Generation of $\rho^0$ cells utilizing a mitochondrially targeted restriction endonuclease and comparative analyses

Alexandra Kukat1,2, Christian Kukat1, Jan Brocher3, Ingo Schäfer1, Georg Krohne4, Ian A. Trounce5, Gaetano Villani6 and Peter Seibel1,*

1Molecular Cell Therapy, Center for Biotechnology and Biomedicine, Faculty of Medicine, Universität Leipzig, Deutscher Platz 5, 04103 Leipzig, Germany, 2Karolinska Institutet, Department of Laboratory Medicine, Division of Metabolic Diseases, Novum, 141 86 Stockholm, Sweden, 3Department of Cell and Developmental Biology, Division of Electron Microscopy, Biocenter of the University of Würzburg, Am Hubland, 97074 Würzburg, Germany, 5Centre for Clinical Neurosciences, The University of Melbourne Department of Medicine, St Vincent’s Hospital, Fitzroy, Victoria 3065, Australia and 6Department of Medical Biochemistry, Biology & Physics, University of Bari, Piazza Giulio Cesare, 11, 70124 Bari, Italy

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ABSTRACT

Eukaryotic cells devoid of mitochondrial DNA ($\rho^0$ cells) were originally generated under artificial growth conditions utilizing ethidium bromide. The chemical is known to intercalate preferentially with the mitochondrial double-stranded DNA thereby interfering with enzymes of the replication machinery. $\rho^0$ cell lines are highly valuable tools to study human mitochondrial disorders because they can be utilized in cytoplasmic transfer experiments. However, mutagenic effects of ethidium bromide onto the nuclear DNA cannot be excluded. To foreclose this mutagenic character during the development of $\rho^0$ cell lines, we developed an extremely mild, reliable and timesaving method to generate $\rho^0$ cell lines within 3–5 days based on an enzymatic approach. Utilizing the genes for the restriction endonuclease EcoRI and the fluorescent protein EGFP that were fused to a mitochondrial targeting sequence, we developed a CMV-driven expression vector that allowed the temporal expression of the resulting fusion enzyme in eukaryotic cells. Applied on the human cell line 143B.TK$^-\rho^0$ cell line was characterized in great detail thereby releasing interesting insights into the morphology and ultra structure of human $\rho^0$ mitochondria.

INTRODUCTION

Mitochondria are organelles that can be found in most eukaryotic cells. Mitochondria harbour critical biochemical processes, such as the Krebs cycle or the aerobic energy supply of the cell. More recently, however, it was shown that they are key players in the ageing process and the programmed cell death. One factor closely linked to all these functions is the genome of the organelle, the mitochondrial DNA. Numerous neurological and neuromuscular diseases presenting with a variety of symptoms have been associated with mutations of the mitochondrial genome (1). Mitochondrial encephalopathies for example are a class of diseases which result from dysfunction of the mitochondria’s oxidative phosphorylation system (OXPHOS). Under regular conditions, this system takes responsibility for cellular respiration and energy production. The associated diseases exhibit disorders manifesting in tissues with high aerobic metabolic demands (e.g. brain, skeletal muscle, heart). The OXPHOS involves five enzyme complexes that assemble from subunits encoded by the nuclear DNA (nDNA) and the mitochondrial DNA (mtDNA). The mitochondrial genome is organized
in a circular fashion, encompassing 16 569 bp (according to 2) and encoding 13 polypeptides involved in OXPHOS, a set of 22 essential tRNAs as well as the large (16S) and small (12S) ribosomal RNA required for mitochondrial translation. Although the mtDNA was the first eukaryotic genome completely sequenced, almost nothing is known about the processes of how nucleus and mitochondria interact, of how mtDNA and nDNA gene expression is coordinated or how mtDNA is maintained within the cell (a process strictly driven by nuclear factors). Resulting from this lack of knowledge, very little is known about the molecular mechanisms leading to human diseases as a result of mtDNA damage. This is due, in part, to our inability to study regulatory and developmental mitochondrial processes under experimental conditions.

One effort to overcome these difficulties was an experiment carried out by King and Attardi (3). A human osteosarcoma cell line (143B.TK⁻, derived from an osteosarcoma) was treated over 4–6 weeks with a low dosage of ethidium bromide. It is known from yeast experiments (4–6) that this reagent interacts with mtDNA to form complexes that interfere with DNA replication. After a series of cell divisions, the endogenous mtDNA is lost, while nuclear DNA is maintained. Since the loss of mtDNA extinguishes the oxidative phosphorylation system after a few days, cells devoid of endogenous mtDNA (so called \( \rho^0 \) cells) need supplementation with nutrients to sustain viability. This is achieved by adding pyruvate and uridine to the growth medium so that the energy demand of the cell is satisfied additionally to the possibility of generating pyrimidines in spite of the inhibition of dihydroorotate dehydrogenase (DHODH) due to loss of respiratory chain electron transfer.

Additionally, other reagents were tested to generate \( \rho^0 \) cells that interfere with mtDNA replication (ditercalinium, ddC, etc.) (7–10). However, all chemicals used presented severe disadvantages, such as mutagenic effects or the induction of mdr (multi-drug resistance) family gene expression so that the inhibitory effect on mtDNA replication was abolished. Moreover, the major limitation of these methods is their inapplicability to different cell lines. For instance, although there are no published accounts of rat cell lines that interfere with mtDNA replication (ditercalinium, ddC, etc.) (7–10), the reagents shown in Table 1 were carried out to characterize the proliferative, metabolic and morphological changes due to a \( \rho^0 \) state of these cells. Therefore a 143B.TK⁻ \( \rho^0 \) cell line generated by incubation with low doses of ethidium bromide and the newly developed 143B.TK⁻ K7 cell clone were compared to wild-type cells for growth rate, glucose consumption as well as lactate and proton production rate.

**MATERIALS AND METHODS**

**Cloning**

Cloning was performed according to standard procedures, and all polymerase chain reaction (PCR) products were verified by sequencing. The plasmid pAN4 [a kind gift of A. Kiss and constructed for the EcoRI gene was used as template in a PCR with following primers: EcoRIR-001-FOR (2) 5'-catggaca getgacaagtgtgtaaatagtaaaac-3', which adds a short sequence complementary to the 3'-end of the EGFP gene and EcoRIR-834-REV (2) 5'-ggccaaatcttagatgtaag-3' which generates a NotI-restriction site. The EGFP gene from the plasmid pEGFP-Mito (Clontech) was amplified with the primers pEGFP-Mito-0597-FOR 5'-ggccaaatcacttagatgtaagctgttc-3' which generates a NotI-restriction site and pEGFP-Mito-1421-REV 5'-cttgta catctgctccgtc-3' which amplifies the EGFP without the stop-codon. Both fragments were utilized in a recombinant PCR and the phosphorylated product was sub-cloned in the PvuII-linearized vector pTRE2hyg. The fusion gene product consisting of EGFP- and EcoRI gene was cloned with Agel and NotI into pEGFP-Mito resulting in the vector pMEE-con.

**Cell culture, transfection and mitochondrial staining**

Human osteosarcoma cells 143B.TK⁻ (ATCC CRL-8303) were cultured under standard conditions in Dulbecco's modified Eagle's medium (high glucose) with GlutaMAX (Invitrogen) supplemented with 10% fetal calf serum, 1% bromdeoxyuridine, 100 U penicillin and 100 \( \mu \)g/ml streptomycin. The \( \rho^0 \) cell line referred to as 143B.TK⁻ \( \rho^0 \) was generated by incubation of 143B.TK⁻ with low doses of ethidium bromide. \( \rho^0 \) cell lines of the parental 143B.TK⁻ and cells after transfection procedure were maintained as indicated with additional supplementation with 100 \( \mu \)g/ml pyruvate and 50 \( \mu \)g/ml uridine.

Transient transfections of 143B.TK⁻ and 143B.DsRed1-Mito (143B.TK⁻ cells stably expressing mitochondrially targeted DsRed1 for co-localization studies with pMEE-con) were performed using Effectene (Qiagen) according to manufacturer's conditions. Geneticin selection with 700 \( \mu \)g/ml was carried out 24 h after transfection for 5 days. An isolated clone named as 143B.TK⁻ K7 was aliquoted and metabolically tested for existing oxidative phosphorylation with DMEM (high glucose) with GlutaMAX, 10% dialysed FCS (3), with 1% bromdeoxyuridine, 100 U penicillin and 100 \( \mu \)g/ml streptomycin without pyruvate and uridine for 30 days.

Two other cell lines were used to extend the applicability of our approach to rodent cells: the mouse line...
LMTK\(^{-}\) (ATCC CCL-1.3), and the rat line NRK52E (ATCC CRL-1571).

MitoTracker CMXRos (Molecular Probes) was used to label mitochondria according to manufacturer’s protocol at a concentration of 100 nM.

FACS analyses
Samples of transfected cells (2 \times 10^6 cells/ml in 0.05\% BSA in PBS) were analysed on a flow cytometer and its software (FACSCAN, FACS Vantage SE, Cell Quest Software, BD Biosciences). Acquisition of data was performed by gating more than 100,000 cells expressing the mitochondrial targeted restriction endonuclease fused to EGFP with the flow cytometer.

Characterization of clones by PCR
To verify the loss of all endogenous mtDNA in the resulting 143B.TK\(^{-}\) K7 cell clone the genomic DNA was isolated and used as a template in a PCR. Controls were accomplished by using the genomic DNA isolated from the 143B.TK\(^{-}\) wild type and a corresponding \(\rho\)\(^0\) cell-line generated by incubation with ethidium bromide as described before. The amplified region of the mtDNA does not contain EcoRI sites. Moreover, the primers selected anneal within the D-loop area encompassing the origin (O\(_{\text{H}}\)) of mtDNA replication. Persisting mtDNA molecules would be expected to contain this region to further possess DNA replication fidelity. The following primers binding in the mtDNA region were utilized: Mmus-Histon 3a-001-For \(5\'-\text{atgagctcatgaccgagaattccacgtccg-3'}\) and Mmus-Histon 3a-411-Rev \(5\'-\text{ttaagcacgttctccgcgtatgc-3'}\). The control-PCR was performed with primers that amplify the human histone H1 gene (Histon H1 5\'- 5\'-atgagctcatgaccgagaattccacgtccg-3' and Histon H1 3\'- 5\'-atccgggcaaacttcttcttgcc-3'). The annealing temperature was 55\(^\circ\)C for 35 cycles with an elongation time of 1 min.

Similar analyses were carried out on rat and mouse DNA extracted from LMTK\(^{-}\) and NRK52E cell lines. Primers utilized for mtDNA amplification were Rnor-15525-For \(5\'-\text{ctataaatcacaacaactgtc-3'}\) and Rnor-1214-Rev \(5\'-\text{ttaaagctcgtacaaagcagactgcc-3'}\) (rat cell line) and Mmus-16033-For \(5\'-\text{tatccg-3'}\) and Mmus-1174-Rev \(5\'-\text{tcaatactttagataaatg-3'}\) (mouse cell line). Histones were amplified utilizing primers Rnor-Histon3H3.3-001-For \(5\'-\text{atggccgaaccaagcagacc-3'}\) and Rnor-Histon3H3.3-539-Rev \(5\'-\text{atggcccgaaccaagcagacc-3'}\) and Mmus-Histon3a-001-For \(5\'-\text{atcccgggcaaacttcttcttgcc-3'}\) and Mmus-Histon3a-411-Rev \(5\'-\text{atggccgaaccaagcagacc-3'}\) (mouse cell line).

Southern blot analysis
Total DNA (15\,\mu g) extracted from cells was digested with the restriction endonucleases BamHI or PvuII. Restricted fragments were separated on a 0.6\% agarose gel, transferred onto a nylon membrane and hybridized with a digoxigenin-dUTP-labelled human mtDNA probe amplified with mtDNA primers annealing to nucleotides 4831–4846 (For) and 5628–5651 (Rev). The membrane was washed, and the fragments were detected with the DIG Nucleic Acid Detection Kit (Roche Applied Sciences) according to the manufacturer’s instructions. Chemoluminescence of the fragments was analysed with a bioimaging system.

Metabolite analyses of culture medium
In a 35-mm dish, 4 \times 10^5 cells were seeded and incubated in 2.5\,ml culture medium supplemented with 100\,\mu g/ml pyruvate and 50\,\mu g/ml uridine under standard conditions. After 24, 40, 48 and 62 h cultivation time the cell number was quantified, the pH value analysed and the glucose and lactate concentrations determined on a Hitachi 917 clinical chemistry analyser (Roche Diagnostics). The pH value was converted into proton concentration in medium by the equation proton conc. (mol/l) = 10^{-pH}, standardized additionally to all other data to 10^6 cells and analysed graphically. The indicated values are averages of four individual measurements or eight independent measurements (pH value of \(\rho\)\(^0\) cells), respectively. Standard deviations are depicted as error bars.

Confocal microscopy
Living cells cultured on glass bottom dishes (MatTek Corporation Ashland) were observed with the inverted confocal laser scanning microscope TCS SP5 (Leica Microsystems). To avoid a cross talk in excitation of multiple stained compounds a sequential scanning mode was used in confocal microscopy exclusively.

Images were acquired with photo multipliers and micrographs were processed and analysed with the software Leica Application Suite Advanced Fluorescence 1.5.1 and Adobe Photoshop CS.

Electron microscopy
Cells grown on cover slips were fixed with a solution of 2.5\% glutaraldehyde, 2\% formaldehyde (made from paraformaldehyde) in 100\,mM cacodylate buffer, pH 7.4 for 1.5 h at 4\(^\circ\)C, washed twice with cacodylate buffer, followed by a fixation with 2\% osmium tetroxide in 50\,mM cacodylate buffer (pH 7.4). Specimens were washed twice with distilled water and stained over night with aqueous 0.5\% uranyl acetate at 4\(^\circ\)C. Cells were dehydrated and flat embedded in Epon 812. Ultra-thin sections were analyzed with a Zeiss EM10 (Oberkochen, Germany). Negatives were digitized by scanning and processed with Adobe Photoshop CS.

RESULTS
Generation of \(\rho\)\(^0\) cells utilizing a mitochondrial targeted restriction endonuclease
The aim of this work was to develop a system for generating cells devoid of any endogenous mitochondrial DNA without using mutagenic substances like ethidium bromide, ditercalium or 2\,3'-dideoxyctydin. The basis of this system was the vector pMEE-con that after transfection in cells constitutively expresses the restriction endonuclease EcoRI as functional unit. EcoRI is known to cleave human mtDNA 3–5 times depending on the cells' background laboratory. Background mouse mtDNA also has
three EcoRI sites, while Rattus norvegicus, the source of the NRK52E cell line, has seven sites. EcoRI was fused to a mitochondrial targeting peptide derived from the human cytochrome c oxidase subunit 8 (COX VIII). Furthermore, this construct was fused to the enhanced green fluorescent protein EGFP (Figure 1A) on the one hand to observe the expression in the cells (Figure 1B) and to have a selection marker suitable for FACS analysis on the other hand so that transfected cells can be selected to purity. Confocal fluorescence microscopy showed that the polypeptide was expressed and presented a mitochondrial localization in punctate structures presumably in close contact to mtDNA in nucleoids (Figure 1B). After selection of transient transfected cells with the antibiotic geneticin for 5 days or alternatively by FACS selection, a clone of 143B.TK\(^{-}\) could be isolated. This cell clone designated as 143B.TK\(^{-}\)K7 was metabolically tested for its dependency on pyruvate and uridine. The cells exhibited only poor proliferation during cultivation in medium without pyruvate und uridine and finally died, strongly suggesting a \(\rho^0\)-state of the cell line.

The analyses of cell proliferation during 62 h of cultivation time (Figure 2) showed an obvious decrease in proliferation rate of the 143B.TK\(^{-}\)K7 cells (dotted lines) in contrast to the wild type (dashed lines). This is consistent with observations seen in a \(\rho^0\) cell line referred to as 143B.TK\(^{-}\) generated by incubation with low doses of ethidium bromide (solid line). Both \(\rho^0\) cell lines exhibit a similar genetic background. The figure indicates that the 143B.TK\(^{-}\)K7 cells possess a similar growth rate as the 143B.TK\(^{-}\)\(\rho^0\) cells.

**Confirmation of \(\rho^0\) state**

Additional confirmations of the depletion of mtDNA were carried out by PCR analyses on genomic DNA isolated from the 143B.TK\(^{-}\)K7 cells with primers that amplify a 1698-bp region located within the D-loop (Figure 3A). Positive controls for the amplification were carried out with genomic DNA of the wild type and as negative control of 143B.TK\(^{-}\)\(\rho^0\) cells. Additional examinations were accomplished with primers amplifying a region of the nuclear gene histone H1 to proof a sufficient amount of template DNA in the samples. In contrast to the wild-type cell line, no amplification product could be observed in 143B.TK\(^{-}\)\(\rho^0\) and 143B.TK\(^{-}\)K7 cells. Furthermore, these results were additionally confirmed by Southern blot testing and showed a complete loss of mtDNA in the 143B.TK\(^{-}\)\(\rho^0\) and 143B.TK\(^{-}\)K7 cells (Figure 3B). Low hybridization signals observed as ‘smear’ could be seen in all lanes, suggesting that these may represent hybridization signals to fragments of mtDNA incorporated into the nuclear genome during evolution. Similar results were obtained in multiple independent clones of the LMTK\(^{-}\} and NRK52E cell lines (Figure 3C).
Additional confirmation of the $\rho^0$ state of the cells was acquired by transfecting the $\rho^0$ cell line with the pMEE-con vector. The rationale behind this experiment is that if mtDNA was completely removed, no nucleoid-like structures should appear upon fluorescence microscopic analyses. As shown in Figure 4, the EGFP-tagged restriction enzyme appears as an evenly green-stained mitochondrial network without any punctate pattern. Thus, the punctate appearance of the EcoRI seen originally in the wild-type cells (Figure 1B) accounts for mitochondrial nucleoids, whereas the absence of nucleoids (mtDNA) leads to an even-stained matrix localization of the restriction enzyme, underlining the $\rho^0$ state of the cells.

To analyse whether or not the plasmid was integrated into the host genome during transient transfection, PCR analysis specific for the coding sequence of the restriction endonuclease was carried out on DNA extracts (see Supplementary Data). As expected, no PCR amplification product could be detected (see Figure S1 in Supplementary Data). Thus, an integration of the EcoRI plasmid into the cells nucleus could be excluded.

During more than 1 year of cultivation, there was no recovery of mtDNA in the 143B.TK−K7 cells tested by PCR and metabolic dependence of pyruvate and uridine.

**Comparative analyses of the culture medium**

The energy production of $\rho^0$ cells completely depends on anaerobic glycolysis. Additionally, all $\rho^0$ cell lines known so far exhibit a dependence on supplementation with pyruvate (3) and uridine (12–14) as well as an increasing acidification of the culture medium by excessive lactic acid production (15). Both characteristic features could be observed in the 143B.TK−K7 cells and the mouse and rat $\rho^0$ clones.

Therefore, the 143B.TK−K7 cells were analysed after 24, 40, 48 and 62 h of cultivation for glucose concentration, lactate concentration and pH value in the culture medium additionally to the determination of cell number. The glucose and lactate concentration as well as the proton concentration were standardized to 10$^6$ cells and graphically illustrated. Reference values were obtained by the examination of 143B.TK− and 143B.TK− $\rho^0$ cells.

Standardized to 10$^6$ cells, the glucose consumption (Figure 5A) of the $\rho^0$ cells (7.5–8.5 g/l, 143B.TK− $\rho^0$, solid line and 143B.TK−K7, dotted line) exceeds the amount of the wild-type cells (5 g/l, dashed line). Furthermore, it is obvious that the glucose consumption decreases during cultivation time (~88% in $\rho^0$ cells and 75% in wild-type cells).

The graphical documentation of lactate production standardized to 10$^6$ cells (Figure 5B) shows a higher lactate production rate of the $\rho^0$ cells (143B.TK− $\rho^0$, solid line and 143B.TK−K7, dotted line) than the wild type. While the values initially increase during 24 until 40 h cultivation, the curve progression resembles a saturation thereafter and then stays at an almost constant level (143B.TK−K7: 17 g/l, 143B.TK− $\rho^0$: 15 g/l, 143B.TK−: 11 g/l).

The pH values of pure medium without addition of cells show only a slight decline of pH 8.0 to pH 7.7 due to saturation of medium with CO$_2$ (Figure 5C, dotted-dashed line). On the other hand, the pH value of the medium of the examined cell lines changes from 7.8 in wild-type cells (dashed line) and 7.5 in $\rho^0$ cells (143B.TK− $\rho^0$, solid line and 143B.TK−K7, dotted line) to pH 7.3 after 48 h cultivation time in all cell lines (Figure 5C). Whereas the 62-h value of the 143B.TK− cells could not be determined due to cell mortality that was caused by high cell density, in the cells depleted of mtDNA the pH value further declined to 6.7–6.8.
Figure 4. 143B.TK−K7 cells were transfected with pMEE-con and analysed by confocal fluorescence microscopy. The EGFP-tagged restriction enzyme (MTS-EGFP-EcoRI, green colour, A1 and A4) co-localizes (A3 and A6) with the MitoTracker Red CMXRos stained mitochondrial network (A2 and A5). When compared to Figure 1B, the punctate appearance (‘nucleoid’ structure) of MTS-EGFP-EcoRI merged into an even-stained mitochondrial network, indicating that the interacting partner (mtDNA) of the restriction enzyme (MTS-EGFP-EcoRI) disappeared thus underlining the ρ0 state of the cell.

Figure 5. Graphic documentation of the glucose consumption (A), lactate production rate (B) and proton production rate (C) in pure culture medium (dash-dotted line) and the culture medium of the cell lines 143B.TK−K7 (dotted lines) 143B.TK−ρ0 (solid lines) and 143B.TK− (dashed lines) at proceeding cultivation times standardized to 10^6 cells. The indicated values are averages of four individual measurements, respectively eight independent measurements (pH value of ρ0 cells). The standard deviations are depicted as error bars. Time scale represents the time intervals proceeded after cell seeding. Standardized to 10^6 cells the glucose consumption as well as the lactate and proton production of the 143B.TK−K7 and 143B.TK− ρ0 cells exceed the amounts of the 143B.TK− wild type. However, the pH value of culture medium in all determined cell lines was nearly similar. The pH value of pure medium without addition of cells shows a slight decline due to saturation of medium with CO2. An especial curve progression could be observed in the analysis of proton production rate where the values initially decline and then increase. It was not possible to determine the 62 h cultivation value of the wild-type cells because of extreme cell density resulting in apoptotic events.
The proton concentration standardized to the cell number at different cultivation times (Figure 5D) exhibits that the proton-discharging rate in the medium of \( r^0 \) cells (143B.TK\(^{-}\)/C0\(_r^0\), solid line and 143B.TK\(^{-}\)/C0\(_K7\), dotted line) highly exceeds the demand of the wild-type cells (2.5–3.5 \( \times \) \( 10^{-8} \) mol/l, dashed line). Besides an especial curve progression could be observed in the \( r^0 \) cells where the values initially decline from 7 \( \times \) \( 10^{-8} \) to 6 \( \times \) \( 10^{-8} \) mol/l and then increase to 1.0–1.5 \( \times \) \( 10^{-7} \) mol/l.

Comparative morphological studies with confocal microscopy

The morphology of the mitochondria in the denoted cell lines was studied with confocal microscopy using MitoTracker Red CMXRos to stain the organelles (Figure 6).

The mitochondria of 143B.TK\(^{-}\) wild-type cells (Figure 6A1) exist as reticular network evenly distributed within the cell. The prevalent mitochondrial structure was elongated and rod shaped.

In the \( r^0 \) form of these cells (Figure 6A2 and A3), the network structure appears disrupted, yielding a distribution of small individual mitochondrial units. This fragmentation is highlighted in the higher magnification images. Furthermore, some of these single organelles seemed to be swollen; this result could not be observed in the wild type.

These observations of an interconnected mitochondrial network in wild-type cells in contrast to punctated mitochondria in \( r^0 \) cells constitute typical changes of mitochondrial morphology due to a loss of mitochondrial DNA (16,17).

Comparative morphological studies with transmission electron microscopy (TEM)

Even more morphological changes of the \( r^0 \) mitochondria could be detected by ultra-structural studies (Figure 7).

Figure 7A1 details a highly interconnected mitochondrial network structure in the wild type. The cross-section through the mitochondrial reticulum in normal 143B.TK\(^{-}\) cells showed a distinct outer and inner membrane with electron-dense matrix full of regular arranged cristae structures. In contrast, the TEM images of the \( r^0 \) cells (Figure 7A2 and A3) show an apparent altered morphology of the mitochondria. The network is degraded in single mitochondrial units, often with swollen appearance. The matrix seems to be electron empty probably due to a dilution of the matrix by increasing influx of water that also explains the swollen appearance. The mitochondria in \( r^0 \) cells still exhibit the distinct outer and inner membranes seen in normal cells; but the cristae displayed gross changes of structure. Most cristae appear curved or as concentric rings consisting of two membranes in close
contact, an appearance denoted as ‘fuzzy onions’ (18,19). Additionally, these membranous rings showed only few contact sites with the mitochondrial border membrane. It could not be distinguished whether these double membranes consist of two inner membranes or one outer and one inner membrane.

These results also are supported by investigations of other groups (20–25).

**DISCUSSION**

Very little is known about the key processes of interaction between nucleus and mitochondria, the coordination of gene expression or how mtDNA is maintained within the cell. Resulting from this lack of knowledge, studies were carried out to shed light on the molecular mechanisms underlying mitochondrial dysfunctions. Considerably, work was done to identify nuclear encoded proteins that act on mitochondrial DNA replication and transcription. It is now clear that for example DNA polymerase γ is one of the key players involved in the generation of mtDNA mutations (26–28). Numerous neurological and neuromuscular diseases have been associated with mutations of the mitochondrial genome (1).

In the past, many efforts to overcome these difficulties were carried out by eliminating the mtDNA as one component in the cross talk between the two genomes and thus to generate cells devoid of endogenous mtDNA (so called ρ0 cells). However, all chemicals used so far (ethidium bromide, ditercalinium, ddC, etc.) presented severe disadvantages, such as mutagenic effects on nuclear DNA or the induction of md family gene expression (multi-drug resistance). Moreover, the major limitation of these methods is their inapplicability to different cell lines or to create transgenic animal models utilizing this pathway.

The methodical approach of mitochondrially targeted restriction endonucleases for the fragmentation of mitochondrial DNA was pursued in the past by two groups. The successful targeting and function of various restriction endonucleases in the mitochondria of human and murine culture cells could be demonstrated (29,30). Cells with heteroplasmic genotypes were used, where mutated mtDNA harbouring a single base replacement co-exists together with wild-type mitochondrial genomes. The mutated site accounts for a newly formed recognition site for a restriction endonuclease that is not present in the wild-type DNA. Hence, by expressing the appropriate restriction enzyme, the mutated DNA can be selectively fragmented. Upon further degradation of the cleaved molecule by endogenous mitochondrial nucleases, a homoplasmic wild-type genotype can be regained. This method was used with the restriction endonuclease PstI in murine cells harbouring the heteroplasmic mutation for NARP (29,31). Additionally, the restriction endonuclease SmaI was applied, to the mitochondria of human cybrids (30) containing the heteroplasmic recognition sequence associated with Leigh disease (32).

Our strategy to destroy endogenous mtDNA in vivo was also based on a restriction endonuclease that was targeted to the matrix of mitochondria thereby cleaving the genomes and allowing endogenous enzymes to fully disintegrate the DNA molecules so that cells devoid of any endogenous mitochondrial genomes (ρ0 cells) can be generated. This goal was reached by selecting the gene for the restriction endonuclease EcoRI that is known to cleave human mtDNA 3–5 times, depending on ethnic background of the mitochondrial DNA. By fusing the EcoRI gene to a mitochondrial targeting sequence (derived from subunit VIII of human cytochrome c oxidase) and to the EGFP gene (enhanced green fluorescent protein) as optical marker, mitochondrial localization could be visually achieved. Moreover, a selection of transfected cells with FACS analyses can be achieved with the aid of EGFP (data not shown). Utilizing this vector in cell culture, we were able to generate ρ0 cells very efficiently on a 143B.TK− background as well as in mouse and rat cell lines. The loss of all endogenous mitochondrial DNA was monitored both by metabolic (ρ0 cells are kept on regular growth media devoid of traces of uridine and pyruvate) and genetic testing (PCR- or Southern blot analysis). Therefore, it could be shown that this method is applicable not only for the 143B.TK− and LMTK− cell lines that are known to be capable of ρ0 generation via the ethidium bromide method, but also for a rat cell line. The latter result is of interest since there are no published rat ρ0 lines despite anecdotal evidence of attempts by various groups using the ethidium bromide method over the past 15 years. It is possible that our method may be more widely applicable to different cell types and species backgrounds.

After derivation of the ρ0 clone 143B.TK−K7, the EGFP fluorescence of the EcoRI fusion protein could not be detected further. Therefore we tested by PCR if the cells had lost the gene for the restriction endonuclease construct after destruction of the mtDNA or if some gene parts were integrated into the nuclear genome so that continually low amounts of the restriction endonuclease were expressed that would render the cell line useless as acceptor cell line in cytoplast fusion experiments. Analyses with different primer pairs amplifying varying regions of the EcoRI gene showed that no amplification products could be observed. Obviously, the gene for EcoRI was not integrated into the nuclear genome of the 143B.TK−K7 cells. In accordance to the data of Tanaka, our results prove that only a short temporal expression of the restriction endonuclease targeted to the mitochondria is sufficient to destroy completely, irreversibly and stably the mitochondrial genome, also indicating that DNA-repair systems in mitochondria are not efficient enough to cope with the DNA damage.

All generated culture cells depleted of mtDNA so far proliferate despite their lack of oxidative phosphorylation in mitochondria. However, 143B.TK− cells with ρ0 genotype exhibit a decrease in proliferation rate compared to wild type (33). This feature was obvious in the newly established ρ0 cell line 143B.TK−K7 due to the exclusive anaerobic energy production via glycolysis. Thus, all ρ0 cell lines depend on supplementation with pyruvate (3) and uridine (12–14). Additionally, they show an increased acidification of the culture medium by lactic acid (15). Within this work, the dependence on supplementation
with pyruvate and uridine was also obvious for the 143B.TK K7 cells under normal growth conditions as well as an acidification of the culture medium by lactic acid fermentation.

Investigation of the culture medium showed that the pH value of the ρ0 cells was substantially decreased, compared to wild-type medium. While it was not possible to determine the 62 h value of the 143B.TK cells due to cell mortality that was caused by high cell density, cells depleted of mtDNA exhibited a further decline of the pH to 6.7–6.8.

Standardized to 10⁶ cells, the glucose consumption as well as the lactate and proton production of the 143B.TK K7 and 143B.TK ρ0 cells were increased. In contrast to the wild-type cells, this results from a complete dependence of the ρ0 cells on glycolysis for ATP production and therefore an acidification by enhanced lactic acid fermentation occurs. It was shown that in MOLT-4 cells the lactate production was increased 4-fold compared with the wild type (20). However, in the present work the lactate acid production of the cells depleted of mtDNA was only increased 1.5–2-fold in contrast to the wild type.

The transport of lactate and other monocarboxylates within and between cells is accomplished by a family of monocarboxylate transport proteins (MCTs). The mitochondrial lactate/pyruvate exchanger probably collaborates with a mitochondrial lactate dehydrogenase that enables the oxidation of lactate in active respiring cells (34). This pathway cannot take place in ρ0 cells, so that the total amount of lactate produced must be released into the surrounding medium. Across the plasma membrane, lactate is co-transported with protons (35) accounting for the decrease in pH value of the culture medium in the ρ0 cells.

The morphology of the mitochondria in the denoted cell lines was studied with confocal microscopy using MitoTracker Red CMXRos to stain the organelles. The cell lines was studied with confocal microscopy using the lactate and proton production of the K7 and 143B.TK ρ0 cell lines under very controlled and mild conditions, avoiding mutagenic effects of chemicals (e.g. ethidium bromide). Thus, ρ0 cell lines exhibiting varying nuclear backgrounds can be used in future experiments to study diseases connected to mitochondrial phenotypes.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR online.

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