Anatomopathological Assessment of the Diaphragm in Formalin-Fixed, Paraffin-Embedded Sections

Avaliação anatomopatológica do diafragma em cortes fixados em formalina e incorporados em parafina

Ricardo Aparecido Baptista Nucci¹,² Wilson Jacob-Filho²,³ Alexandre Leopold Busse²,³ Laura Beatriz Mesiano Maifrino⁴,⁵ Romeu Rodrigues de Souza⁶,⁷

¹ Department of Pathology, Faculty of Medicine, Universidade de São Paulo, São Paulo, SP, Brazil
² Postgraduate Pathology program, Faculty of Medicine, Universidade de São Paulo, São Paulo, SP, Brazil
³ Department of Geriatrics, Faculty of Medicine, Universidade de São Paulo, São Paulo, SP, Brazil
⁴ Laboratory of Morphoquantitative Studies and Immunohistochemistry, Department of Physical Education, Universidade São Judas Tadeu, São Paulo, SP, Brazil
⁵ Postgraduate Physical Education program, Universidade São Judas Tadeu, São Paulo, SP, Brazil
⁶ Department of Anatomy, Institute of Biomedical Sciences, Universidade de São Paulo, São Paulo, SP, Brazil
⁷ Postgraduate Aging Sciences program, Universidade São Judas Tadeu, São Paulo, SP, Brazil

Address for correspondence Ricardo Aparecido Baptista Nucci, PhD, Departamento de Patologia, Faculdade de Medicina da Universidade de São Paulo, Avenida Dr. Arnaldo, 455, 01246-903 São Paulo, SP, Brazil (e-mail: r.aparecido.nucci@uol.com.br).

Abstract

Introduction The analysis of frozen muscle biopsies has become a routine method in the evaluation of muscle structure in health and disease. However, the technique for frozen muscle specimens is not widely available in countries with limited medical facilities. The present study aimed to elucidate a reproducible formalin-fixed and paraffin-embedded (FFPE) method for this type of analysis in postmortem muscles.

Methods Diaphragm muscle was obtained within 1 hour of sudden death. Diaphragm strips were washed in saline solution, fixed in 10% formalin, frozen at 4°C in a refrigerator, and stored for 24 hours. Then, the tissue samples were processed into paraffin-embedded blocks. Transversal sections were cut from each paraffin block and stained with hematoxylin and eosin, Picrosirius red, Verhoeff-Van Gieson, and Congo red for the qualitative analysis.

Results Our analysis indicated a well-preserved muscle.

Conclusion In summary, we demonstrate a simple technique for a reproducible FFPE method in postmortem muscle tissues.
Introduction

The diaphragm (DIA) is the primary muscle of inspiration. Therefore, its uncompromised function is essential to support the ventilatory and gas exchange demands. Many patients with chronic obstructive pulmonary disease (COPD) or emphysema show high levels of DIA activity both at rest and during exercise, in which the DIA may experience maximal activation leading to irreversible muscle injury, mechanical failure, and death. The analysis of frozen muscle biopsies has become a routine method in the evaluation of muscle structure and function in health and disease. However, the analysis of frozen muscle specimens is not widely available in countries with limited medical facilities.

Materials and Methods

Autopsy verification is mandatory in Brazil to define the cause of death for most individuals who die of natural causes. The São Paulo Autopsy Service (Serviço de Verificação de Óbito [SVO], in the Portuguese acronym) is the major morgue serving the metropolitan area of São Paulo, Brazil. The present study was approved by the ethical committee of the Faculdade de Medicina of Universidade de São Paulo under the registration number 2.209.383. The division of pathology from the Instituto do Coração (INCOR) provided the material for the present study exclusively for the sake of methodological standard in future projects.

Muscle Samples

Specimens from the right lobule of the DIA muscle was obtained within 1 hour of sudden death from a previously physically healthy 51-year-old man who died of myocardial infarction.

Diaphragm strips (~5 × 5 × 5 mm) were removed at 4 to 6 cm from the central tendon (midcostal) to avoid the muscle fibers that radiate toward this tendon insert. The DIA strips were gently washed in saline solution, placed in a standard histological cassette (SWINGSETTE™ Tissue Processing/Embedding Cassettes, Histocell Soluções em Anatomia Patológica Ltda., São Paulo, SP, Brazil) to re-establish the initial length, and immediately fixed in 10% formalin, frozen at 4°C in a refrigerator, and stored for 24 hours. Then, the tissue samples were processed into paraffin-embedded blocks.

Histological Staining

Transversal sections of 6 μm thick were cut from each paraffin block for histological staining. The sections were stained with the hematoxylin and eosin (H&E) standard method for the general structure of the sample. The sections were also stained with standard methods for: (a) Picrosirius red (collagen fibers); (b) Verhoeff-Van Gieson (elastic fibers); and (c) Congo red (amyloid deposits).

The qualitative evaluation of the tissue was performed by photographing 40 fields selected at random for each staining technique. The images were captured using a Zeiss Microscope (Binocular microscope Axio Lab.A1 with phototube, Carl Zeiss, Thornwood, NY, USA) with the specific software AxioVision, Version 4.8 (Zeiss, Thornwood, NY, USA).

Results

The H&E stain positivity indicated a well-preserved muscle. The Picrosirius red stain gave superior results under a polarized filter, in which different collagen fiber types were observed in the endomysium, in the perimysium, and in the epimysium. Additionally, we could observe a weak positive stain for elastic fibers and amyloid.

Discussion

With death, a series of postmortem events initiate, including the loss of enzyme activities. The reliability of postmortem muscle samples depends on the extent of these cellular alterations.

Fig. 1 Cross-sectional images of the diaphragm (DIA) muscle stained with hematoxylin and eosin (A) or picrosirius red (B) (100x magnification). A) Red arrow: muscle fiber; yellow arrow: inflammatory process. B) Picrosirius red stain in polarized microscopy showing type I collagen fibers (red); intermediate fibers (yellow/orange); and type III collagen fibers (green).
Eriksson et al.15 demonstrated that muscle samples stored for a maximum of 10 days in controlled temperature (4°C) are reliable as nitrogen stored samples. Van Ee et al.16 had similar results, indicating that lower temperatures may maintain the integrity of the muscle fibers. We have frozen the samples at 4°C, which maintained its integrity.

Nevertheless, it is reported that muscles stiffen as they enter rigor mortis.18 An experimental study by Fitzgerald19 reported that the elastic compliance decreased 95%, and that the viscous compliance decreased 98.3% in relation to the living muscle tissue values at ~6.5 hours after death. Van Ee et al.16 reported that a period of 0.5 hours postmortem (prerigor) had only a modest effect on the mechanical properties of the muscle, while at 48 hours postmortem (postrigor), the response changed greatly. We have chosen to freeze the samples 1 hour postmortem to maintain the prerigor integrity.

A pilot study of an autolytic change in rat diaphragms demonstrated that, up to 96 hours postmortem, the only postmortem morphological artifact was a decrease in the intensity of the H&E staining.20 This artifact was not observed in the present study. The Picrosirius red stain was successful because it specifically labels collagen molecules without relying on the recognition of antigens, which may be degraded postmortem or during fixation.15,17 Although we have observed a decrease in the intensity of the Verhoeff-Van Gieson staining, Rodrigues et al.21 observed that the viscoelastic properties of the DIA decreases with aging, which may explain our finding. Finally, the Congo red staining was efficient to demonstrate amyloid deposits, as has been shown by previous studies with FFPE in the nervous system.22,23

Recently, Suriyonplengsaeng et al.6 successfully developed an immunohistochemistry (IHC) technique, with heat-mediated antigen retrieval, for FFPE muscle biopsy specimens. This study may encourage more researches to use IHC on FFPE muscles in autopsy studies as well.

The present study has some weaknesses that need to be acknowledged for proper interpretation. The main limitation of the present study is the small sample size, which was limited to one patient. However, the small sample size would not explain our positive findings. Due to the limited literature, we could not carry out an extensive discussion about our findings. Nevertheless, we have highlighted in the present study a simple and valid methodological approach for the analysis of postmortem muscle that could be improved and discussed in depth in further studies.

Conclusion

In summary, we demonstrate a simple and reproducible technique for FFPE in postmortem tissue. We suggest that this method could become a valuable tool for the diagnosis of anatomopathological changes in the DIA muscle.

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