Phytopathology laboratories regularly prepare microscope slides in order to observe and identify reproductive and/or vegetative structures from plant pathogenic fungi and those used in biological control. Cotton blue (methylene blue) and Trypan Blue are the most commonly used dyes in the preparation of temporary slides for phytopathological studies (MAFIA & ALFENAS, 2016). Lactophenol Cotton Blue (LPCB) is the most commonly used mounting fluid for preparation of microscopy slides. Phenol is the main component of LPCB, being well known as a mutagenic, tumorigenic and toxic to man and the environment (IPCS, 2017). BASAVA et al. (2016) demonstrated that Iodine-glycerol has great potential to replace LPCB as a slide mounting liquid.

Another alternative dye, phloxine B is used in the study of basidiomycotina (RYVARDEN, 1991). However, phloxine B is harmful to the user and the environment and requires costly procedures for waste disposal, whereas Iodine-glycerol is known to be a hazardous substance, and moreover, both dyes are difficult to acquire and also more expensive than azo dyes in the Brazilian market.

In Brazil, the Ministry of Health, through the Agência Nacional de Vigilância Sanitária-ANVISA, allows the use of azo dyes in the food industry for coloring foodstuff and juices. Azo dyes are the main group of synthetic pigments in the world and widely used in the industry (SINGH et al., 2015, CHUNG, 2016). ROCHA et al. (2005)
studied the efficiency of azo dyes for staining phytonematodes and found that dyes containing bordeaux, red bordeaux, and the mixture of bordeaux and 1% indigotin blue, consistently stained in red color the egg masses, eggs, juveniles and females of Meloidogyne incognita, inside the root, compared to the dyes phloxine B and acid fuchsin. However, studies regarding the use of artificial dyes for staining fungal structures are scarce. The objective of this work was to evaluate the efficiency of artificial dyes, used in the industry of juices, at different concentrations or in mixture with glycerol for staining some hyaline, reproductive or vegetative structures, useful for the identification of Oidium sp., Albugo ipomoeae-panduratae (pathogenic fungi in several crops), Pochonia chlamydosporia (nematophagous fungus) and Phytophthym helicoides (important soilborne oomycete that causes damping-off).

The commercial product Docile®, strawberry flavor, was used to prepare solutions containing the artificial dyes sunset yellow and bordeaux S. Solutions were prepared in distilled water at concentrations of 0.5, 1.0, 1.5, 2, 3 and 5% (w/v). These solutions were used as a mounting liquid for the staining of fungal structures. The first assay was performed to determine the best concentration of the dyes for spore staining (SANTANA et al., 2017). In this study powdery mildew (Oidium sp.) samples obtained from leaves of Peplonia axillaris were used. Spores were collected from the leaf lesions by scraping and the slides were mounted by the wet mount slide method with the staining solutions at different concentrations. The slides were covered with glass coverslips. Slides mounted with the cotton-blue dye were used as reference standard (control). Photographs of the spores were obtained with a light microscope at 40X magnification and a 13 megapixel camera. In the second assay, the effect of adding glycerol at concentrations of 0.25, 0.5 and 1.0% (w/v) into the staining solutions (1, 3 and 5%) for staining sporangium of A. ipomoeae-panduratae, hyphae of P. helicoides and chlamydospores of P. chlamydosporia, was studied. The addition of glycerol in the staining solution was based on previous studies reporting better visualization of the fungal structures on slides (VIGNESH et al., 2013). Leaves of Ipomoea batatas with symptoms of white rust (A. ipomoeae-panduratae) and an isolate of P. helicoides obtained from lettuce seedlings were used to mount the slides for staining the sporangia and hyphae, respectively. For staining chlamydospores, the commercial product Rizotec (5.2x10^7 chlamydospores/g), formulated with P. chlamydosporia var. chlamydosporia isolate Pc-10, was used. The chlamydospores were obtained according to the technique of COOLEN & D’HERDE (1972) with modifications (rotation 1,000 rpm for 2 minutes) and placed in test tubes containing the staining solutions or heated at 80°C for 5 minutes, followed by assembly of the slides and acquisition of the images.

Slides mounted with cotton-blue dye stained the spores of Oidium sp. in blue and those with the artificial dyes sunset yellow and red bordeaux S stained the spores with color intensity varying from orange red to magenta (Figures 1A-G). The increasing concentration of the dyes provided greater contrast in the color of Oidium sp. spores. Starting at a concentration of 2%, a stronger staining intensity of the fungal spores was observed when compared to the cotton-blue dye. The strongest color intensity was observed in spores of Oidium sp. stained with the dyes at 5% concentration (Figure 1G). The results of the staining intensity of A. ipomoeae-panduratae sporangia were similar to those of Oidium sp. (Figures 1J and K). The addition of glycerol in the staining solutions did not interfere with the intensity of sporangia staining but resulted in a higher contrast when compared to those stained only with the staining solution and cotton blue (Figures 1H, L, M and N).

In previous studies performed in our laboratory we also observed a similar result while staining hyphae, sporangiophores and sporangia of P. helicoides (SANTANA et al., 2017). In this work we observed the mixing of 2% concentration of both dyes (sunset yellow and red bordeaux) resulted in better staining and contrast of the fungal structures. However, in the present study we observed a higher contrast in the color intensity of P. helicoides hyphae using 3% staining solution, however, when we added 1% glycerol to the 5% solution there was better sharpness when compared to cotton blue (Figures 1O, P and Q). Therefore, glycerol in addition to acting as a hygroscopic agent, avoiding desiccation and changes in the morphological structure of the fungal structures, also increases the longevity of the slide and the contrast of the fungal structure when observed under a light microscope. However, in this work we only evaluated the color intensity of the artificial dyes, color uniformity and their fixation to the fungal spores for two weeks. During this period it was observed a reduction in color intensity, but the contrast of fungal structures remained excellent. Chlamydospores of P. chlamydosporia showed higher intensity and color uniformity at 5% concentration of the staining solution heated with 1% glycerol, compared to the staining solution without
Staining fungal structures with artificial dyes used in the industry of juices.

Figure 1 - Spores of *Oidium* sp. staining with cotton blue (A) and artificial dyes, sunset yellow and red *bordeaux* S, at concentration of 0.5 (B), 1.0 (C), 1.5 (D), 2 (E), 3 (F) and 5% (G). H-K: Stained sporangia of *Albugo ipomoeae-panduratae* with standard dye and artificial dyes at different concentrations. H- Cotton blue. I- Dye solution at 1%. J- Dye solution at 3%. K- Dye solution at 5%. L-N: Staining with 5% dye solution and glycerol in different concentrations. L- Glycerol at 0.25%. M- Glycerol at 0.5%. N- Glycerol at 1.0%. O-T: Staining of *Phytophthium helicoides* and chlamydospores of *Phochonia chlamydospora* with standard dye and artificial dyes. O-Q: Staining of hyphae of *P. helicoides*. O- Cotton blue. P- Dye solution at 3%. Q- Dye solution at 5% and glycerol at 1%. R-T: Staining of chlamydospores of *P. chlamydospora*. R- Cotton blue. S- Dye solution at 3%. T- Dye solution at 5% and glycerol at 1%.
heating and the control (Figures 1R, S and T). At 3% and lower concentration solutions, there was poor uniformity of coloration of the chlamydospores, especially in the staining solutions without heating. Heating the dye can accelerate the fixation reaction and/or increase the penetration of the dye in the thick-walled chlamydospore (BOEDIJN, 1956; EVANS & KIRK, 2017). Sunset yellow and red bordeaux S (amaranth) are classified as azo dyes, since they are synthesized from various aromatic amines derived from tar (BAFANA et al., 2011; AL-RUBAIE & MHESSN, 2012). Thus, as the dyes used in this study are water soluble, their use in the laboratory may be a safer alternative than the traditional LPCB in the preparation of temporary fungal slides. Although azo dyes, including sunset yellow and red bordeaux S, are used in the food industry in Brazil, these dyes exhibit certain toxicity, but their use in the staining of microscopic samples causes less health problems and environmental risks compared to LPCB. Furthermore, azo dyes are cheaper making them of great utility for routine utilization in mycological studies.

ACKNOWLEDGEMENTS

The first two authors thank Universidade Federal de Minas Gerais (UFMG) and Fundação de Amparo à Pesquisa do Estado de Minas Gerais (FAPEMIG) for the scholarship and the opportunity to carry out this research.

CONFLICTS OF INTEREST

The authors declare no conflict of interest. The founding sponsors had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, and in the decision to publish the results.

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