides for the delivery and removal of gene expression cassettes to the porcine genome for the purpose of cellular transgenesis, selection cassette recycling and conditional gene expression based on transposons and recombinases. In these studies, the Tol2 and PB transposon systems were more efficient than SB, which was more efficient than PP at mediating TnT in PEGE cells, although these relative efficiencies should not be over-interpreted. Although we used favorable conditions for each transposon system by our selection of promoters and transposase/transposon ratios, our focus here was on testing their function, not on determining their relative activities in PEGE cells, an immortalized cell line unsuitable for generating pigs by SCNT.

Indeed, it is well established that the rate of TnT in any cell type is likely to depend not only on intrinsic transposase activity, but also on the presence or absence of cellular cofactors and DNA repair enzymes, the method of DNA introduction, and the amount of transposase produced/provided in the specific cell type. Transposon activity varies not only between cells from different species, but also between different cell types from the same species [26,27,34]. Future studies will focus on the efficiency of different transposon systems and recombinases in pig fibroblasts (applicable to SCNT), pig stem cells (for functional genomics and SCNT), and porcine embryos (for direct transgenesis by PNI).

In addition to potential differences in efficiency, the integration behavior of each transposon may be an important factor in determining the appropriate transposon system for a specific task. For instance, PB appears to preferentially integrate into transcription units [32,61]. Consistent with this observation, in our limited examination of integration sites in the pig genome, flanking sequence from two of seven PB insertions matched porcine cDNAs. In addition, PB primarily leaves no footprint when remobilized [33]. Therefore, PB may be most suitable for functional genomics studies in pigs or pig cells, where mutations due to the interruption of genes, and the potential for precise transposition-based rescue is desirable [62-65]. SB does not integrate into transcription units at a rate much higher than what would be expected by random integration [66], so it may represent a better choice for animal transgenesis, transposon-based DNA vaccination, or other somatic therapies. Alternatively transposon systems engineered to target specific genomic locations may be developed and could provide the safest choice for these applications [34,67,68]. The integration profiles of Tol2 and PP are not well characterized in any organism or cell type, and the integration predilections of any transposon system remains to be addressed in specific swine cells being considered for engineering.

Advantages of transposition for pig transgenesis and genetics

There are several advantages of transpositional versus unguided transgenesis. First, the enzymatic activity of the transposase increases the efficiency of transgene integration (Fig. 1 and 2). Secondly, transposase-mediated transgenesis precisely integrates a single copy of the transposon into one or more locations in the genome. Consequently, transposition avoids the integration of G/C-rich prokaryotic elements of the vector and avoids transgene concatemerization, both of which can lead to shutdown of gene expression [5,6]. In addition, concatemerization is problematic for selection cassette recycling (Fig. 5) and the implementation of more complex genetic rearrangements with recombinases. We propose the use of transposon sys-
Figure 7

Conditional gene-activation of integrated transposons. Colonies from the transfection of pTC-loxPTK-G with pKUb-SBII (Fig 5C) were expanded in selective media containing puromycin. DNA from these transgenic colonies was isolated and analyzed by Southern hybridization. A) A schematic of pKT2C-loxPTK-G that shows the Asel restriction sites and the location of the PTK hybridization probe (diagonal lined rectangle) used for Southern analysis. B) A Southern blot of pKT2C-loxPTK-G colonies. The clones were analyzed without Cre excision, so integrants that result from transposition should be equal to or greater than the transposon size of 4.9 kb (open arrow). Whereas, bands associated with concatemer formation are found at 6.0 kb (vertical line arrow). The positions of the DNA marker bands of the 1 kb Quanti-Marker from ISC Bioexpress (Kaysville, Utah), are indicated by black dots on the right of each blot with sizes of 12, 10, 8, 6, 5, 4, 3, 2.5, and 2 kb shown. C) pKT2C-loxPTK-G colonies were transfected with pPGK-nlsCre and plated under gancyclovir selection. Clones with PTK eliminated by recombination became gancyclovir resistant and were counted. Cre-activation of all clones was determined to be significant (p < 0.5).
tems for transgenesis of porcine cells prior to their use for the creation of pigs by SCNT to enable increased efficiency, better precision, reliable expression, and selection cassette recycling. In addition, SB and PB dramatically improved the transgenesis rate in mice by PNI [32,69], providing a clear rationale for improving the efficiency of transgenic pig production via this method.

Recombinases in swine genetics - selection-cassette recycling and conditional alleles

There are several immediate applications for recombinases in swine genetics. First, as shown in Figs 5, 6, 7, recombinases can be combined with a positive/negative selectable marker like PTK for selection cassette recycling [52]. Currently, most, if not all transgenic animals produced by SCNT contain a selectable marker (e.g. neoR, puroR, GFP) in addition to an experimental transgene. This selectable marker is useful for the proper identification of nuclear donor cells, but generally is undesirable in the transgenic animal. This could be particularly important for removal of xenogenic elements after gene knock-out or manipulation preceding the introgression of engineered germplasm into agricultural production herds. The flanking of selectable markers with RRS provides the opportunity to eliminate them in culture or by breeding to Cre expressing pigs, leaving behind only a single 34-basepair RRS footprint.

Recombinases also permit the creation of conditional alleles for activation or inhibition of gene function in response to Cre or Flp recombinase activity, as illustrated in Figures 6, 7[51]. In addition, the effectiveness of homologous recombination constructs can be improved to allow selection cassette recycling, thereby avoiding 'selection cassette interference', whereby the exogenous regulatory elements in the selection cassette can interfere with the expression of genes in the vicinity of the targeted mutation [52]. As has been elegantly demonstrated in mice, recombinases can also be used to create conditional knock-outs in pigs when tissue specific ablation is desired, or when traditional knockout results in embryonic lethality. The availability of an assay of transposon and recombinase systems should also permit serial cellular transgenesis and recombination to achieve complex genomic rearrangements in the pig. Serial transgenesis provides a direct method for the production of pigs that express several gene products. Given the dramatic long-range conservation of synteny between pig and human genomes, far more extensive than for mouse and human, engineered chromosomal rearrangements between serially provided RRS in the pig could provide superior large animal models of human congenital and cancer related chromosomal abnormalities [53].

Conclusion

Pork represents the single most economically important meat product in the world and pigs are playing an increasingly critical role in biomedicine. An armamentarium of effective genetic tools will be required to capture the value and potential of this species for human nutrition and health. Here we have tested four transposon and two recombinase systems for activity in pig cells. SB, PP, Tol2, and PB and transposons are capable of precise transpositional transgenesis of porcine cells, increasing efficiency by 4–28 fold. We have also demonstrated that Cre and Flp recombinases function efficiently in the nucleus of pig cells for selection-cassette recycling and conditional regulation of transgene expression. The combination of these tools will significantly improve the efficiency and sophistication of porcine genetic manipulation for enhancing pig production and human nutrition, as well as modeling and treating human disease.

Methods

Vector construction

Sequence information, maps, and material requests for these constructs can be found on our web site [70].

\textbf{pT2-FloxP-PTK-} To generate a multiple cloning sequence flanked by \textit{FRT} and \textit{loxP} recombinase recognition sequences (\textit{FRT-loxP MCS}), two oligonucleotides with overlapping sequence (shown in bold) were designed, \textit{FRT-loxP Upper} [ATACCGGCGAGGTTCCCTATCCCAAGTTCCATTTCTCT AGAAAATGATAGAAATCATAAATGATATGCTATACGAGATCAGAATTCGCCGGAG-GCCCTACTAGT], and \textit{FRT-loxP Lower} [GTATTCATGAGAATGCTATACCTTTCAGAGAAATTAGG AAATCGGAAATAGAATCATAAATGATATGCCTATACGAGATCAGAATTCGCCGGAGGG CCGGAAA]. These oligonucleotides were annealed and elongated by PCR using Pwo polymerase. The 218 base pair PCR fragment was cloned into pCR4 using the ZERO Blunt TOPO PCR Cloning Kit (Invitrogen, USA) to create pCR4 \textit{FRT-loxP MCS}, and its sequence was verified. \textit{FRT-loxP MCS} was subsequently excised with EagI and BspHI and cloned into pT2/BH [71] cleaved with EagI and NcoI to produce pT2-\textit{FRT-loxP MCS}. Finally, a completely filled XhoI fragment, containing the mouse PGK promoter, the PTK fusion protein, and bovine growth hormone poly(A) signal from YTC37, a kind gift from the laboratory of A. Bradley [54], was cloned into Sim1 cleaved pT2-\textit{FRT-loxP MCS} to produce pT2-FloxP-PTK.

\textbf{pKUb-SB11-} A 1.0 kb fragment of the SB11 transposase from pCMV-SB11 [72], which had been amplified with CDS-SB11-F1 [CACCATGGGAAATCAGAATTCAGCCT] and CDS-SB11-R1 [GGATCCCAATTTAAG-GCAATGCTACCAAATACTAG] primers and subcloned
into an intermediate vector adding a 5' BglII site and the sequence [AGATCTGAT], was cloned into the BamHI site of pKUb to make pKUb-SB11. pKUb was made by cloning nucleotides 3561–4771 of the human UbC gene (genbank accession D63791), which contains the UbC promoter, non-coding exon 1, and intron 1, into pk-SV40(A) between intact BglII and NheI restriction endonuclease sites. p-SV40(A) was made by cloning a single copy of the SV40 poly(A) signal amplified by PCR with oligos KJC-SV40(A)-F1 [CATGATTGAGTTGGACACACACACCA] and KJC-SV40(A)-R1 [ACCACATTTGAGGTTCATTTGC] into pk-A10 opened with XmnI. pk-A10 was made by cloning KJC-Adapter 10 [CTGATCTTAAGCTAGCACCGATCCAGAACATTCATTGCAATCC] into pk digested with PvuII creating a multiple cloning site with PvuII, BglII, AflII, NheI, BamHI, EcoRI, XmnI, and PvuII recognition sites. pk was made by joining an 0.8 kb PCR product of pBluescriptSK- (Stratagene), containing the pUC-ORI amplified with oligos KJC-pUC-ORI-F1 [CTGTTCCGCT-TCCAGGCTCATGACT] and KJC-pUC-ORI-R1 [AAAAGGACATGAGTTGAAGATCCATTTC] to a 0.9 kb PCR product of pEFR-D-TOPO (Invitrogen), which contains the kanamycin resistance gene amplified by oligos KJC-KanR-F1 [CTGCATCATGAACAATAAAACTGTCTGCT] and KJC-KanR-R1 [TTCCGCTCACTGACTGAGTTTGGACAAACCACA]. The junction of ORI-F1 to KanR-R1 created a single blunt end GTTAACTT to TTGAACTCTAGA including a modified sequence at the 3' end GTTAACTT to GTTAACTGCTACATTGCAAAATTTCATTTAAAT creating a small multiple cloning site with PvuII, BamHI, BglII, KpnI, and Swal sites. pk-PK-PTK-ΔA was constructed as indicated [74] from previously described materials [75].

pk-cr-Tol2-PTK was made by cloning a 2.7 kb PvuII fragment from pKP-PTK-ΔA into pk-cr-Tol2-PTK opened from EcoRI and BglII. cr-Tol2-PTK was made by cloning a 2.3 kb BglII fragment from pSBT-mCAG [73] into pK-A10 opened from XmnI. pk-Tol2-PGKΔ was made by cloning EGFR as a 1.4 kb PCR product of pBT-mCAG opened from BglII to KpnI fragment of pSBT-mCAG [73] into PK-A10 opened from EcoRI and BglII. PK-Tol2-PTK-ΔA was made by cloning a 2.3 kb PvuII fragment from pSBT-PTK-ΔA into pk-cr-Tol2-PTK opened from EcoRI and BglII. cr-PK-PTK-ΔA was made by cloning a 2.3 kb BglII fragment from pSBT-mCAG [73] into pk-cr-Tol2-PTK opened from EcoRI and BglII. cr-PK-PTK-ΔA was made by cloning a 2.3 kb BglII fragment from pSBT-mCAG [73] into pk-cr-Tol2-PTK opened from EcoRI and BglII. cr-PK-PTKΔ was made by cloning a 2.3 kb BglII fragment from pSBT-mCAG [73] into pk-cr-Tol2-PTK opened from EcoRI and BglII. cr-PK-PTKΔ was made by cloning a 2.3 kb BglII fragment from pSBT-mCAG [73] into pk-cr-Tol2-PTK opened from EcoRI and BglII. cr-PK-PTKΔ was made by cloning a 2.3 kb BglII fragment from pSBT-mCAG [73] into pk-cr-Tol2-PTK opened from EcoRI and BglII. cr-PK-PTKΔ was made by cloning a 2.3 kb BglII fragment from pSBT-mCAG [73] into pk-cr-Tol2-PTK opened from EcoRI and BglII. cr-PK-PTKΔ was made by cloning a 2.3 kb BglII fragment from pSBT-mCAG [73] into pk-cr-Tol2-PTK opened from EcoRI and BglII. cr-PK-PTKΔ was made by cloning a 2.3 kb BglII fragment from pSBT-mCAG [73] into pk-cr-Tol2-PTK opened from EcoRI and BglII. cr-PK-PTKΔ was made by cloning a 2.3 kb BglII fragment from pSBT-mCAG [73] into pk-cr-Tol2-PTK opened from EcoRI and BglII. cr-PK-PTKΔ was made by cloning a 2.3 kb BglII fragment from pSBT-mCAG [73] into pk-cr-Tol2-PTK opened from EcoRI and BglII. cr-PK-PTKΔ was made by cloning a 2.3 kb BglII fragment from pSBT-mCAG [73] into pk-cr-Tol2-PTK opened from EcoRI and BglII. cr-PK-PTKΔ was made by cloning a 2.3 kb BglII fragment from pSBT-mCAG [73] into pk-cr-Tol2-PTK opened from EcoRI and BglII.
CCTATATGTAGTCGTATTA] and T3-RevComp [TCTCTCCTTATGTAGGGTTATTA] primers. pPBT was made by cloning the PB LTR1 and LTR2 into pKT2-SE from KpnI to BamHI. LTR1 and LTR2 from PB were amplified from pKC-PB, a kind gift of Malcolm Fraser (Notre Dame University), using PB-LTR1-F1 [TGGATCCCAATCCITTATCTACATGACCGAGTA- CATG] and PB-LTR2-F1 [TGGATCCCAATCCITTATCTACATGACCGAGTA- CATG] and PB-LTR2-R1 [TGGTACCTAGTTAACCCTAGAAG- CATG] and PPTN-TK-F2 [TTAGCTACATGACCGAGTA- CATG] and PPTN-TK-R1 [ATAACTTCGTATACTACGTA- CATG] into pKT2C-EGFP (Invitrogen). pPBT was made by cloning a 1.0 kb SmaI to EcoRI fragment of p3XP3-DsRed, a kind gift of Dr. Malcolm Fraser, University of Notre Dame, containing the PB transposase coding sequence.

pKUb-PTs1 was made by replacing the SB11 gene from pKUb-SB11 with PTs by cloning a 1.0 kb BamHI to NheI fragment from pCR4-PPTs1B into pKUb-SB11 from NheI to BamHI. pCR4-PPTs1B was made by cloning a PCR fragment of pBluKS-PPTN4 [29], a kind gift of Dr. Michael Leaver (University of Stirling, UK), amplified with primers CDS-PPTs-F1 [AAAGCTAGCATGAACAGCAGGATCT- CACC] and CDS-PPTs-R1 [AAAGCTAGCATGAACAGCAGGATCT- CACC] into pCR4-TOPO (Invitrogen).

pKT2C-loxPTK-G was made by cloning a 2.3 kb PvuII fragment of pK-PPT-TS into the MscI site of pKT2C-lox-GFP. pK-PPT-TS was made by cloning a 1.9 kb BglII to EcoRI fragment of pCR4-PPTK into the MCS of pSV40(×2) opened with EcoRI and BglII. pCR4-PPTK was made by cloning a 1.9 kb PCR product of pT2-FloxP-PTK using oligos PuroΔTK-F2 [TTAGCTACATGACCGAGTA- CATG] and PuroΔTK-R1 [TTAGCTACATGACCGAGTA- CATG].

**Cell Culture and transposition/recombinase assays**

Pig fibroblasts were isolated from 43 day old embryos. The tissue was dissociated using a collagenase/DNase I treatment as well as mechanical disruption. The cells from the female piglet #8 were cultured in DMEM enriched with 10% FBS and 2× antibiotic/antimycotic solution (Gibco #15240-022). The cells were passaged in DMEM high glucose media enriched with 10% FBS, 2 mm L-glutamine, 1 × P/S until spontaneously establishing line FP8. A subpopulation of porcine endometrial gland epithelium cells [76] were spontaneously immortalized, strain PEGE. The PEGE cells were maintained in DMEM supplemented with 10% FCS, 1 × Pen/Strep, 10 μg/ml Insulin (Sigma, USA), and 1 × L-Glutamine.

For transposition assays cells were plated in each well of a six well plate to achieve 60–80% confluence within 6–24 hours. Cells were transfected using TransIT-LT1 (Mirus Bio Corporation, WI) transfection reagent according to the manufactures instructions with a ratio of 3:1 lipid:DNA. Each transfection contained a total of 1.15 to 1.5 μg of plasmid DNA. Wells 1–3 contained transposon plus transposase, well 4 contained transposase with no transposase, well 5 contained SB plus SB transposase and well 6 contained pKT2C-EGFP only. Molar amounts of each transposon were fixed at 1.5 × 10−13 moles of transposon product.
(0.75 × 10⁻¹³ Moles for Tol2) while transposase plasmid was added at a molar ratio of 1:1 for SB, Tol2, and PB, and 1:0.5 for PP. The choice of the promoters and transfection ratios for SB and PP was based on the highest transposition activity observed in human HT1080 cells (data not shown). Strong promoters (CMV & miniCAGs) and transfection conditions for Tol2 and PB were selected based on previously published data and the observation that these transposon systems seem less susceptible to overexpression inhibition than SB and PP.[34,61,74] Total DNA weight was adjusted using pCMV-β plasmid. Forty-eight hours after transfection cells were trypsinized, and two replicates of 60,000 cells were plated onto 100 mm plates in media containing 0.3 μg/ml puromycin and selected for 9–12 days. Colonies were visualized by methylene blue staining and counted. A minimum of two six-well plates were transfected for each experiment. The mean colony number and standard error are shown in figures.

Southern hybridizations
Several independent puromycin resistant PEGE foci for each transposon were aspirated and grown to confluence on a 100 mm plate. Genomic DNA was extracted using standard methods and approximately 10 μg was digested with SspI (Tol2 clones) or AseI (SB, PB, and PP) clones. Digested DNA was separated on 0.7% agarose gel and transferred to positively charged nylon membranes (GE Osmotics, USA). Membranes were probed with a random primed 1524 bp Xmal fragment of pKP-PTK-TS that contained the bulk of the PTK gene and visualized by autoradiography or phosphor imaging.

Cloning transposon junctions
Genomic DNA was isolated from pooled, fixed, and stained puromycin resistant clones for each transposon. For splinkerette PCR DNA was cut with Sau3AI or NlaIII and junctions were cloned as described [69]. For blocked linker-mediated PCR, DNA was cut with NspI for Tol2 and SB, and a cocktail of enzymes including XbaI, AvrII, NheI and Spel for PB and PP. The NspI digested DNA was ligated to the blocked linker-SphI that was created by annealing primerette-long [CCTCCACTACGCTACTCAGGCGAAGCTGCTTACAAACCATG] and blink-SphI [5’-GTTGTAGGACTGCTTGC-3’]. Whereas the DNA digested with the cocktail was ligated to the blocked linker-XbaI that was produced by annealing primerette-long to XbaI [5’-CTAGCATGTTGTAGGACTGCTTGC-3’]. Following ligation the junction sequences were amplified by nested PCR. The primary PCR used the common primer primerette-short [CCTCCACTACGCTACTCAGGCGAAGC] with transposon-specific primers SB-IRDR(L)-O1 [ATTTTCAAGCTTTAAAGGCACAGTCAAC], Tol2(L)-O1 [AATTAAACCTGGCATCGCAGAAT], and PB-LTR(R)-O1 [ACAGACCGATAAACACATGGCTCAA], and PTn-IRDR(R)-O1 [GGGTGAATACTTATGCAACCCACAAGATG]. The secondary PCR reactions used the common primer primerette-nested [GGGCAAGCGAGCTCTAACAACCATG] with transposon-specific primers SB-IRDR(L)-O2 [GACTTGTGTCATGCAAAAGTAGATTGCTCT], Tol2(L)-O2 [GCGAAITCAATTGTTGCGTAATTGC], PB-LTR(R)-O2 [TCTCTAATGACAGCGACGGATC], and PTn-IRDR(R)-O2 [CAGTACATAATGCGAATAAGTCCAGGG]. To generate unique sequences serial dilutions (1:50 and 1:500) of the ligation reaction were used as template for the primary PCR. The primary PCR was diluted 1:50 and used as template in the secondary PCR reaction. The PCR fragments were shotgun cloned and sequenced.

Abbreviations
PEGE Porcine endometrial glandular epithelium
GFP Green fluorescent protein
RRS Recombinase recognition site
SCNT Somatic cell nuclear transfer
PNI Pronuclear microinjection
ITR Inverted terminal repeat
SB Sleeping Beauty
PP Passport
PB piggyBac
TnT transpositional transgenesis

Authors’ contributions
KJC designed experiments and transposon vectors and together with DFC constructed transposons, performed experiments, and conducted molecular analysis. LKF optimized and conducted tissue culture experiments. BWK performed preliminary transfections in pig cells and was mentored by DNF. SCF conceived the study and mentored KJC, DFC, and LKF in experimental design and data analysis. KJC drafted the manuscript and along with DFC and SCF completed manuscript preparation. All authors read and approved the final manuscript.

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