

New insights into the *in situ* microscopic visualization and quantification of inorganic polyphosphate stores by 4',6-diamidino-2-phenylindole (DAPI)-staining

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Abstract

Inorganic polyphosphate (PolyP) is a biological polymer that plays important roles in the cell physiology of both prokaryotic and eukaryotic organisms. Among the available methods for PolyP localization and quantification, a 4',6-diamidino-2-phenylindole(DAPI)-based assay has been used for visualization of PolyPrich organelles. Due to differences in DAPI permeability to different compartments and/or PolyP retention after fixation, a general protocol for DAPI-PolyP staining has not yet been established. Here, we tested different protocols for DAPI-PolyP detection in a range of samples with different levels of DAPI permeability, including subcellular fractions, free-living cells and cryosections of fixed tissues. Subcellular fractions of PolyP-rich organelles vielded DAPI-PolyP fluorescence, although those with a complex external layer usually required longer incubation times, previous aldehyde fixation and/or detergent permeabilization. DAPI-PolyP was also detected in cryosections of OCT-embedded tissues analyzed by multiphoton microscopy. In addition, a semi-quantitative fluorimetric analysis of DAPI-stained fractions showed PolyP mobilization in a similar fashion to what has been demonstrated

with the use of enzyme-based quantitative protocols. Taken together, our results support the use of DAPI for both PolyP visualization and quantification, although specific steps are suggested as a general guideline for DAPI-PolyP staining in biological samples with different degrees of DAPI and PolyP permeability.

Introduction

Inorganic polyphosphate (PolyP) is a widely distributed biological polymer composed of phosphate residues linked by high-energy phosphoanhydride bonds, and found in every tested organism.1 Several biological functions have been described for PolyP, including a role as phosphate reservoir, metal homeostasis, regulation of transcription factors and transcription fidelity, as well as regulation of several enzymatic activities.2 More recently, studies on the role of PolyP in vertebrate models revealed the participation of PolyP stores in platelet activation and intrinsic coagulation.3-6 cancer metastasis,7,8 activation of the fibroblast growth factor (FGF) signaling and a role in stem cell differentiation.9,10

Although extensively demonstrated in several subcellular compartments such as nuclei, mitochondria, as well as in the cytoplasm of different cells, millimolar concentrations of PolyP are commonly found inside PolyP-rich organelles. A wide variety of PolyP-rich organelles has been described, but recent studies have pointed that at least some of these compartments probably derive from a common evolutionary step and are similar to the protozoan acidocalcisome,11 the best studied PolyPrich organelle. It has been suggested that PolyP stored in acidocalcisomes have a role in protozoan virulence^{12,13} as well as in the maintenance of osmotic balance14 and metal homeostasis.15 Together with protozoan acidocalcisomes, bacterial volutin granules are also considered one of the most studied types of PolyPrich structures. Although volutin granules have been known for a longer period, it was only after the first description of the acidocalcisome in trypanosomes that the concept that this organelle might have a membrane unit containing specific transporters was established in bacteria. 16,17 After these initial studies, several other organisms were shown to contain PolyP-rich acidocalcisome-like organelles, including eggs of invertebrates such as insects18,19 and sea urchin,20 and vertebrates (chicken),²¹ models where the isolation of subcellular fractions of acidocalcisomes was readily demonstrated.

The significant biological relevance of PolyP in cell metabolism has lead to the establishment of different protocols for PolyP visualiza-

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tion and quantification, an important step towards a better comprehension of the roles played by this polymer in different biological models. In this regard, the use of 4',6-diamidino-2-phenylindole (DAPI), first developed as a trypanocide agent and later largely used for the detection of nucleic acids,22 has progressively become a widely accepted tool for the detection of PolyP-rich compartments. When used as a nucleic acid sensor, both in spectrofluorimetric and microscopic analysis, DAPI-DNA complexes exhibit a maximum fluorescence emission at around 450 nm (blue), whereas when bound to PolyP, a bathochromic shift in the emission to 525-550 nm take place, leading to a greenish-vellowish color.23 DAPI-PolyP interaction was shown to be remarkably specific and to generate a proper quantum yield for microscopic observation.^{21,24-26} Nevertheless, reports concerning the feasibility of using DAPI as a PolyP probe have remained frag-





mented among several publications that suggested a number of possible routines for DAPI-PolyP labeling and visualization.^{23,27,28}

Information available in most catalogs and product data sheet classify DAPI (different varieties) as compounds slightly permeable to cell membranes, although it has been generally recognized that it may permeate cell membranes less efficiently in living cells, making them less susceptible to DAPI-staining.29 DAPI permeability to cell walls has been reported,23 although not systematically studied in different cell types. The degree of permeability presumably depends on the structural characteristics of the different cell surfaces in different organisms. In this regard, although a few studies have shown the spectroscopic properties of analytical grade sodium PolyP-DAPI fluorescence for quantification purposes, 28,30 quantitative analysis of PolyP mobilization using DAPI as a probe in biological samples has not vet been reported. In the present study, we evaluated the effect of different routines of DAPI incubation in subcellular fractions, whole cells (containing or not cell walls), and OCT-embedded semi-thin sections in order to ascertain a unified guideline for PolyP labeling and detection. A semi-quantitative fluorimetric assay for the analysis of PolyP mobilization in PolyP-enriched extracts obtained from biological samples is also shown and provides a fast protocol for PolyP quantification

Materials and Methods

Animals, cell culture and strains

Trypanosoma cruzi epimastigote forms (Y strain) were grown axenically in liver infusion tryptose (LIT) medium supplemented with 10% fetal calf serum as previously described. Tells were collected by centrifugation at 350 g after four days of cultivation. Trypanosoma brucei procyclic forms (strain 427) were maintained at 27-28°C in SDM-79 medium supplemented with 10% fetal bovine serum. Cells were harvested by centrifugation at 350 g after two days of cultivation.

The obtainment of *Eimeria spp* oocysts from chickens was approved by the Ethics Committee of the Biology Institute of the State University of Campinas (UNICAMP) under the protocol number 1084-1, according to the Brazilian federal law (11.794/2008, Decree n. 6.899/2009) that is based on the Guide for the Care and Use of Laboratory Animals prepared by the National Academy of Sciences, USA, and the Australian Code of Practice for Care and Use of Animal for Scientific Purpose. All animals received humane care in compliance with

the above-mentioned guides used by the Ethics Committee to approve the protocol. Briefly, four-week-old chickens were inoculated with 1×10^4 sporulated oocysts of *Eimeria acervulina* (Cu strain) and 5×10^3 Eimeria tenella (Pa strain) oocysts, as previously described. Ocysts were isolated from feces and sporulated after one week of inoculation. Sporozoites were obtained from oocysts by mechanic rupture, enzymatic treatment, and purification by anion-exchange chromatography using DE-52 cellulose columns.

Euglena gracilis wild strain was cultivated axenically at room temperature and constant illumination as previously described.34 Rhodnius prolixus Stahl, 1859 (Hemiptera, Reduviidae) were reared in a colony maintained at 28°C and 70-80% relative humidity. The insects were fed with rabbit blood in an artificial apparatus as described before.35 Total egg homogenates (TEH) were prepared by disrupting the laid eggs and a fraction enriched in acidocalcisomes was obtained as described by Ramos et al. 19 The final pellet was used as acidocalcisome-enriched fraction (acidocalcisome fraction). Anticarsia gemmatalis specimens were obtained as described;36 briefly, a colony was kept about 27°C and 70% relative humidity. Adults were maintained in a plastic cage and paper sheets were added for eggs deposition. After 24 h, eggs were transferred to a plastic box and left for egg hatching and larvae development. Larvae were fed as previously described,37 until they reached the fifth instar around the 10th or 11th day after hatching as detected by visual inspection.

PolyP staining in whole cells

For PolyP detection, living cells were incubated with 50 µg/mL DAPI for 30 min, or different times where indicated, at room temperature. Alternatively, Eimeria sporocysts were first fixed with 4% formaldehyde for 5-20 min at room temperature and permeabilized with 0.3% Triton X-100 for 5 min prior to DAPI incubation. Mechanically disrupted sporocysts were also prepared as described.³³ Following DAPI incubation, samples were mounted on glass slides and observed with a Zeiss Axioplan epifluorescence microscope using a personalized filter set with a bandpass excitation maximum at 350 nm and a 500 nm long pass emission filter, combined with the appropriate dichroic mirror or a FITC-optimized filter set. Following image acquisition, deconvolution was performed using a no-neighbor algorithm.

Isolation of PolyP-rich organelles and PolyP staining

Acidocalcisome-like organelles from

Rhodnius prolixus were fractionated and enriched as described. 19,21,38 Isolated acidocalcisomes were incubated with 2-10 µg/mL DAPI at room temperature, mounted on a glass slides and observed under a Zeiss Axioplan epifluorescence microscope using a personalized filter set as above.

Midgut optimum cutting temperature embedding and PolyP staining in semi-thin sections

Midguts from 5th instar larvae of the insect Anticarsia gemmatalis were dissected and fixed for 3 h at room temperature using 4% formaldehyde, 0.1% glutaraldehyde, 0.1 M sodium cacodylate pH 7.2. The tissue was cryoprotected in 10% sucrose overnight and in 30% sucrose for 24 h before Tissue-Tek optimum cutting temperature (OCT) immersion and freezing in LN2. Semi-thin sections were obtained in a cryostat and left at room temperature to dry for 60 min, then incubated with 0.1 ug/mL DAPI, washed with PBS, and mounted on slides using n-propyl gallate. The sections were observed under a Zeiss Axioplan epifluorescence microscope using a fluorescein filter set. After image acquisition, deconvolution was performed using a no-neighbor algorithm. Alternatively, samples were observed with a Zeiss LSM 510 Meta NLO multiphoton microscope using a two photon fluorescence excitation of 780 nm. A wavelength scan was performed in order to detect the most intense fluorescence wavelengths.

Fluorimetric analysis of PolyP

Time-lapse analysis of PolyP fluorescence was evaluated by washing cell suspensions with PBS and incubating with 100 µg/mL DAPI. After addition of DAPI, samples were immediately transferred to a microplate reader and the fluorescence intensity was registered over the course of 60 min, using excitation/emission wavelength of 350/450 (nucleic acid signal) and 350/550 (PolyP signal). Water-soluble PolyP fractions (WSF) were prepared from Periplaneta americana and R. prolixus eggs as described26 and both WSF, sodium phosphate glass - PolyP75+ (Sigma Aldrich, St. Louis, MO, USA) - and plasmidial DNA were incubated with 20 µg/mL DAPI for 30 min at room temperature. Samples were analyzed with a spectrofluorometer at the excitation wavelength of 354 nm. Relative quantification of PolyP in R. prolixus eggs was performed as described.26 For comparison, PolyP samples were extracted as described and incubated with an excess of scPPX for 15 minutes at 37°C.39 Resulting Pi levels were analyzed with malachite green.





Results

PolyP-DAPI staining of subcellular isolated compartments and single cells

Subcellular fractions of PolyP-rich organelles were obtained as previously described19 and verified by electron microscopy analysis of whole mounts adhered to formvar-coated grids, showing the typical electrondense characteristic of acidocalcisomes (Figure 1A). Additional confirmation was obtained by electron-probe X-ray microanalysis where high amounts of phosphorus and oxygen, bound to a number of cations, were detected (Figure 1B). PolyP stores in these fractions were rapidly stained (2-3 min at room temperature) using a range of 2-10 µg/mL DAPI and observed using a filter set optimized for DAPI-PolyP visualization (Figure 1C). Similar results were obtained using acidocalcisome fractions isolated from other organisms, such as PolyP-rich granules of chicken eggs (data not shown) and are consistent with those obtained from iodixanol differential sedimentation from other models.19-

DAPI-PolyP was also detected in whole cells of Trypanosoma cruzi (Figure 2A-B), a protozoan where both the size and number of T. cruzi PolyP-rich organelles (acidocalcisomes) are well known. 41,42 Preparations of whole cell mounts seen in TEM showed electron dense organelles in the intracellular milieu (Figure 2A), presenting morphological similarities with the acidocalcisome-rich fractions depicted in Fig. 1A. X-ray microanalysis spectra confirmed the high content of phosphorus, oxygen and several cations (Figure 2A, inset), as previously described. 43,44 When DAPI-PolyP staining was carried out in whole cells, higher concentrations of DAPI and longer incubation times were usually required. Incubation of intact cells with DAPI at the same conditions used for staining subcellular fractions (2-10 ug/mL or 5 min) did not result in satisfactory staining (data not shown). In contrast, 50 ug/mL DAPI for 10-20 min showed an intense staining pattern compatible with the presence and distribution of acidocalcisomes (Figure 2B) in these cells. This result suggested that the presence of external membranes or additional layers with different degrees of DAPI permeability may potentially diminish the penetration of the dye, therefore requiring slight modifications in the staining protocol. To verify whether these additional layers would actually interfere in PolyP-DAPI staining, we performed a PolyP detection assay under similar conditions in a range of unicellular organisms, i) containing plasma membrane with different properties, or ii) presenting additional cell or

cvst walls. We first analyzed detection of PolyP stores in Euglena gracillis, known to present unique plasma membrane morphology supported by a dense fibrillar layer, the pellicular cortex.45 In Euglena cells, intracellular PolyP stores (acidocalcisome-like) were usually detected using higher concentrations of DAPI (50-200 μg/mL), following a 30 min incubation period at room temperature, without any further modifications in the protocol (Figure 3A). To verify whether the presence of a cell wall would diminish DAPI penetration into the cells, we incubated Candida albicans cells with different concentrations of DAPI and followed the staining pattern over time. In general, although the additional cell wall layer seemed to reduce DAPI penetration in some cells, PolvP stores were readily visible using 50 ug/mL DAPI (Figure 3B). Eimeria parasites have also been shown to contain PolyP-rich organelles.33 These cells provided an interesting model for testing PolyP detection with DAPI, since the sporozoites (infective cells that contain only a limiting plasma membrane as a surface barrier) are found inside sporocysts that contain a cyst wall (each sporocyst contain two zoites). In contrast to yeast cells, PolyP detection in Eimeria sporozoites protected by a cyst wall inside sporocysts was not possible using the above mentioned protocols. In an attempt to enhance the access of DAPI to the intracellular milieu, sporocysts were submitted to i) gentle mechanical disruption/permeabilization as previously reported46 (Figure 3 C,D) or ii) Triton X-100 permeabilization followed by aldehyde fixation (Figure 3 E,F), before incubation with DAPI. Results showed an intense staining of PolyP-rich organelles (acidocalcisomes) inside the sporozoites. Although a previous step of aldehyde fixation has not significantly altered PolyP staining properties of DAPI in Eimeria cells, the same procedure seemed to impair DAPI-PolyP staining in other organisms, either by excluding DAPI penetration across the fixed cell wall or causing leaking of PolyP from the target organelles after chemical fixation (Figure 3B, inset). Nevertheless, it is important to note that in other yeast cell types such as the pathogenic fungi Cryptococcus neoformans, suc-

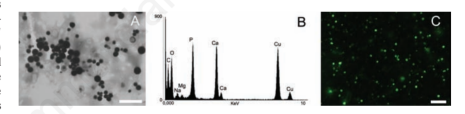


Figure 1. PolyP detection on subcellular stores in fraction of PolyP-rich organelles obtained from *R. prolixus* eggs. A) Electron micrograph where the stores appear as electron dense granules. B) X-ray microanalysis elemental composition. C) PolyP-rich granules after DAPI staining as observed by epifluorescence microscope with a long pass 500 nm emission filter. Scale bars: A, 2 μ m; C, 2.5 μ m.

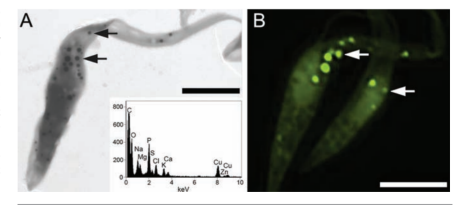


Figure 2. PolyP staining of *T. cruzi* whole cells. A) transmission electron micrograph of the electron dense acidocalcisomes whose elemental composition was determined by X-ray microanalysis (A, inset). B) PolyP-rich composition of acidocalcisomes was confirmed by fluorescence DAPI staining (50 μg/mL DAPI for 10-20 min). Arrows: acidocalcisomes. Scale bars: A, 3 μm; B, 4 μm.





cessful PolyP-DAPI staining has been carried out without the previous step of aldehyde fixation (data not shown), suggesting that the structural organization of the cell wall may be important for the permeation of the dye. Additional time-lapse analyses showed that the staining intensity was time-depended and varied with the cell type and the number and type of surface barriers. Fluorescence microscopy analysis of C. albicans incubated for different times showed acidocalcisomes became visible after 30 min and fluorescence intensity reached a plateau within 50 min (Figure 4A). Time-scale spectrofluorimetric analysis showed that in contrast to C. albicans, this plateau was reached in T. cruzi and T. brucei within 15-20 min, both for nuclei and PolyP staining (Figure 4 B,C).

DAPI staining in tissues

Several studies have shown that the inorganic content of PolyP stores is usually washed during different steps used in sample preparation for electron microscopy, resulting in empty organelles observed in transmission electron microscopy images from several models.^{24,47,48} This suggested that the preparation of semi-thin sections of PolyP-containing tissues for light microscopy would potentially require protocol adaptations to preserve the PolyP content. In this regard, preparation of semi-thin cryosections of OCT-embedded tissues allowed on one hand minimal sample handling and, on the other, the obtainment of sections containing the whole diameter of the PolyP-rich organelles, avoiding the leaking of PolyP from sectioned organelles, as observed in TEM preparations. Analysis of semi-thin sections of the midgut of the insect A. gemmatalis showed the presence of PolyP stores in some apocrine vesicles at the apical region of columnar cells (Figure 5A) as well as in vesicles occurring mainly around the goblet cell cavities (Figure 5B), as previously described.36,49

Results provided by different groups have shown that in situ DAPI-PolyP staining of PolyP granules may result in a slight variation of fluorescence emission peaks or colors (from green to yellow) (Figure 3).50,51 To check whether different organelles in the same organism might yield different emission peaks, a localized in situ wavelength scan of different PolyP-rich apocrine secretions in the midgut of A. gemmatalis was carried out using multiphoton microscopy. Results showed that different organelles generated similar fluorescence emission profiles, ranging from 516 to 592 nm, with a maximum around 528-549 nm (Figure 6, lines 1-3), which in general corresponded to DAPI-bound commercial sodium PolyP75 emission wavelengths (Figure 7A). Wavelength scans of con-

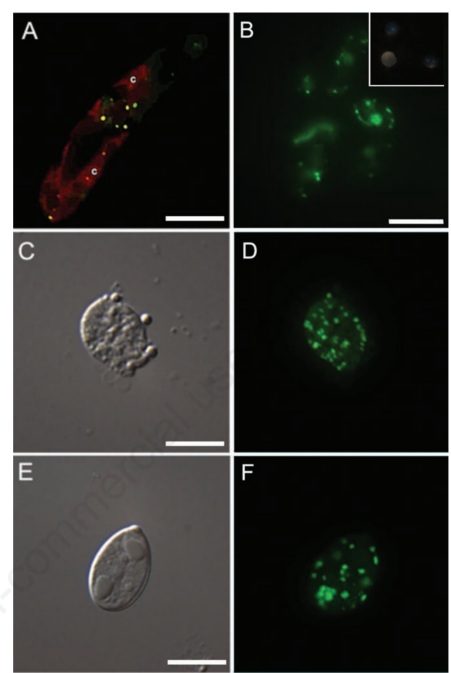


Figure 3. Neither prior fixation nor permeabilization was needed for detecting PolyP by DAPI staining (50 μg/mL for 30 min) in intracellular stores in living cells from (A) *E. gracillis* presenting c: chlorophyll autofluorescence, or (B) *C. albicans*. Formaldehyde fixation prevented DAPI staining of the PolyP granules in *C. albicans* (B, inset). On the other hand, formaldehyde fixation followed by either mechanical lysis (C,D) or Triton X-100 permeabilization (E, F) was needed to visualize subcellular PolyP stores in *Eimeria* cells by DAPI staining. Scale bars: A, 10 μm; B, 10 μm; C, 5 μm; E, 5 μm.



trol areas did not generate similar fluorescence spectra (Figure 6, line 4).

Quantitative fluorimetric analysis of DAPI-PolyP staining in dynamic events

Spectrofluorimetric analysis of analytical grade sodium polyphosphate (PolyP₇₅) (Figure 7A) and PolyP extracted from P. americana WSF (Figure 7B) confirmed fluorescence spectra qualitatively different from DNA-rich (plasmid) samples (Figure 7B), supporting the use of fluorimetric methods for the quantitative analysis of DAPI-PolyP staining. Although some reports have suggested similar approaches with minor variations, 28,30,52 a semi-quantitative measurements of PolyP mobilization in biological samples using DAPI as a probe were yet to be compared against results from more widely accepted methodologies such as scPPX quantification following PolyP binding onto glassmilk particles. 14,39 Here, we measured PolyP mobilization by following DAPI-PolyP fluorescence levels in samples containing PolyP extracted from eggs of the insect R. prolixus during the early stages of development. Results showed that a strong mobilization of PolyP takes place during this period (Figure 7C) and were validated after a comparison against the well-established protocol for PolyP quantification based on the PolyP binding to silica powder in suspension followed by enzymatic PolyP hydrolysis into Pi by recombinant yeast exopolyphosphatase, 14,26,39,53 which showed similar profiles of PolyP mobilization during the early days of development.

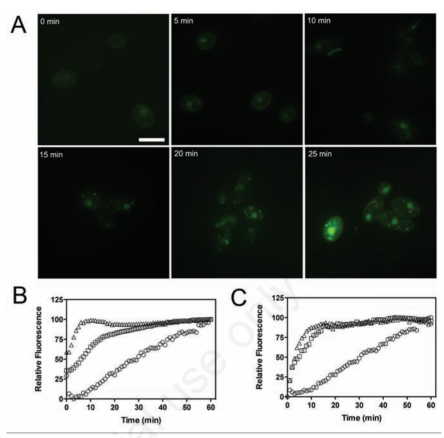


Figure 4. Time-lapse analysis of PolyP DAPI fluorescence. (A) Development of PolyP store labeling in *C. albicans* after staining with 50 μg/mL DAPI in PBS. (B,C) The influence of cell permeability to DAPI for the signal yield was compared over time using samples of *T. cruzi* (squares), *T. brucei* (triangles) and *C. albicans* (circles) incubated with 100 μg/mL DAPI at room temperature. Normalized fluorescence intensities were recorded by a microplate fluorimeter using an excitation/emission wavelength of 350/450 B: nuclear staining) or 350/500 (C: PolyP staining). Scale bar: 10 μm.

Discussion

PolyP-rich organelles comprise the major PolyP reservoir in several eukaryotic models such as yeast and protozoans. 11,54-56 PolyP stores were first detected with metachromatic reactions using basic dyes such as toluidine blue, methylene blue or neutral red, and were, thus, named metachromatic granules. 57-62 These reactions generally depend on a previous step of specimen fixation, thus limiting experiments in live cells. Polymeric compounds other than PolyPs, such as nucleic acids, have also been shown to be stained with basic dyes, 63,64 dragging the attention to the poor specificity of some dyes and the necessity for the use of markers with higher specificity. Tetracycline has been alternatively used to indirectly stain bacterial PolyP.65 Nevertheless, fluorescence mechanism involves binding of tetracycline to divalent cations (usually chelated by PolyP) within PolyP-rich organelles, pro-

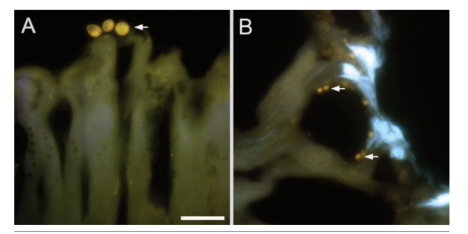


Figure 5. In situ detection of PolyP in midgut cells of A. gemmatali. Cryostat sections after incubation with 0.1 µg/mL DAPI. Apocrine secretions containing PolyP stores were detected emerging from columnar cells (arrows in A). PolyP granules concentrate around goblet cells (arrows in B). Scale bar: 25 µm.





viding signals of low specificity. Alternative methods such as the use of PolyP-binding domain of the *E. coli* exopolyphosphatase (PPBD) fused with an Xpress tag⁶⁶ for the indirect detection of PolyP with the use of secondary antibodies have been reported. 67,68 As yeast exopolyphosphatase (scPPX) is highly specific to PolyP, the method provides both specificity and sensitivity, although as PPBD is not commercially available, in lab production of PPBD requires protein engineering, bacterial strain transformation and protein expression and purification. In the last years, the use of DAPI as a PolyP-rich organelle sensor has gained popularity.^{21,24-26} It yields fluorescence at 525-550 nm after PolyP binding, displays a greenish-vellowish color²³ and is both commercially available and present a fairly good sensitivity and specificity for the detection of PolyP-rich stores. In the present report, we evaluated the effect of different routines for PolyP detection and demonstrated the feasibility of using DAPI as a probe for the detection of PolyP-rich organelles. We further demonstrated that a set of straightforward treatments steps allows a wide range of biological samples to be analyzed following sample-optimized protocols. DAPI was used to detect PolyP stores in biological models after a relatively simple incubation step. Nevertheless, DAPI concentrations needed for PolyP detection were generally above those used for nuclei staining and usually demanded longer incubation times. Whether those conditions might not be optimal for in vivo studies, they are still the least invasive technique for PolyP visualization at present. Nevertheless, it was observed that DAPI intake could not be performed in vivo in some models. This might be the result of both the inability of DAPI to surpass complex cell layers or outward DAPI pumping mechanism of living cells. In those cases, we have been able to observe PolyP stores by using both pre-incubation cell fixation and permeabilization routines. Nevertheless, special care must be taken as we have observed that aldehyde fixation could either prevent or allow visualization of PolyP granules by DAPI.

Subcellular fractionation has been the method of choice for the isolation and posterior characterization of PolyP storage compartments. 20,25,40,48 In those cases, a fast and reliable method of PolyP detection is essential as it allows identification of PolyP-enriched granules among the several resulting fractions. The subcellular compartments we tested did not need previous treatments, and smaller DAPI concentrations (2-10 μ M DAPI) were successfully used. Additionally, DAPI staining in semithin sections have not been widely used and scarcely reported. 49 In part, this is due to a loss of inorganic components during sample preparation, resulting in empty compartments. In

this work, we have used an OCT-embedding protocol avoiding extensive sample manipulation and the steps of alcohol dehydration and medium substitution found in paraffin embedding protocols.

Several microscope settings have been previously used for *in situ* detection of PolyP stores by DAPI likely resulting in a confusing scenario for the outcomer researcher. As PolyP shifts DAPI emission from 450 nm to 525-550 nm,²³ the fluorescence hue changes to a yellowish-green color. Thus, fluorescence emitted from nuclei can theoretically be discerned from fluorescence from PolyP using regular

DAPI filters (data not shown), if color sensitive cameras are used. Nevertheless, emissions from nuclei often become too intense to enable differentiation from the weaker PolyP emissions. The DAPI filter system has thus to be modified and set to block wavelengths shorter than 500-515 nm. This will not prevent nuclei greenish emission area from being detected, yet it should block the most intense blue wavelengths and even allow detection by a monochromatic detection system. During our analyses, these settings more commonly provided optimal results suggesting that it should be used as a default condition for DAPI-PolyP

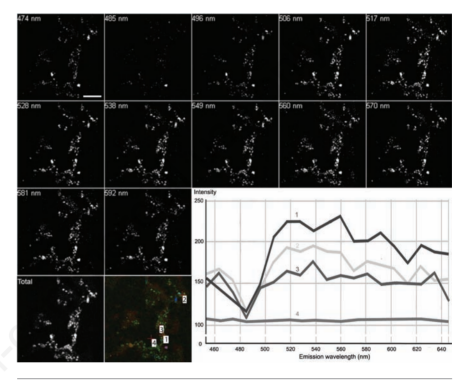


Figure 6. Spectral analysis of PolyP detection in situ. Cryostat sections of the midgut of A. gemmatalis after incubation with 0.1 µg/mL DAPI as observed by a multiphoton microscope at 780 nm excitation wavelength. PolyP stores fluorescence was more intense around 520-550 nm (lines 1-3). The background signal was negligible and did not show wavelength specificity (line 4). Scale bar: 100 μ m.

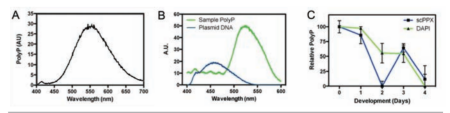


Figure 7. Spectroscopic analysis of PolyP fluorescence after staining with 20 μg/mL DAPI for 30 min at room temperature. The fluorescence intensities were measured by a fluorimeter at 354 nm excitation wavelength. Spectra were obtained from (A) 50 ug PolyP₇₅ on (B) the water soluble fraction from *P. americana* (green line); plasmid DNA was also measured (blue line). (C) normalized DAPI fluorescence at 550 nm for PolyP samples of *R. prolixus* eggs at different development days (green line); the results were compared against an exopolyphosphatase (scPPX)-based assay (blue line), as a control.





detection. Alternatively, previous reports suggested that the use of excitation wavelengths around 400-415 nm could similarly favor PolyP signals.²⁸ A similar adjustment was used with OCT-embedded sections with good results. Nevertheless, care must be taken as DAPI excitation is performed over the maximum 350 nm excitation wavelength and low signals might be generated, hindering proper detection.

Quantification of PolyP levels have been done either with the use of its metachromatic properties⁶⁹ or using enzymatic assays mainly based on the hydrolysis of PolyP by scPPX14,70 or incorporation of Pi from PolyP into ADP and synthesis of ATP by a polyphosphate kinase (PPK).39 Again, the same issues described for detection of PolyP stores are present and the methods either lack specificity and sensitivity, or require recombinant enzymes that are not commercially available. Nevertheless, biotechnological applications of PolyP such as the removal of phosphorous and heavy metals from wastewater demands a faster and cheaper method of quantification that still maintain its reliability.50 In this regard, some authors have approached on a method for DAPI-PolyP quantification.^{26,28,30,50,71} Here we performed an evaluation of relative PolyP levels to ascertain PolyP mobilization on a time scale using fluorescence levels generated by DAPI as a probe. Again, fluorescence levels were measured using the maximum DAPI excitation wavelength (~350 nm). It has been suggested that DAPI-DNA and unbound DAPI signal are significantly removed by using 415 nm excitation, which is imperative to discern PolyP specific signals in conditions of low PolyP content. Our present results were compared with the scPPXbased routine, which remains the most widely accepted PolyP quantification method, and demonstrated no major discrepancy between both methods. This suggests that when significant levels of PolyP are present, therefore giving a proper specific PolyP signal when compared to unbound or DNA/RNA-bound DAPI signals, quantification using excitation wavelengths around 350 nm are allowed. Nevertheless, it has been demonstrated that smaller polymers (<5 Pi residues) do not yield DAPI fluorescence.30 Interference by DAPI binding to contaminants may also account for imprecise quantification. In this regard, DAPI binding to RNA has been described,64 and also shown that treatment of PolyP extracts with RNAse into has a strong influence on DAPI-PolyP signals under specific circumstances.⁷² Analysis of the level of interference of DAPI-RNA on the samples should help to establish how absolute quantification of PolyP by DAPI can be accomplished.

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Correlation between PARP-1 immunoreactivity and cytomorphological features of parthanatos, a specific cellular death in breast cancer cells

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Abstract

In parthanatos, a PARP-1 (poly (ADP-ribose) polymerase 1)-mediated cell death, dissipation of mitochondrial membrane potential, largescale DNA fragmentation and chromatin condensation were observed. In contrast to apoptosis, it does not cause apoptotic bodies formation. Although PARP-1-mediated cell death presents loss of membrane integrity similar to necrosis, it does not induce cell swelling. The purpose of the study was to correlate the immunohistochemical parameters of PARP-1 reactivity and the selected cytomorphological features of parthanatos: the lack of apoptotic bodies and the absence of necrosis in breast cancer (BC) specimens. Immunohistochemistry for PARP-1 was performed on 83 BC specimens. Correlations between parameters of PARP-1 expression and sub-cellular localisation and the presence of apoptotic bodies and necrosis were evaluated. High expression of PARP-1 (immunoreactive score ≥6) was associated with the lack of apoptotic bodies (P=0.013) and with the absence of necrosis (P=0.002). The presence of apoptotic bodies was correlated with re-distribution of PARP-1 from the nucleus to cytoplasm in BC cells (P=0.029). Additionally, a tendency was observed between necrosis and loss of nuclear PARP-1 expression (P=0.049). Our study suggests that PARP-1 may play a crucial role in induction and regulation of specific type of cellular death called parthanatos.

Introduction

PARP-1 has an important role in DNA damage repair and maintaining genome integrity by repairing DNA single strand breaks (SSBs) by base excision repair (BER). It is also involved in chromatin remodelling, regulation of transcription, proliferation processes and differentiation as well as specific cell death called parthanatos. The so-called damage sensor model is proposed in which activated PARP-1 identifies DNA breaks and temporarily binds to the ends of the damaged DNA.1-3 PARP-1's enzymatic activity can increase 500fold on its binding to the damaged DNA, which results in quick elongation of PAR chain.4 ADPribose polymers situated on PARP-1 molecules attached to the damaged DNA function as specific enticements which facilitate repair by recruiting repair enzymes to the site of damage. Additionally, ADP-ribosylation of XRCC1 (vital for BER) greatly enhances its affinity for BER pathway main enzymes, i.e. DNA ligase III and AP endonuclease.2,5

Current knowledge and results of recent studies suggest that the initiation of death cell called parthanatos is closely related with the accumulation of PAR in response to DNA damage.⁶ The polymers function as specific signals and cytophysiological messengers informing about DNA damage that is too extensive to be repaired by the cell itself. Their excess induces parthanatos to prevent and protect against further loss of energy for ineffective DNA repair.^{6,7}

PARP-1-mediated cell death is morphologically and molecularly distinct from other known forms of cell death such as apoptosis, necrosis or autophagy. Parthanatos has been shown to involve mitochondrial transmembrane potential dissipation, chromatin condensation and large DNA fragmentation.8 Unlike apoptosis, it does not cause apoptotic body formation or small scale DNA fragmentation. In vitro experiments with cell lines and caspase inhibitors (z-VAD-fmk, boc-aspartyl-fmk) have conclusively confirmed that the process is caspase-independent and it is not regulated by Bcl-2 proteins.^{5,9} It is worth noting that PARP-1-mediated cell death involves loss of membrane integrity similar to necrosis, yet it does not induce cell swelling.10 Parthanatos is distinct also from autophagy as it does not involve autophagic vacuoles formation or lysosomal degradation.11,12

PARP-1 was widely examined in some types of human tumors, 13,14 but it must be stressed that there are no reports that would describe cytomorphological features of parthanatos in clinical material obtained from breast cancer (BC) patients in correlation with overexpression of PARP-1 as the main protein involved in this type of cell death. The purpose of the study

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Key words: parthanatos, cell death, PARP-1, immunohistochemistry, breast cancer.

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was to correlate the immunohistochemical parameters of PARP-1 reactivity and the selected cytomorphological features of parthanatos, namely the presence of apoptotic bodies and necrosis in BC specimens.

Materials and Methods

Patients

Tissue samples were obtained from 83 patients treated radically for stage II ductal BC, diagnosed between 1993-1994 in the Lower Silesian Oncology Centre in Wroclaw, Poland. The mean age of the patients was 55.2 years. The patients were selected based on the availability of tissues. All patients underwent surgery (Madden mastectomy) with or without adjuvant treatment. Following the applied treatment, the patients were subjected to permanent control in the Lower Silesia Oncology Centre. The study was approved by the Institutional Review Board of the Wroclaw Medical University, Poland.

Tumor samples and immunohistochemistry

Tumor specimens were fixed in 10%

