Optimization of methods to assess mitochondrial DNA in archival paraffin-embedded tissues from mammary canine tumors

Otimização dos métodos para avaliar o DNA mitocondrial obtido a partir de tumores mamários caninos incluídos em parafina

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In this study we describe the alterations used to extract and amplify mitochondrial desoxyribonucleic acid (DNA) from formalin-fixed paraffin-embedded samples of canine mammary tumors. The epithelial and mesenchymal components (chondromyxoid and chondroid) of each tumor, as well as the normal mammary gland tissues, were manually microdissected from 19 mixed canine mammary tumors (10 benign mixed tumors and nine carcinomas arising in mixed tumors). DNA was extracted by Invisorb® Spin Tissue Mini Kit, with protocol changes proposed by the manufacturer. A 273-bp fragment was amplified by polymerase chain reaction (PCR) and submitted to automatic sequence analysis. The fragment was successfully analyzed in 100% of the samples. However, an additional lysis step, the reduction of volume in buffer solutions and PCR, a higher annealing temperature and an increase in the number of PCR cycles were required. The initial PCR products were diluted and re-amplified in six

Neoplasia
Dog
Mesenchymal

A presente comunicação descreve as modificações usadas para extrair e amplificar o DNA mitocondrial obtido de amostras de tumores mamários caninos fixados em formol tamponado a 10% e incluídos em parafina. Os componentes epiteliais e mesenquimais (condromixóide e condróide), bem como a mama normal adjacente, foram microdissectados manualmente de 19 tumores mamários (10 tumores mistos benignos e nove carcinomas em tumores mistos). O DNA foi extraído utilizando-se o Invisorb® Spin Tissue Mini Kit com modificações do protocolo proposto pelo fabricante. Um fragmento de 273-bp foi amplificado por reação em cadeia da polimerase (PCR) e sequenciado em sequenciador automático. O fragmento foi analisado em 100% das amostras, entretanto modificações como lise adicional, redução do volume das soluções de extração e PCR, aumento da temperatura de anelamento e do número de ciclos de amplificação foram necessárias. Em seis amostras os produtos iniciais de PCR foram diluídos e re-amplificados para obtenção de sucesso.

Neoplasia
Cão
Mesenquimal

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In general, desoxyribonucleic acid (DNA) recovered from formalin-fixed paraffin-embedded tissues is highly degraded and may be unsuitable for most molecular techniques such as polymerase chain reaction (PCR). Although some studies have shown that PCR can be successfully performed with DNA recovered from formalin-fixed paraffin-embedded tissues, results have still been limited in some laboratories.

In this study, we describe our experience concerning standardization of mitochondrial DNA (mtDNA) extraction techniques, amplification and sequencing of 10% formalin-fixed paraffin-embedded tissues, microdissected from mixed canine mammary tumors.

Nineteen canine mammary tumors (10 benign mixed tumors and nine carcinomas arising in mixed tumors) were retrieved from the files of the Laboratory of Comparative Pathology, Department of General Pathology, Instituto de Ciências Biológicas of Universidade Federal de Minas Gerais (UFGM). The tissues used in this study had been fixed in 10% neutral buffered formalin (without knowledge of total time fixation) and paraffin-embedded.

Seven 10 µm-thick sections were cut from formalin-fixed paraffin-embedded tumors and mounted on polylysine-coated slides. The microtome used to cut the sections was kept clean and excess paraffin and tissue fragments were wiped from the blade holder with xylene between blocks. The sections were dewaxed twice, with two changes in xylene (15 min each) and washed with 100% and 70% ethanol (twice, for 5 min).

Microdissection target areas were previously marked on hematoxilin-eosin stained slides. The epithelial (neoplastic epithelial proliferations) and the mesenchymal (chondromyxoid and chondroid) components of each tumor and the normal residual glandular parenchyma were manually removed from each of the sections using a scalpel, under stereomicroscopic observation. We verified that the addition of deionized water to the the area to be microdissected facilitates tissue recovery of the material. The tissues were scraped into 1.5-ml tubes and submitted to digestion and extraction using an Invisorb® Spin Tissue Mini Kit (Invitrek). The procedures were performed according to the manufacturer’s protocol with the following modifications: the suggested volumes of reagents (lysis, binding and wash buffer, proteinase K) were reduced to half, the elution buffer volume was reduced to 33 µl, an additional lysis step was performed, and mixtures were homogenized by gentle pipetting instead of vortexing.

The digestion included the incubation of samples in a lysis buffer (200 µl) and 20 µl proteinase K (10 µl/ml) at 56°C, with gentle agitation in a thermomixer. An overnight incubation was usually enough to obtain total digestion of the tissue, indicated by a clear solution. In the cases where the samples were still turbid and tissue fragments were visible, an additional lysis step was performed by a new addition of 20 µl proteinase K for another eight hours at 56°C.

After tissue digestion, the tubes were centrifuged for 2 min at maximum speed in order to precipitate all non-lysed material, and 100 µl binding buffer T were added. The solutions were homogenized with gentle pipetting to minimize DNA breakage. The suspensions were transferred onto spin columns placed in 2-ml receiver tubes and incubated for 1 min. The spin columns were centrifuged at 12,000 rpm for 2 min and the filtrate was discarded. The column was placed back in the receiver tube, 200 µl wash buffer were added and centrifuged as in the previous step. An additional centrifugation was performed to remove any residual ethanol. For DNA elution the spin columns were placed in new 1.5-ml receiver tubes and 33 µl of pre-heated elution buffer D were added and incubated for 5 min. A final centrifugation was performed for 2 min at 10,000 rpm. A total volume of 33 µl of eluted DNA was recovered from each sample.

The DNA extract was stored at 4°C for 24 hours and used for PCR amplification.

For successful amplification of degraded DNA samples it is recommendable that the amplicon size be relatively small. However, numerous known polymorphisms strongly constrain primer design in this region. The primers designed in this work for the amplification of a 273-bp fragment of the hypervariable mitochondrial D-loop represent a compromise for obtaining the smallest and most potentially informative amplicons.

PCR amplifications were optimized by varying the number of cycles and the annealing temperature. The annealing temperatures tested were 56°C, 57°C, 59°C and 60°C. Reactions were performed with 30, 35 and 40 cycles. PCR optimization reactions were tested for each possible combination of annealing temperature and number of cycles. The best results were obtained with an annealing temperature at 60°C with 35 cycles.

The optimized PCR was performed in a 25-µl total volume, using 11 µl of Quiagen® Multiplex PCR Kit; 11 µl of H2O; 1 µl of each primer (2.5 mM); 1 µl of DNA template; and 1 µl of sterile deionized water. The PCR program consisted of an initial denaturation step at 95°C for 15 min followed by 35 cycles at 95°C for 5 min; 60°C
for 1.3 min and 72°C for 1 min; and a final extension step at 72°C for 10 min.

DNA isolated from the peripheral blood of a dog was used as positive control, and sterile deionized water was used as negative control. The PCR products were run in 6% polyacrilamide electrophoresis gel and visualized after silver staining. A DNA ladder was run as a molecular marker in each gel.

Electrophoresis showed that the 273-bp fragment of the mitochondrial canine control region was amplified in 52 out of 57 samples (91%). Five samples (three chondroid, two epithelial and one normal mammary tissue) yielded low PCR product, and a second round of PCR amplification was performed. One microliter from a 100-fold diluted aliquot of the initial PCR product was used as template. This reaction was performed following the same procedure as in the first round. The use of products from the first PCR round as starting template produced bands that could be visualized in the electrophoresis gel (Figure 1A) and were sufficient for subsequent sequencing.

The PCR products were analyzed by automated sequencing (Figure 1B). In order to clean the PCR product from primers and nucleotides, PCR products were purified using a Sephacryl HR300 column (Pharmacia, Biotech, Saclay, France), and sequenced using BigDye Terminator v 3.0 sequencing kit. The sequencing reactions were performed in both directions using 2.6 µl of the purified PCR products, 0.25 mM of primer, 0.6 µl of sequencing kit and deionized water to complete 5 µl of total reaction volume. The thermocycler program comprised an initial denaturation step at 96°C for 2 min, followed by 35 cycles at 96°C for 15 s, 50°C for 9 s, 60°C for 2 min and a final extension step at 60°C for 10 min. Sequencing reaction products were purified using Sephadex G-50 fine columns (Pharmacia) and eluted in deionized formamide. DNA sequencing was carried out in an ABI Prism 3130 Genetic Analyzer (Applied Biosystems Inc., Foster City, CA, USA) using POP-6, and was used for the capillary electrophoresis separation matrix.

The sequences obtained were compared to the reference sequence, indicated as haplotype A19 (GenBank accession entry: NC_002008) and confirmed the amplification of the predicted 273-bp fragment of the hypervariable D-loop control region in 48 cases (Figures 1B and 1C). Difficulties in deducing the sequences were observed in nine samples, which were re-submitted to sequencing reactions with higher and lower amounts of purified PCR product, according to the intensity of the band previously detected in polyacrilamide gel electrophoresis. A total of 2.4 µl and 2.6 µl were used for sequencing in samples with high and low intensity bands, respectively. In all the samples it was possible to successfully confirm the DNA sequence.

The analysis of mtDNA offers advantages over nuclear DNA due to a larger number of copies per cell, what potentiates the recovery of DNA from difficult or degraded materials. The literature describes several successful protocols for analysis of shorter and longer mtDNA fragments recovered from archival fixed paraffin-embedded human tissues. In our series of cases, the success of mtDNA analysis was achieved by the use of a DNA extraction kit with modifications in several steps of the process: lysis (additional lysis step), extraction (reduction of buffer volumes and homogenization by gentle handling), amplification (high-annealing temperature and increase in the number of PCR cycles) and sequencing (adjustment of the volume of the PCR product).

The samples had been fixed in 10% buffered formalin, but no information on the handling of specimens before tissue fixation (tissue amount, degree of autolysis), fixation-related factors (temperature, and duration of fixation,) and post-fixation procedures storage (temperature and duration of storage) was available. Several of the samples may well have been subjected to conditions that contributed to the partial degradation of the nucleic acids.

**Figure 1** - A: Evaluation of DNA amplification by polyacrilamide gel electrophoresis. C+= positive control (peripheral blood of a dog); C-= negative control; 1, 2, 3, 4, 5 and 6 = PCR products (273bp) from second round PCR, M, molecular weight marker. 1.2 microliters of PCR products were loaded in each lane; B and C: Confirmation of amplification by direct sequencing of PCR product. B = Sample 1; C = Positive control. Aliquots of the amplicons were purified and sequenced as describe under Material and Methods. Picture in the ‘5 → 3’ orientation is the sense strand of the amplicon from mesenchymal component (case 5).
The major limitation in this analysis was the difficulty in obtaining high quality DNA in some samples (mainly from cartilaginous component), probably due to lower cell content present. In addition to the improvements in the extraction procedure, the re-amplification step using the initial PCR product as template was crucial to achieving success.

Improvement and routine control of pre-, during and post-fixation procedures should be considered in medicine veterinary laboratories so that, in the future, tissue banks may consistently offer high-quality DNA.

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