Introduction of a plasmid and a protein into bovine and swine cells by water-in-oil droplet electroporation

Takeshi ISHINO1, Hirofumi KURITA2, Rikio KIRISAWA3, Yoshinori SHIMAMOTO4, Rika NUMANO2 and Hiroshi KITAMURA1*

1)Laboratory of Veterinary Physiology, Departments of Veterinary Medicine, School of Veterinary Medicine, Rakuno Gakuen University, Ebetsu, Hokkaido 069-8501, Japan
2)Department of Applied Chemistry and Life Sciences, Graduate School of Engineering, Toyohashi University of Technology, Toyohashi, Aichi 441-8580, Japan
3)Laboratory of Veterinary Virology, Departments of Veterinary Medicine, School of Veterinary Medicine, Rakuno Gakuen University, Ebetsu, Hokkaido 069-8501, Japan
4)Laboratory of Animal Therapeutics, Department of Veterinary Science, School of Veterinary Medicine, Rakuno Gakuen University, Ebetsu, Hokkaido 069-8501, Japan

ABSTRACT. Instrument cost is a major problem for the transduction of DNA fragments and proteins into cells. Water-in-oil droplet electroporation (droplet-EP) was recently invented as a low-cost and effective method for the transfection of plasmids into cultured human cells. We here applied droplet-EP to livestock animal cells. Although it is difficult to transfec plasmids into bovine fibroblasts using conventional lipofection methods, droplet-EP enabled us to introduce an enhanced green fluorescent protein (EGFP)-expressing plasmid into bovine earlobe fibroblasts. The optimal transfection condition was 3.0 kV, which allowed 19.1% of the cells to be transfected. For swine earlobe fibroblasts, the maximum transfection efficacy was 14.0% at 4.0 kV. After transfection with droplet-EP, 69.1% of bovine and 76.5% of swine cells were viable. Furthermore, droplet-EP successfully transduced Escherichia coli recombinant EGFP into frozen-thawed bovine sperm at 1.5 kV. Flow cytometry analysis revealed that 71.5% of spermatozoa exhibited green fluorescence after transfection. Overall, droplet-EP is suitable for the transfection of plasmids and proteins into cultured livestock animal cells.

KEY WORDS: bovine cell, electroporation, sperm, swine cell
recombinant protein to bovine and swine cells by using droplet-EP.

**MATERIALS AND METHODS**

**Bovine ear-derived fibroblasts and swine ear-derived fibroblasts**

Bovine ear fibroblasts have been previously reported [19]. Pieces of swine earlobes were obtained from the Rakuno Gakuen Field Education Center (Ebetsu, Japan) when pigs (aged 3 days) were ear-notched. The swine earlobe tissue blocks were cut into fragments, and vigorously washed with phosphate-buffered saline (PBS) containing 200 U/ml penicillin, 200 µg/ml streptomycin, and 500 ng/ml amphotericin B (Nacalai Tesque, Kyoto, Japan). Subsequently, the tissues were treated with 2 mg/ml collagenase type I (Wako Pure Chemical, Osaka, Japan) in Tyrode’s solution and cut into small pieces using scissors. The tissue slices were further digested for 30 min at 37°C with strong agitation. After extensive washing with PBS and centrifugation, the pellet was resuspended in Dulbecco’s modified Eagle medium (DMEM) containing 10% fetal calf serum (FCS), 200 U/ml penicillin, 200 µg/ml streptomycin, and 500 ng/ml amphotericin, and filtered with a cell strainer (70 µm; Corning, New York, NY, USA) to remove debris. The cells were cultured in DMEM supplemented with 10% FCS at 37°C, 5% carbon dioxide, and 95% humidity.

**Enhanced green fluorescent protein (EGFP)-encoding plasmid**

The pEGFP-N2 plasmid was purchased from Takara Bio (Kusatsu, Japan). The plasmid was prepared from *Escherichia coli* (E. coli) DH5α using a FavorPrep Plasmid DNA Extraction Midi Kit (Favorgen, Ping-Tung, Taiwan).

**Water-in-oil droplet EP of EGFP-encoding plasmid to bovine or swine fibroblasts**

Droplet-EP was performed as previously reported with some modifications [24, 25]. One well of a 24-well cell culture plate was layered with 1 ml silicone oil (KF96-1; Shin-Entsu Chemical, Tokyo, Japan) on 1.5 ml of fluorocarbon oil (Fluorinert FC-96; 3M, Maplewood, MN, USA). The anode and cathode were set at a 6 mm interval. The parameters for each experiment are shown in Table 1. Bovine and swine fibroblasts were detached by treatment with trypsin, and washed with DMEM supplemented with 10% FCS twice. After counting cell number, the cell were resuspended in DMEM without FCS. To date, we employed DMEM as a vehicle of droplet-EP when we collected basic transfection data of mouse and human cells. Therefore, we used DMEM for transfection of the plasmid to bovine or swine fibroblasts. A droplet including the cells and transfected materials was gently added into the silicone oil phase in the well. Through a strong charge applied using a DC High Voltage power supply (HAR-30R10; Matusada Precision, Kusatsu, Japan), the droplet was elongated and attached to both electrodes, resulting in the generation of a short circuit. Subsequently, the droplet was retrieved and transferred to the culture medium.

**Microscopic observation**

Fibroblasts seeded on a collagen-coated cover glass slide were fixed with 4% paraformaldehyde for 10 min at room temperature. After a thorough wash with PBS, the cells were stained with Hoechst33342 (Thermo Fisher Scientific, Waltham, MA, USA) for 5 min at room temperature. The cover glass was embedded on the cell-seeded glass slide with 50% glycerol, and the cells were observed using a C2 confocal microscope (Nikon, Tokyo, Japan). Scanned images were analyzed using ImageJ software [42]. Transfection efficiency was determined as the ratio of the number of EGFP-positive cells to Hoechst33342-positive cells.

**Cell viability**

Cell viability was evaluated immediately after droplet-EP or conventional EP. Dead cells were stained with 0.5% trypan blue (Nacalai Tesque). Cell viability was calculated as the ratio of the number of non-stained cells to total cells.

**Transfection of EGFP-encoding plasmid to bovine or swine fibroblasts using NEPA21**

As a reference of gene transfection, we performed transfection of the EGFP-encoding plasmid using a NEPA21 electroporator (Nepagene, Chiba, Japan) according to the protocol provided by the manufacturer. Briefly, 1.0 × 10⁶ trypsinized bovine or swine fibroblasts and 10 µg of pEGFP-N2 plasmid in 100 µl of DMEM (Thermo Fisher Scientific) were placed in a 2 mm-gap cuvette (Nepagene) and pulsed using NEPA21. The parameters for the transfection pulse were as follows: poring pulse voltage: 275 V; pulse length: 2.5 msec; pulse interval: 50 msec; number of pulses: 2; decay rate: 10%; polarity +; and transfer pulse voltage: 20 V; pulse length: 50 msec; pulse interval: 50 msec; number of pulses: 5; decay rate: 40%; and polarity +/- of DMEM containing 10% FCS.

**Table 1.** Transfection condition

| Transfected cell | # of Cell (+10⁶) | Plasmid/Protein | Size of plasmid/protein (µl) | Amount of plasmid/protein (pmol) | Solvent | Droplet volume (µl) |
|------------------|-----------------|-----------------|-----------------------------|---------------------------------|--------|-------------------|
| Bovine fibroblast| 1.0             | pEGFP-N2        | 4.7 kb                      | 0.121                            | DMEM   | 3.0               |
| Swine fibroblast | 1.0             | pEGFP-N2        | 4.7 kb                      | 0.121                            | DMEM   | 3.0               |
| Murine sperm     | 5.0             | EGFP protein    | 27 kDa                      | 0.605                            | Opti-MEM® | 10.0            |
| Bovine sperm     | 5.0             | EGFP protein    | 27 kDa                      | 0.605                            | Opti-MEM® | 10.0            |

DMEM, Dulbecco’s Modified Eagle Medium; EGFP, enhanced green fluorescent protein.

doi: 10.1292/jvms.19-0475
Transfection of EGFP plasmid to bovine fibroblasts by lipofection

Bovine earlobe fibroblasts were transfected with pEGFP-N2 using Lipofectamine 2000 (Thermo Fisher Scientific), Lipofectamine LTX (Thermo Fisher Scientific), Fugene 6 (Promega, Madison, WI, USA), or Fugene HD (Promega) according to the instructions provided by the manufacturer. Briefly, 0.2, 0.1, 0.1, and 0.1 µg of the EGFP-encoding plasmid was encapsulated with Lipofectamine 2000, Lipofectamine LTX, Fugene 6, and Fugene HD, respectively. The liposome-and DNA complexes were subsequently supplemented to 1 × 10⁶ adherent bovine fibroblasts. After 4 hr incubation, the medium was exchanged to DMEM supplemented with 10% FCS. The cells were observed using a fluorescence inverted microscopy IX71 (Olympus, Tokyo, Japan) with a cooled CCD camera DP73 (Olympus) at 24 hr after the transfection.

Preparation of recombinant EGFP

Recombinant EGFP was prepared as previously described [48]. The pET-His6-GFP-TEV-LIC plasmid was obtained from Addgene (Cambridge, MA, USA). The plasmid was introduced to E. coli BL21 (DE3) (BioDynamics Laboratory, Tokyo, Japan). The plasmid-introduced E. coli clone was treated with 1 mM isopropyl β-D-1-thiogalactopyranoside (IPTG, Nacalai Tesque) for 3 hr. After centrifugation at 3,000 × g for 20 min, the bacterial pellet was lysed with 20% Triton X-100 on ice, followed by sonication using a VP-050 sonicator (Taitec, Koshigaya, Japan). After centrifugation at 13,000 × g for 30 min at 4°C, the supernatant was collected. For the purification of EGFP, HisPur Cobalt Resin (Thermo Fisher Scientific) was used according to the instructions provided by the manufacturer.

Droplet-EP of recombinant EGFP protein to mouse or bovine sperm

Mature (aged 12–15 weeks) male C57BL/6J mice (SLC, Hamamatsu, Japan) were sacrificed through cervical dislocation under anesthesia with 100 mg/kg of sodium pentobarbital. The sperm of mice was collected from the cauda epididymides. Frozen bovine sperm in straws was kindly gifted by Genetics Hokkaido Association (Sapporo, Japan). To recover sperm activity, both mouse and bovine sperm in Opti-MEM (200 µl), covered with liquid paraffin (Wako Pure Chemical), was pre-incubated at 37°C for 1.5 hr [45]. After centrifugation at 1,000 × g for 5 min, the sperm was resuspended in a drop of 10 µl Opti-MEM. To minimize lot difference, we selected Opti-MEM for a pre-conditioning and transfection medium. A droplet (3 or 10 µl) including the sperm and EGFP was gently floated in the silicone oil layer in the well. Immediately after droplet-EP was performed, the droplet was retrieved. The parameters for droplet-EP of EGFP protein to the sperm are listed in Table 1. The procedures of the animal experiments were approved by the Animal Ethics Committees of Rakuno Gakuen University (Permit Number: VH19A8).

Bovine sperm viability

Viability of bovine sperm was determined by the eosin-nigrosin staining method [4]. Briefly, the sperm was stained with 0.5% eosin Y (Muto Pure Chemical, Tokyo, Japan) for 15 sec, and subsequently stained with 0.5% nigrosin (Wako Pure Chemical) for 15 sec. The sperm was smeared on a glass slide and then air-dried. Since dead sperm was dyed red, sperm viability was calculated as the ratio of the number of non-stained cells to total cells. Viability was determined immediately after transfection.

Flow cytometry analysis

The transfected sperm was analyzed by flow cytometry to evaluate the efficacy of droplet-EP for the transfection of EGFP into sperm. The sperm was fixed in 4% paraformaldehyde for 10 min at room temperature. After washing twice with PBS, they were analyzed using a FACS Verse system (BD Biosciences, San Jose, CA, USA). The data were further analyzed using Flow Jo 10 (FlowJo, Ashland, OR, USA).

Statistical analysis

Statistical analysis was performed via one-way ANOVA corrected for multiple comparisons with the Holm-Bonferroni method, using Kaleida Graph software (Hulinks, Tokyo, Japan).

RESULTS

Droplet-EP efficiently transfects EGFP plasmid DNA into bovine fibroblasts

We previously reported that the transfection efficiency of plasmids into bovine fibroblasts was extremely low using commercially available lipofection reagents [19]. Hence, we initially attempted to transfect a mammalian expression plasmid into bovine fibroblasts using the droplet-EP technique. As previously reported [24, 25], we floated a droplet including the cells and plasmids in silicone oil (Fig. 1). Under the lower current, the droplet frequently moved back and forth between the electrodes. An increase in the electric field caused morphological changes to the droplet (flattening). Eventually the droplet attached to both electrodes and created a short circuit. For the transfection into bovine fibroblasts, we encapsulated 1.0 × 10⁶ cells and 0.12 pmol plasmid in a 3.0 µl droplet, and applied a voltage of 2.0, 3.0, 3.5, or 4.0 kV. Previously, we transected plasmids into human embryonic kidney cells using droplet-EP [25]. In this series, we found that droplet size is negatively correlated with transfection efficiency when voltage, cell number, and DNA amount were constant. Since 2–3 µl droplet was most effective for transfection of plasmids into 1.0 × 10⁶ cells, we first tried to transfect the EGFP-encoding plasmid using a 3.0 µl droplet. Electric current <2.0 kV failed to generate a short circuit, and did not result in transfection (Fig. 2A). In contrast, 3.0 kV caused a short circuit, followed by the expression of EGFP in 27.3 ± 9.7% of live cells (Fig. 2A and 2B). Both 3.5 and 4.0 kV transduced the plasmid into the cells to a significant
degree, despite the inversely proportional relationship between the transfection efficiency and voltage. Notably, cell viability was also inversely proportional to the voltage (Fig. 2C). Cell viability was 82.2 ± 2.5, 69.0 ± 4.1, 57.9 ± 0.9, and 49.4 ± 1.8% when the cells were subjected to droplet-EP at 2.0, 3.0, 3.5, and 4.0 kV, respectively. In a further step, we multiplied the efficiency and viability to calculate the yields of transfections.

Fig. 1. Configuration of a droplet-electroporation (EP) apparatus. (A) Positive and negative electrodes were connected to a high voltage direct current (DC) power supply. (B) A medium droplet containing cells and plasmids or recombinant proteins was floated in silicone oil and placed between the electrodes.

Fig. 2. Efficiency, viability, and yields of transfection of the pEGFP-N2 plasmid into bovine fibroblasts by droplet-EP. A total of 4 × 10^5 bovine earlobe fibroblasts were subjected to droplet- electroporation (EP) at the following settings: 2.0, 3.0, 3.5, or 4.0 kV. Cell viability was determined immediately after charging the cells through droplet-EP. Subsequently, the transfected cells were cultured for 24 hr. After staining the nuclei with Hoechst33342, the transfection efficiency was determined using a laser-scanning microscope. (A) Transfection efficiency was determined based on the proportion of enhanced green fluorescent protein (EGFP)-expressing cells in Hoechst33342-positive cells. (B) A representative image of EGFP-transfected cells. Scale bar=50 µm. (C) Cell viability was determined using the trypan blue staining method. (D) The yields of transfections were determined by multiplying the efficiency and cell viability obtained at each voltage. “Intact” indicates that droplet-EP was not performed. Values are means ± one standard deviation over three wells. *P<0.05 vs. intact cells.
cell viability at each voltage to more precisely determine the effectiveness of droplet-EP. These indices, namely transfection yield, resulted in transfection at each setting. As shown in Fig. 2D, 3.0 kV resulted in the greatest yield (19.1 ± 7.9%, n=3), followed by 3.5 kV and 4.0 kV (14.4 ± 3.7% and 9.5 ± 1.8%, respectively). On the other hand, transfection yield and cell viability of EP using the commercially available electroporator NEPA21 at the optimal setting was 40.2 ± 5.0 and 76.9 ± 2.6%, respectively. We also transfected the EGFP-encoding plasmid to the bovine fibroblasts by lipofection. Neither Lipofectamine 2000, Lipofectamine LTX, Fugene 6, nor Fugene HD effectively transfected the plasmid to the cells: Transfection efficacies of Lipofectamine 2000, Lipofectamine LTX, Fugene 6, and Fugene HD were 0.72 ± 0.87% (n=5), 0.00 ± 0.00% (n=5), 0.16 ± 0.38% (n=5), and 1.63 ± 0.79% (n=5), respectively. Collectively, these results indicate that the optimal condition of droplet-EP effectively transfects plasmid DNA into bovine fibroblasts at a comparable level with that reported using a commercially available electroporator.

**Droplet-EP efficiently transfects EGFP plasmid DNA into swine fibroblasts**

We subsequently attempted to transfet plasmids into swine cells. For this purpose, we isolated fibroblasts from pieces of swine earlobes using a method previously described [1]. Morphologically, swine earlobe fibroblasts were flatter than bovine earlobe fibroblasts, and were proliferative for ≥20 passages (Fig. 3A). In accordance with bovine fibroblasts, swine fibroblasts were not transfected with the pEGFP-N2 plasmid when droplet-EP was performed at the 2.0 kV setting (Fig. 3B). On the other hand, transfection efficiency was increased in a directly proportional manner to the voltage at ≥4.0 kV. At 4.0 kV, 18.3% of cells were transfected with the plasmid. Even at this intense voltage setting, >75% cells were alive, although cell viability was slightly higher at lower voltage settings (Fig. 3C; 90.5 ± 2.3%, 87.9 ± 1.3%, and 82.6 ± 2.2% at the 2.0, 3.0, and 3.5 kV settings, respectively). Given that viability of bovine earlobe fibroblasts was ~50% at the 4.0 kV setting, swine earlobe fibroblasts were resistant to relatively higher voltage. Figure 3D exhibits the yields of transfection to swine earlobe fibroblasts. The 4.0 kV setting resulted in the highest yield (14.0 ± 5.9%), followed by the 3.0 and 3.5 kV settings (11.3 ± 6.7% and 12.7 ± 6.2%, respectively). The transfection yields of the NEPA21 at the optimal condition were ~3-fold higher (40.6 ± 9.0%) than those obtained with droplet-EP. Meanwhile, cell viability (78.3 ± 3.8%) after EP using a NEPA21 was slightly less than that of droplet-EP at 4.0 kV setting. These results show that droplet-EP sufficiently transfected a plasmid into swine fibroblasts but that transfection efficacy was lower than that obtained with the commercially available electroporator.

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**Fig. 3.** Efficiency and cytotoxicity of transfection of the pEGFP-N2 plasmid into swine fibroblasts by droplet-electroporation (EP). Swine fibroblasts were isolated from earlobe skin, and subjected to transfection by droplet-EP. (A) Morphology of swine earlobe fibroblasts. Scale bar=50 µm. (B–D) Transfection efficiency (B), cell viability (C), and yields of transfection (D) were determined (as shown in Fig. 1). Values are means ± one standard deviation over three wells. *P<0.05 vs. intact cells.
Droplet-EP introduces EGFP protein into mouse and bovine spermatozoa

In the dairy scientific fields, gonadal cells were recently manipulated by gene editing [6, 46]. This requires that exogenous proteins or nucleotides are introduced into the cells. Notably, sperm cannot survive under cell culture conditions for >1 day [7]. Considering that cells typically require 24 hr to produce a sufficient amount of recombinant proteins, the introduction of a plasmid may not be applicable to sperm. Therefore, we attempted to introduce a purified protein into sperm using the droplet-EP approach. We are aware of no previous study investigating the transfection of recombinant protein to cells through droplet-EP. The present study was the first to introduce recombinant EGFP into mouse sperm which have previously been subjected to the introduction of proteins through conventional EP [40]. Droplet-EP at the 3.0 kV setting, which is the lowest voltage producing a short circuit using a 3.0 µl of sperm suspension, killed 91.3 ± 3.3% of total sperm (n=15) although 77.0 ± 9.1% (n=15) of sperm was alive without EP. To reduce the voltage producing a short circuit, we increased the size of the droplet to 10 µl, which included 2.7 µg of purified recombinant protein and 5.0 × 10⁵ spermatozoa. In this case, 1.5 kV sufficed to generate a short circuit through droplet-EP. Reduction of voltage to 1.5kV increased sperm viability to 55.2 ± 15.6% (n=15). Figure 4A shows a microscopic image of a sperm after transfection at this setting. A number of spermatozoa exhibited an intense green florescent signal in the head, middle piece, and tail part, suggesting the successful transfection of EGFP through the droplet-EP method. In agreement with this finding, flow cytometric analysis demonstrated that 24.9% of the cells showed a stronger green fluorescent signal compared with untransfected cells (Fig. 4B).

We subsequently examined whether droplet-EP may be applicable to protein transfection into bovine sperm. As with the transfection into mouse sperm, we encapsulated frozen-thawed bovine sperm (5.0 × 10⁵ cells) and 2.7 µg of recombinant EGFP in 10 µl of Opti-MEM, and applied a voltage (1.5 kV). Similar to the mouse sperm, a proportion of spermatozoa exhibited bright green fluorescence after droplet-EP with the EGFP (Fig. 4C). Moreover, flow cytometry demonstrated that 71.5% of the transfected cells showed a green fluorescent signal (Fig. 4D), and 24.9% showed intense green fluorescence, suggesting that the transfection efficacy was not constant among the sperm samples. Nevertheless, droplet-EP may be applied to the transfection of proteins into animal sperm.

![Image](image.png)

**Fig. 4.** Introduction of recombinant enhanced green fluorescence protein (EGFP) to mouse and bovine sperm. *E. coli* recombinant EGFP was transduced into mouse (A, B) and bovine (C, D) sperm by droplet- electroporation (EP) with a 10 µl droplet at the 1.5 kV setting. The nuclei were counterstained with Hoechst33342. (A, C) Representative images of mouse (A) or bovine (C) sperm transfected with recombinant EGFP. Arrowheads indicate EGFP-transfected spermatozoa. Scale bar=10 µm. (B, D) Histograms of the flow cytometric analysis of EGFP in mouse (B) or bovine (D) sperm. Mouse or bovine sperm were mixed with EGFP followed by droplet-EP (blue-filled area, Droplet-EP) or mixed with EGFP without droplet-EP (red-filled area, Intact). Representative data from three experiments are shown.
DISCUSSION

Transfection of gene fragments or recombinant proteins is currently widely performed in veterinary basic and clinical research. EP is one of the most effective methods for transfection. It is notable that EP allowed us to transfect plasmids into bovine cells, whereas lipofection methods were scarcely able to introduce them into the cells [19]. A major problem hindering the utilization of EP is cost. For example, the cost of NEPA21, one of the most widely used instruments for EP in livestock animal cells [17], is approximately 20,000 USD. Other popular instruments also typically cost 10,000–25,000 USD. The manipulation of cells with exogenous genes or proteins through EP is thus limited to research laboratories with relatively substantial funds. In contrast, droplet-EP only requires a DC power supply, which costs approximately 2,000 USD. Considering that droplet-EP does not require any special expendables, the present study provides basic information regarding a novel direction for transfection with high efficiency and low cost.

Although the proliferation of most somatic cells is determined by the Hayflick limit [15], numerous fibroblast-lined cells are still able to divide after 10 passages under appropriate culture conditions [23]. Moreover, fibroblasts express a relatively wide variety of signaling molecules. Thus, they are frequently used as atypical models for research into intracellular signaling cascades. For example, mouse derived-NIH3T3 and Swiss3T3 cells have been used to investigate the insulin, calcium, and Notch signals [32, 39, 43]. In these experiments, gene manipulations (e.g., overexpression, gene knockdown, gene knockout, and gene editing) were conducted by introducing exogenous gene fragments or recombinant proteins [3, 22]. On the other hand, non-negligible variations in cell signaling exist, which are derived from differences between species [2, 33]. Therefore, it is necessary to establish transfection methods for several species-derived fibroblasts. In this study, we determined the optimal setting of droplet-EP for bovine and swine fibroblasts; both are difficult to transfect through conventional lipofection [20, 34], and droplet-EP thus facilitates the study of cell signaling events in bovine and swine cells.

In previous studies, droplet-EP introduced plasmids into human fibroblasts cells at a relatively low voltage without generating a short circuit [25]. In contrast, we found that higher voltage was required for transfection into bovine and swine fibroblasts in the present study. Transfection of the plasmid was only observed in droplets which generated a short circuit between the electrodes. Thus, the generation of a short circuit rather than high voltage appears to be the crucial factor in introducing DNA and protein to livestock animal cells. We also observed the effective transfection of recombinant protein into mouse and bovine sperm by generating a short circuit at a low voltage setting, supporting this notion.

So far, a conventional electroporation was reported to effectively introduce recombinant Cas9 protein into mouse sperm [46]. Since transfection efficiency as well as viability of sperm have not been described in this report, we are not able to compare effectiveness of conventional EP and droplet-EP regarding protein transfection to sperm. In this report, 55% spermatozoa were alive after droplet-EP. Given that 23% sperm died during a freeze-thawing process, droplet-EP virtually damaged less than only one-third of the cells using an optimal setting (1.5kV). Additionally, we could introduce recombinant EGFP protein into significant proportion (mouse, 24.9%; cattle, 71.5%) of sperm. Nevertheless, we have to evaluate whether efficacy of droplet-EP is enough to proceed downstream applications such as genome editing and fertility treatments.

One of the disadvantages of droplet-EP might be limitation of cell number for transfection. This limitation is attributed to size of droplet. We had to set less than 10 µl/droplet between the electrodes at a 6 mm interval. In the droplet, we concentrated 1.0 × 10^5 fibroblasts, which is comparable to the number of cells for lipofection at 35 mm dish scale. Other popular instruments also typically cost 10,000–25,000 USD. The manipulation of cells with exogenous genes or proteins through EP is thus limited to research laboratories with relatively substantial funds. In contrast, droplet-EP only requires a DC power supply, which costs approximately 2,000 USD. Considering that droplet-EP does not require any special expendables, the present study provides basic information regarding a novel direction for transfection with high efficiency and low cost.

The reasons for the markedly lower transfection efficacy observed in bovine cells remain unclear [37]. Transfection efficiency is highly dependent on membrane permeability, which is determined by the size and number of pores on the plasma membrane [41]. In addition, the duration of high permeability is also a critical factor influencing transfection [44]. These factors are highly dependent on membrane flexibility, which is determined by the lipid constituents of the plasma membrane. Previous reports showed that the content of cholesterol is somewhat higher in bovine cells than in human and mouse cells [8, 27]. Cholesterol confers inflexibility to the plasma membrane [38]. Therefore, temporary deprivation of cholesterol may improve the transfection efficiency in bovine cells by droplet-EP as well as other transfection techniques.

Addition of electrolytes or dimethyl sulfoxide is considered another option for improving the transfection efficiency by droplet-EP [5, 36]. Alternatively, repeated generation of a short circuit may increase the transfection efficiency in livestock animal cells. Since swine fibroblasts are relatively resistant to high voltage, the repeated generation of short circuits appears to be particularly suitable for these cells. Further improvements may increase the effectiveness of nucleotide and protein transfection into livestock animals (e.g., bovine, swine, and chicken) by droplet-EP.

In this study, we transfected bovine and swine cells with an EGFP-expressing plasmid. The molecular weight of the EGFP-expressing plasmid was ~4.7 kb. Transfection efficiency is critically influenced by the size of the introduced materials [12]. For example, introduction of a 3.5 kb and a 10.9 kb plasmid to mesenchymal stem cells through conventional EP yielded a 42% and 5% transfection, respectively [26]. Further studies are warranted to optimize the transfection of larger products into bovine and swine cells.

In Japan, livestock animals are bred mainly through artificial insemination. A decrease in the rate of conception is one of the
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