PHOTOMICROGRAPHY: COMMON GROUND FOR SCIENCE AND ART

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Figure 1. Title: none. Description: A single exposure of molecular weight calf thymus DNA in a slightly hydrated condition. Awards: Honorable mention in the 1987 Nikon Small World Competition. Exposure 1: A 900 milligram/milliliter mixture of DNA and saline buffer was sandwiched between a microscope slide and coverglass and sealed with a polymethylmethacrylate mounting medium. The sample was observed with a first order retardation plate (530 nm) inserted between the sample and the analyzer. Exposure was colour corrected with a 20 CC magenta filter. The 10x objective was utilized with polarized light for the 0.17 second exposure.

Figure 2. Title: DNA Liquid Crystals. Description: A single exposure of Smectic-like liquid crystalline DNA. Awards: Photographs of the type have been published in numerous periodicals including Nature, the British Journal of Photography, Second Opinion, The Bethesda Research Laboratories catalog, Microscopy and Analysis, and Functional Photography. Exposure 1: A 350 milligram/milliliter solution of nucleosome core length (approximately 500 Angstroms) DNA in a buffered ammonium acetate (0.3M Ammonium acetate, 0.01M Sodium Cacodylate, and 0.01M EDTA adjusted to pH 6.5 by addition of solid Cacodylic Acid) spontaneously forms smectic-like liquid crystals. The sample was sandwiched between a coverslip and a microscope slide and sealed with a polymethylmethacrylate mounting medium before examination. The exposure was taken with a 10x objective for 0.12 seconds with a 10 CC magenta colour correction.

Figure 3. Title: Nebraska. Description: A multiple (4) exposure of ascorbic acid (the wheat in the foreground), stretched polyethylene (the morning sky), polybenzyl-1-glutamate spherulites (the stars), and the field diaphragm defocused (the sun).
Figure 4. *Title*: Neptune Beach, Florida. *Description*: A multiple (5) exposure of the antibiotic, chloramphenicol (the grassy foreground), stretched polyethylene with a blue filter (the morning sky and ocean), the field diaphragm (the morning sun and its reflection), and polybenzyl-1-glutamate (the stars).

Figure 5. *Title*: Ascorbic Acid, Idaho. *Description*: A multiple (5) exposure of ascorbic acid (the wheat in the foreground), Xanthan gum base (the snow-covered mountains), stretched polyethylene (the morning sky), polybenzyl-1-glutamate spherulites (the stars), and the field diaphragm image defocused (the sun).

Figure 6. *Title*: Nuclear Sunrise. *Description*: A multiple exposure (4) of ascorbic acid (the desert foreground), stretched polyethylene (the morning sky), polybenzyl-1-glutamate (the stars), and the field diaphragm defocused (the rising sun).
PHOTOMICROGRAPHY

various textures formed by DNA as it entered the liquid crystalline state (illustrated in Figure 11-4). To this end, we have used both light and electron microscopy; however, this article will be dedicated to our colour photomicrography with a polarising light microscope.

In December 1986, we first observed a highly birefringent, beautifully coloured, fan-textured smectic (2-dimensionally ordered) liquid crystalline state of DNA (Figure 24) which we attempted to photograph using standard E6 transparency films processed by commercial laboratories. The results that we obtained were disappointing because the contrast and colour saturation of our transparencies were of an inferior quality to the specimens that we observed first-hand in the microscope. To correct this, we initiated a series of tests using various daylight and tungsten-balanced films which we began

The escalating technology in optics and optical coatings, coupled to the development of high quality microscopes and deeply colour-saturated photographic films has, in part, led to an explosion in the utilisation of photomicrography for numerous fields. Many years ago, the microscope was an exclusive tool of biologists who spent countless hours observing and drawing various specimens of biological interest. Today, however, the microscope has found a home in disciplines as diverse as Chemistry, Physics, Geology, Psychology and Materials Science, to name a few.

Our interest in photomicrography originated from a need to view, at relatively high magnification, the various textures formed by DNA as it entered the liquid crystalline state (illustrated in Figure 11-4). To this end, we have used both light and electron microscopy; however, this article will be dedicated to our colour photomicrography with a polarising light microscope.

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**Figure 7.** Title: Sulphur Canyon. Description: A multiple (4) exposure of recrystallized sulphur (the canyon), polybenzyl-1-glutamate (the stars), and the field diaphragm defocused (the moon) with a blue filter (the sky).

**Figure 8.** Title: Sunday. Description: A multiple (3) exposure of the antibiotic chloramphenicol (the grassy foreground), hardened epoxy resin (the purplish sky), and the field diaphragm defocused with a red filter (the rising sun).

**Figure 9.** Title: Cety Alpha 5. Description: A multiple (4) exposure of smectic liquid crystalline DNA (the foreground), polybenzyl-1-glutamate spherulites (stars), and the field diaphragm defocused with a blue filter.

**Figure 10.** Title: Tornadoland. Description: A multiple (6) exposure of ascorbic acid (Vitamin C-the desert foreground), cholesteric liquid crystalline DNA (the tornado), and Cibachrome bleach crystals (the clouds and dust) with a blue sky.
**Figure 11.** *Title:* Westworld. *Description:* A multiple (2) exposure of epoxy resin (the clouds and the mountains) and the field diaphragm defocused (the red sun).

**Figure 12.** *Title:* The Stand. *Description:* A multiple (4) exposure of ascorbic acid (the wheat in the foreground), polybenzyl-1-glutamate (the stars), the field diaphragm defocused with a mask (the new moon), with a blue sky.
processing in-house with commercially available Kodak E6 kits. After many months of tests and comparisons, we finally settled on Fujichrome 64 T and have been using this film almost exclusively, with the exception of Polachrome HC 35 mm instant colour transparency film, which we employ for the annual Polaroid International Instant Photomicrograph Competition.

Our entry into the photomicrograph competition circuit started with the 1987 Nikon Small World contest in which we received 6th and 9th prizes and two honorable mentions for photomicrographs of liquid crystalline DNA samples. The Nikon contest heralded a new era for our photomicrography because one of the winning micrographs was a multiple exposure which we intended to resemble an alien landscape. Photomicrographs of this type have now won many competitions and we have extended our emphasis to make these photographs more life-like. We now term these types of photomicrographs, Microscapes. Several examples of our most recently fabricated Microscapes are illustrated in Figures 3-12.

**FABRICATION OF MICROSCAPES**

Nikon’s UFX-II digitally controlled exposure monitor allows for a double or multiple exposure mode which is employed in the construction of multiple exposure Microscapes. Our microscope is a Nikon Optiphot-pol in which light intensity and distribution are regulated by the field diaphragm (a leaf-type shutter) and emitted through a lens in the base of the microscope. By carefully masking a portion of the field diaphragm lens, selective areas of film can be exposed to light captured through the microscope.

The sun or moon is added by closing the field diaphragm almost completely and defocusing the image until the individual leaves merge to form a complete circle. Next, an orange or red filter is inserted into the light pathway and the substage condenser is decentered, by realignment of the adjustable centering pins, to move the diaphragm image to the appropriate location and an exposure is recorded. Long exposures yield a bright, white

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**Figure 13.** *Title:* Vitamin C-horse. *Description:* A single exposure of ascorbic acid recrystallized from the melt. Exposure 1 (see Figure 16).

**Figure 14.** *Title:* The Many Faces of Vitamin C. *Description:* A single exposure of ascorbic acid recrystallized from the melt. Awards: Honorable mention in the 1988 Nikon Small World Competition. Exposure 1 (See Figure 16).

**Figure 15.** *Title:* Man in the Moon-Greencheese. *Description:* A single exposure of ascorbic acid recrystallized from the melt. Exposure 1 (See Figure 16).
Figure 16. **Title:** The Ghost. **Description:** A single exposure of ascorbic acid (Vitamin C) recrystallized from the melt. Exposure 1: A 20 milligram sample of ascorbic acid was sandwiched between a microscope slide and a cover slip and heated until melted (with some decomposition) in a bunsen burner. Upon crystallization (1 to 9 weeks), the image was photographed with a 10x objective with a 20 CC magenta colour correction filter for a period of 0.14 seconds.

Figure 17. **Title:** Swirlaway. **Description:** A single exposure of Niacin (a member of the B complex) recrystallized from the melt. Exposure 1: A 20 milligram sample of niacin was sandwiched between a microscope slide and a coverglass and was heated in a bunsen burner until melted. Upon recrystallization, the sample was photographed under polarized light with the 10x objective and a 20 CC magenta colour correction filter for a period of 0.10 seconds.

Figure 18. **Title:** Welcome to the Jungle. **Description:** A single exposure of ascorbic acid (Vitamin C) recrystallized from the melt. Exposure 1 (see Figure 17).

Figure 19. **Title:** Sunflowers. **Description:** A single exposure of ascorbic acid (Vitamin C) recrystallized from the melt. Exposure 1: A 40 milligram sample of ascorbic acid was sandwiched between a coverslip and a microscope slid and heated until melted with a bunsen burner. After recrystallization the sample was photographed with the 4x objective after a 530 nm retardation plate had been inserted between the sample and the analyser. Colour correction was 20 CC magenta and the exposure time was 0.07 seconds.
centered sun or moon, while short exposures give more colour-saturated images with a yellow-to-red transition from centre to edges. The new moon illustrated in Figure 12 is created by placing the tip of a ballpoint pen in the light path after closing down the field diaphragm and defocusing.

The mountains are masked exposures of unrefined Xanthan-gum dissolved in aqueous solution and allowed to concentrate while sandwiched between a microscope slide and a cover slip. This polysaccharide tends to undergo a transition to cholesteric liquid crystalline states, in concentrated solutions, which resemble mountainous formations.

The morning sky is created by cutting a rectangular portion of a polyethylene film (plastic storage bag, thickness~100 micrometers) and stretching it in a uniform manner before pressing onto a microscope slide. Upon stretching, the polyethylene molecules align to a form birefringent diffraction gradient which casts a yellow-to-red-to-blue visible light spectrum to simulate a morning sky. After masking previous exposures, the film is exposed to a masked section of the diffraction spectrum in order to obtain the effect desired.

Stars are created by imaging small cholesteric liquid crystalline spherulites of polybenzyl-l-glutamate dissolved in dimethyl formamide. Previously exposed areas of the film are masked for these exposures in order to eliminate burning of the spherulite image.

The clouds present in Figure 9 are generated by defocusing colourless birefringent crystals of Cibachrome bleach.

Various foregrounds are obtained by photographing crystalline formations of a wide spectrum of drugs, vitamins, and other assorted biochemicals. For instance, the foregrounds in Figures 3 and 4 are exposures of ascorbic acid (Vitamin C) and chloramphenicol (an antibiotic) respectively, recrystallized after sandwiching the pure biochemical between a microscope coverslip and slide and heating until melted. In some instances, the chemicals recrystallize rapidly within a few minutes. However, many take weeks or even months to completely recrystallize. The composition of the foreground in each photomicrograph is discussed in the respective figure legends.

Generally, after the selected foreground is exposed, the exposed area is carefully masked by placing a black card over a large enough area of the field lens to completely stop any additional light from reaching the exposed portion of the film. Next, a second exposure portion of the film, Next, a second exposure is made usually either with Xanthan
THE MANY FACES OF UBIQUITOUS VITAMIN C

Our various photomicrograph portfolios are each constructed around a central theme as described for the Microscapes collection in detail above. Additional portfolios include a vitamin collection, an antibiotic collection, as well as selected collections of other chemicals and biochemicals.

However, probably the most unusual and visually exciting collection is our vitamin collection (Figures 13-20 are selections from this group). A sub-portfolio from the vitamin collection has been named The Many Faces of Vitamin C due to the fact that many of the wide spectrum of crystalline morphologies displayed by recrystallized ascorbic acid resemble faces in one respect or another. Figure 16 is an example of an unusual morphology which possesses a haunting appearance. The pattern illustrated in Figure 13 was titled Vitamin C-Horse by Steven Rosenbaum of Modern Photography due to its striking resemblance to a sea horse. Likewise, surrealistic faces can be found in Figures 14 and 15. Figure 15, a photomicrograph of an isolated crystallite of ascorbic acid has been titled The Man in the Moon-Greencheese due to its uncanny resemblance to the fictional Man in the Moon on one side of the inner circle and to green Swiss cheese on the other side. Currently, we have about 40 members in the Faces of Vitamin C portfolio and constantly search out additional morphological formations from recrystallized ascorbic acid samples.

The needle-like morphology demonstrated in many recrystallized samples of ascorbic acid can be further enhanced in appearance by the addition of a 530 nanometer retardation plate inserted. It strongly resembles a boiling solution with chips at the bottom. These are only a small sampling from The Vitamin Collection intended to illustrate the inherent beauty to be found in polarized light photomicrographs of recrystallized vitamins.

CONCLUSION

By coupling a well-developed photographic methodology to a creative imagination, there is virtually no limit to the types of photomicrographic artwork possible with the light microscope. Here we have demonstrated a few examples of the wide spectrum of possibilities available to the microscopist who is willing to take the time and effort necessary to compose micro-art.

EXPERIMENTAL METHODS

All photomicrographs were composed using a Nikon Optiphot-pol polarizing light microscope in either the brightfield or polarizing mode. A UFX-II digitally controlled exposure monitor measures light intensity through a 30 per cent central portion of the viewfield by employing a photomultiplier and calculates exposure time based on this reading. Illumination is provided by a tungsten-halide 12 volt bulb operating at 8.5-9 volts. Images were recorded on Fujichrome 64 T, a 3200 Kelvin colour-balanced (tungsten) transparency film operating at approximately ISO 64. Exposures were usually made at 1 to 3 f-stops below the recommended exposure of the UFX-II system and Kodak E-6 processing (done in-house) was extended 25-50 per cent in the first developer. Slight modifications were made to the...
chemical composition of the first colour developers to enhance contrast and colour saturation.

ACKNOWLEDGEMENTS
The authors wish to thank Kaye Merchant for her enthusiasm and inspiration during the early days of this study. In addition, we would like to acknowledge the assistance of Tom Fellers (FSU Department of Physics) for advice concerning photographic methods and David Van Winkle (FSU Department of Physics) for advice on liquid crystal behavior. This work was supported, in part, by the NIH.

REFERENCES


5. The authors collectively have won prizes in the following competitions: **Nikon Small World Contest**, 1987, held by Nikon Inc. Instrument Group, 623 Stewart Avenue, Garden City, New York 11530. Prizes: 6th, 9th, 2 honorable mentions.


**Polaroid International Instant Photomicrography Competition**, 1987, held by the Polaroid Corporation, 575 Technology Square, 9P Cambridge, Massachusetts 02139. Prize: 3rd place.


**American Society of Clinical Pathology 1988 Medical Photograpy Competition Sponsored by Nikon, Inc.**, held by the American Society of Clinical Pathologists, 2100 West Harrison Street, Chicago, Illinois 60612. Prize: 1st in the Micro Division.


**Polaroid International Instant Photomicrography Competition**, 1988, held by the Polaroid Corporation, 575 Technology Square, 9P Cambridge, Massachusetts 02139. Prizes: 1st and Honorable Mention in colour transparency category, and overall best of competition grand prize.


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