Telomere is a nucleoprotein structure at the ends of chromosomes that helps to protect the ends of chromosomes from being fused with other chromosomes. Knockout of histone methyltransferases Suv39h1 and Suv39h2 increases the telomere length in murine cells, whereas downregulation of SUV39H1 and SUV39H2 genes decreases the telomere length in human cells, suggesting that telomere biology is different among mammalian species. However, epigenetic regulation of the telomere has not been studied in mammals other than the human and mouse. In the present study, the effect of knockdown of SUV39H1 and SUV39H2 genes on telomere length was examined in porcine embryonic stem-like cells (pESLCs) and porcine embryonic fibroblasts (PEFs). The telomeres in SUV39H1 and SUV39H2 knockdown (SUV39KD) pESLCs (37.1 ± 0.9 kb) were longer (P<0.05) compared with those of the control (33.6 ± 0.7 kb). Similarly, SUV39KD PEFs had longer telomeres (22.1 ± 0.4 kb; P<0.05) compared with the control (17.8 ± 1.1 kb). Telomerase activities were not different between SUV39KD pESLCs (10.4 ± 1.7) and the control (10.1 ± 1.7) or between SUV39KD PEFs (1.0 ± 0.3) and the control (1.0 ± 0.4), suggesting that telomerase activities did not contribute to the telomere elongation in SUV39KD pESLCs and SUV39KD PEFs.

Relative levels of trimethylation of histone H3 lysine 9 and expressions of DNMT1, DNMT3A and DNMT3B were decreased in SUV39KD cells, suggesting that telomere shortening in SUV39KD pESLCs and SUV39KD PEFs might be not only related to the loss of histone modification marks but also linked to the decrease in DNA methyltransferase in pigs.

**Key words:** Embryonic stem-like cells, Fibroblasts, Knockdown, Pig, Telomere length

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**Abstract.** Telomere is a nucleoprotein structure at the ends of chromosomes that helps to protect the ends of chromosomes from being fused with other chromosomes. Knockout of histone methyltransferases Suv39h1 and Suv39h2 increases the telomere length in murine cells, whereas downregulation of SUV39H1 and SUV39H2 genes decreases the telomere length in human cells, suggesting that telomere biology is different among mammalian species. However, epigenetic regulation of the telomere has not been studied in mammals other than the human and mouse. In the present study, the effect of knockdown of SUV39H1 and SUV39H2 genes on telomere length was examined in porcine embryonic stem-like cells (pESLCs) and porcine embryonic fibroblasts (PEFs). The telomeres in SUV39H1 and SUV39H2 knockdown (SUV39KD) pESLCs (37.1 ± 0.9 kb) were longer (P<0.05) compared with those of the control (33.6 ± 0.7 kb). Similarly, SUV39KD PEFs had longer telomeres (22.1 ± 0.4 kb; P<0.05) compared with the control (17.8 ± 1.1 kb). Telomerase activities were not different between SUV39KD pESLCs (10.4 ± 1.7) and the control (10.1 ± 1.7) or between SUV39KD PEFs (1.0 ± 0.3) and the control (1.0 ± 0.4), suggesting that telomerase activities did not contribute to the telomere elongation in SUV39KD pESLCs and SUV39KD PEFs.

Relative levels of trimethylation of histone H3 lysine 9 and expressions of DNMT1, DNMT3A and DNMT3B were decreased in SUV39KD cells, suggesting that telomere shortening in SUV39KD pESLCs and SUV39KD PEFs might be not only related to the loss of histone modification marks but also linked to the decrease in DNA methyltransferase in pigs.

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Knockdown of *SUV39H1* in human cells, however, causes telomere shortening [14]. These data suggest that telomere biology is rather different among mammalian species. However, no study on epigenetic regulation of telomeres has been performed in mammals other than the mouse and human. In the present study, we investigated the impact of *SUV39H1* in human cells, however, causes telomere shortening [14].

### Materials and Methods

#### Cell culture

Two pESLC lines were produced from *in vitro* fertilization blastocysts according to Haraguchi et al. [15]. The cells were subcultured at 37 C in a 5% CO₂ atmosphere on mouse embryonic fibroblasts as feeder cells in ES medium consisting of KnockOut DMEM (KO-DMEM, Gibco, Grand Island, NY, USA) supplemented with 2 mM GlutaMAX, 1% MEM nonessential amino acids, 20% KnockOut Serum Replacement (KSR), 1% antibiotic-antimycotic liquid (all from Invitrogen, Carlsbad, CA, USA), 20 ng/ml porcine recombinant LIF (pLIF) (made in-house) and 0.1 mM β-mercaptoethanol (Sigma, St. Louis, MO, USA). The reference gene was α-tubulin (*TUBA*).

#### RNA interference

Four sets of siRNA were designed by Cosmo Bio (Tokyo, Japan) for double knockdown of *SUV39H1* and *SUV39H2* genes (SUV93KD). The sequences of the two siRNA sets designed to target porcine *SUV39H1* were 1) sense 5'-ggaaacgacagcagcagcagcagtctcc-3' and antisense 5'-gagaagccacagcagcagcagaagcagtctcc-3, and 2) sense 5'-ggaaacgacagcagcagcagcagtctcc-3 and antisense 5'-gagaagccacagcagcagcagcagtctcc-3'. The sequences of the two siRNA sets designed to target porcine *SUV39H2* were 1) sense 5'-ggaaacgacagcagcagcagcagtctcc-3 and antisense 5'-gagaagccacagcagcagcagcagtctcc-3 and 2) sense 5'-gagaagccacagcagcagcagcagtctcc-3 and antisense 5'-gagaagccacagcagcagcagcagtctcc-3. One day prior to transfection, pESLCs and PEFs were passaged, and the cell concentration was adjusted to a density of 30–50% confluency. Before transfection, 80 nM of each siRNAs were mixed and incubated together with 0.4% lipofectamine X-tremeGENE siRNA Transfection Reagent (Roche, Mannheim, Germany) in culture media without serum and antibiotics (transfection media) for 15–20 min at room temperature. The transfection media were then supplemented with each 10% FBS for PEFs or 20% KSR for pESLCs, and transfection was then carried out in culture media for 24 h or 48 h for pESLCs or PEFs, respectively. The transfections were performed for three successive passages, and the cells were collected after the third transfection. Non-siRNA-treated pESLCs and PEFs (at the same passage as SUV93KD pESLCs and SUV93KD PEFs, respectively) were also cultured in parallel with SUV93KD pESLCs and SUV93KD PEFs to serve as controls.

#### Extraction of total RNA and real-time PCR

Total RNA was isolated by using an RNeasy Micro kit (Qiagen, Hilden, Germany) according to the manufacturer’s instructions. cDNA synthesis was performed using a PrimeScript RT reagent kit (TaKaRa, Tokyo, Japan) by following the manufacturer’s protocol. PCR reactions contained approximately 5 ng cDNA, 0.25 μM of forward and reverse primers for each transcript and 10 μl of LightCycler 480 SYBR Green I Master (Roche Applied Science, Penzberg, Germany). The reference gene was α-tubulin (*TUBA*).

#### Table 1.

Primer sequences used for gene expression or chromatin immunoprecipitation (ChiP) assays

| Locus | Forward primer | Reverse primer |
|-------|----------------|----------------|
| Chr2start | ttcaacccaaaatgttttgc | tggactgcaaggaagcaca |
| Chr8start | atgcagccgtaactgccttct | cactgcggccagtaagtatc |
| Chr12start | tcctcagcactcaccttctg | gagctgcctctgcctttcta |

Knockdown of *SUV39H1* in human cells, however, causes telomere shortening [14]. These data suggest that telomere biology is rather different among mammalian species. However, no study on epigenetic regulation of telomeres has been performed in mammals other than the mouse and human. In the present study, we investigated the impact of *SUV39H1* and *SUV39H2* on telomere length in porcine ES-like cells (pESLCs) and porcine embryonic fibroblasts (PEFs).

### Extraction of genomic DNA and determination of telomere length by TRF assays

Genomic DNA was extracted using a DNeasy Blood & Tissue Kit (Qiagen) by following the manufacturer’s protocol. Mean telomere length was determined by mean terminal restriction fragment (TRF) length analysis using a TeloTAGGG Telomere Length Assay Kit.

Knockdown of *SUV39H1* in human cells, however, causes telomere shortening [14]. These data suggest that telomere biology is rather different among mammalian species. However, no study on epigenetic regulation of telomeres has been performed in mammals other than the mouse and human. In the present study, we investigated the impact of *SUV39H1* and *SUV39H2* on telomere length in porcine ES-like cells (pESLCs) and porcine embryonic fibroblasts (PEFs).
Telomerase activity assays

Telomerase activity assays were examined by using a TRAPEZE Telomerase Detection Kit (EMD Millipore, Billerica, MA, USA) following the manufacturer’s protocol. Briefly, samples were resuspended in 200 μl of 1× CHAPS Lysis Buffer. The suspensions were then incubated on ice for 30 min and centrifuged at 12,000 × g for 20 min at 4 C. About 160 μl of the supernatant was transferred into a fresh tube for determination and adjustment of protein concentration. The remaining extract was then quick-frozen on dry ice and stored at −80 C until use. The relative telomerase activities were determined using the SYBR Green real-time quantitative telomeric repeat amplification protocol (RQ-TRAP) assay according to Wege et al. [16] with some modifications. Briefly, the SYBR Green RQ-TRAP assay was conducted with 2 μl extracts, 0.1 μg of telomerase primer TS, 0.05 μg of anchored return primer ACX and 12.5 μl of LightCycler 480 SYBR Green I Master (Roche Applied Biosystems). Primer sequences were as described by Kim and Wu [17]. Inactivated (heat treated) samples and lysis buffer were also assayed on every plate as a negative control for each sample and primer-dimer/PCR contamination control, respectively. Samples were incubated for 20 min at 25 C and amplified in 35 PCR cycles of 30 sec at 95 C and 90 sec at 60 C (two-step PCR). All PCRs were performed twice and in triplicate on each plate with a LightCycler 480 Instrument (Roche Applied Science). Standard curves were generated from telomerase-positive cell extract (positive telomerase extract control supplied in the TRAPEZE Telomerase Detection Kit), which was serially diluted with PCR-graded water by 2-fold per dilution to produce five concentrations of DNA. The relative telomerase activities of cell samples were normalized to the relative quantification of non-siRNA-treated PEFs by the standard curve method.

Chromatin immunoprecipitation (ChIP) assay

For ChIP analysis, 2×10^6 pESLCs or PEFs were used per condition. ChIP assays were performed based on Boyd and Farnham [18] with some modifications. In preparation for the chromatin preclear step, Pansorbin Cells (EMD Millipore) were washed with 2 mM EDTA and 50 mM Tris-Cl (pH 8.0), blocked with 10 mg/ml calf thymus (Sigma) and 10 mg/ml BSA and stored at 4 C before use. Cells were treated with 1% formaldehyde for 10 min to cross-link histones to DNA at room temperature on a shaking platform. The cross-linking reaction was stopped by adding glycerol to a final concentration of 0.125 M. The cross-linked cells were washed twice with cold PBS and then lysed in 100 μl of 1% SDS (Sigma), 50 mM Tris-HCl (pH 8.0), 10 mM EDTA and 1% protease inhibitor cocktail (Nacalai Tesque, Kyoto, Japan) for 10 min on ice. The lysate was sonicated for 5 min (4 pulses for 15 sec with a 1-min rest interval) by a Bioruptor sonicator (Cosmo Bio) to obtain chromatin fragments <1 kb. The chromatin fragments were then centrifuged at 14,000 × g for 10 min at 4 C, and the supernatant was collected. About 10 μl of the chromatin fragments were cleaned using a QIAquick PCR Purification Kit (Qiagen) and run on a gel to check the sonication. The remaining chromatin fragments were precleared by adding 2 μl blocked/washed Pansorbin Cells and 1 mM phenylmethanesulfonyl fluoride (PMSF, Nacalai Tesque) and incubated on a rotating platform for 10 min at 4 C. The fragments were diluted 1:3 with 1.1% Triton-X100, 1.2 mM EDTA, 167 mM NaCl, 16.7 mM Tris-HCl (pH 8.0), 0.01% SDS, 1 mM PMSF and 1% protease inhibitor cocktail. The fragments were then incubated with 5 μl of rabbit polyclonal antibody to 3MeH3K9 (EMD Millipore) or IgG from rabbit serum (for negative control; Sigma) at 4 C overnight on a rotating platform. After that, 2 μg of blocked/washed Pansorbin Cells and 1 mM PMSF were then added and the lysate was incubated on a rotating platform for 15 min at room temperature. The immunoprecipitated pellets were then washed twice with 0.2% SDS, 2 mM EDTA and 50 mM Tris-HCl (pH 8.0) and thrice with 0.5 M LiCl, 1% Nonidet P-40 (Sigma), 1% sodium deoxycholate, 100 mM Tris-HCl (pH 9.0) and 1 mM PMSF. The complexes of antibody-protein-chromatin were then eluted by incubation with 100 μl 1% SDS and 50 mM NaHCO3 for 15 min at room temperature with rotation. Cross-links were reversed by adding 4 μl of 5 M NaCl and incubating samples for 4 h at 65 C. Fragment DNAs were then purified using a QIAquick PCR Purification Kit (Qiagen) following the manufacturer’s protocol. The presence of subtelomeric sequences of chromosomes 2, 8 and 12 in ChiP samples was detected using RT-PCR with the primer sets Chr2start, Chr8start and Chr12start shown in Table 1. The primer sequences were designed was detected using RT-PCR with the primer sets Chr2start, Chr8start and Chr12start shown in Table 1. The primer sequences were designed was detected using RT-PCR with the primer sets Chr2start, Chr8start and Chr12start shown in Table 1. The primer sequences were designed was detected using RT-PCR with the primer sets Chr2start, Chr8start and Chr12start shown in Table 1. The primer sequences were designed was detected using RT-PCR with the primer sets Chr2start, Chr8start and Chr12start shown in Table 1. The primer sequences were designed was detected using RT-PCR with the primer sets Chr2start, Chr8start and Chr12start shown in Table 1. The primer sequences were designed.
USA). Data are expressed as means ± SEM.

Results

Efficiency of siRNA knockdown of SUV39H1 and SUV39H2

SUV39KD pESLCs and SUV39KD PEFs were obtained by siRNA knockdown. The efficiency of siRNA knockdown was evaluated by the expression levels of SUV39H1 and SUV39H2 genes in SUV39KD pESLCs and SUV39KD PEFs in comparison with their respective controls. The primer sets for SUV39H1 and SUV39H2 were designed so that they amplified the regions that contain the sites siRNAs targeted. Therefore, the expressions of these two genes in effectively knocked-down cells were expected to be remarkably decreased compared with controls. The expressions of SUV39H1 (0.02 ± 0.01) and SUV39H2 (0.02 ± 0.01) in SUV39KD pESLCs decreased (P<0.001) approximately 50 and 14 times compared with those of the control (1.00 ± 0.14 and 0.28 ± 0.08, respectively) (Fig. 1A). Likewise, the expressions of SUV39H1 (0.82 ± 0.14) and SUV39H2 (0.49 ± 0.03) in SUV39KD PEFs decreased (P<0.05) approximately 3.5 and 12 times compared with those of the control (2.82 ± 0.62 and 5.89 ± 0.43, respectively) (Fig. 1B).

Gene expression in SUV39KD cells

Following the downregulation of SUV39H1 and SUV39H2 genes, the expressions of DNMT1, DNMT3A and DNMT3B also decreased in SUV39KD cells. Significant decreases (P<0.05) in expressions of DNMT1 (0.03 ± 0.01), DNMT3A (0.05 ± 0.01), and DNMT3B (0.03 ± 0.01) in SUV39KD pESLCs was observed compared with the control (0.45 ± 0.06, 1.13 ± 0.3 and 0.48 ± 0.18, respectively) (Fig. 1A). In contrast, expression levels of ATP5A1 (1.08 ± 0.15) and CMOS (0.47 ± 0.11) in SUV39KD pESLCs did not differ from those of the control (1.60 ± 0.37 and 0.37 ± 0.13, respectively) (Fig. 1A). Similar to the case in pESLCs, significant decreases (P<0.05) in expressions of DNMT1 (0.12 ± 0.01), DNMT3A (0.30 ± 0.01) and DNMT3B (1.00 ± 0.01) were found in SUV39KD PEFs compared with the control (7.14 ± 0.19, 3.51 ± 0.07 and 1.93 ± 0.02, respectively) (Fig. 1B). In opposition to that, expression levels of ATP5A1 (5.94 ± 1.03) and CMOS (1.09 ± 0.04) in SUV39KD PEFs were similar to those of the control (3.42 ± 0.56 and 0.75 ± 0.03, respectively) (Fig. 1B).

Telomere length in SUV39KD cells

SUV39KD pESLCs (37.1 ± 0.9 kb) and SUV39KD PEFs (22.1 ± 0.4 kb) had longer telomeres (P<0.05) compared with their respective controls (33.0 ± 0.7 kb and 17.8 ± 1.1 kb, respectively) (Fig. 2A, B).

Telomerase activity in SUV39KD cells

Relative telomerase activities were not different between SUV39KD pESLCs (10.4 ± 1.7) and the control (10.1 ± 1.7). Likewise, SUV39KD PEFs had similar relative telomerase activities (1.0 ± 0.3) compared with the control (1.0 ± 0.4).

Trimethylation of H3K9 in subtelomeric regions

The relative levels of 3MeH3K9 at subtelomeric regions of chromosome 2 (18.3 ± 3.4), chromosome 8 (4.5 ± 1.8) and chromosome 12 (2.5 ± 0.4) in SUV39KD pESLCs were significantly lower (P<0.05) than those of the control (46.3 ± 6.9, 11.0 ± 2.0 and 8.0 ± 0.6, respectively) (Fig. 3A). Similar to the case in pESLCs, significant decreases (P<0.05) in the 3MeH3K9 levels at subtelomeres of chromosome 2 (11.0 ± 1.3), chromosome 8 (1.1 ± 0.3) and chromosome 12 (4.8 ± 1.8) were observed in SUV39KD PEFs compared with the control (39.6 ± 6.5, 4.2 ± 1.0 and 11.2 ± 2.5, respectively) (Fig. 3B).

Discussion

In the present study, we successfully performed double knockdown of SUV39H1 and SUV39H2 by siRNA transfection in pESLCs and PEFs. In a preliminary experiment, we found that telomere length in SUV39KD pESLCs and SUV39KD PEFs did not significantly change compared with the control when the transfection was performed for only one passage (see Suppl Fig. 1: online only), although...
significant downregulations of *SUV39H1* and *SUV39H2* were found in SUV39KD cells compared with the controls. The possible cause for this observation is that although the siRNA treatment showed its effects on gene expressions in the very first trial, it might take a little longer to show effects on telomere length, since the cells were harvested only 24 h or 48 h after siRNA treatment for pESLCs and PEFs, respectively. However, SUV39KD PEFs started to show slow growth speed and changes in morphology after the third transfection. The siRNA knockdown of *SUV39H1* and *SUV39H2*, therefore, was carried out for three consecutive passages. To check the efficiency of siRNA knockdown, we designed primers for *SUV39H1* and *SUV39H2* so that they amplified the regions that contain the targeted sites of designed siRNAs. Therefore, the expression of *SUV39H1* and *SUV39H2* genes in effectively knocked down cells were expected to be remarkably decreased compared with controls. The remarkably low expressions of these two genes in SUV39KD pESLCs and SUV39KD PEFs compared with controls suggested that *SUV39H1* and *SUV39H2* were effectively knocked down.

We found that the knockdown of *SUV39H1* and *SUV39H2* in pESLCs and PEFs significantly increased the length of the telomere in pigs. This result is in agreement with a previous study in the mouse [8] and inconsistent with a study in humans [14]. This might suggest that the telomere biology of the pig is more similar to that of the mouse in respect to epigenetic/genetic regulation of telomere length. In the present study, the telomere in PEF control was 17.8 kb in length. This length is comparable with that of the eGFP-transduced fetal fibroblast cell line in a previous report, which had a telomere length of 18.5 kb [19]. We found that, in comparison with the PEF control, the telomeres in SUV39KD PEFs increased approximately 24%. Meanwhile, an approximate 12% increment in telomere length was observed in SUV39KD pESLCs compared with the pESLC control. These increments are comparable to those in the previous study in the mouse (from 11% to 28%) [8].

Fig. 2. Telomere length in (A) *SUV39H1* and *SUV39H2* knockdown (SUV39KD) porcine embryonic stem like cells (pESLCs) compared with the control; and (B) SUV39KD porcine embryonic fibroblasts (PEFs) compared with the control. Telomere lengths were determined by using a TeloTAGGG Telomere Length Assay kit. SUV39KD pESLCs and SUV39KD PEFs had longer telomeres compared with controls. A: Control DNA, DNA from immortal human cell lines provided in the kit; Control pESLCs, non-siRNA-treated pESLCs. B: Control DNA, DNA from immortal human cell lines provided in the kit; Control PEFs, non-siRNA-treated PEFs.

Fig. 3. Relative level of trimethylation of Histone H3 Lysine 9 (3MeH3K9) in the subtelomeres of chromosomes 2, 8 and 12 in (A) *SUV39H1* and *SUV39H2* knockdown (SUV39KD) porcine embryonic stem-like cells (pESLCs) compared with the control; and (B) SUV39KD porcine embryonic fibroblasts (PEFs) compared with the control. The relative levels of 3MeH3K9 in subtelomeric regions were determined by using a chromatin immunoprecipitation (ChIP) assay followed by RT-PCR analysis. A: The relative levels of 3MeH3K9 in the subtelomeric regions of chromosome 2, chromosome 8 and chromosome 12 in SUV39KD pESLCs were significantly lower than those of the control. B: Similar to this, significant decreases in 3MeH3K9 levels in the subtelomeres of chromosomes 2, 8 and 12 in SUV39KD PEFs were observed compared with the control. Different letters denote statistically significant differences between groups.
SUV39h1 and SUV39h2 are 3MeH3K9-specific HMTs in mammals [13]. Thus, a decrease in the 3MeH3K9 level was expected as SUV39H1 and SUV39H2 genes were downregulated. We examined the relative level of 3MeH3K9 in the telomere-adjacent regions on chromosomes 2, 8 and 12. These are GC-rich regions based on the Sscrof10.2 database. Significant reductions in the 3MeH3K9 level in these subtelomeric regions were observed in pESLCs and PEFs. The decreases in 3MeH3K9 at subtelomeric regions might be responsible for telomere lengthening, since this reduced chromobox proteins binding to the telomere [8] and Heterochromatin Protein 1, the chromobox homolog in *Drosophila*, plays an important role in telomere capping [20]. Further study on chromobox proteins is needed to confirm this hypothesis.

In mammals, DNA methylation requires 3MeH3K9 by SUV39h1 and SUV39h2 HMTs, suggesting that histone methylation directs DNA methylation at the pericentric chromatin [21, 22]. Downregulation of the genes SUV39H1 and SUV39H2, therefore, might also affect DNA methyltransferases. We revealed that the expressions of DNMT1, DNMT3A and DNMT3B were decreased in both SUV39KD pESLCs and SUV39KD PEFs compared with controls. The downregulation of DNMT1, DNMT3A and DNMT3B that occurred as a consequence of SUV39KD might be due to the fact that, at the enzyme levels, SUV39H1 and its interacting partner bind to and direct Dnmt1 of SUV39KD might be due to the fact that, at the enzyme levels, SUV39H1 and SUV39KD PEFs compared with controls. The downregulation were decreased in both SUV39KD pESLCs were found in SUV39KD cells in comparison with controls.

CMOS telomerase activity also contributes to the abnormal elongation of cells are needed to confirm this hypothesis. The possibility that a decrease in DNA methyltransferases might also be responsible for the abnormal lengthening of telomeres in SUV39KD cells in pigs. Further study on the protein levels of DNA methyltransferases and the changes in DNA methylation at subtelomeres in SUV39KD cells are needed to confirm this hypothesis. The possibility that telomerase activity also contributes to the abnormal elongation of telomeres is excluded, as we found that telomerase activities did not differ between SUV39KD pESLCs and the control or between SUV39KD PEFs and the control. It should be noted that no significant differences in expressions of the housekeeping genes ATP5A1 and CMOS were found in SUV39KD cells in comparison with controls. This result confirms that SUV39KD of pESLCs and PEFs seemed to specifically downregulate the expression of SUV39H1, SUV39H2, DNMT1, DNMT3A and DNMT3B but do not significantly alter the expressions of other essential genes in our study.

It also should be noted that we obtained similar results with another pESLC line and two other PEF cultures (data not shown). This may eliminate the possibility that all the observations might only apply to specific cell cultures.

In conclusion, double knockdown of SUV39H1 and SUV39H2 results in elongation of telomeres in pigs. This elongation might be not only related to the decrease of trimethylation of histone H3 lysine 9, but also linked to the reduction of DNA methyltransferases.

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