Experience of Using Antifungal Rocima GT for Protection of Paper from Biological Damage Caused by Fungi

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Abstract: This study proposes a method for the chamber disinfection of paper with biocide in the form of fine, volatile droplets using antifungal Rocima GT. This method provides a fungicidal effect, and within a short exposure time, a fungistatic one. At a concentration of 5% Rocima GT solution, the minimum treatment time to ensure the complete disinfection of paper was 15 min. The proposed method of disinfection by a chemical mist was less harmful to paper than disinfection using a swab saturated in a biocide solution. It was noted that when using Rocima GT at insufficient concentrations to inhibit fungal growth completely, Rocima GT can induce, as well as suppress, organic acid produced by Aspergillus niger depending on the concentration.

Keywords: Aspergillus niger; biodeterioration; organic acids; paper; biocides; Rocima GT

1. Introduction

Biodeterioration caused by microscopic fungi is a significant problem affecting different construction materials and buildings, but the threat of biological destruction is especially alarming for the preservation of the historical and cultural heritage of museum archives and libraries [1–5]. Due to their enormous enzymatic activity and their ability to grow at low water-activity values, fungi are able to inhabit and decay paintings, textiles, paper, parchment, leather, oil, casein, glue, and other materials used for historical art objects [6]. The great majority of these fungal species are already able to develop at water activity levels near 0.80, which qualifies them as xerotolerants. Regarding temperature, the optimum growth levels are about 20–25 °C, a common interval for human comfort in libraries, archives, and museums [5].

Most of the micromycete species found in archives and libraries are ubiquitous species, but they are able to grow in conditions different from in nature. When placed in favorable conditions, their germination and development can occur very quickly. The combination of the high bio-receptivity of paper with inappropriate storage conditions or extreme situations associated with water damage makes the material very susceptible to microbial contamination, mainly by fungi [3,7,8]. The bio-receptivity of paper for fungi is considered high due to its hygroscopicity and composition (cellulose, hemicellulose, adhesives, and sizing), which is an abundant source of carbon for fungi [3,7].
The fungi can produce cellulolytic enzymes which break the cellulose polymer into smaller units, weakening the paper structure. Acidic metabolites can be excreted by these microorganisms, promoting acid hydrolysis in the paper fibres. In addition, they can excrete coloured metabolites with several different hues, having the potential to blemish the documents and artworks where they dwell. The release of all these extracellular enzymes and/or pigments acts chemically on the substrate [5].

Paper-based cultural heritage kept in museums and libraries can show notorious signs of deterioration, including foxing stains, caused by fungal colonization [9].

*A. niger* is one of the very viable and harmful species of micromycetes frequently found in archives and libraries and in museums on paper and cellulose textiles [5,6]. Moreover, *A. niger* is known as a producer of organic acids [10], which cause the rapid and deep destruction of various materials [5,11–15]. The overproduction of citric acid is a characteristic feature of *A. niger*. In addition to citric acid, *A. niger* also produces a number of other acids [10,16,17].

In addition to the harmful effect on materials, a high degree of contamination of premises with fungal spores can cause allergic reactions, dermatoses, mycoses, and SBS (Sick Building Syndrome), and some species can induce serious lung diseases [5,18–21]. The fungi that most often cause allergic symptoms belong to the genera *Alternaria*, *Cladosporium*, *Aspergillus*, *Candida*, *Penicillium*, *Mucor*, *Fusarium*, and *Rhizopus* [22], as well as *Aspergillus niger* and some other species [23].

The level of contamination is naturally correlated to dust accumulation, and one can argue that this can be prevented with maintenance [5]. Preventive measures are fundamental in the field of biological document security [24]. In the storage areas of libraries and museums, certain precautions should be taken, such as limited lighting, a low temperature (16–18 °C), and relative humidity in the range of 40–55% [6,25,26].

However, there are situations when documents are seriously damaged and require urgent disinfection processing. Moreover, fungal growth mostly happens between shelves with little aeration or near walls with temperatures below the dew point. Micro-niches are often also created by the wrapping of single objects into extremely tight boxes, not allowing an exchange of air and vapor [6]. There is a wide selection of antifungal substances with different chemical compositions and principles of action. A proper antifungal treatment method of materials should have a broad activity spectrum, good chemical stability, low cost, no adverse effects on the treated material, and be non-toxic to humans. [3,25,27].

Traditional antifungal methods usage may increase the risks of developing drug resistance. Therefore, the development of non-toxic and safer antifungal strategies to treat the fungal biodeterioration of paper products is still an important task and remains a challenge [27].

This issue is most problematic in the case of the mass disinfection of archival collections; when a large number of documents need to be processed, fumigation methods are preferable. Substances traditionally used for the fumigation of documents (formaldehyde, ethylene oxide, methyl bromide, and thymol) have a number of disadvantages that make further use inadvisable, with toxicity to humans and negative effects on document materials being the main ones [3]. In France, for some years, thermal fogging with an alkyl dimethylbenzylammonium chloride solution has been employed successfully for the cleansing of an atmosphere contaminated by fungi [28]. Over the last 10 years, chemical misting methods using disinfection based on quaternary ammonium salts [26], as well as silver nanoparticles [29], have shown their effectiveness in museums, archives, and libraries for protection against microbiological contamination.

Nevertheless, there is little research on chemical misting today, and its usage in conservation is limited. It is important to note that the heating involved in most of these developments may adversely affect the properties of some biocides.

This study proposes a method for the chamber disinfection of paper by the antifungal Rocima GT using ultrasonic action without additional heating.
Rocima GT is an antifungal used for different types of materials, including plasters, textile, and paper. This antifungal is a mixture of chlormethyl-/methylisothiazolone, a quaternary ammonium compound, formaldehyde, and a formaldehyde donor. Rocima GT provides very good bacterial and fungal control [30]. The studies of Rocima GT for use in archives and libraries indicate its efficacy and safety for historical papers. Rocima GT is relatively safe for humans and has a pronounced antifungal activity at a minimum concentration of 0.5% [31].

The aim of this work was to evaluate the effectiveness of Rocima GT against Aspergillus niger using an application of a traditional swab treatment and the new means of aerosol disinfection in chambers. An analysis of the effect of Rocima GT on the production of organic acids was one of the tasks of this study.

2. Materials and Methods

2.1. Fungal Strain Isolation and Identification

For the experiments, we used Aspergillus cf. niger strain P3.17. It was isolated from paper-based documents in 2014 in Saint-Petersburg. The strain was stored in test tubes cultured on Czapek Dox Agar at 12 °C.

Morphological species identification was carried out with the use of guidebooks [32,33]. Molecular identification was performed by Sanger DNA sequencing of the locus TEF1α. To isolate genomic DNA, we took fungal spores by cotton swabs and processed them with GeneJET Plant Genomic DNA Purification Mini Kit (Thermo Fisher Scientific, Lithuania) according to the manufacturer’s protocol. DNA amplification was performed with the use of EF1-1018F (5′-GAYTTCATCAAGAACATGAT-3′) and EF1-1620R (5′-GACGTTGAADCCRAYCRTTGTC-3′) primers [34]. The amplification program was 94 °C for 30 s, 48 °C for 45 s, 72 °C for 1 min. The PCR products were purified by precipitation in acidified isopropanol with the addition of glycogen and sequenced on both strands using BigDye Terminator v3.1 Cycle Sequencing Kit (Thermo Fisher Scientific, USA) on a 3500 Genetic Analyzer (Thermo Fisher Scientific, Waltham, MA, USA).

2.2. Fungal Cultivation and Paper Inoculation

Before undergoing Rocima GT treatment, a sample of experimental paper was inoculated with A. niger suspension (0.1 mL, conidia concentration 1 × 10⁴).

Three types of test papers were taken for all experiments:

B-5—82 g/m², 100% sulfite cellulose, ash content—5% (filler-kaolin);
B-19—82 g/m², 100% sulfate cellulose, ash content—5% (filler-kaolin);
B-25—85g/m², 100% bleached cotton half-mass, ash content—9% (filler-kaolin).

In the case of studying the biocidal effect by the method of swab and treatment in a chamber, in addition to experimental paper samples with specified compositional parameters, commercially produced, typographical printing paper was also used. Typographical printing paper contains equal parts cellulose and wood pulp and 15% kaolin. The physical and mechanical parameters of this paper have not been tested.

After treatment in the chamber or by a swab saturated in Rocima GT solution, samples of paper with spores of A. niger were placed in Petri dishes on the Czapek-Dox nutrient medium. The samples were incubated at 25 °C for 1 month. The radial growth rate was assessed by the diameter of the colony. The experiments were carried out in triplicate.

In order to investigate the effect of Rocima GT on the fungal resistance of paper during storage, the paper samples exposed to a cold mist of Rocima GT were stored in paper envelopes at 40% relative humidity and room temperature. After 2 months, the samples were placed on a nutrient Czapek-Dox medium in Petri dishes, inoculated by A. niger, and incubated at 25 °C for 15 days.

In order to determine the composition of organic acids in the cultural liquid, the cultivation of fungus was carried out on a liquid Czapek medium (g/L: glucose −20; NaNO₃−2.0; KH₂PO₄−1.0; MgSO₄•7 H₂O−0.5; KCl−0.5; FeSO₄•7 H₂O−0.01) in static conditions at 25 °C (volume—50 mL). The cultivation time was 8 days.
The effect of Rocima GT on the growth of *A. niger* in liquid culture was determined by weighing of dry biomass of mycelium.

### 2.3. Rocima GT Treatment

#### 2.3.1. The Treatment of the Inoculated Paper Samples with Rocima GT

Paper samples were treated by Rocima GT solution by two means: traditional methods using a swab and by cold mist in a special chamber.

**Swab Treatment**

The samples of paper were carefully blotted with a gauze swab moistened with a biocide solution. Aqueous solutions of a biocide concentration of 0.5 and 2% were used.

**Treatment in Chamber**

The Rocima GT aqueous solution was aerosolized into a cold mist using ultrasonic action to avoid negatively affecting the active substance with heat. This method allows the aqueous solution to be converted into a state of fine droplets due to high-frequency vibrations. For the experimental study of Rocima GT, the cold mist’s antifungal activity, and its effect on the mechanical properties of paper, a special chamber (volume 80 L) was constructed. This chamber was connected by an adapter tube to a vessel filled with Rocima GT 5% aqueous solution (1 L), in which ultrasound generators were placed. Three generators (power 24 W) were used at the same time. The chamber was filled with Rocima GT solution mist for three minutes. In this case, the approximate amount of liquid transferred to the state of mist was 50 mL, which corresponded to approximately 625 µL/L.

In order to select the optimal disinfection time, paper samples contaminated by *A. niger* spores were fumigated by Rocima GT for 5, 10, 15, 20, 25, and 30 min.

#### 2.3.2. The Cultivation of *Aspergillus niger*, Medium Enriched with Rocima GT

To study the effect of Rocima GT on the organic acids produced by *A. niger*, the biocide was added to the medium in concentrations 0.01%, 0.05%, and 0.1%.

### 2.4. Organic Acids Analysis

#### 2.4.1. Sample Preparation

In order to determine the content of organic acids in the culture broth of *A. niger*, an aliquot of the culture broth (10 mL) was acidified with 0.1 M HCl to pH 1.0 to dissolve insoluble salts. Then, this aliquot was diluted by distilled water and was passed through a cation exchanger (KU–2–8 C) to free the acids from their salts. The resulting aqueous solution of organic acids was evaporated at 40 °C. The dried extracts were dissolved in pyridine (30 µL) and BSTFA (N,O-bis—3-methyl-silyl-3-F-acetamide) (30 µL) and incubated at 100 °C for 15 min.

#### 2.4.2. Gas Chromatography–Mass Spectrometry

The derivatized samples were analyzed by gas chromatography–mass spectrometry (GC-MS) on a Maestro instrument (Interlab, Moscow, Russia) with an Agilent 5975 mass-selective detector (Column HP-5MS, 30 m × 0.25 mm × 0.25 µm). Chromatography was performed with linear temperature programming from 70 °C to 320 °C in the mode of constant carrier gas flow through the column (1 mL/min); the carrier gas was helium. Mass spectra were scanned in the range of 50–750 m/z with a frequency of 1.6 scans/s. Chromatograms of the samples were recorded using the total ion current. Mass spectrometric information was processed and interpreted using the AMDIS program [35], standard NIST2005 library, and the library of standards compounds of BIN RAS. Quantitative interpretation of chromatograms was performed by the internal standard method using tridecane in the UniChrom program [36].
2.5. Estimating the Physical Characteristic of Paper

The effect of the Rocima GT treatment on the mechanical and physical properties of the paper was assessed by the change in the values of the fracture resistance index. The measurements were made on the device “I-1-3” (Russia) for determining the fracture strength with multiple kinks [37]. The tests were carried out in 10 repetitions in the longitudinal and transverse directions. Three types of test papers were the same as the ones used for inoculation by fungi (B-5, B-19, B-25). Heat aging of the test paper samples was carried out in the thermostat «EKROS ES-4610» at 105 °C for 72 h [38,39].

The measurements of whiteness were made on the device “photoelectric spherical photometer FMSh-56M” (Russia) [40]. The tests were carried out in 10 repetitions.

2.6. Statistical Analysis

Statistical analysis was performed via the Student’s *t*-test using Microsoft Excel and Origin Pro software.

3. Results

3.1. Fungal Species Identification

The obtained TEF1α sequence of *A. niger* isolate P3.17 was deposited in a GenBank with the accession number MH029907. It was used for a MegaBLAST search against the fungal sequences of the NCBI Nucleotide database. The top score hits included the *A. niger*, *A. welwitschiae*, and *A. awamori* sequences. To refine the identification, we aligned the TEF1α sequence of the isolate P3.17 with sequences of well-annotated Aspergillus strains [41,42]. The studied sequence was identical in *A. niger* ITEM 4501 (FN665404) and *A. welwitschiae* CBS 139.54b (QQZQ01000062.1:113618-114218). Based on the combined results of morphological and cultural studies, the strain was identified as *Aspergillus niger Tiegh*.

3.2. Evaluation of the Viability of *A. niger* after Treatment by Rocima GT

First, four types of experimental paper, B5, B 19, and B 25, and typographical printing paper inoculated by *A. niger*, were treated with a 0.5 and 2% Rocima GT solution by swab. A treatment by swab provided a good antifungal effect at both concentrations for 15 days. On the 30th day, weak growth of *A. niger* was observed on only one sample of the typographical paper treated by the 0.5% Rocima GT solution. In the control samples without treatment, the growth of mycelium began on the second day after the inoculation of the spore suspension.

In the case of the chamber treatment, after a week of incubation of the paper samples on a nutrient medium, the growth of the mycelium of *A. niger* was not observed in any of the experimental paper samples (Figure 1). On the 15th and 30th days (Figure 2), growth was observed on all samples (B5, B 19, and B 25, and the typographical printing paper) treated for 5 min and in one sample treated for 10 min (growth rate was 30–50% less than in the control). There was no significant difference in the growth rate of fungi depending on the type of paper.

After treatment by the Rocima GT and storage for two months, the paper samples were reinoculated by *A. niger*; the antifungal effect of Rocima GT was absent for all types of paper.

3.3. The Influence of Rocima GT on the Physical Properties of Paper

The comparison of the effect of the Rocima GT mist on the strength and whiteness of paper with an aqueous solution (applied by swab) of 0.5% and 2% (Table 1) showed that the greatest negative effect on the mechanical strength of the paper appeared to be the treatment with a 2% water solution of Rocima GT. The treatment of paper by a 0.5% aqueous solution of Rocima GT, as well as treatment by a mist of 5% aqueous solution of Rocima GT, affected the strength of paper made of sulphite cellulose (B-5) and cotton (B-25) in the same way. For paper made of sulphate cellulose (B-19), the average value of the fracture resistance after treatment with the 0.5% aqueous solution of Rocima GT decreased by 8% more than after treatment with Rocima GT “vapors.”
Figure 1. Antifungal effect of aerosol treatment by Rocima GT, depending on the exposure time (typographical printing paper): (a) control (treatment was not carried out); (b)—5 min; (c)—10 min; (d)—15 min; (e)—20 min; (f)—30 min.
Figure 2. Samples of paper inoculated by *A. niger* on the 30th day of incubation after aerosol treatment by Rocima GT (typographical printing paper): (a) control (treatment was not carried out); (b) treatment by 5 min; (c) treatment by 15 min; (d) treatment by 30 min.

After thermal aging, a decrease in the fracture resistance index was observed in the treated samples of paper made of sulfate cellulose (B-5) up to 30%. Samples from the cotton half-mass (B-25) showed a tendency towards a slight increase in this indicator. For specimens made of sulfate cellulose (B-19) that were subjected by thermo-aging, the value of the fracture resistance index practically did not change. This applies to paper samples that have undergone either an aqueous treatment with a Rocima GT solution (by using swab) or treatment in the Rocima GT “vapors.” The whiteness of all three types of paper after various treatments either decreased insignificantly or remained unchanged. The decrease in the whiteness of various types of paper after heat aging did not depend on the antifungal treatment method and corresponded to the whiteness of paper without treatment after heat aging.
### Table 1. Indicators to breakage resistance and whiteness of paper after various types of treatments.

<table>
<thead>
<tr>
<th>Type of Paper</th>
<th>Treatment</th>
<th>Break Resistance (h.p.)</th>
<th>Whiteness</th>
<th>Thermo-Aging</th>
<th>Whiteess</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Longitudinal</td>
<td>Transverse</td>
<td>Longitudinal</td>
<td>Transverse</td>
</tr>
<tr>
<td>B-5</td>
<td>without treatment</td>
<td>11.6 ± 2.4</td>
<td>32.6 ± 3.9</td>
<td>71.8 ± 0.2</td>
<td>8.9 ± 1.7</td>
</tr>
<tr>
<td>B-19</td>
<td>distressed water</td>
<td>21.4 ± 3.4</td>
<td>30.1 ± 5.0</td>
<td>71.2 ± 0.3</td>
<td>16.8 ± 6.2</td>
</tr>
<tr>
<td>B-25</td>
<td>distilled water</td>
<td>8.9 ± 0.6</td>
<td>6.5 ± 0.5</td>
<td>83.0 ± 0.0</td>
<td>10.3 ± 0.9</td>
</tr>
<tr>
<td></td>
<td>0.5% aqueous solution of Rocima GT</td>
<td>16.3 ± 3.2</td>
<td>24.4 ± 4.7</td>
<td>69.0 ± 0.6</td>
<td>15.1 ± 3.5</td>
</tr>
<tr>
<td>B-19</td>
<td>2% aqueous solution of Rocima GT</td>
<td>46.4 ± 7.3</td>
<td>24.7 ± 2.9</td>
<td>69.5 ± 0.0</td>
<td>62.5 ± 17.3</td>
</tr>
<tr>
<td>B-25</td>
<td>2% aqueous solution of Rocima GT</td>
<td>4.9 ± 0.3</td>
<td>4.0 ± 0.0</td>
<td>84.0 ± 0.0</td>
<td>5.8 ± 0.4</td>
</tr>
<tr>
<td></td>
<td>water mist</td>
<td>18.7 ± 2.6</td>
<td>34.3 ± 7.3</td>
<td>69.0 ± 0.0</td>
<td>10.6 ± 2.7</td>
</tr>
<tr>
<td>B-19</td>
<td>mist of 5% aqueous solution of Rocima GT</td>
<td>168.5 ± 54.0</td>
<td>30.2 ± 7.0</td>
<td>70.0 ± 0.0</td>
<td>152.5 ± 49.1</td>
</tr>
<tr>
<td>B-25</td>
<td>mist of 5% aqueous solution of Rocima GT</td>
<td>7.4 ± 0.6</td>
<td>5.2 ± 0.4</td>
<td>84.0 ± 0.0</td>
<td>10.3 ± 1.2</td>
</tr>
</tbody>
</table>

3.4. Effect of Rocima GT on Organic Acids Produced by A. niger

Rocima GT added in the liquid Czapec-Dox medium suppressed the growth of A. niger. For the A. niger growth suppression by 10%, the Rocima GT concentration was 0.01%; for growth suppression by 50%, the Rocima GT concentration was 0.05%, and for growth suppression by 80%, the Rocima GT concentration was 0.1%.

The sporulation of A. niger was inhibited under all concentrations of Rocima GT used. At high concentrations of Rocima GT, 0.05% and 0.1%, the growth of the mycelium occurred mainly by pellets, and the surface mycelium film was not formed.

On the eighth day of growth, oxalic, citric, succinic, fumaric, malic, and gluconic acids were identified in the culture liquid of A. niger grown on a medium without biocides. The effect of Rocima GT on the production of organic acids by A. niger was manifest in different ways, depending on the concentration of the fungicide in the nutrient medium (Table 2). At 0.01% Rocima GT suppressed the formation of oxalic acid and had virtually no effect on the formation of other organic acids. At 0.05% Rocima GT, the oxalic acid content increased by 1.6 times. The amount of citric acid under the influence of the fungicide and in the control did not change significantly, while for fumaric, succinic, and malic acids, tendencies to decrease were observed. At 0.1% Rocima GT, the production of all organic acids was inhibited.
Table 2. Effect of Rocima GT on organic acid production by *A. niger*.

<table>
<thead>
<tr>
<th>Rocima GT Concentration, %</th>
<th>Growth Suppression, %</th>
<th>The Amount of Organic Acids, mg/g Mycelium</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Oxalic Acid</td>
</tr>
<tr>
<td>0%(control)</td>
<td>0</td>
<td>125.0 ± 34.8</td>
</tr>
<tr>
<td>0.01%</td>
<td>10</td>
<td>20.0 ± 6.2</td>
</tr>
<tr>
<td>0.05%</td>
<td>50</td>
<td>194.8 ± 28.7</td>
</tr>
<tr>
<td>0.1%</td>
<td>80</td>
<td>- *</td>
</tr>
</tbody>
</table>

*"- acids not found.

4. Discussion

The results indicate the method based on chemical misting of paper using an aqueous solution of Rocima GT by ultrasonic fogging provides a good antifungal effect and, with a short exposure time, a fungistatic effect. The fungistatic effect of the Rocima GT solution mist is observable after 5 min of treatment. However, a longer exposure is required for the fungicidal effect. At the selected concentration of 5% Rocima GT solution, the minimum treatment time that ensures the complete disinfection of paper is 15 min.

At the same time, the treatment of paper made of sulphite cellulose and cotton half-mass with a cold mist of a 5% aqueous solution of Rocima GT leads to a lower decrease in the paper’s strength than treatment with an aqueous solution of 0.5% and 2%. This shows the advantage of this method compared to disinfection by the application of a swab saturated in a biocide aqueous solution.

The scope of the biocide is very wide, but the use of Rocima GT for the disinfection of cultural heritage is not very common today. Nevertheless, the studies [31], as well as our study, have shown the relative efficacy of the use of Rocima GT for paper disinfection.

According to our data, using this biocide should be related to Rocima GT having both a stimulating and an inhibitory effect on the acid production activity of *A. niger*, depending on the concentration. The *A. niger* strain cultivated on a Chapec-Dox media produces oxalic, citric, succinic, malic, and fumaric acids.

Organic acids are among the most aggressive fungal metabolites in relation to many materials, including paper. As paper is highly susceptible to acidic hydrolysis, acid production represents an important biodeterioration potential [5,15]. The acids’ production of fungi can damage paper not only due to a low pH; among all the acids secreted by fungi, the strongest oxalic acid is able to form insoluble salts-oxalates with di and trivalent metal. There is evidence that CaCO$_3$-containing paper susceptible to fungal growth accumulates calcium oxalate, along with the disappearance of carbonate [43].

In general, the biocide suppressed the formation of organic acids by the fungus. However, IC$_{50}$ Rocima GT (0.05%) had a stimulation effect on the productivity of oxalic acid by *A. niger*. Nevertheless, it should be kept in mind that we are talking about the productivity of acid formation, i.e., when the growth of mycelium is suppressed by 50%, the total amount of acid released is less than in the control. Perhaps, in this case, oxalic acid can help maintain the integrity of the membranes and, accordingly, greater resistance to the action of a biocide, which is consistent with the hypothesis of the role of oxalic acid in maintaining membrane stability [44]. Most likely, the decrease in the production of citric, malic, fumaric, and succinic acids in fungi under the action of Rocima GT is associated with their general inhibitory effect on the growth and processes of the energy metabolism of fungi.

It should be noted that the production of organic acids by fungi is a very labile process by default and is influenced by a lot of different factors. The change in acid production under various influences, including biocides, will differ mainly depending on the species and even the strains of fungi [17].
In addition to the possible increase in the oxalic acid production under the action of Rocima GT, the effect of low concentrations of biocides, including Rocima GT, contributes to a fairly rapid formation of fungal resistance to the fungicide. The formation of resistant strains requires an increase in the used concentrations of biocides or the use of new ones [25].

Thus, the choice of the optimal concentration sufficient for the complete antifungal effect is one of the most important factors for the successful disinfection of paper and other materials.

When disinfecting with a cold mist of Rocima GT aqueous solution, fungal spores die; however, upon further storage, the fungicidal effect is lost. Disinfection by swab shows the same trend. The lack of preservation of the fungicidal effect over time could be problematic.

The issue of the short-term effect of some biocides was also noted in other studies, including the development of the chamber methods of disinfection by a chemical mist of quaternary ammonium salts [26]. If proper storage conditions are not observed, treatment does not protect against secondary contamination of fungal spores and subsequent damage to documents.

The disinfection should be used only on the basis of a preliminary mycological examination of documents and an estimation of the threat to document safety. Preventive treatment by biocide, less prolonged fumigation, and at a lowered concentration of the active substance are not desirable.

5. Conclusions

The results of this study indicate that the method of the chamber fumigation of paper by a Rocima GT solution mist is promising. This method will allow the antifungal treatment of a large volume of archival papers. In addition, the chemical mist treatment does not deform the paper. Research of this method of disinfection using Rocima GT and other water-soluble biocides will be continued for introduction in the practice of conservation and restoration measures in archives and libraries.

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