Blemish Formation in Processed Microfilm

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Oxidizing and reducing agents generated by paper cartons during storage may react with the image silver to form blemishes. The type of fixing bath, concentration of chlorine in the wash water, washing time and composition of the storage cartons can be factors that accelerate or retard the attack of the peroxide on the image silver. Microfilm washed after fixation in distilled water or chlorine-free tap water formed blemishes when exposed to peroxide paper. A trace of silver chloride in the image silver of processed microfilm augmented the formation of the natural type blemishes when exposed to peroxide paper, but high concentrations of residual silver chloride inhibited blemish formation. During storage, some storage cartons evolved formaldehyde, formic acid, and ammonia which created a chemical environment favorable for blemish formation in microfilm in the presence of peroxide. A correlation was found between the incidence of blemishes and the brand of paper storage cartons. A procedure was developed for testing processed microfilm to determine its susceptibility to blemish formation. According to the present theory, peroxide reacts with the silver in the image, forming colloidal silver which imparts a yellowish or reddish color to the blemishes.

Key Words: Archival record film, blemishes due to aging, colloidal silver, microfilm, microfilm storage cartons, permanent record film, peroxides.

1. Introduction

Blemish formation in stored microfilm has been under investigation for some time [1, 2, 3]. The nature of the problem and the types of blemish formation have been described [1, 2]. It has been shown that oxidizing and reducing agents, probably peroxides, emitted by paper cartons in which the microfilm is stored attack the silver in the image, causing yellowish or reddish blemishes [4]. In the present study, the processing procedure was investigated to determine its effect on blemish formation. In this paper the term “blemish” includes various types of spots and edge attack on characters and other image boundaries.

2. The Role of the Processing Procedure in Blemish Formation in Microfilm

In the investigation of processing variables, four fine grain 16 mm microfilms, made by three manufacturers, were used. The processing solutions and times of developing, stopping, and fixing are given in the appendix. The microfilm samples were processed in beakers and agitated with a glass rod. The chlorine concentration of the wash water was determined by a Taylor chlorine analyzer [7] using the o-tolidine dihydrochloride reagent.

The peroxide source for testing the microfilm was a chromatographic paper, 0.03 cm thick, which had been treated in aqueous hydrogen peroxide. The test procedure is described in section 5. From 2 to 4 cm² of the peroxide paper formed spots in areas of microfilm containing natural spots. All microfilms were dried at least 1 day after processing or other treatment before exposure to peroxide paper.

2.1. Effect of the Fixing Bath and Wash Water

A study was made to determine if processed microfilm formed blemishes when the thiosulfate was removed by washing in chlorine-free water. The fixing baths tested were a neutral bath, a nonhardening acid bath, and an acid hardening bath. The wash waters were distilled water and chlorine-free tap water which had a pH of about 8. Chlorine-free tap water was obtained by storing cold tap water until the chlorine disappeared or by adding sodium sulfite to the tap water to neutralize the chlorine. For a chlorine concentration of 1.2 ppm, 50 ml of a sodium sulfite solution containing 0.500 g per liter was sufficient for 4 liters of tap water. About 50-cm lengths of the microfilms were exposed to a test pattern having various kinds of information, developed, fixed and washed in 1-liter baths of the wash water in 1.5-liter beakers. Three baths of the wash water were used for a 3-min wash and for longer washes a bath was used for each additional 5 min of washing. The sample was trans-
exposed to peroxide paper if the processing procedure
was such that the residual thiosulfate was reduced to
results show that micro film developed blemishes when
had a residual thiosulfate concentration of 10
were almost contiguous to each other for microfilm
had a very low level. A higher concentration of residual
and 8 cm² of the peroxide paper. No blemishes de-
were washed for 30 min in running tap water at 27 °C,
amounts of silver chloride enhance blemish formation but larger concentrations inhibit it.
Water from city taps generally contains 0.1 to 1 ppm
Tests were made of the effect of chlorine in the
had a residual thiosulfate concentration of 10 μg per
process was such that the residual thiosulfate was reduced to
fer from beaker to beaker and agitated with a
glass rod. The temperature of the processing solutions
and wash water was 26 °C.
The processed microfilms developed spots and, in
some cases, edge attack when exposed to peroxide
paper. The microfilm fixed in the neutral and the non-
hardening baths and washed in distilled water de-
veloped much larger spots than microfilm fixed in the
nonhardening and the acid hardening baths and
washed in chlorine-free tap water. The spots in the
latter two fixing procedures were more numerous and
had sharper edges than those for the first two fixing
procedures with distilled water washing. The spots
were almost contiguous to each other for microfilm
fixed in the acid hardener bath. The spots from all
fixing baths appeared white when they first developed
but turned yellow or yellowish as the peroxide attack
continued. The edges of the spots were slightly fuzzy.
No significant differences were observed between
blemishes formed after a 3-min wash and a 15-min wash.
The results of the tests are shown in table 1, where
the observed effect for each processing procedure and
the amount of peroxide paper necessary to induce
blemish formation are indicated. The residual thio-
sulfate in the microfilm samples that formed blemishes
ranged between 0 and 3 μg per 6 cm² of the image
area. Microfilm D retained more residual thiosulfate
than the other microfilms and exhibited only slight
blemish formation for the neutral and the nonharden-
ing baths and none for the acid hardening bath when
washed in chlorine-free water. Microfilm A, fixed in
the acid hardening bath and washed in distilled water,
had a residual thiosulfate concentration of 10 μg per
6 cm², making it resistant to peroxide attack. The
results show that microfilm developed blemishes when
exposed to peroxide paper if the processing procedure
was fixed in an acid hardening bath. Samples
were washed for 30 min by placing them in a succession of tap water baths containing 0.3 ppm
of chlorine for 2, 3, 5, 10, and 10 min, respectively.
Exposure to 4 cm² of peroxide paper produced some

<table>
<thead>
<tr>
<th>Processing procedures</th>
<th>Microfilm</th>
<th>Neutral fix and washed in distilled water for 3 min</th>
<th>Nonhardening acid fix and washed in distilled water or chlorine-free tap water for 3 min</th>
<th>Acid hardening fix and washed in chlorine-free tap water for 3 min</th>
<th>Acid hardening fix and washed in distilled water for 40 min</th>
</tr>
</thead>
<tbody>
<tr>
<td>A...</td>
<td>Many spots and edge attack.</td>
<td>Many spots and edge attack.</td>
<td>Many spots but no edge attack.</td>
<td>No spots or edge attack.</td>
<td></td>
</tr>
<tr>
<td>B...</td>
<td>Many spots and edge attack.</td>
<td>Many spots and edge attack.</td>
<td>Many spots but no edge attack.</td>
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</tr>
<tr>
<td>C...</td>
<td>Many spots and edge attack.</td>
<td>Many spots and edge attack.</td>
<td>Many spots but no edge attack.</td>
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</tr>
<tr>
<td>D...</td>
<td>Few spots but edge attack.</td>
<td>Few spots but edge attack.</td>
<td>No spots or edge attack.</td>
<td></td>
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<tr>
<td>Amount of peroxide paper used.</td>
<td>1 to 2 cm²</td>
<td>2 cm²</td>
<td>4 to 8 cm²</td>
<td>8 cm².</td>
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</table>
spots but a considerable resistance to blemish formation was noted when compared to samples washed in chlorine-free tap water.

After fixation, microfilm A was washed in running tap water at 27 °C, containing 1.2 ppm chlorine for 3, 6, 9, 15 and 30 min. A sample was removed at the end of each washing time. The processed microfilm was exposed to 4 cm² of the peroxide paper. The microfilm samples fixed in the acid hardening bath developed a high incidence of yellow spots for the 3-min wash, a few in the 6-min wash, and none in the 9, 15, and 30-min washes. The microfilm samples fixed in the nonhardening bath developed many light yellow spots for the 3-min wash, with slight yellowing of the image silver, a decrease in number of spots and no yellowing for the 6-min wash, a few spots for the 9-min wash, and none for the 15 and 30-min washes.

Thus a high concentration of chlorine in the wash water formed silver chloride in the image silver making it resistant to yellowing and spot formation when exposed to peroxide. Furthermore, resistance to blemish formation increased with the washing time for 1.2 ppm of chlorine in the tap wash water and 10 to 15 min was required to impart complete resistance. At the end of the 3-min wash, the thiosulfate had reached a low concentration and the chlorine began to form silver chloride. As the washing progressed, more silver chloride was formed and it finally reached a concentration which inhibited blemish formation. The spots which formed after fixation in the acid hardening bath and a 3-min wash were more yellow than those obtained from the nonhardening bath and a 3-min wash. This would be the expected effect of acid carried over into the wash water from the acid bath, since chlorine reacts more rapidly in the presence of acid to form silver chloride. Since the tap water had a pH of about 8, a high concentration of chlorine and a considerable washing time was necessary to form sufficient residual silver chloride to inhibit blemish formation. Tap water containing low concentrations of chlorine, such as 0.3 ppm, may not form a sufficient amount of silver chloride to inhibit blemish formation even after a 30-min wash but the tests described above for 0.3 ppm of chlorine in wash water showed that some resistance to blemish formation was obtained.

Microfilm containing sufficient silver chloride to make it resistant to blemish formation loses this resistance if treated for 10 min in a 0.2 percent solution of ammonium hydroxide to remove the silver chloride. Microfilm so treated showed only an even yellowing of the image silver when exposed to 2 to 4 cm² of the peroxide paper.

It is not known how long silver chloride formed from chlorine in the tap water will impart resistance to blemish formation from peroxide attack. But in section 3 it is shown that chemicals given off by the storage cartons do react with the image silver, causing blemish formation in the presence of peroxide. Evidently the silver halide, if present, is partially reduced by the peroxide.

2.3. Effect of Iodide in the Fixer

It has been shown [12] that the silver in a photographic image is partially sulfided during fixation. If iodide is added to the fixing bath most of the sulfiding is prevented, leaving a layer of silver iodide on the silver grains.

Henn, Wiest, and Mack [14] have reported that collections of microfilm processed in iodide-containing developers or fixing baths to be relatively free from blemishes. They recommend the use of 0.2 g of potassium iodide per liter of fixing solution to give microfilm greater resistance to blemish formation.

Tests were made on four microfilms by adding 0.2 g of potassium iodide to each liter of the acid hardening fixing solution. After development, the samples were fixed and washed in chlorine-free tap water. The chlorine-free tap water was used to avoid the blemish inhibiting effect of the chlorine. The samples were exposed to 2 to 8 cm² of the peroxide paper. The microfilms showed more even yellowing of the image silver than the controls without iodide, indicating that iodide had made the image silver more susceptible to attack by peroxide. (If natural blemishes were caused by peroxide from the boxes, this finding seemed inconsistent with the field observations.) The resistance to spot formation was about the same as the controls.

However, if the microfilms fixed with iodide in the fixer were washed in the chlorine-free tap water to which 1 ppm of chlorine was added, they, like the controls, resisted blemish formation and did not show an even yellowing. Old microfilms which had been processed with iodide in the fixing bath formed blemishes when exposed to peroxide at low levels of 2 to 4 cm² of the peroxide paper.

To simulate actual practice, a study was then made of the effect of storing processed microfilm in fresh storage cartons. The films in cartons were placed in a desiccator jar for 10 days at 86 percent relative humidity and 26 °C and then the samples were exposed to 2 to 4 cm² of the peroxide paper. The samples processed with potassium iodide in the fixer developed a few blemishes but samples processed without potassium iodide in the fixer developed many more. These tests confirmed the field observations and suggested that chemicals in addition to peroxide were generated in the storage carton and played a part in the chemistry of blemish formation.

2.4. Natural-Type Blemishes Obtained With Acid and Silver Chloride in the Image

A small amount of silver chloride was formed in the image silver of old films and freshly processed films. When these films were exposed to peroxide, sharp-edged blemishes formed which resembled very closely those found in nature, even duplicating the ring formation and edge attack. The films were exposed to about 25 cm² of the peroxide paper at 76 percent relative humidity and 26 °C. A higher relative
humidity did not produce the natural-type blemishes. Although these tests may not be applicable to microfilm as normally processed and stored they do suggest that a trace of silver chloride contributes to blemish formation under natural storage conditions.

For these experiments, processed films were treated 10 min in a solution containing 0.025 g of sodium chloride and 3 ml of glacial acetic acid per liter, dried and exposed to peroxide paper. (Formic acid may be substituted for the acetic acid.) The 0.025 g of sodium chloride per liter gave a solution containing 15 ppm of the chloride ion which is close to the median value of 13 ppm in the water supplies of large cities [6]. The oxygen dissolved in the sodium chloride-acetic acid solution oxidized the image silver to silver ion which combined with the chloride ion to precipitate silver chloride in the image silver. The silver chloride formed in microfilm by the sodium chloride-acetic acid solution may be removed by treating the film for 10 min in a 0.2 percent solution of ammonium hydroxide. After such treatment, the image silver showed only an even yellowing after exposure to peroxide paper. When microfilm was treated in a solution of 3 ml of acetic acid per liter without the sodium chloride and exposed to peroxide paper, no spot or edge blemishes formed in the image silver, only an even yellowing.

The microfilms A, B, C and D, reported in section 2.1, which readily formed blemishes when exposed to peroxide paper were treated in the sodium chloride-acetic acid solution and dried. Peroxide exposure produced natural type spots, some with ring structure, and edge attack. There was no even yellowing, indicating that silver chloride was formed throughout the image. The edges of the spots were sharp and the spots were deep yellow, showing the effect of silver chloride. The number of spots was much less than before the sodium chloride-acetic acid treatment since too much silver chloride had been formed at most points in the image for spot induction.

Blemish formation was induced in old processed microfilms by treatment in chlorine gas or an acid solution of chlorine and subsequent exposure to peroxide. The films were exposed to chlorine gas by suspending the samples over a chlorine solution in a 1-liter closed glass bottle. The chlorine solution was prepared by dissolving 0.250 g of technical grade calcium hypochlorite containing 70 percent calcium hypochlorite in 1 liter of water and adding 5 ml of glacial acetic acid. Approximately 400 ml of this solution was added to the bottle. The film samples were soaked in distilled water for 10 s, squeegeed with a dry glass rod and suspended immediately in the chlorine gas above the solution for 20 to 60 s, dried overnight and exposed to peroxide paper. This method produced spot and edge attack but each brand of microfilm required a different time of exposure to the chlorine gas.

To determine whether silver chloride was formed in the image silver by the chlorine gas, microfilm samples were treated in a 0.2 percent ammonium hydroxide solution for 5 min to remove any existing silver chloride, dried overnight, treated 10 s in distilled water, squeegeed, treated to the chlorine gas for 30 s and exposed to peroxide paper. These films had spot and edge attack.

Microfilm samples were immersed for 5 s in the chlorine solution described above, dried and exposed to peroxide paper. These microfilms showed spot and edge attack but were more resistant to peroxide because more silver chloride had been formed in the image silver.

The findings with respect to chlorine and chloride ion raised a question about the effects of the other halides, silver bromide and silver iodide. Old processed microfilm samples, blemish free, were treated 10 min in solutions of sodium bromide containing 0.045 g/l and 3 ml of glacial acetic acid, and potassium iodide containing 0.060 g/l and 3 ml glacial acetic acid. Strong spot and edge attack were obtained in most samples when exposed to peroxide paper. The amount of blemish formation decreased as the solubility of the corresponding silver halides decreased.

A method was developed for estimating the free silver in processed microfilm. "Free silver" as here defined includes silver ion, silver oxide and silver halides but the method does not show how much of the silver is in the form of halide salt. This method indicated as much free silver present in old processed microfilm which had been treated in a solution containing 3 ml of acetic acid per liter as in microfilm treated in a solution containing 3 ml of acetic acid and 0.025 g of sodium chloride per liter. However, when they were exposed to peroxide paper, the samples given the acetic acid and sodium chloride treatment resisted background yellowing though they had natural-like spots and edge attack while those given the acetic acid treatment showed only an even yellowing of the image silver. Apparently traces of silver in the image are oxidized by oxygen dissolved in the water, in the presence of acid, and if the chloride ion is present, silver chloride is formed.

The procedure for determining the free silver is outlined as follows. The free silver was extracted from 12 cm² of processed microfilm with 10 ml of a 0.2 percent solution of ammonium hydroxide containing 1 g of sodium chloride per liter. Two 6 cm² pieces of microfilm were cut, folded emulsion in and placed back to back in a glass vial containing the test solution and allowed to stand 15 min. The samples were removed by a glass rod having a hook at one end and 0.10 ml of a potassium bromide solution containing 100 g/l was added and mixed. Then the solution was acidified by adding 1 ml of concentrated nitric acid diluted 1 to 5 and mixed, allowed to stand 1 hr and the turbidity measured with a nephelometer. A calibration curve was prepared from known amounts of silver added to the standard solution as silver nitrate. One-fourth microgram of silver as silver nitrate gave a measurable turbidity. The values obtained are relative but were helpful in showing changes in the free silver content of microfilm after various treatments. The free silver content of old processed microfilms containing blemishes ranged from 0.2 to 1 μg for 6 cm² of film.
3. Formaldehyde, Formic Acid, and Ammonia
From Paper Cartons

Microfilms in which blemishes had formed during natural storage were studied to determine if more blemishes would develop in the laboratory peroxide test. Several rolls of microfilm having blemishes on the outer convolutions but none on the remainder of the roll were tested. Samples taken from blemish-free areas were tested and no blemishes developed, but samples from areas where blemishes had formed during storage did develop blemishes like those already present. The blemishes present before the test were rinsed with a stylus to differentiate them from blemishes produced by the peroxide. Evidently chemicals in addition to peroxide had evolved from the paper carton during storage and created a chemical environment conducive to blemish formation in the presence of peroxide.

Formaldehyde, formic acid, and ammonia have been detected in the atmosphere within many of the paper storage cartons when stored at 100 percent relative humidity at 26°C. A glass petri dish, 60 mm in diameter and 13 mm high, containing 10 ml of distilled water was placed in the paper carton to absorb evolved chemicals. The carton was stored in a desiccator jar containing 25 ml of distilled water for 1 to 3 days, one day being generally sufficient. This extraction procedure was used in testing for the presence of formaldehyde, formic acid, and ammonia. Formaldehyde-urea and formaldehyde-melamine resins have been widely used as wet-strengthening agents in paper manufacture for over 20 years. The chemicals detected were probably formed by hydrolysis of these resins at high relative humidities.

3.1. Detection of Formaldehyde, Formic Acid, and Ammonia

a. Formaldehyde

Two ml of the water solution which had been stored in the paper carton were added to 5 ml of a 1 percent solution of chromotropic acid (4,5-dihydroxy, 2,7-naphthalene disulfonic acid) in concentrated sulfuric acid. Chromotropic acid readily detects formaldehyde in concentrations of 1 ppm (15). Chromotropic acid gave a pink to dark purple color, a positive test for formaldehyde. The less sensitive resorcinol test gave a positive indication of formaldehyde from some cartons when the same water solution was stored successively in 3 cartons for a period of 3 days each. This test was made by adding 0.1 ml of 1 percent solution of resorcinol to 5 ml of the water solution and pouring the mixture carefully down the side of a test tube containing 5 ml of concentrated sulfuric acid to form two layers. A pink color formed at the junction of the two layers. The formaldehyde was also identified by microwave absorption analysis.

b. Formic Acid

The water stored in the paper cartons had a pH range of 3.5 to 4.5 after storage for 3 days at 100 percent relative humidity at 26°C. Titration with 0.001 N sodium hydroxide solution indicated that the water stored in the cartons was about 0.001 N with respect to formic acid for some storage cartons. The water stored in the cartons decolorized a light pink solution of potassium permanganate, indicating that formic acid was present. When 5 ml of the water solution was added to 5 ml of saturated mercuric chloride and warmed in hot tap water for 10 min, a precipitate formed, indicating reduction of the mercuric chloride. This test indicated the presence of formic acid. A precipitate of reduced silver formed when 2 ml of silver nitrate solution containing 10 g per liter were added to 10 ml of the water solution from the paper cartons and warmed 15 min in hot tap water. The reduced silver indicated the presence of formic acid which is capable of reducing the silver ion. This procedure detected 5 ppm of formic acid in a known solution. The formic acid is evidently formed by oxidation of the formaldehyde by the peroxide generated in the storage cartons and possibly by aerial oxidation of formaldehyde, catalyzed by the silver in the image. A water solution of formaldehyde was found to increase in acidity when stored over silver dust for 4 hr showing that formic acid was formed by the oxidation of formaldehyde.

c. Ammonia

About 10 ml of the water stored in the paper cartons were placed in a 50-ml beaker and made alkaline with sodium hydroxide. The beaker was covered with a watch glass having a strip of moistened red litmus paper attached to the underside and the beaker was warmed in hot tap water. After 30 to 60 min the litmus paper turned blue suggesting the presence of ammonia. A control test run on the distilled water was negative.

3.2. Tests of Old and New Storage Cartons

Some new storage cartons readily evolved formaldehyde when stored for 3 hr in the drawer of a metal laboratory desk. A strong indication of formaldehyde was obtained after a new carton was stored at 100 percent relative humidity at 3°C for 1 day, indicating that a low temperature does not stop the evolution of formaldehyde if the relative humidity is high. Most of the old storage cartons tested evidenced formic acid evolution.

Storage cartons from one collection of blemish-free microfilms evolved no formaldehyde or only a very small amount. Cartons from a second collection where blemishes have not been found evolved an appreciable amount of formaldehyde, but this collection is located in an arid climate which would minimize the evolution of formaldehyde and the formation of formic acid. All the tests of paper storage cartons from collections where blemishes have been found, even collections that were ten years old, gave a strong indication of formaldehyde.

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1 Obtained from Matheson Coleman and Bell, Cat. No. P5425, East Rutherford, New Jersey.
3.3. Reactions of Formaldehyde, Formic Acid, and Ammonia on Silver and Gelatin

During natural storage, blemishes generally form on the outside leader and edges of the microfilm roll, which are the areas most accessible to gaseous chemicals evolved by the storage carton. Formaldehyde will harden gelatin by reacting with the amino groups attached to the amino acids in the gelatin. Blemish formation was inhibited in a blemish forming microfilm after an exposure of 24 hr to the gaseous phase of a 2 percent solution of formaldehyde. This severe test may not be relevant to natural storage conditions but suggests that formaldehyde, alone, may inhibit blemish formation. Formaldehyde and formic acid neutralize the alkalinity of the amino groups making the gelatin more acidic.

Formic acid and chloride ion reacted with the image silver at high relative humidities to form silver chloride in the same manner as described in section 2.4 for the sodium chloride-acetic acid solution, making the high-density uniform image more resistant and the edges of the blemishes sharp, when the film was exposed to peroxide evolved by the storage cartons.

In section 2.2 it was noted that tap wash water having as much as 1 ppm of chlorine formed a sufficient amount of silver chloride in the microfilm to make it resistant to blemish formation. The formic acid from cartons stored at high relative humidities appears capable of reducing the silver chloride concentration to a level favorable for blemish formation. Formic acid readily reduced the silver ion in an aqueous solution of silver nitrate, forming a gray precipitate of silver. Solid silver chloride in a solution of formic acid turned slightly gray after standing in the dark for 7 days. A control without formic acid did not exhibit this effect. The gray color was attributed to reduced silver.

Experiments were performed to see whether or not residual silver chloride in the image silver of processed microfilm is susceptible to reduction by formic acid and peroxide. To incorporate silver chloride into clear (image-free) processed microfilm, the film was washed 20 min in distilled water, treated 10 min in a solution containing 5 g of silver nitrate per liter, treated 10 min in a second solution containing 5 g sodium chloride per liter, washed 20 min in distilled water and dried. When exposed to 24 cm² of paper treated with formic acid, samples of this microfilm developed a yellow color in the gelatin layer, indicating the reduction of the silver chloride. The test was made at 86 percent relative humidity and 26 °C and in the same type of desiccator jar used for peroxide tests. The paper was prepared by treating Whatman No. 3MM filter paper in a 10 percent formic acid solution for 30 min drying it 3 hr at 26 °C and storing it at approximately −10 °C. Microfilm placed in a new storage carton for 12 days at 86 percent relative humidity and 26 °C showed evidence of reduction of silver chloride. Samples exposed to 6 cm² of the peroxide paper also evidenced such reduction. Thus residual silver chloride reduction in stored microfilm should decrease the blemish resistance.

It was found that the silver-chloride-ammonia complex was more easily reduced by formic acid and peroxide than silver chloride precipitated in the gelatin layer as described above. Tests were made on clear (image-free) processed microfilm treated 5 min in a solution containing 0.5 g of silver chloride dissolved in one liter of water containing 8 ml of concentrated ammonium hydroxide. The results indicated that silver chloride formed in microfilm stored paper cartons which evolve relatively large amounts of formaldehyde and formic acid.

4. The Chemistry of the Formation of Blemishes in Processed Microfilm

4.1. Theory

Paper cartons in which microfilm is stored undergo degradation at high relative humidity and high temperature to form peroxide. The silver in the image serves as a catalyst to decompose the peroxide, forming oxygen which oxidizes the image silver to the silver ion. The chloride ion catalyzes the oxidation of silver by peroxide. The peroxide [11] and formic acid reduce the silver ion to yellow colloidal silver.

Processed microfilms exposed for 2 min to the gas phase of 20-percent hydrochloric acid formed spot blemishes when exposed to peroxide paper. The hydrochloric acid reacted with the image silver to form silver chloride [10]. Many white spots formed, a few with a sharp ring structure, but they were colorless because the gelatin was too acid for the formation of the yellow to red colloidal silver found in naturally occurring spots. Under these conditions only oxidation of the silver occurred but concomitant oxidation and reduction must occur to form yellow to red colloidal silver. When the samples with the white spots were given a printout exposure and treated in a developer solution the spots were still colorless. This indicated that the silver ion had not remained in the colorless area but had migrated to the edge of the spots.

The silver ion may be reduced in an aqueous solution to form colloidal silver [9]. The reduction of silver ion in the gelatin layer of processed film by peroxide forms yellow colloidal silver. A proprietary product containing 19 to 23 percent silver in the colloidal state in gelatin is known in the trade as “Solargentum” [13]. Processed microfilm having no silver image was washed in distilled water, treated in a dilute solution of silver nitrate or silver acetate and dried. When samples were exposed to peroxide at 86 percent relative humidity and 26 °C, the gelatin layer turned yellow. Reducing agents evolved by paper storage
cartons formed the same yellow color in the gelatin of other samples stored in the cartons for 10 days at 86 percent relative humidity and 26 °C.

There is another type of colloidal silver formed when ionic silver is reduced in the presence of an insoluble silver salt such as silver chloride. This type of colloidal silver appears to be silver dispersed in a silver halide and gives colors ranging from pink, red, yellow, brown to gray when formed in an aqueous solution but yellow or reddish when formed in the gelatin layer of microfilm. Carey Lea claimed that this colloid was a mixture of silver chloride and a subhalide of silver. Mees [5] has pointed out that it may be regarded as a mixture of silver in silver halide.

Tests showed that the blemishes contained ionic silver in addition to colloidal silver. The black specks in the centers of many spots appear to be reduced silver since they are readily dissolved by strong nitric acid. In section 2 it was noted that large spots, with light yellowing and even attack on the image silver, were obtained when the microfilm was fixed in a neutral fixer or a nonhardening fixer and washed in distilled water. In these processing procedures no chloride or bromide ions should be present to form silver halide and the yellow color would be attributed to the silver-gelatinate type of colloidal silver. However, when the microfilm was fixed in the nonhardening fixer or the acid-hardening fixer and washed in chlorine-free tap water the spots were sharper, more yellow, attended by no edge attack or even attack on the image silver, and required more peroxide for spot formation. The chloride ion introduced by the tap water formed silver chloride which led to the formation of the silver-silver chloride type of colloidal silver in spots and prevention of even attack on the image silver. The silver-silver chloride colloidal silver probably predominates in blemish formation.

4.2. Formation of Silver-Silver Halide Colloidal Silver in Solution and on Silver Strips

It was found possible to form the silver-silver chloride type of colloidal silver in water solution. Thirty percent hydrogen peroxide was added to a solution containing varying concentrations of sodium chloride in the presence of solid silver oxide. A small amount of silver oxide dissolved making the solution alkaline. Reduced gray silver accumulated in the bottom of the beaker, leaving the solution clear when no chloride ion was present. When chloride ion was added, a colloidal suspension formed, ranging in color from pink to yellow and sometimes a reddish precipitate formed along with the gray reduced silver. Treatment of the reaction mixture with nitric acid dissolved the reduced silver and left a yellow or reddish colloidal suspension and a reddish precipitate. The pink, yellow and reddish color in the colloidal suspension resembled the color of natural spots. The optimum condition for producing the colloidal silver was a 60-ml solution containing 0.05 g of sodium chloride and about 0.5 to 1 g of solid silver oxide. The hydrogen peroxide was added dropwise with stirring.

Equal mol concentrations of potassium bromide (0.102 g) and potassium iodide (0.142 g) equivalent to 0.05 g of sodium chloride were used in the above procedure to test for the formation of colloidal silver. The potassium bromide solution produced a light yellowish colloidal suspension with a trace of reddish color and some reddish precipitate. The potassium iodide solution gave no visual evidence of the formation of colloidal silver in the solution although a faint trace may have been present. A small amount of sodium chloride in the potassium iodide solution did produce a slightly yellowish colloidal solution indicating that colloidal silver will form in the presence of potassium iodide and the chloride ion.

The silver-silver chloride type of colloidal silver was also formed on silver strips. The silver strips were cleaned with nitric acid, washed in distilled water and immediately immersed in a 50-ml solution containing 0.05 g of sodium chloride, to which 30 percent hydrogen peroxide was added dropwise. A heavy reddish coating of colloidal silver formed on the silver strip showing that the silver on the strip was both oxidized and reduced by the peroxide. When no sodium chloride was added, the silver strip remained bright and served only as a catalyst to decompose the peroxide into water and oxygen. There was no precipitate of reduced silver. These results showed that the chloride ion played an important part in the oxidation of the silver by the peroxide.

5. Test for the Formation of Blemishes in Processed Microfilm With Peroxide Paper

5.1. Preparation of the Peroxide Paper

It was found that paper treated in a solution of hydrogen peroxide was a good source of gaseous peroxide for the formation of blemishes in processed microfilm. All of the types of blemishes described in NBS Handbook 96 [2] have been produced by exposing films to such paper. The paper used was Whatman No. 3MM filter paper which is used in chromatography. The paper was treated 30 min in a 5-percent solution of hydrogen peroxide at room temperature with occasional stirring, hung in a ventilating hood for 2 hr, dried for 24 hr in a desiccator over anhydrous calcium sulfate at 26 °C and stored in a sealed glass bottle in a refrigerator at -10 °C.

5.2. Apparatus

The tests were made in a desiccator jar containing a saturated salt solution to control the relative humidity. The jar had an inside diameter of 160 mm at the ground flange and a volume of approximately 2300 ml. The air and salt solution were not stirred but the relative humidity value reported is that recorded in the literature for equilibrium conditions. The jar was covered with a flat glass plate about 0.6 cm thick and sealed with petroleum jelly. A porcelain plate perforated with small holes was located just above the saturated salt
solution. Saturated solutions of sodium chloride and potassium chloride were used to obtain relative humidities of about 76 and 86 percent, respectively, at 26°C.

The test samples, about 6 cm in length, were fastened at each end with black photographic masking tape to a thin glass plate about 8 × 10 cm in size. Four or five samples could be mounted on a plate. The glass plate was then fastened at two opposite edges to the underside of the cover with a strip of the tape, so that the emulsion faced downward. A glass dish, 60 mm in diameter and 13 mm high, was centered in the middle of the porcelain plate to hold the peroxide paper. The mounted microfilm samples were about 8.5 cm from the peroxide paper. In testing microfilms for blemish formation the peroxide paper was added to the glass dish and allowed to react 24 hr at 26°C ± 1°C. Two cm² of peroxide paper was sufficient for some microfilms. If no blemishes formed, 2 cm² more of peroxide paper was added and the test was continued. Most of the blemish forming microfilms required 2 to 4 cm² of the peroxide paper.

6. Summary

Microfilms fixed in a neutral or nonhardening acid bath and washed in distilled water or chlorine-free tap water, formed blemishes when exposed to peroxide paper. Microfilms fixed in an acid hardening bath and washed in chlorine-free tap water, also formed blemishes when exposed to peroxide paper but no blemishes formed after washing in distilled water. The distilled water wash did not reduce the concentration of the residual thiosulfate in the microfilm sufficiently to permit blemish formation. The silver sulfide formed in microfilm during fixation affords some protection against oxidation of the image silver by peroxide but its concentration is not large enough to prevent oxidation of the silver after washing in distilled water. Residual thiosulfate reacts with the image silver in processed microfilm to form additional silver sulfide, giving a greater resistance to peroxide attack. Tap wash water containing a chlorine concentration of 1 ppm formed a sufficient amount of silver chloride in the image silver to make microfilm resistant to blemish formation when exposed to peroxide paper. One microfilm required a 10-min wash after fixing in a nonhardening acid bath to inhibit blemish formation. The resistance to blemish formation by the silver chloride in the image silver can be removed by treating the processed microfilm in a 0.2 percent solution of ammonium hydroxide. The image silver showed an even yellowing after this treatment and exposure to peroxide paper. Some resistance to blemish formation was found when microfilms were washed 30 min in tap water containing a chlorine concentration of 0.3 ppm. A trace of silver chloride in the image silver promoted blemish formation. Thus the type of fixing bath, concentration of the chlorine in the wash water, and the washing time after fixation can be factors in the formation of blemishes in processed microfilm stored in paper cartons. Formaldehyde-urea and formaldehyde-melamine type resins are incorporated in microfilm paper storage cartons as wet-strengthening agents. These resins hydrolyze at high relative humidity to give formaldehyde and ammonia. Formic acid may form by the oxidation of the formaldehyde by the peroxide evolved by the storage carton and aerial oxygen in the presence of silver in the image. Microfilm containing sufficient residual silver chloride to prevent blemish formation loses this resistance when exposed to formic acid evolved in storage cartons. It was found that formic acid was capable of reducing silver chloride and the silver chloride ammonia complex in the gelatin layer of microfilm. The silver chloride ammonia complex was more easily reduced than silver chloride, suggesting that ammonia evolved by storage cartons may sensitize the reduction of the silver chloride in the image silver.

The present theory is that peroxide oxidizes the silver in the image and also reduces the silver ion. The reduced silver may form colloidal silver of the silver-gelatinate type or combine with a silver halide, forming the silver-silver halide type of colloidal silver.

7. Appendix. Formulas for Developing and Fixing the Silver Image in Microfilm

7.1. Developer

Monomethyl p-aminophenol sulfate.................. 2.0 g.
Sodium sulfite, anhydrous.......................... 90.0 g.
Hydroquinone........................................ 8.0 g.
Sodium carbonate, monohydrate...................... 52.5 g.
Potassium bromide................................... 5.0 g.
Water to make...................................... 1 liter.
The development time was 3 min.

7.2. Stop Baths

No. 1 for Neutral Fixing Bath:
Treated sample 1 min in each of two 2-liter lots of distilled water.
No. 2 for Nonhardening Acid Fixing Bath:
Sodium bisulfite ........................................ 60.0 g.
Water to make ........................................... 2 liters.
Treated sample 2 min.

No. 3 for Acid Hardening Fixing Bath:
Glacial acetic acid ...................................... 30 ml.
Water to make ........................................... 2 liters.
Treated sample 30 secs.

7.3. Fixing Baths

Neutral Fixing Bath:
Sodium thiosulfate (Na$_2$S$_2$O$_3$) ........... 245.0 g.
Water to make ........................................... 1 liter.
Fixed sample 4 min.

Nonhardening Acid Fixing Bath:
Sodium thiosulfate (Na$_2$S$_2$O$_3$) ........... 240.0 g.
Sodium bisulfite ......................................... 25.0 g.
Water to make ........................................... 1 liter.
Fixed sample 5 min.

Acid Hardening Fixing Bath:
Sodium thiosulfate (Na$_2$S$_2$O$_3$) ........... 300.0 g.
Sodium sulfate ........................................... 10.0 g.
Glacial acetic acid ..................................... 10 ml.
Boric acid .................................................. 5.0 g.
Potassium aluminum sulfate ....................... 10.0 g.
Water to make ........................................... 1 liter.
Fixed sample 4 min.

7.4. Chemical Specifications

All of the solutions were made up with distilled water. The following chemicals were of reagent quality meeting ACS specifications: sodium sulfite, acetic acid, boric acid, potassium bromide, and potassium iodide. The monomethyl p-aminophenol sulfate, hydroquinone, sodium carbonate, sodium thiosulfate, and potassium aluminum sulfate conformed to the USA Standards Institute Specifications for Photographic Grade Chemicals. The sodium bisulfite was analytical reagent grade.

8. References

[15] Eugene Sawicki, T. R. Hauser, and Sylvester McPherson, Spectrophotometric determination of formaldehyde and formaldehyde-releasing compounds with chromotropic acid, 6-amino-1-naphthol-3-sulfonic acid (J acid), and 6-amino-1-naphthol-3-sulphonic acid (phenyl J acid), Anal. Chem. 34, 1460 (1962).

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