Increased efficiency of transfection of murine hybridoma cells with DNA by electropermeabilization *

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Dispase-treated murine hybridoma cells (SP2/0-Ag14) were transfected with the G418 resistance gene bearing plasmid pSV2-neo by electropermeabilization with a high degree of efficiency. The cells were subjected to intermittent multiple high-voltage short duration (5 μs) DC pulses at intervals of 1 min in a weakly conducting medium followed by selection in G418-containing medium. The transfection medium, temperature, pulse duration, and voltage were empirically determined by preliminary electropermeabilization experiments. Increasing the number of pulses resulted in a higher percentage of transfected cells, but a decrease in the number of viable cells, with the optimal transfectant yield resulting when five pulses of 10 kV/cm were administered. This method allows the rapid and efficient injection of DNA into mammalian cells, and permits the rapid production of stable, drug resistant hybridoma cell lines for use in subsequent fusion experiments.

Key words: DNA transfection; Electropermeabilization; Eukaryotic cell; Hybridoma; Drug resistance

Introduction

The injection of foreign DNA into eukaryotic cells, and its subsequent expression in these cells is a powerful research tool with wide application in cellular and molecular biology. Using this technique, it is possible to rapidly render otherwise susceptible cell lines resistant to antibiotics by injection and subsequent genomic integration and expression of bacterial antibiotic resistance genes. One important use for such drug resistant cells is the production of ‘fusomas’ or hybrid hybridomas (Milstein et al., 1983; Semenenko et al.; 1985, Suresh et al., 1986) which may then be conveniently selected after somatic cell hybridization in appropriate antibiotic-containing medium (Lanza­vecchia et al., 1987). A number of such drug resistance genes, incorporated in well characterized plasmid vectors are now readily available and functional in mammalian cells after integration.

We have recently refined and optimized the methodology for insertion of the plasmid vector containing the bacterial drug resistance gene, neomycin phosphotransferase (pSV2-neo) (Southern and Berg, 1982) into the non-secreting mammalian hybridoma cell line, SP2/0-Ag14 (Sp2/0) by means of electropermeabilization, thus enabling the selection of transfectants in growth medium supplemented with the antibiotic G418.

The crucial step in this method is the production of reversible electrical breakdown of the cell
membrane, which is achieved when the cell is exposed to a field pulse of high intensity (kV/cm range) and short duration (ns to \( \mu \text{s} \) range) (Zimmermann et al., 1974a,b; Zimmermann, 1986). When the total membrane potential exceeds a critical value of approximately 1 V at 20°C or 2 V at 4°C the membrane is transiently 'permeabilized' permitting the influx of DNA, proteins, or other substances into the cell (Zimmermann et al., 1976, 1981; Zimmermann, 1982; Stopper et al., 1985, 1987). Longer duration and/or higher intensity pulses will result in irreversible electrical breakdown with consequent cell lysis (Zimmermann, 1982, 1986). The molecular events that follow electrical breakdown, culminating in membrane permeabilization are poorly understood and opinion remains divided regarding the possible formation of transient channels or 'pores' in the cell membrane as a consequence of electrical field application, i.e., so-called 'electroporation' (Neumann et al., 1982; Zimmermann, 1982; Sowers, 1985; Stopper et al., 1987). The presence of such channels alone could not account for the passage of whole cells through the cell membrane and it seems more likely that the application of the electrical field temporarily disrupts the orderly arrangement of the phospholipid bilayer and other membrane components, i.e., permeabilization of the membrane in areas where the field strength is sufficiently high (Zimmermann, 1983). Nevertheless, existing knowledge permits cogent manipulation of the process such that a variety of substances may be introduced into many cell types (Auer et al., 1976; Zimmermann and Pilwat, 1976; Zimmermann et al., 1980; Schussler and Ruhnstroth-Bauer, 1984).

We have previously shown that this method may be used for the transfection of yeast (Broda et al., 1987) and murine fibroblasts (Stopper et al., 1975; Zimmermann et al., 1976), with an efficiency greater than that reported for chemical transfection methods (Graham and Van der Eb, 1973; Pellicer et al., 1980; Berman et al., 1984). In the experiments reported here, high numbers of stable transformants were obtained in the presence of very low plasmid concentrations by electroporation using a series of 5-10 pulses with the cell suspension maintained at 4°C, each pulse being administered at intervals of 1 min. This delay is required to allow sufficient rescaling of the electropermeabilized cell membrane such that the cumulative toxicity of multiple pulsations does not result in excessive cell death. With some modifications, this method was used to efficiently generate large numbers of stable G418 resistant murine hybridoma cells, SP2/0.

**Materials and methods**

The non-secreting murine hybridoma cell line, SP2/0 (SP2/0-Ag14) was obtained from ATCC (Bethesda, MD) and cultured in complete growth medium (CGM) as follows: RPMI 1640 (Boehringer, Mannheim) supplemented with L-glutamine 2 mM (Boehringer, Mannheim), 10% fetal bovine serum (Boehringer, Mannheim), and 5 \( \times \) 10^{-3} M 2-mercaptoethanol. The cells were grown at 37°C in an incubator supplemented with 5% CO₂. A subclone of these cells chosen for the subsequent experiments was found to die rapidly in G418 (Gibco, Grand Island, NY) supplemented medium at concentrations of less than 100 μg/ml. Cells were harvested in log growth phase at concentrations of 3 \( \times \) 10^5/ml and incubated for 1 h in complete growth medium supplemented with dispase (grade I) 0.1 mg/ml (Boehringer, Mannheim). The cells were then washed once in chilled (4°C) CGM followed by 1–2 washes in pulse medium, the composition of which is as follows: inositol (Serva, Heidelberg) 0.22 M, KCl 30 mM, KH₂PO₄/K₂HPO₄ 1.1 mM, pH 7.2 then resuspended in pulse medium at a final concentration of 0.5–1.0 \( \times \) 10^6/ml.

The plasmid pSV2-neo was linearized by digesting a 50 μg aliquot for 3 h at 37°C with 3 \( \times \) 50 U of EcoRI (BRL, Bethesda, MD) added at 0, 1 and 2 h. The linearized plasmid was added to the final cell suspension at a concentration of 1.0 μg/ml excepting those experiments which were performed in the absence of plasmid DNA to serve as controls. 1.2 ml of the cell suspension was then loaded into the chamber of a high voltage electric discharge apparatus which was maintained at 4°C.

In brief, the apparatus used consists of two flat stainless steel electrodes mounted in parallel in a plexiglass chamber at a distance of 3.3 mm and cooled by a Peltier element. The total volume of
the chamber is 1.2 ml. The chamber is connected to a capacitor by a switch, which is, in turn, connected to a voltage generator. When the switch is closed, the capacitor is discharged exponentially (Zimmermann, 1983).

A varying number (0–10) of short duration, high voltage pulses (10 kV/cm for 5 μs) were then applied at intervals of 1 min. After pulsing, the cells were gently flushed from the pulse chamber and transferred into warm phenol-free RPMI 1640 medium (Biochrom, Berlin) and allowed to 'reseal' for 10 min at 37°C. The cells were then washed and resuspended in CGM, examined microscopically, counted and distributed into the wells of 24-well culture plates (Greiner, Nurtingen) at a concentration of 10^4/ml. Each well of these plates is marked by a raised 4 x 4 grid to prevent 'cross-contamination' of G418-resistant colonies and to facilitate recording of colony numbers. Numbers of viable cells were determined by microscopic examination using a phase contrast microscope and cell counts were determined using a hemocytometer.

48 h after pulsing, G418-supplemented CGM was added such that the final concentration of G418 was 250 μg/ml. In preliminary experiments, this dose was determined to inhibit the growth of non-transfected SP2/0-Ag14 cells, while minimally inhibiting the growth of transfected cells. The culture medium was changed twice weekly thereafter with the same dose of G418 being maintained. After 12 days the plates were examined with an inverted phase-contrast microscope and the number of viable colonies were counted and the efficiency of transfection, i.e., the percent viable G418-resistant colonies obtained were calculated. Some of these colonies were grown for a longer period in G418 (more than 28 days) and continued to exhibit G418 resistance.

### Results

The number of cells surviving pulsing and the percent of resulting stable G418-resistant colonies from a representative set of experiments are given in Table I. With the voltage and pulse duration held constant at 10 kV/cm and 5 μs, respectively, it will be seen that as the number of pulses administered increases, the number of cells surviving immediately after pulsation diminishes, while the percent of stable G418-resistant cells (as a function of the number of surviving cells) rises (Fig. 1). The absolute number of resistant colonies diminishes beyond five pulses, however, owing to the smaller number of surviving cells (Fig. 2).

There was some interexperimental variation in the optimal yield of resistant clones (varying from 0.4 to 4.8% in a total of five experiments) depending upon the condition of the cells at the time of

**TABLE I**

**RESULTS OF A TYPICAL TRANSFECTION EXPERIMENT**

<table>
<thead>
<tr>
<th>Number of pulses administered</th>
<th>Plasmid added</th>
<th>Cells before pulsing (x10^5)</th>
<th>Cells after pulsing (x10^5)</th>
<th>% Surviving cells post-pulse</th>
<th>Number of G418-resistant colonies</th>
<th>% G418-resistant colonies</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>–</td>
<td>9.0</td>
<td>8.8</td>
<td>97.8</td>
<td>0</td>
<td>0</td>
</tr>
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<td>7.5</td>
<td>83.3</td>
<td>36</td>
<td>0.005</td>
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<td>+</td>
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<td>4.1</td>
<td>64.1</td>
<td>96</td>
<td>0.023</td>
</tr>
<tr>
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<td>3.7</td>
<td>61.6</td>
<td>401</td>
<td>0.11</td>
</tr>
<tr>
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<td>2.8</td>
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<td>0.24</td>
</tr>
<tr>
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<td>658</td>
<td>0.30</td>
</tr>
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<td>1.5</td>
<td>25.0</td>
<td>766</td>
<td>0.51</td>
</tr>
<tr>
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<td>+</td>
<td>6.0</td>
<td>0.6</td>
<td>10</td>
<td>474</td>
<td>0.79</td>
</tr>
</tbody>
</table>

* Total number of viable cells (as determined by phase contrast microscopy) flushed from the pulse chamber after administration of the indicated number of pulses.

b Number of colonies of SP2/0-Ag14 cells observed after 12 days of culture in CGM supplemented with G418.

c G418-resistant colonies/total viable cells post-pulse administration (a) x 100.
Fig. 1. Percent of cells surviving electroporation as determined by microscopical examination immediately following application of varying numbers of 10 kV/cm pulses, and percent stable G418-resistant transfectants obtained after selection in G418-supplemented CGM (as a percentage of surviving cells initially cultured).

Fig. 2. Absolute numbers of stable G418-resistant colonies selected in CGM supplemented with G418 in a typical experiment after the application of varying numbers of 10 kV/cm pulses. The optimal number of colonies (766) were obtained when five pulses were administered.

Some of the cells bearing nonintegrated pSV2-neo are able to temporarily survive in G418 due to the transient synthesis of neomycin phosphotransferase. However, for most purposes, only those cells which are truly resistant to the selecting drug, i.e., stable transfectants are of interest. It is likely that cells exhibiting active growth in high concentrations of G418 beyond 12 days post transfection had integrated the plasmid into their genome and were therefore resistant.

It is possible that higher transfection efficiencies might have resulted if higher concentrations of plasmid were used but the addition of carrier DNA did not increase the efficiency of transfection in preliminary experiments (data not shown). However, those cells undergoing sham transfection in the presence of plasmid DNA had a much lower survival as compared with these cells undergoing sham transfection in the absence of plasmid DNA (Table I) suggested that increasing the DNA concentration may have had deleterious consequences.

A small number of G418-resistant clones were consistently found in the experiments wherein pulses were applied in the absence of plasmid DNA (Table I). This occurred much less frequently in the control groups that were not pulsed, although it was seen when plasmid was added to cell suspensions that were sham pulsed. This latter observation is explained by the spontaneous transfer of plasmid across the cell membrane. The increased frequency of G418 resistant cells after puling in the absence of plasmid may be the result of an increased frequency of mutation induced by electrical field interactions with DNA, although there is no experimental evidence to support this to date.

Discussion

A number of methods for DNA transfection of a variety of mammalian cell lines have been previously described including the use of DEAE dextran (McCutchan and Pagano, 1968), calcium phosphate coprecipitation (Graham and Van der Eb, 1973), direct transfer by bacterial protoplast fusion (Schaffner, 1980), microinjection (Yamamoto et al., 1982), liposome mediated transfer
(Schaeffer-Ridder et al., 1982), retroviral vectors (Gruber et al., 1985), laser-mediated transfer (Kurata et al., 1986) and electroinjection methods (Zimmermann, 1974a). Most of these methods were developed for use with adherent cells in monolayer culture and their efficiency is markedly reduced when they are applied to lymphoid cells in suspension (Rice and Baltimore, 1982; Oi et al., 1983). Some of the more exotic techniques requiring lasers, liposomes or microinjection apparatus are technically demanding and have not enjoyed widespread application, while retroviral transfer requires extensive DNA manipulation.

The electropermeabilization method which was first introduced more than 12 years ago by Zimmermann and coworkers (1974a,b) has been consistently more efficient than chemical methods for the transfection of mammalian cells, is reproducible and is technically easy. Previous applications of this method required relatively high concentrations of DNA (Pellicer et al., 1980; Zimmermann, 1986), have produced suboptimal efficiencies of stable transfectants (Pellicer et al., 1980; Toneguzzo et al., 1986) and have not been reported to be successful for the transfection of hybridomas. The results reported here demonstrate that when appropriate pulse medium, temperature and field conditions are maintained, the electropermeabilization method may be used to derive large numbers of stable transfected hybridoma cells for use in subsequent experiments, and that relatively small amounts of DNA may be used.

The electrolyte concentration of the pulse medium is of particular importance. The addition of KCl 30 mM facilitates an asymmetrical breakdown of the cell membrane, such that electropermeabilization of only one ‘hemisphere’ of the pulsed cells results (Zimmermann and Stopper, 1987). Thus, the restoration of the original membrane impermeability following breakdown is greatly enhanced, and this is especially so at low temperatures. In this manner, cell survival increases greatly, without compromising entry of DNA.

The application of pulses in an iterative fashion with sufficient interval time for membrane recovery (1 min) to occur, while maintaining the cells at 4°C is key to obtaining optimum yields. Several pulses of slightly supercritical field strength are required in order to permeabilize sufficiently large areas of membrane to allow entry of plasmid DNA owing to the angular dependence of the breakdown voltage (Zimmermann et al., 1974a, 1980). Multiple pulse application in this fashion is best performed with relatively long interval times between consecutive pulses since Brownian motion and cell rotation will allow exposure of different cell membrane areas to electropermeabilization, increasing efficiency of transfection, while reducing cell injury.

Moreover, without sufficient time for resealing to occur between pulses, there would not be an adequate increment in the cell membrane potential in response to subsequent pulses. This would result in the flux of current through the still permeabilized area of the membrane, injuring the cell.

Maintenance of the cell suspension at relatively low temperature allows maximal duration of the permeabilized membranes areas after pulsing. This facilitates the entry of DNA into the cell, while diminishing premature cell resealing.

It should be noted that the best growth conditions for particular cell lines selected for transfection, as well as the optimal concentration of the selecting drug must be carefully chosen if the maximum number of transfected clones are to be obtained while minimizing the number of cells lost to nonspecific toxicity, and inhibiting the growth of non-transfected cells. This is usually best accomplished by using cells in log growth phase, and by performing a preliminary experiment wherein the lowest concentration of selecting drug that inhibits the growth of the cell line is determined.

It will be seen that the optimal number of resistant colonies is obtained when five pulses are administered (Fig. 2). It will be noted that a small number of resistant colonies were found after sham electroinjection when the cells are suspended in the presence of plasmid, but the number of transfectants is, of course, low. A small number of resistant colonies were also found in the multiple pulse sham transfection, even though no plasmid was present in the chamber. This might have been the result of the appearance of spontaneous mutants resistant to G418. The fact that fewer or
no G418-resistant cells were found among those cells which were not pulsed, and had no exposure to plasmid suggests that electric field interactions with DNA may cause an increased mutation frequency.

The addition of plasmid to the final cell suspension consistently resulted in a somewhat greater cell loss even when pulses are omitted. This might be a result of the general increased 'cell stickiness' that results when DNA is present in the cell suspension. The cell clumping that results may lead to increased cell susceptibility to electrical pulse toxicity. Alternatively, the presence of higher amounts DNA on the cell surface may render the cells more susceptible to electrical field toxicity or increase the likelihood of DNA integration into sites which are not compatible with survival. We have previously noted that the use of higher concentrations of DNA resulted in reduced yields of murine fibroblasts (Stopper et al., 1985).

As previously noted by our group (Stopper et al., 1985) and others (Pellicer et al., 1980) the use of linearized DNA is several fold more efficient than circular, i.e., 'native' plasmid DNA. This might be due to increased efficiency of DNA injection, integration or both. Our results would indicate that the use of carrier DNA is not mandatory when µg/ml quantities of plasmid DNA are available.

From the foregoing discussion, it is clear that one must carefully consider the interaction of the various parameters which come into play during the process of electroinjection, if one hopes to obtain high efficiency yields. The lower transfection efficiencies reported by other authors using various electroinjection protocols (Potter et al., 1984; Toneguzzo et al., 1986) may in part be attributed to lack of attention to one or more of the aforementioned considerations. Only Chu et al. (1987) have reported stable transfection rates as high as 1%, but this was achieved in a variety of adherent monolayer cells by applying very long duration pulses (7 ms) of relatively low voltage and using DNA concentrations of 20 µg/ml with 500 µg/ml carrier DNA. Such long pulse durations would be expected to result in irreversible membrane breakdown and to result in the death of many otherwise viable cells. The fact that many of the transfectants derived from these experiments survived only transiently in the selection medium may suggest that they sustained injury which was ultimately lethal.

With appropriate application of the method described here, we believe that it should be possible to efficiently obtain large numbers of stable transfected mammalian cell clones to be used in subsequent fusion experiments or for other purposes, even when comparatively small numbers of cells and small amounts of DNA are available, and possibly where other methods have failed.

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