Progress toward generating a ferret model of cystic fibrosis by somatic cell nuclear transfer
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Abstract
Mammalian cloning by nuclear transfer from somatic cells has created new opportunities to generate animal models of genetic diseases in species other than mice. Although genetic mouse models play a critical role in basic and applied research for numerous diseases, often mouse models do not adequately reproduce the human disease phenotype. Cystic fibrosis (CF) is one such disease. Targeted ablation of the cystic fibrosis transmembrane conductance regulator (CFTR) gene in mice does not adequately replicate spontaneous bacterial infections observed in the human CF lung. Hence, several laboratories are pursuing alternative animal models of CF in larger species such as the pig, sheep, rabbits, and ferrets. Our laboratory has focused on developing the ferret as a CF animal model. Over the past few years, we have investigated several experimental parameters required for gene targeting and nuclear transfer (NT) cloning in the ferret using somatic cells. In this review, we will discuss our progress and the hurdles to NT cloning and gene-targeting that accompany efforts to generate animal models of genetic diseases in species such as the ferret.

Introduction
Until recently, the generation of gene-targeted animal models has primarily relied upon homologous recombination following direct introduction of transgenes into embryonic stem cells (ES cells). While this technique has been successful for animal modeling in the mouse, it has thus far proven significantly more difficult in larger species. To date, the most exciting and promising research in transgenesis involves the use of fetal and adult somatic cells to produce genetically identical animals through nuclear transplantation \([1,2]\). Successful production of cloned animals derived from somatic cells was first demonstrated in sheep \([3,4]\) and has more recently been demonstrated in mice \([5]\), cattle \([6]\), goats \([7]\), pigs \([8]\), cats \([9]\), rabbits, \([10]\) and mules \([11]\). Transgenic calves \([12]\), gene-targeted sheep \([13]\), and \(\alpha\)-1,3-galactosyltransferase knockout pigs \([14,15]\) have also been obtained by nuclear transfer from somatic cells. These successes have made animal modeling using nuclear transfer in less-studied species, such as the ferret, more feasible. Since somatic cell nuclear donors can be easily maintained \(in vitro\) and readily targeted for gene mutations, somatic cell-based embryo cloning is undoubtedly the future method of choice for generating genetically modified larger animals. Two major steps are required to clone genetically defined animals (Figure 1). First, gene targeting must be achieved in a somatic cell type appropriate for nuclear cloning, and karyotypically normal clonal cell lines must be isolated. Second, the nucleus from the mutant somatic cell must be used to reprogram enucleated recipient oocytes. The reconstructed embryos must then be implanted in foster mothers to generate cloned genetically defined animals.
CF is a recessive inherited genetic disease in the Caucasian population, with a frequency of about 1 in 3,000 newborns [16]. CF is caused by a defect in an epithelial chloride channel called the cystic fibrosis transmembrane conductance regulator (CFTR) [16]. CF patients suffer from recurrent bacterial infection in the lung, leading to bronchiectasis, compromised lung function, and ultimately death. Substantial efforts have been made to generate mouse models capable of reproducing the lung pathology seen in CF patients. However, due to differences in lung biology between mice and humans, CFTR-deficient and mutant mice do not develop spontaneous lung disease as seen in humans [17,18]. This lack of appropriate CF animal models has hindered progress in the development and testing of therapies for this disease.

The domestic ferret, Mustela Putorius Furos, has proven to be an excellent animal model for studying CFTR lung biology. Several aspects of ferret lung biology make this species an attractive model for CF lung disease. First, in contrast to mice, the ferret has marked similarities to humans in terms of lung physiology, airway morphology, and cell types [19–24]. Second, the expression of CFTR in the ferret airway epithelium and submucosal glands is identical to that in humans [25,26]. Third, amino acid identity between ferret and human nucleotide binding domain 1 (NBD1) of CFTR is a striking 97% [25], which is just as high as for non-human primates (96%, Macaca nemestrina) and is significantly higher than for rodents (80%, rat and mouse). Fourth, the ferret had been a useful model for viral and bacterial lung infections seen in humans [27–32]. Lastly, the ferret, with a gestation period of 42 days and 6 months to sexual maturity, has obvious advantages over larger species for animal modeling.

Strategy and Progress in Cloning a CF Ferret
Unlike sheep and cattle, for which NT cloning procedures are well established, cloning in the ferret presents numerous challenges since experimental parameters for embryo manipulation have not been defined. Over the past few years, several of the critical steps required for NT cloning in the ferret have been established [33–35]. This review will focus on progress in the various steps highlighted in Figure 2. Furthermore, we will discuss approaches currently under investigation to facilitate efficient gene targeting in somatic cells, a process required to ultimately clone a CF ferret.

In Vitro Culture of Ferret Embryos and Successful Production of Offspring from Implanted Embryos
A critical first step toward genetic manipulation in a new species is defining the parameters required for embryo culture and adoptive transfer into pseudopregnant recipient females. Optimal superovulation in the ferret (19.3 ± 0.6 oocytes and/or embryos per female) was achieved by a combination of hormonal injections, including 100 IU eCG and 150 IU hCG at 72-hr intervals [33]. This ovulation rate is more than double that induced by mating. Mating with a male immediately following hCG injection did not significantly alter the ovulated number of oocytes and/or embryos, indicating that mating is not required for superovulation in ferrets. Of embryos harvested at the one-cell stage, 64.5% and 47.1% developed into blastocysts when cultured in vitro in CZB or TCM-199 plus 10% fetal bovine serum (FBS) media, respectively. In contrast, only 17.1% of embryos cultured in vitro in NCSU-23 developed to the blastocyst stage. Both freshly retrieved and in vitro cultured embryos from cinnamon coat–colored parents produced live young when transferred at the 8-cell stage into albino coat–colored, pseudopregnant recipients. The percentage of kits delivered relative to embryos transferred was 61% when freshly retrieved embryos were used and 32% when in vitro cultured embryos were used. The most important achievement of these studies was the successful birth of cinnamon coat–colored pups from an albino female. These results demonstrate successful embryo transfer in the ferret and open the door to animal modeling with this species following embryo manipulation [33].

Conditions for In Vitro Maturation and Parthenogenetic Activation of Ferret Oocytes
One of the most critical and difficult steps in successful cloning of any species is the reprogramming of oocytes to
divide in the absence of fertilization. Optimization of this step, termed parthenogenetic activation, is very empirical, and it varies greatly between species. To increase the yield of oocytes for NT cloning, ovaries are often harvested and immature oocytes extracted for use. This method requires a step involving in vitro maturation (IVM), which must also be optimized prior to activation. The success of activation is closely linked to the quality of IVM oocytes. To this end, we have optimized conditions for in vitro maturation and parthenogenetic activation of ferret haploid oocytes.

Immature oocytes (cumulus-oocyte complexes) harvested from ovaries of superovulated ferrets have been evaluated for in vitro maturation conditions using several types of media [34]. The optimal media for maturation of ferret oocytes consisted of TCM-199 containing 10% FBS, 10 IU/ml of eCG, and 5 IU/ml of hCG. In this media, the maturation rate of ferret oocytes was 72% at 24 hrs of IVM. Optimization of oocyte activation was evaluated using both electrical and chemical stimuli individually or in combination. Treatment with cycloheximide (5 μg/ml, 5 min) and 6-dimethylaminopurine (6-DMAP, 2 mM/ml, 4 h) following electrical stimulation (an alternating current pulse of 3 V for 5 secs, followed by one direct current pulse of 180 V/mm for 30 μsec) resulted in 43% of the oocytes developing to the blastocyst stage. Such an activation rate represented a significant improvement over those obtainable under other tested conditions, including individual treatment with electrical pulses (10%),

Figure 2
The important steps involved in somatic cell nuclear cloning are shown schematically. Progress in each of the boxed methodologies will be discussed in detail in subsequent sections [33–35]. Italicized labels indicate descriptive markings, not methodological steps per se.
cycloheximide (3%), or 6-DMAP (5%). Blastocysts derived from in vitro activation appeared to be normal morphologically and were composed of an appropriate number of both inner cell mass (10.3 ± 1.1) and trophectoderm (60.8 ± 2.9) cells when they were examined using a technique [36] that differentially stains the inner cell mass (ICM) and trophectoderm (TE) layer [34].

Developmental Capacity of Ferret Embryos by Nuclear Transfer Using G0/G1-Stage Fetal Fibroblasts

With the ultimate goal of establishing experimental protocols necessary for cloning ferrets, we have begun to examine parameters for ferret cloning by nuclear transfer using G0/G1-stage donor fetal fibroblasts [35]. Cumulus-oocyte complexes were harvested from ovaries of superovulated ferrets and cultured in a maturation medium for 24 hrs. Following removal of cumulus cells, oocytes with normal morphology, uniform cytoplasm, and a first polar body were enucleated with a 25 µm (inside diameter, ID) glass pipette. Critical to successful embryo reconstruction by nuclear transfer is the complete removal of oocyte chromosomal DNA. Fluorescent DNA dye (such as Hoechst 33342) can be used to practice and evaluate enucleation efficiencies. However, these stains cannot be used during cloning since they affect the viability of the reconstructed embryo. Since the removal of too much of the oocyte cytoplasm also leads to poor cloning efficiencies, it is critical that the individual performing the work optimizes this step. The proportion of completely enucleated ferret oocytes was 80.8% ± 2.6 (n = 82) in the studies discussed below.

Following optimization of enucleation procedures, ferret fetal fibroblast cells (serum-starved for 14–16 hrs prior to NT) were injected directly into enucleated oocytes with a 10 µm (ID) PiezoDrill glass pipette. Reconstructed embryos were then activated by a combination of electrical pulses and chemical stimulations. Subsequently, the reconstructed and activated embryos were either cultured in vitro or transferred to pseudopregnant ferrets to evaluate their development capabilities in vitro and in vivo. Our results demonstrated that 56.3% of reconstructed embryos cleaved, while 26.0% and 17.6% developed to morula and blastocyst stages in vitro, respectively [35]. The blastocysts derived from NT embryos demonstrated normal morphology by differential staining and also contained cell numbers appropriate for normal blastocysts developed in vitro. In vivo developmental studies at 21 days post-transplantation demonstrated 8.8% of reconstructed embryos implanted into the uterine lining of recipients, while 3.3% formed fetuses (Figure 3). However, reconstructed embryos failed to develop to term (42 days). These results demonstrate that donor nuclei of G0/G1-stage fetal fibroblast cells can be reprogrammed to support the development of reconstructed ferret embryos in vitro and in vivo [35].

One facet of cloning that has been highly variable between species is the narrow window of post-mating uterine development that is receptive for implantation of NT-reconstructed embryos. These differences are due to the developmental delay caused by NT reconstruction of embryos. To solve this problem, researchers have utilized asynchronous breeding schedules of the oocyte donors.
and the female recipients used for implantation of NT-reconstructed embryos [10]. Current efforts attempting to clone ferrets to live birth are optimizing asynchronous breeding schedules to improve survival in vivo.

Allele-Specific Targeting of Single Base-Pair Changes in Somatic Cells with Chimeric RNA/DNA Oligonucleotides

In addition to embryo manipulation procedures in the ferret, efficient strategies for introducing mutations into the ferret CFTR gene must be developed in order to generate appropriate somatic cell donors for NT cloning. Several criteria are important when considering the appropriate targeting strategy for somatic cells. First, the strategy cannot be dependent on expression of the target gene (i.e., gene targeting using promoter-trapping) since the CFTR gene is not expressed in fibroblasts. Second, since low-passage somatic cells are preferred for NT cloning, the targeting strategy should be efficient and optimally not require selective pressure. Third, a highly sensitive and efficient screening method must be developed to isolate gene-targeted clonal cell lines at early passage. One advantage of NT cloning for generating genetically defined animal models is that theoretically, relatively few somatic cell donors are required to carry out the procedure. Hence, we have focused on developing methods based on nuclear injection of gene-targeting agents, followed by high-throughput screening of single-cell clones in 96-well plates for successful gene alterations. One suitable targeting agent we are currently evaluating utilizes chimeric RNA/DNA oligonucleotides (also called chimeraplasts).

To establish our ability to target the CFTR gene and rapidly screen large numbers of clonal cell lines, we began evaluating chimeraplasts in cell systems already established to target at high efficiencies. **Chimeraplasts** are self-folding RNA/DNA duplexes that create hairpin ends with flanking homology (10–30 bp) to the target base. Chimeraplasts have been reported to introduce single base-pair changes at the β-globin locus at efficiencies between 1 and 5% in Hela cells [37] and as high as 10% in other systems [38,39]. Since chimeraplasts are most efficient at introducing single base-pair alterations, we have evaluated the ability of chimeraplasts to target the G551D mutation to the CFTR gene that requires only a single-base alteration. With the ultimate goal of generating ferret fibroblast cell lines heterozygous for the G551D mutation, we first sought to develop targeting and screening methodologies in Hela cells. Since such targeting methods are non-selective, it is imperative that the targeting efficiency be ≥ 0.1% so that suitable somatic clones can be identified from a pool of fewer than 1000 cells.

Initially, we based our chimeraplast design on the original reports describing this technology [38,39]. The non-coding strand in the oligonucleotide contains two 10–15-nt long, 2′-O-methyl RNA stretches, flanking a 5 bp DNA sequence, with the mutation to be generated in the central position. A 68-mer chimeric oligonucleotide, shown in Figure 4a, was synthesized and purified by IDT (Iowa City, IA) and transfected into Hela cells using standard Lipofectamine (Gibco)-mediated methods. Although direct nuclear injection was used in similar targeting strategies for primary ferret fetal fibroblasts, transfection efficiencies in Hela cells were high enough to work out the methodologies of screening. The efficiency of transfection, which was tested with a FITC-labeled oligonucleotide of similar length, was demonstrated to be >95% (data not shown). Specifically, 1.8 µg of chimeraplasts were transfected into 50% confluent 5 × 10⁴ Hela cells in 200 µl serum-free DMEM medium/lipofectamine reagent at RT for 30 min in a 24-well plate. The cells were allowed to grow to confluence following the addition of 10% FBS. They were then trypsinized and serially diluted into 96-well plates for cloning. The cell concentration was determined before seeding so that only 50% of the wells would lead to the clonal outgrowth of a single cell. After the wells reached confluence, approximately 5000 cells were removed by scraping with a multi-channel pipette. The remaining cells were fed fresh media and allowed to grow back to confluence. Harvested cells from each clone were then pelleted and lysed directly in 60 µl of denaturing buffer containing 500 mM NaOH, 2.0 M NaCl, and 25 mM EDTA. After boiling for 5 min, 6 µl of 1 M Tris-Cl was added to cell lysates for neutralization. Nested PCR was then performed with 6 µl of the neutralized lysates as templates, using the following primers:

1st round 5′-ACATTAAGAGGAGATGTGCC-3′/5′-GTGCCCTTCAATCTCAGATG-3′

2nd round 5′-GGGCACAGATTTGAGTAACC-3′/5′-AATGTGATTTCTAACCACCTAGGC-3′

One tenth of each PCR product was loaded onto a Nylon membrane using a slot-blotting apparatus and screened for mutations by allele-specific oligonucleotide (ASO) hybridization. The sequences for the wild-type and mutant ASO primers were as follows: 1) wild-type, GAGTGGCCTTTCAAATTCAGATTGAGC-3′; 2) G551D mutant, GAGTGCAAGATGTGCC-3′. Primers were end-labeled with [γ-32P]ATP, and hybridization was performed overnight in a solution containing 3 M tetramethylammonium chloride (TMAC), 0.6% SDS, 1 mM EDTA, 10 mM Na₃PO₄ (pH 6.8), 5 × Denhardt, and 40 µg/ml yeast DNA. The blots were washed in 3 M TMAC, 1 mM EDTA, 10 mM Na₃PO₄ (pH 6.8), and 0.6% SDS twice at RT, followed by 2 more washes at 53 °C. Membranes were then exposed to film. Since the G to A mutation leads to the loss of a HindIII restriction site and a gain of DpnI and DpnII sites, positive

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clones detected by ASO were confirmed by restriction digestion analysis following amplification of genomic DNA.

Since gene targeting efficiencies were expected to be fairly low, we initiated this line of research by developing methods for mutant screening that were amenable to analyzing 1000 cell clones. We settled on using a 96-well plate to screen crude cell lysates by PCR. These plates could also be replicated easily by simply scraping cells from one plate into another without the need for trypsinization. This approach will be of considerable benefit when applied to ferret fibroblasts since it requires minimal amplification of cell clones, and will thus likely increase cellular competence for NT experiments. A variety of lysis buffers were tested for efficient amplification of the target by nested PCR. The lysis buffer discussed above allowed successful PCR from as few as 1,000 cells. ASO screening of 44 single-cell clones generated from chimeraplast-transfected Hela cells resulted in 3 positive clones that strongly hybridized to the G551D mutant oligonucleotide (Figure 4b and 4c). Each of these targeted clones demonstrated the expected restriction site changes caused by a G → A conversion. In summary, we feel that this preliminary data has provided the means for rapid effective screening of targeted primary fibroblasts or any other cell type useful for NT cloning. The 7.6% targeting efficiency for the human CFTR gene found in this study also suggests that chimeraplasts may be a useful targeting strategy in the generation of mutant ferret somatic cells.

Although studies with Hela cells have helped establish the screening protocols for selecting CFTR-targeted clonal cell lines, approaches for targeting primary fetal ferret fibroblasts have proven more difficult due to low transfection efficiencies. To this end, a single-cell injection approach was adopted in which targeting agents can be injected directly into the nucleus of primary ferret fibroblasts. Studies utilizing a mutant EGFP(Y66S) fluorescent recovery assay were established to help define the optimal conditions for gene targeting with chimeraplasts in ferret fetal fibroblasts [40]. Both episomal mutant EGFP plasmids and mouse transgenic fetal fibroblasts expressing the mutant EGFP gene were used as targets for chimeraplasts. Since the mutant EGFP protein is non-fluorescent, the simple index of fluorescent recovery could be used to assess the efficiency of gene targeting. Results from this study demonstrated that chimeraplasts could efficiently (~1–2%) target correction of mutant episomal DNA targets. However, they failed to correct an integrated mutant EGFP gene in fetal fibroblasts from transgenic mice. Although these results are disappointing, several advances in small oligonucleotide targeting may help to improve the efficiency of this approach. Recent reports have suggested that the polarity of the DNA segment in the

Figure 4
Chimeraplast targeting of the G551D mutation to the human CFTR gene in Hela cells. (A) The 25 bp segment homologous to CFTR is bolded, with the mutated nucleotide (G → A) indicated by asterisks. This structure is based on the traditional design with a 5 bp DNA (capitalized and underlined) core region flanked by two 10-nt 2’-O-methyl RNA stretches (lowercased). Only one base-pair change exists in parentheses for generating such an oligo for ferret CFTR. (B, C) Cell lysates prepared with 44 targeted Hela cell clones after G551D chimeric oligonucleotide transfection were amplified by 2 rounds of PCR and analyzed by ASO hybridization against (B) wild-type CFTR and (C) G551D mutant oligonucleotide probes. Three of these cell clones (2b, 3c, and 6a) were highly positive for both the wild-type and the G551D genotype, as indicated by the asterisks. Positive plasmid cDNA controls for the wild-type (1e) and the G551D (1f) CFTR sequences were also run as standards. PCR blanks are shown in 1 g and 1 h. All other wells contain experimental Hela cell clone PCR material. Wells that showed no hybridization to either probe probably contained no DNA material, which was likely due to insufficient cells for efficient PCR.
targeting construct affects the efficiency of gene targeting by more than 1000-fold [41]. We have currently only tested for base alteration targeted to the antisense strand of the target gene (i.e., chimeraplast DNA segments encode the mutation in the sense strand). However, the alternative design, placing mutations in the anti-sense strand of the targeting DNA oligonucleotide, has proven significantly more effective [41]. Such adaptations to the described approach may be used to increase targeting efficiencies for integrated genes in primary fibroblasts.

**Future Challenges in Mammalian Cloning of Genetic Disease Models**

Mammalian cloning has been accomplished in several mammalian species by nuclear transfer of somatic cells. However, widespread use of this technology has been limited due to low efficiencies of cloning to live births. At present, cloning efficiency – as determined by the proportion of live offspring developed from all oocytes that receive donor cell nuclei – is no more than 3%, regardless of the developmental age of the donor cell or the type of cell used [42].

The low efficiency associated with cloning may be attributed to many factors that are not fully understood, such as the oocyte-donor cell interaction [3], the stage of the donor cell cycle [4,43–46], the type of donor cell used [47,48], and inappropriate or incomplete nuclear reprogramming following nuclear transfer [49,50]. In addition, technical skill greatly contributes to the cloning success rate. Even the slightest damage to the donor cell (cytoplasm and/or nucleus) may render the nuclei incapable of participating in normal embryo development.

Changes in DNA methylation patterns may also account for the low efficiency of present cloning approaches. DNA methylation is highly dynamic in cleavage-stage embryos of a number of mammalian species. Failure to properly recapitulate pre-implantation DNA methylation patterns in embryos derived by nuclear transfer may contribute to the low efficiency of nuclear transfer in producing live offspring [49]. It is natural to speculate that oocyte cytoplasm has ‘special ingredients’ that reprogram epigenetic imprinting from the somatic state to the zygotic state. Attempts to increase the efficiency of cloning by increasing the exposure time of donor nuclei to the oocyte’s cytoplasm have met with some, though by no means dramatic success [45,51–53]. It was also somewhat surprising that using the so-called ‘totipotent’ ES cells for cloning was not as successful as expected [54,55]. However, more recent studies using out-bred F1 ES cell lines have demonstrated higher efficiencies in reconstructing oocytes capable of developing to live-born pups and increasing post-natal survival [56].

In conclusion, although mammalian cloning is still in its infancy, it is likely to change the face of animal modeling in the near future. As new methods for embryo manipulation and NT cloning merge with highly efficient gene targeting approaches, the ability to generate innovative larger animal models of genetic disease will significantly increase. Such efforts will greatly benefit the field of molecular medicine.

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**References**


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