CYTOLOGICAL AND CYTOCHEMICAL STUDIES
OF GREEN MONKEY KIDNEY CELLS
INFECTED IN VITRO WITH SIMIAN VIRUS 40

ROBERT LOVE, M.D., and MARIO V. FERNANDES, V.M.D.

From the Department of Pathology, Jefferson Medical College, and The Wistar Institute, Philadelphia

ABSTRACT

Cytological and cytochemical studies of green monkey kidney cells infected with SV40 virus indicated that the type of lesion produced was influenced by the multiplicity of infection and that the lesions appeared later and progressed more slowly when the inoculum was diluted. The earliest change consisted of enlargement of ribonucleoprotein-containing spherules in the nucleolus (nucleolini). This was followed by rarefaction, with or without condensation, of the chromatin and the appearance of one or more homogeneous masses of inclusion material containing DNA, RNA, and non-histone protein which eventually filled the nucleus. In some instances the chromatin appeared to be directly transformed into inclusion material. In the later stages of infection, the ribonucleoprotein of the nucleolini was no longer stainable and material resembling the nucleoprotein of the intranuclear inclusions was found in the nucleolar vacuoles and in the cytoplasm. The nucleic acids in the inclusions were stained by toluidine blue, toluidine blue-molybdate, the Feulgen stain, and by methyl green. The stainable material was extractable by nuclease digestion or by hot trichloroacetic acid. Green or yellowish green staining by acridine orange was apparently due to binding of dye by protein and not by nucleic acids since the staining reaction was not reduced by extraction of nucleic acids by hot trichloroacetic acid. Extraction with pepsin in combination with ribonuclease or deoxyribonuclease removed practically all the inclusions from the cells; consequently they could not be stained with acridine orange. The cytochemical studies suggest that the use of pepsin together with nuclease is not a meaningful technique.

INTRODUCTION

Several reports of the cytological, cytochemical, and immunocytochemical changes produced by infection of green monkey kidney cells in vitro with SV40 virus (34) have been published (2, 5, 11, 12, 16, 19, 27, 28, 31, 36). The present study combines more extensive cytochemical investigations with observations by phase-contrast microscopy and immunofluorescent techniques and reveals some new aspects of the cytopathology. In particular, the cytochemical changes in the nucleolus and the nature of the inclusion material in the nucleus have been elucidated and the effect of variable multiplicity of the viral inoculum on the evolution of the lesions has been determined.

MATERIALS AND METHODS

Virus

The virus was kindly provided by Dr. M. R. Hilleman and maintained by passage in green monkey kidney cells.
monkey (Cercopithecus aethiops) kidney cultures at the Wistar Institute. The preparation of virus pools and the method of infecting monolayers have been described (30). The pool used in the present experiments contains 10^4 TCID<sub>50</sub> per 0.1 ml. Replicate cultures were infected with undiluted virus (high multiplicity), or 10<sup>-4</sup> and 10<sup>-5</sup> (low multiplicity) dilutions of the pool. The undiluted inoculum corresponded to an approximate input multiplicity of 1 TCID<sub>50</sub> per cell.

Cells

Primary monolayer cultures of African green monkey kidney cells from Microbiological Associates, Bethesda, Maryland, were subcultured into 60 mm Petri dishes containing coverslips and grown at 37°C in a humid atmosphere of approximately 5 percent carbon dioxide in air. The medium was Eagle's basal medium (5) containing 2 mM glutamine and 10 percent filtered calf serum with added bicarbonate (5.6 percent NaHCO<sub>3</sub>, 25 ml per liter) and antibiotics (penicillin, 10<sup>4</sup> units; and streptomycin, 10<sup>4</sup> μg, or aureomycin, 5 × 10<sup>4</sup> μg, per liter). Coverslips were removed for examination at 4 and 24 hours and at daily intervals up to 18 days after infection.

Cytological and Cytochemical Studies

Infected and uninfected cultures were observed by phase-contrast microscopy, ordinary, and ultraviolet light microscopy. The following cytochemical techniques were employed: the Feulgen reaction, periodic acid-Schiff (PAS) (29), toluidine blue-molybdate (TBM, methods A, B, C, D, and E) (25), dilute toluidine blue after nitration for 18 hours (22), acridine orange as described by Mayor et al. (28) a modification of Pauly's method for tyrosine and histidine (32), the ninhydrin-Schiff and dinitrofluorobenzene methods for protein-bound amino groups (3), and the alkaline fast green method for histones (1). The methods of fixation were those described in the references, except that a mixture of two parts ethanol and one part glacial acetic acid was substituted for the regular Carnoy fixative. Methods of digestion, so that analysis of the effects of these procedures was impossible. Preparations were also fixed for 10 minutes at 37°C in formal sublimate and, after treatment with Lugol's iodine solution and sodium thiosulfate (25), stained with pyronin-methyl green (29). Various types of pretreatment after fixation and before staining were employed as described in the results. Deoxyribonuclease (DNase, crystallized, Worthington Biochemical Corporation, Freehold, New Jersey) and ribonuclease (RNase, 5 times crystallized, Worthington) were used as 0.01 percent solutions in 0.02 M trishydroxymethylaminomethane (Tris) buffer containing 0.045 M MgCl<sub>2</sub>-6H<sub>2</sub>O and 5 mM CaCl<sub>2</sub>, anhydrous, brought to pH 7.3 by addition of 1 N HCl (22). Preparations were digested for 2 hours at 37°C. Pepsin (3 times crystallized, Nutritional Biochemical Corporation, Cleveland) was used as a 0.2 percent solution in 0.02 N HCl for 10 minutes at 20 to 25°C (28). Control preparations were treated with buffer or HCl for the same time and at the same temperature as the enzyme-treated preparations. Cells were also treated for 30 minutes at 90°C with 5 percent trichloroacetic acid to remove nucleic acids (33).

Immunofluorescence

Preparations were rinsed briefly in acetone at 4°C, followed by fixation in acetone for 30 minutes at −20°C and stored at this temperature until stained. Rabbit anti-SV40 gamma globulin, labeled with fluoresceinisothiocyanate, was kindly provided by Drs. W. and G. Henle, of the Children's Hospital, Philadelphia. The method of staining has been described (30).

RESULTS

Morphological Changes

Despite the fact that specific changes were observed in almost every cell in the later stages of infection of cultures inoculated with a high multiplicity (undiluted) inoculum, the rate of development of the lesions varied considerably from cell to cell in the same culture (Figs. 6, 8, 9 and 16). The pathogenesis was, therefore, assessed in part from the time sequence of events, but also by analysis of increasing disorganization of nuclear structure in living cells under the phase microscope and in fixed preparations stained to demonstrate the nucleoproteins of the cell. It should be noted that, by virtue of the type of DNP or RNP that they contain, certain structures are optimally stained by each of the five TBM methods (25). Thus, the morphology of the cell appears to be different in each preparation. The structure of the nucleolus in preparations stained by method C is essentially the negative image of that stained by method B (cf. Figs. 3 and 6). In preparations of uninfected green monkey kidney cells stained by method B, the nucleolus appeared to be solid spherical bodies embedded in the lightly staining
Changes in the nuclei of green monkey kidney cells infected with SV40 virus. The drawings represent a composite picture of preparations stained to demonstrate nucleolini by TBM method B and chromatin and inclusions by TBM method C. After the initial nuclear change (8), the lesions evolve along three different pathways, A, B, and C.

In preparations stained to demonstrate the RNP of the nucleolini (method B), the center of some of the enlarged nucleolini was devoid of stainable material (Fig. 4). The enlarged nucleolini appeared light under the phase microscope (Fig. 5). The size of the nucleolus increased, presumably as a result of the enlargement of nucleolini. The nucleus of the affected cell also increased in size. The subsequent development of the lesions appeared to depend upon the rates at which two different processes occurred. One process was the appearance of a newly formed homogeneous material in the nucleus. The other process involved the chromatin and consisted of three types of change: (a) rarefaction, (b) condensation, and (c) replacement by the newly formed homogeneous material. The evolution of the lesions varied according to which of the three types of chromatin change was predominant and also depended upon...
whether the chromatin changes proceeded more quickly or contemporaneously with the appearance of the new homogeneous material in the nucleus. In one series of lesions (Fig. 1, series A), rarefaction of the chromatin was accompanied by replacement of the remaining chromatin by material that appeared to be homogeneous under the phase microscope and in suitably stained preparations (Fig. 1, 3A, and Fig. 6). As the lesions progressed, increasingly dense homogeneous material appeared in the rarefied areas (Fig. 1, 4A, and Fig. 7) and the transformation of chromatin continued until the entire nucleus (with the exception of the nucleolus) was homogeneous (Fig. 1, 5A, and Fig. 6). Subsequently, the homogeneous material became more dense and stained more intensely at first in localized areas (Fig. 1, 6A; and Figs. 5 and 8) and finally throughout the entire nucleus.
(Fig. 1, 7A, and Fig. 9). While these changes were occurring, the enlarged nucleolini appeared to fuse and became less numerous (Fig. 10). In the final stage that preceded karyorrhexis (Fig. 1, 7A, and Fig. 9), the deeply staining homogeneous nuclei shrank, the characteristic stainable material of the nucleolini was no longer detectable, and a vacuolated pars amorpha persisted. This series of changes was more frequent in the low multiplicity infections where the onset of detectable cytological changes was delayed to 96 hours, and the lesions evolved more slowly than with high multiplicity infection.

In a second series of lesions (Fig. 1, series C), the appearance of localized or widespread areas of rarefaction in the nucleus was accompanied by granular condensation of the remaining chromatin (Fig. 1, 3C, and Fig. 11). Homogeneous material, usually with an ill defined periphery, but occasionally quite circumscribed, appeared in the areas of rarefaction (Fig. 1, 4C, and Fig. 12) and subsequently increased in amount (Fig. 1, 5C, and Fig. 13). Further progression of these changes led to irregular margination and condensation of the chromatin and the production of shrunken nuclei filled with deeply staining homogeneous material (Fig. 1, 6C, and Fig. 14). The changes in the nucleolini were similar to those in the A series of lesions except that the enlargement was more irregular. One or two nucleolini were often dis-

\textbf{FIGURE 5} Phase-contrast photograph of an infected cell. The enlarged nucleolini appear as less dense structures in the nucleolus. Note three small ill defined dense inclusions (arrow) near the nucleolus and extensive cytoplasmic vacuolation. \( \times 1600 \).

\textbf{FIGURE 6} Two stages in the transformation of chromatin into homogeneous inclusion material. The chromatin of the cell on the left has lost its normal granularity and the rest of the nucleus is uniformly rareded. The chromatin of the two other cells is no longer recognizable and the nucleoplasm appears structureless. The enlarged nucleolini are very weakly stained in this preparation and appear as vacuoles in the nucleolus. TBM, method C. \( \times 1000 \).
proportionately enlarged, with apparent displacement of the others (Fig. 1, 3C and 4C, and Figs. 15 and 16). In the later stages (Fig. 1, 5C and 6C), the nucleoli fused and, finally, were no longer stainable by method B. Fragments of pars amorpha usually persisted (Fig. 14) until disintegration of the cell by karyorrhexis. This pattern of cytopathic change was characteristic of high multiplicity infection and was first detectable 24 hours after application of virus. The changes in the cells also progressed fairly rapidly and advanced lesions (Fig. 1, 5C and 6C) were observed before the earliest detectable abnormality in the low multiplicity infection.

A third series of changes (Fig. 1, series B), which appeared to be intermediate between A and C, was also observed. Following the usual stage of enlargement of nucleoli, localized or widespread areas of rarefaction of the chromatin were seen. The remaining chromatin was not condensed (Fig. 1, 3B, and Fig. 11). Masses of homogeneous material then appeared in the areas of rarefaction (Fig. 1, 4B, and Fig. 17) and, in some instances, around, or adjacent to, the nucleoli (Fig. 1, 4B, and Fig. 18). It was not possible to distinguish any separate series of cells indicating a further specific type of progression of these changes. It must be assumed that, as the amount of homogeneous material increased, the chromatin was replaced or condensed and the cells underwent the late changes of series A or C. The B series of nuclear disorganization was found in high, intermediate, and low multiplicity infections.
There was little evidence for extrusion of material from the nucleolus into the nucleoplasm, similar to that which has been described with other viruses (23, 26). Occasional ribonucleoprotein bodies in the nucleoplasm with cytochemical properties similar to those of the nucleolini were observed rarely, although more frequently in the infected than in uninfected control preparations. Protrusions of nucleolini from the body of the nucleolus were not seen.

Figure 10 Reduced number of enlarged nucleolini in an infected cell. Note the light staining center (arrow) of one of the nucleolini. TBM, method B. X 1400.

The characteristic cytoplasmic vacuolation that was the first indication of the existence of infection of monkey kidney cells with SV40 virus (34) was noted in the virus-infected cultures (Fig. 5). It was also found, although less commonly and to a lesser degree, in the control cultures that had not been exposed to virus. The degree of cytoplasmic vacuolation bore no relationship to the development of the nuclear changes, and striking cytoplasmic vacuolation was frequently observed without any detectable nuclear abnormality, and vice versa (Fig. 14). Cytoplasmic vacuolation was invariably associated with the formation of syncytia. In the later stages of infection, cytoplasmic inclusions that were cytochemically indistinguishable from the homogeneous intranuclear material were frequently seen (Figs. 13, 17, and 19). A possible origin of some of these inclusions by some form of nuclear budding is suggested by the appearance of the nucleus shown in Fig. 9.

Figure 11 Rarefaction of the chromatin with some irregular condensation around the nuclear membrane and elsewhere in the nucleus of the largest cell. Note the large vacuoles in the nucleolus. There is widespread rarefaction of the chromatin of the two smaller cells, without condensation of the chromatin. TBM, method C. X 1500.

Cytochemistry

The results of cytochemical staining procedures with and without appropriate extraction procedures are summarized in Tables I and II. The chromatin and the intranuclear homogeneous material (inclusions) were stained positively by the Feulgen method and colored purple by TBM, method C, greenish blue by toluidine blue and TBM, method B, and green by pyronin–methyl green (Table I). Staining of the chromatin and
inclusions by these methods was due to the presence of DNA since they could not be stained after digestion with DNase. Hydrolysis by hot trichloroacetic acid prevented all staining, except for weak pink coloration by the pyronin–methyl green procedure (Table I). Digestion with RNase led to a slight reduction in the intensity of staining of the chromatin by TBM (method C). An accurate assessment of any slight reduction in the intensity of staining of the inclusions by TBM was impossible because of the considerable variability in staining from one cell to another in the same preparation. The presence of RNA in the inclusions was none the less established by the fact that digestion with RNase prevented staining by TBM, method B, and by dilute toluidine blue after nitrosation (Table I). Nothing was stained by TBM after digestion with RNase and DNase.

In almost all procedures involving the use of pepsin, alone or in combination with nucleases, the results were variable, not only from one
experiment to another, but also from one area or cell to another, in the same preparation. This was particularly true when pepsin was used in combination with a nuclease (Tables I and II), and in many experiments the cells were considerably distorted or disrupted by the enzyme treatment. Digestion with pepsin resulted in a change in distribution of nuclear chromatin with blurring of the outline of the chromatin and the inclusions, suggesting that some diffusion of stainable material had taken place. Pretreatment with pepsin gave rise to a variable reduction in the intensity of staining of the chromatin and the inclusions by the Feulgen method. Pepsin followed by RNase removed most of the cells from the glass. There was, again, variable weak staining of
The cytochemical properties of chromatin and SV40 virus-induced inclusions were examined using various pretreatment staining methods. The table below summarizes the staining results for chromatin inclusions:

<table>
<thead>
<tr>
<th>Pretreatment</th>
<th>Staining methods</th>
<th>Chromatin</th>
<th>Inclusions</th>
</tr>
</thead>
<tbody>
<tr>
<td>None or buffer</td>
<td>Feulgen, TBM, TB, PMG*</td>
<td>++</td>
<td>++</td>
</tr>
<tr>
<td>DNase</td>
<td>Feulgen, TBM, TB, PMG*</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>RNase</td>
<td>Feulgen, PMG</td>
<td>+++</td>
<td>+++</td>
</tr>
<tr>
<td>RNase, RNase</td>
<td>TBM</td>
<td>+</td>
<td>See text</td>
</tr>
<tr>
<td>RNase, RNase, and versa</td>
<td></td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Hot trichloroacetic acid</td>
<td>Feulgen, TBM, TB, PMG*</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>DNase, RNase, and versa versa</td>
<td></td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Pepsin</td>
<td>Feulgen</td>
<td>Variable ++</td>
<td>Variable ++</td>
</tr>
<tr>
<td>Pepsin, RNase</td>
<td>Feulgen</td>
<td>Variable +</td>
<td>Variable +</td>
</tr>
<tr>
<td>RNase, pepsin</td>
<td>TBM</td>
<td>++</td>
<td>++</td>
</tr>
<tr>
<td>Pepsin, RNase, and versa versa</td>
<td>TBM</td>
<td>Variable</td>
<td>0</td>
</tr>
<tr>
<td>0.2 N HCl</td>
<td>Feulgen, TBM</td>
<td>+++</td>
<td>+++</td>
</tr>
<tr>
<td>Pepsin, DNase</td>
<td>TBM</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>DNase, pepsin</td>
<td>TBM</td>
<td>Absent or 0</td>
<td>Absent</td>
</tr>
</tbody>
</table>

TBM, toluidine blue-molybdate, method C; TB, dilute toluidine blue after nitrosation; PMG, pyronin-methyl green; DNase, deoxyribonuclease; RNase, ribonuclease, 0, unstained. + to ++++ reflects intensity of staining.

* Pale pink color persisted.
‡ Most of the cells were removed from the glass.
§ A very occasional inclusion was weakly stained.

When pepsin was used after RNase, the intensity of staining by the Feulgen method was reduced more uniformly (Table I). Treatment with pepsin led to an increase in staining by TBM, as previously reported (21). Digestion with pepsin and ribonuclease, irrespective of the order of use, produced a variable reduction in staining of the chromatin by TBM and prevented staining of almost all the inclusions. No significant effect was produced on staining by the Feulgen or TBM methods by treatment with 0.2 N HCl. The effects of pepsin were, therefore, due to activity of the enzyme and not to the hydrochloric acid in which it was dissolved. Pepsin and DNase in any order prevented staining of the chromatin and the inclusions by TBM. When the pepsin treatment followed the DNase, all the inclusion material and most of the chromatin disappeared and could no longer be seen by phase microscopy. The inclusions were not stained by TBM, methods A, D, and E. The inclusions stained green to greenish-yellow by acridine orange, as described by others (28). The chromatin of the nuclear membrane and that associated with the nucleolus appeared to stain more yellow than the inclusions. If the preparations were kept for some time in the light, the red color faded and the chromatin then lost the apparent yellow tinge and was colored green. The yellowish green color of the chromatin would appear therefore to be, at least in part, due to the superimposition of green upon red. After the various extraction procedures, the amount of yellow in the green-stained structures was variable and no constant agreement on the color could be reached by four independent observers. For this reason, the varying shades of yellowish-green are referred to as green in Table II. It was, however, unanimously agreed that after extraction of both nucleic acids with hot trichloroacetic acid the color of the acridine orange-stained material was yellowish green.

The results of the various extraction procedures on staining with acridine orange are summarized in Table II. Digestion with nuclease clearly indicated that staining of the chromatin was in part due to DNA. After pretreatment with DNase, the chromatin was colored very weakly red.
FIGURE 18 Juxtanucleolar masses of inclusion material. TBM, method C. X 1500.

FIGURE 19 Nucleus containing a large, ill defined inclusion and granular remnants of chromatin. Two inclusions are present in the cytoplasm. There is a large vacuole in the nucleolus containing DNA. TBM, method C, after ribonuclease digestion. X 1500.

FIGURE 20 An infected cell treated with deoxyribonuclease and stained by TBM, method C. There is no staining of chromatin or inclusion material. Vacuoles in the nucleus contain homogeneously staining RNA. X 1700.

Subsequent digestion of such preparations with RNase resulted in weak green staining of chromatin. Extraction of both nucleic acids with hot trichloroacetic acid had the same effect as digestion with both nucleases. The results of digestion with RNase indicated that the red coloration of the cytoplasm and the nucleolus was due to the presence of RNA. In contrast to the normal constituents of the cell, the intensity of staining of the inclusions was not reduced by any of the procedures designed to remove one or both nucleic acids. Extraction with hot TCA did not produce any fluorescence in preparations that were not immersed in acridine orange.

Procedures involving digestion with pepsin suffered from the disadvantages that have been
TABLE II
Staining of Green Monkey Kidney Cells and SV40 Virus-Induced Inclusions with Acridine Orange

<table>
<thead>
<tr>
<th>Pretreatment</th>
<th>Chromatin</th>
<th>Inclusions</th>
<th>Cytoplasm</th>
<th>Nucleus</th>
</tr>
</thead>
<tbody>
<tr>
<td>None or buffer</td>
<td>Green ++++*</td>
<td>Green ++++</td>
<td>Red ++++</td>
<td>Red ++</td>
</tr>
<tr>
<td>DNase</td>
<td>Red +</td>
<td>Green ++++</td>
<td>Red ++++</td>
<td>Red ++</td>
</tr>
<tr>
<td>RNase</td>
<td>Green ++++</td>
<td>Green ++++</td>
<td>Green +</td>
<td>Green ++++</td>
</tr>
<tr>
<td>Hot trichloroacetic acid</td>
<td>Green ++</td>
<td>Green ++++</td>
<td>Green +</td>
<td>Green ++</td>
</tr>
<tr>
<td>RNase, DNase, and pepsin</td>
<td>Green ++</td>
<td>Green ++++</td>
<td>Green +</td>
<td>Green ++++</td>
</tr>
<tr>
<td>vice versa</td>
<td>Pepsin</td>
<td>Green ++++</td>
<td>Red ++++</td>
<td>Red ++</td>
</tr>
<tr>
<td>RNase, pepsin</td>
<td>Green +</td>
<td>Absent</td>
<td>Green +</td>
<td>Green ++++</td>
</tr>
<tr>
<td>Pepsin DNase</td>
<td>Variable green or red</td>
<td>Absent§</td>
<td>Variable green</td>
<td>Variable green</td>
</tr>
<tr>
<td>DNase, pepsin</td>
<td>Absent</td>
<td>Absent</td>
<td>Red ++++</td>
<td>Red ++</td>
</tr>
<tr>
<td>0.2 N HCl</td>
<td>Green ++++</td>
<td>Absent</td>
<td>Red ++++</td>
<td>Red ++</td>
</tr>
</tbody>
</table>

Abbreviations as in Table I.

* The green color often appeared to have a yellowish tinge (see text).
§ Probably due to nucleolar-associated chromatin.
§ Rarely present and stained green +.

methods A and B. In the early stages of infection, the contents of the vacuoles no longer stained after digestion with RNase, but were not significantly affected by DNase (Fig. 20). In the later stages of infection, the contents of the largest vacuoles became Feulgen-positive and assumed the same cytochemical properties as the inclusion material in the rest of the nucleus; they were digested by DNase but staining by TBM, method C, was not detectably altered by RNase (Fig. 19).

Immunofluorescence
The results of immunofluorescent staining were essentially similar to those described by Mayor et al. (28), Diderholm (5), and Levinthal and Shein (19) and need not be described in detail. Considerable asynchrony was noted in the development of the antigen. Thus, diffuse nuclear staining, perinucleolar staining and staining of the cytoplasm were sometimes observed in the same preparation. Antigen was observed in particulate form in the nucleolus in the late stages of infection, when DNA became detectable in the nucleolar vacuoles.

DISCUSSION
The changes in the nucleolus correspond very closely to those observed by Granboulan et al. (12)
by electron microscopy and by Prunieras (31) by time-lapse cinematography. Nucleolar changes are illustrated diagrammatically in Fig. 2 of the paper by Granboulan et al. (12). The less dense vacuolar structures in the nucleoli that are surrounded by a zone of denser material correspond to the nucleolini. These vacuoles become larger in the infected cells and the amount of dense material increases and is often somewhat eccentrically placed at the edge of the vacuole, cf. Fig. 15 and 16 of this paper. Later in the infection Granboulan et al. found virus in the less dense vacuoles at the time when the vacuoles observed in the present study contained material that was identical to that of the inclusions. The likelihood of virus being responsible for the DNA-containing material in the vacuolar structures is suggested by the present observations, and by those of others (5, 19, 28) that antigen is seen in small particulate points in the nucleolus. The observation of antigen and inclusion material around the nucleoli in occasional cells also suggests that virus may be formed there and subsequently pass outwards into the nucleoplasm. A process of extrusion of material from the nucleolus has been observed by Prunieras (31), whose studies envisage the transportation of material from nucleolus to cytoplasm. Extrusion of RNP from enlarged nucleolini into the nucleoplasm has been observed with herpes and polyoma virus where large RNP bodies or “B-bodies” were formed in the nucleoplasm (23, 26). This was only rarely observed with SV40 virus. On the other hand, there was no suggestion of extrusion of DNA-containing material from the nucleoli of cells infected with polyoma or herpes virus (23, 26). Nor is there definite evidence for the formation of virus or antigen in the nucleoli of cells infected with these viruses (13, 37).

An increase in the size of nucleolini has been observed with all intranuclear DNA virus infections that have been studied (23). In electron micrographs, dense bodies which appear to correspond to enlarged nucleolini have been noted in cells infected with varicella (35) polyoma (2) and SV40 viruses (12). The nucleolini were unaffected in cells infected with vaccinia or Newcastle disease virus (23). Enlargement was also noted when DNA synthesis was inhibited by treatment with 5-fluorodeoxyuridine (FUDR) (24). The changes produced by FUDR can be prevented by incorporation of thymidine in the FUDR-containing medium (24). Thus, when the conversion of deoxyuridylic to thymidylic acid was by-passed by the introduction of exogenous thymidine, DNA synthesis could take place and the resultant changes in the RNP of the nucleolini were prevented. The factor common to all the circumstances in which enlargement of nucleolini occurs is some abnormality of DNA metabolism, most probably either a focal or general inhibition of synthesis. The exact mechanism by which a disturbance of DNA metabolism results in changes in the RNP of the nucleolini is not yet clear. It is of interest, however, that there is some evidence that the nucleolini may reflect the presence of a DNA-RNA hybrid, or at least that the structural integrity of the nucleolini is dependent upon the presence of intact DNA (22).

The results of cytochemical studies using toluidine blue, TBM, pyronin–methyl green and the Feulgen method, and selective extraction procedures for nucleic acids are quite clear. The inclusions in the virus-infected cells contain DNA and RNA. They also contain non-histone protein. The DNP and RNP of the inclusions respond in the same way as cellular nucleoprotein to digestion with nucleases and extraction with hot TCA. As reported by others (28), the inclusions were colored green or greenish yellow by acridine orange and their staining properties were unaffected by digestion by nucleases. The inclusions also stained after extraction of both nucleic acids by hot TCA. The procedures for nuclease digestion must have been effective since they removed all nucleic acids stainable by the TB, TBM, Feulgen and pyronin–methyl green methods. Residual green staining of cellular structures and the inclusions after extraction or digestion was not therefore due to the staining of nucleic acids. In the absence of acid mucopolysaccharides which stain metachromatically with TB, or which, by virtue of accompanying sugars, are PAS-positive (18), the residual staining by acridine orange must be attributed to binding of the dye by carboxyl groups of protein that were charged at pH 4.0. Herrman et al. (14) have shown that, under the conditions of their experiment, the basic dye toluidine blue binds protein as well as nucleic acid above pH 4.2. However, the pH above which a basic dye will bind the carboxyl groups will vary with the type of fixation (17), concentration of dye, temperature, and many other factors (21). Green staining of the inclusion body in tissue culture cells infected with rabies virus...
and stained with acridine orange has also been shown to be due to protein and not to the presence of nucleic acids (20). Since digestion with pepsin and nuclease produced extensive cell damage, diffusion of material within the nucleus and green or even red staining of chromatin such digestion procedures would seem to be of doubtful value. Furthermore, there was no selective specificity of nuclease action since DNase and RNase, in combination with pepsin, were almost equally effective in removing the inclusions or preventing their coloration by dye. Furthermore, treatment with pepsin produced a reduction in the intensity of staining of DNA by the Feulgen method and, in combination with RNase, still further reduced the intensity of staining.

There is considerable evidence that the nucleic acid of unfixed viruses is not susceptible to digestion with nucleases (4, 10). The biological activity and therefore the nucleic acid of a number of unfixed riboviruses is not affected by digestion with ribonuclease (4). Furthermore, the nucleic acid phosphoryl groups, which bind cationic dyes such as toluidine blue or acridine orange, are often masked in unfixed preparations of cells or viruses (21, 27). After suitable fixation, however, the nucleic acid core of riboviruses (7) and deoxyriboviruses (8, 9) becomes susceptible to digestion with nucleases. Similarly, some viruses stain more strongly with acridine orange after fixation (27). Digestion with nucleases prevents staining of fixed purified virus preparations containing RNA or double or single-stranded bacteriophage DNA. However, the yellow-green staining of fixed virus preparations containing double-stranded DNA is not prevented by digestion with DNase unless the preparation has been previously treated with pepsin (27). Epstein has shown that the core of adenovirus and herpes virus, two viruses containing double-stranded DNA, is susceptible to digestion by deoxyribonuclease after fixation by permanganate (8, 9). In the light of these observations and those reported here, it seems unlikely that acridine orange staining of the SV40 virus-induced inclusions was due to binding of dye by the DNA of the virus.

The authors wish to thank Joyce Sills Shahn for the drawings, and Oksana Bohachevsky and Kathy Wannemacher for valuable technical assistance. This work was supported by United States Public Health Service Grants CA-05402-04, RG-813-03, A-102954-05, and CA-04534-06. 

Received for publication, July 28, 1964.

R E F E R E N C E S

3. BURSTONE, M. S., J. Histochem. and Cytochem., 1955, 3, 32.
24. LOVE, R., and WALSH, R. J., J. Histochem. and Cytochem., 1963, 11, 188.