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*J Immunol* published online 28 August 2015
http://www.jimmunol.org/content/early/2015/08/28/jimmunol.1501132

Supplementary Material
http://www.jimmunol.org/content/suppl/2015/08/28/jimmunol.1501132.DCSupplemental

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Not All SCID Pigs Are Created Equally: Two Independent Mutations in the Artemis Gene Cause SCID in Pigs

Emily H. Waide,* Jack C. M. Dekkers,* Jason W. Ross,* Raymond R. R. Rowland,† Carol R. Wyatt,‡ Catherine L. Ewen,‡ Alyssa B. Evans,* Dinesh M. Thekkoot,* Nicholas J. Boddicker,‡ Nick V. L. Serão,* N. Matthew Ellinwood,* and Christopher K. Tuggle*

Mutations in >30 genes are known to result in impairment of the adaptive immune system, causing a group of disorders collectively known as SCID. SCID disorders are split into groups based on their presence and/or functionality of B, T, and NK cells. Piglets from a line of Yorkshire pigs at Iowa State University were shown to be affected by T−B−NK+ SCID, representing, to our knowledge, the first example of naturally occurring SCID in pigs. In this study, we present evidence for two spontaneous mutations as the molecular basis for this SCID phenotype. Flow cytometry analysis of thymocytes showed an increased frequency of immature T cells in SCID pigs. Fibroblasts from these pigs were more sensitive to ionizing radiation than non-SCID piglets, eliminating the RAG1 and RAG2 genes. Genetic and molecular analyses showed that two mutations were present in the Artemis gene, which in the homozygous or compound heterozygous state cause the immunodeficient phenotype. Rescue of SCID fibroblast radiosensitivity by human Artemis protein demonstrated that the identified Artemis mutations are the direct cause of this cellular phenotype. The work presented in the present study reveals two mutations in the Artemis gene that cause T−B−NK+ SCID in pigs. The SCID pig can be an important biomedical model, but these mutations would be undesirable in commercial pig populations. The identified mutations and associated genetic tests can be used to address both of these issues. The Journal of Immunology, 2015, 195: 000–000.

Severe combined immunodeficiency is a diverse group of primary immunodeficiencies that involve impaired development or function of both B and T cells (1). Phenotypes causing SCID can be classified based on the presence or absence of B, T, and NK cells and further categorized by the genetic defect that causes the disease. Mutations in genes necessary for V(D)J recombination at the TCR or Ig clusters, which are required for receptor maturation and T and B cell survival, respectively, lead to T−B−NK+ SCID (2). T−B−NK+ SCID defects can be split into two groups based on sensitivity of affected cells to ionizing radiation. Lesions in the RAG1 and RAG2 genes result in immunodeficient animals that are resistant to radiation (3). Cells with defects in DNA-PKcs, DNA ligase IV, Cernunnos, or Artemis are radiosensitive, as they are unable to repair damage to DNA caused by ionizing radiation (4).

SCID animals are valuable biomedical models and have been used to study many aspects of human immune function, disease progression, as well as vaccine and drug development and testing. SCID mice, in particular, have been widely used as a vehicle to study the human immune system and play important roles in preclinical studies of the stability and function of human stem cells and their differentiated counterparts as potential therapies (5, 6). To improve SCID mice as biomedical models, genetic engineering has been used to overcome some of the differences between the immune environment of mice and humans (7). Much progress has been made in creating immunodeficient mice models used in biomedical research (7, 8) through selection of appropriate mouse strains and conditioning regimens (9). However, regardless of these advances, the translatability of rodent immunology and inflammatory studies to humans is imperfect (10, 11) and other models may be beneficial (12, 13).

Anatomical, genetic, and immunological similarities underpin the usefulness of swine as a large animal model for humans (13, 14). Recently, transgenic methods have been used to create SCID pigs, including mutations in the X-linked IL2RG gene (15, 16) and mutations in RAG1 or in both RAG1 and RAG2 (17, 18). Recently, our group (19) reported, to our knowledge, the first identification of a naturally occurring form of SCID in a line of purebred Yorkshire pigs that had been selected for increased feed efficiency at Iowa State University (20). Necropsy of four piglets that died abnormally early in a viral challenge study showed that they lacked functional adaptive immune systems. In subsequent work, we (21) demonstrated that...
these immunodeficient pigs failed to reject human cancer cells, and Ewen et al. (22) reported minimal circulating B and T cells but normal amounts of NK cells in a preliminary analysis of these SCID pigs. In this study, to our knowledge we present the first report of the characterization of the genetic lesions that cause the SCID phenotype in this pig population. Quantitative real-time PCR (qRT-PCR) and flow cytometry assays confirmed that the SCID pigs lack B and T cells, but do have NK cells; we also demonstrate T cell maturation defects in the thymus. Irradiation of fibroblasts from these SCID pigs and normal controls showed that the genetic defect is in the group of genes involved in DNA damage repair. Genetic mapping, phasing of genetic markers, and targeted sequencing revealed two independent mutations in the *Artemis* gene that cause SCID in this line of pigs. We confirmed that mutations in *Artemis* are responsible for the radiosensitivity of fibroblasts by rescuing this phenotype with expression of human *Artemis* in fibroblasts derived from the SCID pigs. These *Artemis*-deficient pigs provide a large animal model to study various aspects of human disease and immune function.

**Materials and Methods**

All animal experiments were performed under a protocol approved by the Iowa State University Experimental Animal Care and Use Committee (Ames, IA).

**Flow cytometry to determine cellular phenotype**

Thymic tissues were collected into ice-cold MEM, supplemented with HEPES, and penicillin/streptomycin. Tissues were minced into 2 × 2-cm² pieces and treated with 100 U/ml collagenase (collagenase I; Life Technologies, Grand Island, NY) in HBSS (plus calcium and magnesium; Life Technologies) for 30 min at 37°C, with gentle agitation. Cells were washed in PBS, counted, and 1 million cells were stained using the following fluorescently conjugated Abs: Alexa Fluor 647–conjugated anti-pig CD8α (clone MAC320; BD Biosciences), or the appropriate isotype controls (BD Biosciences), or a with FITC-conjugated mouse anti-pig CD3ε (clone PPT3; SouthernBiotech, Birmingham, AL). Cells were washed and resuspended in 400 μl 0.25% BSA/PBS solution prior to acquisition on an EC800 flow cytometer (Sony Biotechnology, Champaign, IL). Data were analyzed using FCS Express 4 software (DeNovo Software, Los Angeles, CA).

For samples used in the homzygous h12 and h16 comparison, an aliquot of 100 μl whole blood, collected in ACD, was transferred to a 12 × 75-mm polystyrene tube. Fc receptors were blocked with 10% serum (Equitech-Bio, Kerrville, TX). Directly conjugated Abs to CD3, the lymphocyte gate were analyzed for the expression of CD3, CD21, or CD28 (Invitrogen, Carlsbad, CA). To examine the entire coding sequence for variation, the full coding region of the *Artemis* cDNA was PCR amplified using primers in the first and last exon (forward, 5'-GGATCCGTG-GCGGCCGCAGAGCTGCCTTTT-3' and reverse, 5'-GGCCGCCGACAGAGTTCTT-ATGGTAGG-3') using an annealing temperature of 60°C and an elongation time of 2 min and 30 s. *Artemis* cDNA PCR products were cloned into a TOPO vector and grown in *Escherichia coli* bacteria on Luria–Bertani plates with ampicillin. Individual colonies were then picked into Luria–Bertani/ampicillin media and PCR amplified to ensure that each colony contained a vector with *Artemis* cDNA. Positive PCR products were cleaned using Exo-Sap and sent to the Iowa State University DNA Facility for sequencing.

**Haplotypes**

Following the results from the association study, genotypes of subjects used for the genome-wide association study for 21 SNPs located in a 1-Mb region surrounding *Artemis* from the SNP60 panel were separated into haplotypes using PHASE 2.1.1 (28). This revealed two haplotypes that segregated with SCID status when present in a homozygous or compound heterozygous state.

**Candidate gene sequencing**

Total RNA was extracted from either ear tissue or fibroblasts cultured from ear tissue using the TRIzol extraction method (Life Technologies, Grand Island, NY). mRNA was reverse transcribed using SuperScript VILO (Invitrogen, Carlsbad, CA). To examine the entire *Artemis* coding sequence for variation, the full coding region of the *Artemis* cDNA was PCR amplified using primers in the first and last exon (forward, 5'-GGATCCGTG-GCGGCCGCAGAGCTGCCTTTT-3' and reverse, 5'-GGCCGCCGACAGAGTTCTT-ATGGTAGG-3') using an annealing temperature of 60°C and an elongation time of 2 min and 30 s. *Artemis* cDNA PCR products were cloned into a TOPO vector and grown in *Escherichia coli* bacteria on Luria–Bertani plates with ampicillin. Individual colonies were then picked into Luria–Bertani/ampicillin media and PCR amplified to ensure that each colony contained a vector with *Artemis* cDNA. Positive PCR products were cleaned using Exo-Sap and sent to the Iowa State University DNA Facility for sequencing.

To examine the genomic region for sequence variation, primers were designed to amplify targeted exons of *Artemis* and portions of each surrounding intron. The genomic region containing exons 7 and 8 was PCR amplified using primers forward, 5'-CTCAATGGGGTATTTAGGCACCT-3' and reverse, 5'-GGCCATCTGATAGGGTTTCCA-3' using an annealing temperature of 58°C and an elongation time of 30 s. The genomic region containing exons 10 and 11 was PCR amplified using primers forward, 5'-GCTAAATGGCCGAGGTATTTCT-3' and reverse, 5'-CAAGAGTCCCAAGCCACTTCT-3' using an annealing temperature of 56°C and an elongation

**Genome-wide association analysis**

Subjects. Genomic DNA from 20 SCID affected pigs, 50 unaffacted littermates, their six parents, and 96 ancestors from the previous seven generations of the experimental residual feed intake (RFI) selection line at Iowa State University (20) was isolated from tail or ear tissue using the Qiagen DNeasy blood and tissue kit (Qiagen, Hilden, Germany). DNA samples were sent to GeneSeek (Lincoln, NE) and genotyped using the Illumina porcine SNP60 beadchip (25). SCID status was determined by plotting lymphocyte against WBC numbers obtained from complete blood cell (CBC) analysis of whole blood from each piglet (data not shown).

**Statistical analysis.** Initial attempts to map the genomic region harboring the causative mutation using runs of homozygosity (26) in the affected piglets were unsuccessful. Associations of single nucleotide polymorphisms (SNPs) with the binary phenotype of affected versus unaffected SCID status were then analyzed using the disease within family (dfam) option of the PLINK toolset (27) to determine the genomic region most strongly associated with the categorical disease phenotype. The dfam option examines differential transmission of alleles from parents to affected versus unaffected offspring. This analysis assumes homogeneity of the associated allele within each family, but not between families.

**Haplotype analysis.** Following the results from the association study, genotypes of subjects used for the genome-wide association study for 21 SNPs located in a 1-Mb region surrounding *Artemis* from the SNP60 panel were separated into haplotypes using PHASE 2.1.1 (28). This revealed two haplotypes that segregated with SCID status when present in a homozygous or compound heterozygous state.
time of 60 s. PCR products were cleaned using Exo-Sap and sent to the Iowa State University DNA Facility for sequencing. Sequences for each haplotype were compared with those of normal littermates and to the reference sequence obtained from Sus scrofa build 10.2.

Rescue of radiosensitive phenotype

Fibroblasts for h12/h16 compound heterozygous SCID (n = 2) and normal animals (n = 3) from three litters were cultured in the same manner as described above. Fibroblasts were transfected with either 5 μg human Artemis plasmid (29) (pExodus CMV-Artemis, which was procured through Addgene [plasmid no. 40211] by Andrew Scharenberg), with 3.45 μg empty vector construct (pEco) (also deposited by A. Scharenberg, Addgene; plasmid no. 39991), or shocked with no plasmid added. Cells were electroporated using electroporation media and conditions described by Ross et al. (23). Briefly, 1 million cells in 200 μl were electroporated using three, 1 ms square-wave pulses of 300 V in a 2-mm gap cuvette using a BTX ECM 2001 electroporator. Transfected cells were incubated overnight in cell culture media. One day after transfection, fibroblasts were subjected to 4 Gy radiation and plated in triplicate as described above. Cells were cultured for 14 d, with one media change 7 d after irradiation, stained with methyl blue, and colonies were determined by visual inspection and counted.

Statistical analysis. The average number of colonies of three replicates for each animal and plasmid was used as observations. Data were analyzed using Proc Mixed in SAS 9.3, including the fixed effects of SCID status, plasmid, and their interaction. Litter and animal within litter were fitted as random effects. Different error variances were allowed for data from normal versus SCID animals.

Results

SCID phenotype segregates as an autosomal recessive trait in Yorkshire pig population

Weaned piglets of the eighth generation of a line of purebred Yorkshires selected for increased feed efficiency (RFI) at Iowa State University (20) were sent to Kansas State University to undergo experimental infection with the porcine reproductive and respiratory syndrome virus to evaluate their ability to respond to an infectious disease stressor (30). Necropsy of four of these piglets that died soon after arrival at Kansas State University showed that they had very low numbers of circulating B and T cells, and their lymph nodes and thymus were small (19). Pedigree analysis of the four litters that these piglets were from showed that the six parents of these four litters were related. To explore the genetic basis of this phenotype, matings between these pigs were shown, each of which produced at least one affected piglet, as expected 25% if the immunodeficiency phenotype was caused by an autosomal recessive mutation.

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SCID fibroblasts are sensitive to ionizing radiation

Human SCID patients with a T− B− NK− (c) phenotype have defects in genes in the somatic recombination pathway required for TCR and Ig maturation. This involves two groups of genes. The first group includes RAG1 and RAG2, whose proteins catalyze the first step of TCR and Ig maturation, creating a dsDNA break (DSDB) in pre-T and pre-B cells (34). The second group of genes repairs these breaks in the process of forming functional TCR and Ig receptors (35). Defects in RAG1 and RAG2 affect somatic DNA recombination, which occurs only in B and T lymphocytes. However, the ubiquitously expressed genes in the second group are involved in the nonhomologous end joining (NHEJ) DNA damage repair pathway, which is important in all cells (36). Cells with lesions in DNA damage repair genes show increased sensitivity to ionizing radiation (37). To determine whether the genetic defect was in the first or second group of genes, we tested the ability of fibroblasts from SCID and normal pigs to repair DSDB caused by ionizing radiation. Fibroblasts cultured from skin tissue of SCID and non-SCID littermates from five litters were gamma irradiated. Fibroblast survival was significantly decreased for cells from the SCID pigs compared with cells from normal littermates at all radiation doses tested (p < 0.0001; Fig. 2). Even at the lowest dose of radiation (2 Gy), the proportion of fibroblasts that survived and developed colonies was significantly lower for those derived from SCID pigs (0.28 ± 0.03) compared with those of normal pigs (0.73 ± 0.03; p < 0.0001). This radiosensitivity in fibroblasts suggested that the mutation causing SCID in these pigs was in one of the ubiquitously expressed genes involved in NHEJ repair of DSDB.

Genetic mapping of the locus mutated in SCID pigs

Genotypes for >60,000 SNPs represented on the Illumina porcine SNP60 panel (25), covering the entire porcine genome, were obtained on the six carrier parents, 20 SCID affected piglets, 50
unaffected littermates of the SCID piglets, and 96 ancestors of these animals. A genome-wide association study was performed to identify the region harboring the causative mutation. We identified a 5.6-Mb region on *S. scrofa* chromosome 10 that was differentially transmitted (*p* = 2.7 × 10^-7 for SNP with strongest association) to SCID affected versus unaffected piglets (Fig. 3). This region was found to contain the porcine *Artemis* gene, which, when mutated in humans (38) or mice (39), causes a T^+^ B^+^ NK^-^ immunodeficiency phenotype. Lesions in *Artemis* have also been shown to cause increased sensitivity to ionizing radiation (40).

To confirm the suspected autosomal recessive manner of transmission, a haplotype analysis was performed using 21 SNPs from the SNP60 panel in the 1-Mb region surrounding *Artemis*. Results showed two distinct haplotypes (h12 and h16), which, when present in a compound heterozygous or homozygous recessive state, were associated with the SCID phenotype. Both of these SNP haplotypes could be traced back to the founder generation of the RFI selection line (20), which was sourced from commercial Yorkshire populations in the Midwest region of the United States (Fig. 4).
missing the second exon of an *Artemis* porcine cDNA sequence isolated from alveolar macrophages that was reported by others (NM_001258445.1). This exon was not present in any of the *Artemis* transcripts sequenced in this study, although it exists in the pig genome (*S. scrofa* 10.2) between exon 1 and what we refer to as exon 2 and may represent a bona fide exon. However, based on BLAST analysis of sequences that are present in dbEST (February 17, 2015), cDNAs with the top six alignments did not contain this exon. Additionally, this exon did not align to any portion of human *Artemis* cDNA sequences (NM_001033855.2). Both genome-guided transcriptome assembly and de novo transcriptome assembly using whole-blood RNA sequencing data from 31 pigs in an unrelated project was found to contain only one *Artemis* transcript, which did not contain this exon (H. Liu and C.K. Tuggle, unpublished data). Based on these findings, we think that the swine reference transcript that contains this extra exon (NM_001258445.1) is a minor transcript. For clarity and consistency with the human literature, we denoted this extra exon as exon 1A and considered the second exon that is present in the transcripts observed in our study as exon 2. Translation of the longest transcript we sequenced from non-SCID pigs would produce a protein containing 712 aa, with 81% sequence identity to the human encoded *Artemis* protein.

*Haplotype 16 has a lesion causing a predicted splice defect.* Most cDNAs sequenced from normal pigs contained exons 1–14. The most complete transcript sequenced from cDNA from the h16 SCID haplotype contained all these exons except for exon 8, resulting in the loss of 141 nucleotides (Supplemental Fig. 1). In fact, none of the cDNA sequences obtained from h16 chromosomes contained exon 8. Genomic sequencing of exon 8 and a portion of the surrounding introns revealed a splice donor site mutation in intron 8 (g.51578763 G→A; Fig. 5B). This G→A mutation was only seen in the h16 genomic sequence, as a G nucleotide at this position was seen in both the h12 and normal sequences; to distinguish h16 from h12 transcripts in compound heterozygous animals, we used a synonymous point mutation in exon 13 that we identified to be present only in h16 transcripts (position g.51587796). This G→A nucleotide conversion disrupts the signals required to correctly splice exon 8 to exon 9, which explains the observed lack of exon 8 in cDNAs expressed from h16 haplotype chromosomes. An *Artemis* transcript with no exon 8 would retain the normal reading frame but would be expected to produce a protein missing 47 aa of the predicted full-length 712-aa *Artemis* protein.

*Haplotype 12 has a nonsense point mutation and a predicted severely truncated protein.* Multiple examples of alternative splicing, including aberrant splicing within exons and one intron, were observed for transcripts from chromosomes that carried the h12 haplotype (Supplemental Fig. 1). All transcripts sequenced from homozygous h12 animals were found to lack the 137-bp-long exon 10, which would cause a frameshift that results in a stop

**FIGURE 2.** Effect of ionizing radiation on fibroblasts from SCID and normal pigs. Fibroblasts from SCID piglets (*n* = 10) and normal littermates (*n* = 10) were exposed to increasing doses of gamma rays. Colonies (≥2 mm) were stained and counted after 14 d. Survival proportion was calculated as the average number of surviving colonies for three replicates at each radiation dose divided by the number of colonies from nonirradiated cells for each animal. Error bars represent the SE of the least squares means. *p < 0.0001, difference between SCID and normal pigs within radiation dose.

**FIGURE 3.** Manhattan plot of the genome-wide association study for SCID status. Results show the −log (*p* value) of the association of ordered SNPs on *S. scrofa* chromosomes 1–18, X, Y, and unknown (U) with SCID status based on the dfam option in PLINK.
The predicted protein translated from transcripts missing exon 10 would be severely truncated; at 277 aa long, it would be 50% of normal length. To investigate the cause of this lack of exon 10 in all h12 transcripts, exons 10 and 11 and portions of the surrounding introns were amplified from genomic DNA of homozygous h12 animals. Signal sequences required for splicing of exon 10 were identical to those seen in the sequence of normal pigs, as well as in the reference sequence for normal pigs.

**Figure 4.** Pedigree of SCID ancestors showing carriers of mutated haplotypes based on the Illumina porcine SNP60 panel. Circles represent females and squares represent males. Green symbols are h16 haplotype carriers and pink symbols are h12 haplotype carriers, with lines of each color tracking the respective SCID haplotype through the pedigree to the founder generation. Beige symbols are pigs that were genotyped with the SNP60 panel and did not carry either SCID haplotype. White symbols in generations 0–7 represent nongenotyped individuals. Boxes in generation 8 give information on the numbers of SCID (Affected) and non-SCID (Unaffected) piglets and the number of piglets that died before their SCID phenotype was determined (Unknown) from each parent pair.

**Figure 5.** Two independent mutations were found in *Artemis*. (A) The coding region of the *Artemis* transcript is indicated by slanted lines and genotypes at mutated positions are shown for chromosomes that carry normal, h12, and h16 haplotypes. Mutant alleles that cause SCID are shown in black lettering. (B) Genomic sequence of the h16 haplotype shows a splice donor site mutation (g.51578763 G→A) responsible for the lack of exon 8 in all h16 transcripts. Capital letters denote exonic sequence whereas lowercase letters denote intronic sequence. (C) A nonsense point mutation (g.51584489 G→A) in exon 10 changes the tryptophan at position 267 to a stop codon in the h12 haplotype.
sequence (Sscrofa 10.2, http://www.ensembl.org/Sus_scrofa/Info/Index). However, sequence analysis of the exon 10 genomic region in homozygous h12 pigs identified a G→A nonsense point mutation at g.51584489 that changes the tryptophan amino acid codon at position 267 to a stop codon (Fig. 5C). Our interpretation of these data is that any transcript that contained exon 10 was not stable enough for cDNA analysis to detect. Furthermore, although not identified as part of any cDNA from homozygous h12 pigs, the predicted translation of this presumably unstable transcript containing exon 10 with the g.51584489 G→A mutation would also produce a severely truncated protein of 266 aa.

**Animals homozygous for either mutation are not different in numbers of specific lymphocytes or in radiosensitivity**

The flow cytometry and radiosensitivity assays described earlier were performed on a combination of compound heterozygous and homozygous h12 or h16 SCID animals. To determine whether homozygosity for either mutation would result in altered severity of phenotypes, flow cytometry of circulating blood and assays of fibroblast irradiation were performed. Peripheral blood from SCID phenotypes, flow cytometry of circulating blood and assays of homozygous h12 or h16 SCID animals. To determine whether were performed on a combination of compound heterozygous and The flow cytometry and radiosensitivity assays described earlier numbers of specific lymphocytes or in radiosensitivity.

Animals homozygous for either mutation are not different in numbers of specific lymphocytes or in radiosensitivity.

**Transfection with human Artemis rescues radiosensitivity of fibroblasts**

To further confirm the role of Artemis in the observed SCID phenotype, we assessed the complementation of the radiosensitivity of SCID pig fibroblasts by expression of human Artemis. Fibroblasts cultured from compound heterozygous SCID (n = 2) and normal (n = 5) piglets from three litters were transfected with a human Artemis-containing plasmid (pArt), with the pExo, or shocked without the addition of either plasmid. Cells were then subjected to 4 Gy radiation and survival was measured. Addition of pArt increased the radiosensitivity of fibroblasts of SCID or non-SCID genotypes.

**Discussion**

Through immunophenotyping, genetic mapping, cDNA and genomic DNA sequencing, fibroblast radiation sensitivity testing, and cDNA rescue of radiosensitivity phenotypes of fibroblasts, we provide substantial evidence that a novel T- B- NK* immuno-deficiency trait in pigs is caused by two independent mutations in the same gene that codes for a DNA repair enzyme, which is known to cause this type of SCID phenotype in human patients. We document that the identified mutations in the Artemis gene are necessary to observe a SCID phenotype in directed matings and are sufficient to cause the radiosensitivity phenotype observed in SCID fibroblasts from this population.

**Novel immuno-deficiency phenotype in pigs is a recessive Mendelian trait that shows defects in thymocyte maturation consistent with flawed somatic recombination**

Ozuno et al. (19) reported the initial analysis of the first cases of naturally occurring SCID in pigs, which was serendipitously found within a line selected for a commercially relevant trait at Iowa State University (20). In our expanded analysis of litters born within the pedigree of this line, a Mendelian and non- sex-linked segregation of SCID affected status in litters born to carrier-by-carrier matings was observed, indicating that the SCID phenotype was an autosomal recessive trait (2). Findings from qRT-PCR of lymphocyte marker expression in whole blood showed that the SCID pigs had very low numbers of B and T cells, whereas NK cells were present. These results suggested that the ability of B and T lymphocytes to recombine genetic material to form functional receptors was compromised in the SCID affected piglets. Flow cytometry of thymus tissue showed an increased population of CD4− CD8α− and γδ− T cells in SCID pigs compared with non-SCID littermates. This suggested an arrest in early thymocyte development and TCR rearrangement in the SCID pigs (31, 33). The increased sensitivity to ionizing radiation in fibroblasts of SCID pigs further narrowed the list of candidate genes to be evaluated for mutations to those involved in the NHEJ DNA damage repair system.

**Two distinct mutations in the Artemis gene found in SCID pigs**

Genomic association analysis of the population segregating the SCID phenotype pointed to a region containing the Artemis gene, whereas SNP genotype phasing results indicated that there were two distinct haplotypes that each carried a mutation postulated to cause the SCID phenotype. Both of these haplotypes could be traced back to the founders of this population, which suggests that these deleterious mutations may be present in commercial Yorkshire pigs in the United States. This is an important implication for the swine industry, as SCID piglets thrive while suckling from their dam, but succumb to opportunistic infections after weaning. Parentage information is generally not tracked into the nursery, at least in part due to cross-fostering practices, the lack of permanent individual identification, and the frequent use of mixed semen from more than one boar for artificial insemination. Therefore, although the death of several littermates represents a concentrated loss for that litter, this clue to a genetic defect is not likely to be
recognized on commercial farms. Additionally, if the frequency of the SCID mutations is low, carrier-by-carrier matings may be rare in industry practice.

Sequencing of *Artemis* cDNA from non-SCID and SCID pigs carrying each haplotype identified multiple examples of alternative splicing. The cause for absence of specific exons in sequences of each SCID haplotype was explored using targeted genomic sequencing. A point mutation at the splice donor site for intron 8 in the h16 genomic sequence explained the lack of exon 8 in all cDNA sequences from the h16 haplotype. A compendium of 48 different causative mutations in *Artemis* in human SCID patients has cataloged six splice site mutations, although none involve exon 8 (41). Translation of the most complete h16 transcript would produce a protein of 665 aa, lacking a 47-aa-long portion of the β-CASP region, which is required for *Artemis* function (42, 43). Genomic deletions of *Artemis* exons 5–8 (38) and exons 7 and 8 (40) have been shown to cause SCID in human patients. For h12, no cDNAs were identified that contained exon 10, but the most complete cDNAs from homozygous h12 cells contained all other exons that were present in the non-SCID cDNA sequence. Because the lack of exon 10 causes a frameshift and the open reading frame stops in exon 11, the longest protein that is predicted to be encoded by the observed cDNAs from haplotype h12 is 277 aa long, containing only 39% of the normal protein. Interestingly, genomic sequence of the h12 haplotype identified a nonsense point mutation in exon 10, which would result in a protein of only 266 aa if this exon 10 were present in mRNA. This severely truncated protein would lack more than half of the total amino acid sequence, including a highly conserved part of the β-CASP motif (42). Therefore, in both observed cDNAs and RNA transcripts predicted from genomic sequence, h12 can encode only a greatly shortened protein, with 277 or 266 aa, respectively. In humans, an 8-bp insertion in exon 14 of *Artemis* that changed a cysteine to a stop codon at position 330 of the protein was also shown to cause SCID (44). Pannicke et al. (41) listed several small deletions causing SCID, most of which lead to a frameshift that introduces novel nonsense codons that cause truncations that are distal to exon 10, in exons 12, 13, and 14. This suggests that either version of the h12 haplotype protein is highly unlikely to be functional in DNA damage repair.

We thus hypothesized that the h12 mutation may cause a more severe immunodeficient phenotype than the h16 mutation, in which only 47 internal amino acids are predicted to be lost. However, flow cytometry and fibroblast irradiation showed no phenotypic differences between SCID pigs that were homozygous for either mutation or compound heterozygous. Although single missense mutations, as well as deletions of single amino acids without a frameshift, can lead to a SCID or Omenn syndrome phenotype in humans (41), examples exist for frameshift mutations producing a truncated protein with partial activity. For example, the D451fsX10 mutation is a large truncation of >200 aa that retains partial nuclease activity (45). In a mouse model, the hypomorphic D451fsX10 mutation is associated with aberrant DNA joining that causes chromosomal rearrangements (46). Thus, underlying cellular differences in DNA repair activity between the h12 and h16 encoded proteins may exist, and further studies should be conducted to determine the functional differences between these two mutations in pigs.

**Value of Artemis SCID pigs in preclinical testing**

Similarities between humans and pigs emphasize the value of the pig as a biomedical model (13), and pigs as large animal models for many diseases are being created (47). Body, tissue, and organ size make them physiologically very similar, allowing surgical and imaging techniques to translate well (48). Size similarity also makes the pig a more suitable model for testing delivery of cells (i.e., heart) and for measuring location/stability of transplanted cells under orthopedic stress (i.e., joint tissue). However, the use of pig models in regenerative medicine has been minimal, due to the heretofore lack of a porcine immunocompromised model that could be used as an xenogenic transplant model. As recently demonstrated for human cancer cells, the SCID pigs we describe in the present study cannot reject a xenograft (21). Immunodeficient porcine models have long been sought, as is evident by the recently reported transgenic SCID pigs with targeted IL2RG (15, 16) and RAG1 and/or RAG2 mutations (17, 18). In the present study, we have described two naturally occurring genetic defects in *Artemis* that cause SCID in pigs. These porcine models will be valuable in studies involving cellular and organ transplantation, human cancer progression and treatment, vaccine development, and many other immunological questions. Specifically, defects in *Artemis* (as well as all genes encoding proteins needed for VDJ recombination) have been shown to be associated with poor outcome of stem cell transplantation in humans (reviewed in Ref. 2). In the case of *Artemis*, affected patients have substantially higher risk of late toxicity following hematopoietic cell transplants as compared with patients with RAG lesions (49). This was attributed to the use of alkyllating agents in conditioning regimens. A separate study also found *Artemis* defects were associated with negative clinical events following hematopoietic cell transplants, which included graft-versus-host disease and a range of infections (50). Whereas Xiao et al. (51) generated an *Artemis*-deficient mouse model that replicates the phenotype observed in human *Artemis*-deficient SCID patients, these *Artemis*-deficient pigs provide an additional and perhaps more relevant preclinical model to test and improve clinical therapies used in treating these patients. Future studies will focus on defining optimal husbandry practices to increase the lifespan of SCID pigs so that such modeling can be performed.

**Acknowledgments**

We sincerely thank G. Kaper and the staff at the Iowa State University swine breeding farm in Madrid, IA. We also thank E. Powell, as well as members of the Tuggle, Dekkers, Ross, Ellinwood, and Laboratory Animal Resources Groups at Iowa State University and the Rowland Group at Kansas State University for assistance with animal care and laboratory work. We acknowledge B. MacPhail, Jaeger Corporation (Omaha, NE), for help with irradiation of fibroblasts. We are grateful to M. Georges for valuable advice regarding genetic mapping and sequencing results of the two mutations.

**Disclosures**


