Transcription of the Influenza Ribonucleic Acid Genome by a Virion Polymerase

III. Completeness of the Transcription Process

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The virion ribonucleic acid (RNA) polymerase of influenza strain A0/WS transcribed at least 81% of the viral genome in vitro. The polymerase is shown to be associated with each of the major size classes of the virion RNA-ribonucleoprotein complexes. Under optimal in vitro conditions, at least 45% of the RNA contained in a population of influenza virions was involved in a repetitive transcription process. The detectable proteins associated with enzymatically active complexes containing RNA, ribonucleoprotein, and polymerase have been identified by polyacrylamide gel electrophoresis.

Influenza virus possesses a virion ribonucleic acid (RNA)-dependent RNA polymerase (10, 14, 20). In vitro assays have demonstrated that even closely related strains of influenza virus (e.g., WSN and WS) apparently possess different amounts of enzyme activity as judged by their rate of uridine monophosphate (UMP) incorporation per milligram of viral protein (10). We have previously examined the properties of the in vitro transcription process using the less active influenza WSN strain. This strain was chosen because it can be grown in tissue culture and the virion RNA can be labeled sufficiently with 3H-uridine to allow its fate to be monitored in product analysis experiments (4, 5). In such experiments, we demonstrated that in 2 hr of in vitro incubation, about 7% of the virion RNA was involved in transcription and at least 14% of the genome was transcribed. Since most of the product RNA was initially associated with the virion 3H-RNA species and all of it could be annealed to the 3H-RNA, these results suggested that the enzyme activity was responsible for transcribing some or all of the viral genome into complementary product RNA species.

In this communication, we have used the more active influenza WS strain and examined the amount of viral RNA involved in transcription, the extent of the transcription process, the association of the virion polymerase with the RNA-ribonucleoprotein complexes, and the protein species associated with these complexes. Which of these proteins constitute the virion polymerase is under study.

MATERIALS AND METHODS

Virus preparations. Influenza A virus, strain WS, was grown in the allantoic membrane of embryonated or de-embryonated chicken eggs by use of a modification of the technique of Bernkopf (1; Stevens and Simpson, in preparation). Each egg received 0.2 mCi of 3H-uridine (specific activity, 22 mCi per pmole) and 1.4 X 106 to 140 X 106 egg infectious units of virus. After incubation at 37 C for 30 hr, the egg fluids were harvested and pooled. These pools, having hemagglutinin titers of 1:4,096, were then centrifuged at 7,500 X g for 30 min at 4 C to remove debris. Virus was obtained from the clarified supernatant fluid by precipitation with polyethylene glycol (Carbowax 6000) and was thereafter purified by a double-cushion centrifugation with the use of 30 and 60% (w/v) sucrose as described previously (9). Virus was harvested from the surface of the 60% sucrose layer, diluted with 0.15 M NaCl, 0.01 M tris(hydroxymethyl)aminomethane (Tris)-hydrochloride buffer (pH 7.4), and then centrifuged to equilibrium in a 20 to 70% (w/v) sucrose gradient prepared in 0.15 M NaCl, 0.01 M Tris-hydrochloride buffer (pH 7.4). The centrifugation conditions were 25,000 rev/min in a Spincos SW27 rotor at 4 C for 17 hr. Virus was collected from the gradient, and sucrose was removed by passage through a 10-ml column of Sephadex G25, equilibrated and eluted in 0.15 M NaCl, 0.01 M Tris-hydrochloride buffer, pH 7.4. The final virus preparations contained 0.8 mg of protein per ml and possessed RNA specific activities of 2 X 106 or 2.5 X 104 counts/min of 3H per mg of RNA—the latter being derived from the WS-infected de-embryonated eggs.

Enzyme assays and purification of the reaction product nucleic acids. Reaction mixtures were prepared containing 3H-α-uridine triphosphate (UTP) to label the product species as described previously (4, 5). In
the experiment designed to determine the extent of the transcription process with WS virus of lower 3H-RNA specific activity, the specific activity of the 32P-UTP was 8 counts per min per pmole (i.e., approximately 6,000 counts per min per μg of product RNA, assuming 25 mole % of UMP in the product). In other experiments, the UTP specific activity was 5,300 counts per min per pmole. Reaction product nucleic acids were purified from protein and triphosphates, etc., as described previously (6).

Annealing of reaction product nucleic acids and RNA polyacrylamide gel electrophoresis. Procedures have been described for annealing reaction product nucleic acids and determining the ribonuclease resistance of the annealed RNA (5). The separation of RNA species by gel electrophoresis in swollen 3.2% polyacrylamide gels by use of Tris-acetate buffer (pH 7.4) containing 0.1% sodium dodecyl sulfate (SDS) has also been described (17). After electrophoresis, the gels were sliced, and the distribution of RNA species was determined by dissolving the slices in 30% (v/v) H2O and counting in a liquid scintillation counter with Aquasol (New England Nuclear Corp., Boston, Mass.).

Isolation of influenza virion RNA-ribonucleoprotein complexes. The procedure used to isolate the virion RNA-ribonucleoprotein complexes of influenza was a modification of that described by Pons (15). A preparation of virus in 1 ml of 0.15 M NaCl, 0.01 M Tris-hydrochloride buffer, pH 7.4, was mixed with Triton N101, sodium deoxycholate (DOC), urea, and mercaptoethanol (at final concentrations of 3 mg/ml, 0.1%, 0.2 M, and 0.01 M, respectively), and was then incubated at 31°C for 5 min; by this procedure, the virus suspension completely lost its turbid appearance. Incubation at 4 or 20°C was insufficient to achieve release of the RNA-ribonucleoprotein complexes from the virions. The mixture was cooled to 4°C and loaded over a 10.5-ml linear gradient of 15 to 30% (w/v) sucrose in 0.15 M NaCl, 0.01 M Tris-hydrochloride buffer (pH 7.5), 0.01 M mercaptoethanol, 0.2 M urea, 0.02% DOC, and 3 mg of Triton N101 per ml, prepared and held at 3°C in a Spinco SW41 tube over a 0.5-ml cushion of 70% (w/v) sucrose. The gradient was centrifuged at 40,000 rev/min at 3°C for 4 hr. Each fraction of the gradient was assayed for content of 3H-RNA and RNA polymerase activity (4) with or without added influenza RNA (0.5 μg per 125-μl reaction mixture), and the indicated fractions were pooled to determine their content of protein or RNA by polyacrylamide gel electrophoresis.

The conditions employed for dissociating virus were selected solely with reference to their effect on the virion transcriptase activity. For example, the rate of enzyme activity was determined in reactions containing increasing concentrations of Triton N101. Maximal enzyme activity was obtained in the presence of 1 to 10 mg of Triton N101 per ml of reaction mixture. In the presence of 3 mg of Triton N101 per ml of virus preparation, pretreatment of the virus at 31°C for 5 min with DOC (pH 7.8) was examined. After pretreatment, the virus was diluted fivefold into the reaction mixture, and the rate of enzyme activity was again determined. It was found that pretreatment with greater than 1% deoxycholate was totally inhibitory for polymerase function. Pretreatment with 0.1, 0.3, and 0.6% DOC inhibited the enzyme activity by 16, 60, and 82%, respectively. With 3 mg of Triton N101 per ml, inclusion of urea in the reaction mixture was also examined. It was found that reaction concentrations of urea of greater than 1 M were totally inhibitory. Reaction concentrations of 0.1, 0.3, and 0.5 M urea depressed the rate of enzyme activity by 15, 40, and 70%, respectively. Pretreatment of virus with Triton N101 (3 mg per ml) and DOC (0.1%) and dilution into a reaction mixture containing 0.2 M urea depressed the enzyme activity by 30% in comparison to a Triton-treated virus preparation. The enzyme activity, however, was linear for only 1 hr after these treatments, in contrast to Triton-treated virus whose enzyme activity is linear for 6 to 8 hr. The reason for this inhibition is not known. Inclusion of mercaptoethanol (0.01 M final concentration) did not inhibit the reaction rate. From these preliminary experiments, the conditions for pretreating virus were devised.

Separation of influenza proteins by SDS polyacrylamide gel electrophoresis. Virus or subviral components were precipitated with 5% (w/v) trichloroacetic acid and held at 4°C for a minimum of 4 hr; the precipitate was collected by centrifugation at 4°C and 20,000 × g for 30 min by use of a Sorvall swinging-bucket HB4 rotor and a Spinco SW41 centrifuge tube. The precipitate was washed by centrifugation with 5% trichloroacetic acid, followed by two washes with absolute alcohol to remove the detergents and acid. After drying, the pellets were dissociated in 50 to 200 μl of 0.01 M sodium phosphate buffer (pH 7.0), 0.1% SDS, 1 M urea, and 0.01 M dithiothreitol for 30 min at 60°C; they were then mixed with 20 μl of glycerol and subjected to electrophoresis in 8% polyacrylamide gels as described previously (8). The gels were stained by Coomasie brilliant blue, and the stained proteins were scanned at 640 nm with an automatic double-beam spectrophotometer (model SD3000, Schoeffel Instrument Corp., Westwood, N.J.). The main protein bands were identified by comparison with parallel electropherograms of SDS-dissolved virus.

RESULTS

Extent of transcription by influenza WS virion transcriptase. In incubations with WSN virus, we have obtained by 2 hr of incubation at least 14% transcription of the virion genome in a transcription process in which at least 7% of the viral RNA was involved (5). Since the linear incorporation of triphosphates ceased between 6 and 7 hr of incubation, in those in vitro reactions, we have not been able to obtain more than 28% transcription involving 18% of the viral RNA. We have not been able to determine with WSN whether this incomplete transcription was due to an inherent property of the transcription process or was caused by the in vitro conditions to which the virus was exposed. However, we have found that product RNA is associated with the five major classes of virion RNA (containing seven
RNA species) suggestive of a transcription process associated with each virion RNA molecule (5). In the present dissertation, we report the extent of transcription by the egg-grown influenza strain A<sub>6</sub>/WS virus which has been shown to contain a more active polymerase (10).

A reaction mixture containing <sup>32</sup>P-α-UTP to label product RNA was incubated at 31 C with <sup>3</sup>H-uridine-labeled WS virus grown in embryo-nated eggs. Samples were withdrawn at intervals, and the reaction product RNA (consisting of <sup>32</sup>P-product and <sup>3</sup>H-virion species) was purified. The ribonuclease resistance of the reaction product nucleic acids was determined before and after an annealing pretreatment (Fig. 1). It was found that through 8 hr of reaction incubation the ribonuclease resistance of the virion <sup>3</sup>H-RNA increased from 0.5% for the zero-time unincubated sample to 40 and 47% for the 6- and 8-hr samples, respectively. This result indicated that in this population of WS viral particles at least 40 to 50% of the virion RNA was involved in the transcription process.

The ribonuclease resistance of the annealed <sup>3</sup>H-RNA increased from 3% for the zero-time sample to 62 and 81% for the 6- and 8-hr samples, respectively, indicating that at least 80% of the virion RNA had been transcribed. Since, however, only 40 to 50% of the viral RNA population became ribonuclease-resistant as a result of transcription, these results suggest either that 40 to 50% of the virions possessed active enzyme whose transcription process was repetitive and at least 80% complete, or that 81% of the virion RNA population was involved in transcription.

The ribonuclease resistance of the <sup>32</sup>P-product RNA species was 70% for the unannealed 1-hr sample and 42% for the unannealed 8-hr sample. After annealing, the ribonuclease resistance of the product species increased to 98 and 70% for the same two samples, respectively. When more WS viral RNA was added to the 8-hr sample (6 μg per ml annealing mixture), the ribonuclease resistance of the annealed product species increased to 97%, indicating that most of the 8-hr product RNA was complementary to the virion genome rather than identical in sequence to the viral RNA. Since the 27% increase in the ribonuclease resistance of the 8-hr <sup>32</sup>P-product RNA was obtained only after addition of WS viral RNA, this result indicated that the transcription process was indeed repetitive.

**Dissociation of influenza WS and separation of the virion RNA-ribonucleoprotein complexes.** A useful procedure has been developed for the dissociation of influenza strain A<sub>6</sub>/WSN and isolation of the virion RNA-ribonucleoprotein complexes (15). We have confirmed these obser-

![Figure 1](http://jvi.asm.org/)
FIG. 2. Dissociation of influenza WS and separation of the virion RNA-ribonucleoprotein complexes. A 1-ml preparation of 3H-uridine-labeled influenza WS virus (2.5 x 10⁴ counts per min of 3H µg of RNA) in 0.15 m NaCl, 0.01 m Tris-hydrochloride buffer, pH 3.4, was dissociated by Triton N101, DOC, urea, and mercaptoethanol at 31 C for 5 min as described in Materials and Methods, and then was cooled to 4 C in an ice bath. The dissociated virus, which visually was completely clear, was then loaded on a 10.5-ml linear gradient of 15 to 30% sucrose containing 0.15 m NaCl, 0.01 m Tris-hydrochloride buffer (pH 7.5), 0.01 m mercaptoethanol, 0.2 m urea, 0.02% DOC, and 3 mg of Triton N101 per ml in a Spinco SW41 centrifuge tube. The gradient, which was made over a 0.5-ml cushion of 70% (w/v) sucrose, was centrifuged at 40,000 rev/min at 3 C for 4 hr. The distribution of 3H-RNA was determined on samples of each fraction, and the indicated fractions were pooled, mixed with sodium dodecyl sulfate, and subjected to polyacrylamide gel electrophoresis to resolve the RNA species. The electrophoretic conditions are described in the text. After electrophoresis, the gels were sliced and dissolved in H₂O₂ as described previously (3, 5, 7); the distribution of 3H-RNA was determined by use of Aquasol. The sucrose gradient profile of the dissociated 3H-WS virus is shown in the first panel. Subsequent panels show the eight RNA gel electropherograms of the indicated gradient fractions.
vations and applied the procedure to answer the question of whether these complexes have enzyme activity and, if so, which of the virion proteins are present in these structures. A preparation of \(^{3}\text{H}\)-labeled virus was dissociated by Triton N101, DOC, and urea, and the products were resolved by velocity centrifugation in a gradient of sucrose as described in Materials and Methods. Each fraction was monitored for labeled RNA (Fig. 2); the indicated fractions were pooled and dissociated with SDS, and their content of RNA was determined by polyacrylamide gel electrophoresis (Fig. 2). The electrophoretic conditions were adapted to overcome the problems engendered by the high ionic content of the samples, as well as their large loading volumes (0.1 to 0.3 ml), by running the gels in the presence of 3E buffer (i.e., 0.12 M Tris-acetate buffer pH 7.4, 0.06 M sodium acetate; see reference 7). Very little label was recovered on the 70% sucrose cushion or in the top fractions of the gradient, indicating that most of the virus had been dissociated and that most of the label was present in the virion RNA-ribonucleoprotein complexes (see below). Examination of the gel electropherograms revealed that sucrose gradient fractions 14 to 16 principally contained the large class of viral RNA species (compare with Fig. 3, which shows the RNA gel electropherogram of the complete virus preparation). Fractions 23 to 25 principally contained the smallest class of virion RNA species. The intermediate fractions contained various mixtures of the five classes of virion RNA molecules as shown in Fig. 2. These results suggest, therefore, that the conditions chosen for dissociation of WS influenza virus were adequate, but the separation between various virion ribonucleoprotein complexes was not very good.

**Association of polymerase and proteins with influenza RNA-ribonucleoprotein complexes.** A preparation of \(^{3}\text{H}\)-labeled influenza WS was dissociated with Triton N101, DOC, and urea, and the RNA-ribonucleoprotein complexes were resolved by sucrose gradient velocity centrifugation as described in the previous section. Each fraction of the gradient was assayed for RNA polymerase activity in the presence or absence (endogenous activity) of added influenza RNA (Fig. 4). The majority of the endogenous polymerase activity was recovered in those fractions which contained the \(^{3}\text{H}\)-RNA-ribonucleoprotein complexes. Addition of influenza RNA only slightly enhanced the enzyme activity of some of these fractions. There was also some templated enzyme activity at the top of the gradient.

The indicated fractions were pooled and precipitated with 5% trichloroacetic acid; the precipitate was collected by centrifugation. After dissociation in SDS (see Materials and Methods), protein gel electrophoresis was performed on each sample (Fig. 5). By comparison with the protein electropherogram of complete virus, it was determined that the virion ribonucleoprotein, NP, and the two minor proteins, P, were present in the gradient fractions which also contained the labeled RNA and endogenous polymerase activity (i.e., gradient pools 3 and 4, Fig. 4). Although the resolution between NP and HA1 and NA (the glycoproteins hemagglutinin subunit 1 and neuraminidase, respectively) was poor, these latter two proteins were apparently absent from the gradient fractions which contained the labeled RNA and endogenous enzyme activity (see Discussion and legend to Fig. 5). The top fractions of the gradient (i.e., gradient pools 6 and 7, Fig. 4) apparently contained all the glycoproteins as well as the membrane (M) protein (see Discussion). From these results, it was concluded that the RNA-ribonucleoprotein complexes of influenza virus contain RNA, polymerase, and probably only the major virion protein NP and minor proteins P. As can be seen from Fig. 5, most of the virion ribonucleoprotein was recovered in the fractions which also contained the RNA and transcriptase enzyme activity. Procedures to dissociate proteins from these complexes are currently being investigated.
**DISCUSSION**

Transcription process of influenza viruses. It has been shown that influenza WS virion transcriptase is able to transcribe at least 80% of the viral genome. The process is repetitive and probably involves all classes of the segmented viral RNA genome. As we have previously shown, the products of the transcription process consist only of complementary RNA (5); therefore, these results suggest that the product RNA species probably have some important function in the infection process of a cell by influenza virus. Such a proposal has been substantiated by the recent finding that polysomal messenger RNA species in influenza-infected cells is complementary to the virion genome (16). Consequently, it would appear that the virion transcriptase is responsible for the synthesis of influenza messenger RNA species. In vitro, the transcription process is slow, even with the relatively active WS virion transcriptase. For the WSN strain, the in vitro process appears even slower, although it is not known whether this relates to the possession of a different polymerase or reflects the lower number of virions in a population of particles exhibiting enzyme activity. To obtain an answer to these possibilities will require obtaining genome recombinants between the two viruses and an examination of the rate of product synthesis as well as the amount of virion RNA involved in transcription; this is currently being undertaken.

Association of transcriptase with the RNA-ribonucleoprotein complexes. When influenza virus is dissociated by appropriate techniques, the virion RNA-ribonucleoprotein complexes can be partially separated from each other by sucrose gradient velocity centrifugation. Since separation of the complexes can be obtained in this manner, these results indicate that the virion complexes...
are either separate from each other or linked through hydrogen-bonding or possibly disulfide bridges between their proteins in the virus particle. It is not known whether the complexes are also linked to other structural components of the virus, e.g., the membrane. We have demonstrated that most of the recovered endogenous RNA polymerase activity is associated with those gradient fractions which contain the $^3$H-RNA-ribonucleoprotein complexes. We have obtained similar results, but with somewhat better separation, using influenza WSN strain (unpublished data).

It is of particular interest to note that the profiles of $^3$H-RNA and RNA polymerase activity of the dissociated virus spread by sucrose gradient centrifugation are not coincident (Fig. 4). We have obtained similar noncoincident profiles with influenza WSN virus. However, it should be remembered that the $^3$H-RNA in each fraction reflects the mass of RNA species present and not their molar ratio. Since the influenza RNA segments vary in molecular weight (from $1.05 \times 10^5$ to $3.5 \times 10^5$ [5]), this means that per unit of $^3$H counts per minute there can be a threefold variation in the number of RNA molecules between the fastest and slowest sedimenting RNA-ribonucleoprotein complexes. The number of active polymerase molecules per gradient fraction could possibly be related either to the RNA species mass or to their molar concentration. In the latter case, if all of the complexes have a similar number of active enzyme molecules asso-

FIG. 5. Protein gel electrophoresis of influenza WS and its dissociated components. The gradient pools from Fig. 4 were mixed in an SW41 centrifuge tube with trichloroacetic acid (5% final concentration) and kept at 4 C for 4 hr. The precipitated proteins were collected by centrifugation at 4 C and 20,000 × g, and were washed once with 5% trichloroacetic acid and then twice with absolute ethanol to remove trichloroacetic acid and the detergents. After drying, the proteins were dissociated and resolved by gel electrophoresis, and the stained proteins were scanned at 640 nm (see Materials and Methods). The proteins of complete influenza WS virus, run in parallel, are also shown. The influenza proteins were identified from the data of Combs et al. [11] and Schulze (18), and are named in accordance with the decision of the influenza workshop held in Madison, Wis., in 1971 [19]. The influenza hemagglutinin glycoprotein HA1:HA2 was completely dissociated under the conditions used, and the subunits were recovered as indicated. The other glycoprotein, neuraminidase (NA), could not be identified, but its presumed position is also indicated (see 11, 18, 21). The two major non-carbohydrate-containing virion proteins, nucleoprotein (NP) and membrane protein (M), were clearly seen as indicated, although the former (the sharp NP band) runs close to the two diffuse glycoproteins (HA1 and NA). Owing to the small amounts of protein in the samples, the whole sample of each pool was loaded on the gels except for gradient pools 3 and 4, of which only one-third of the protein was loaded. No stained protein was observed in gradient pool 5 despite the scan obtained, and this was probably because of irregularities.
ciated with them, then one would expect to have less enzyme activity on a 3H-RNA mass basis in the leading fractions. Such is the observation in the results presented in Fig. 4, where it is apparent that the variation in enzyme activity across the RNA-ribonucleoprotein complexes profile is in the range predicted by these arguments. However, this result does not preclude the possibility that one or more of the RNA species do not have associated transcriptase enzyme. A conclusive decision on this point will have to await better resolution of the various complexes.

Proteins associated with the RNA-ribonucleoprotein complexes of influenza virus. The identification of the influenza virion proteins (Fig. 5) has been made by reference to the published protein electropherograms of influenza A virus (11, 18, 19), with the use of the nomenclature adopted at the influenza workshop held at Madison, Wis., in 1971. Under the conditions of dissociation we have used, the hemagglutinin complex (HA1: HA2) has been completely dissociated into its subunits HA1 and HA2 (11, 18, 21).

We have not been able to identify or differentiate the two glycoproteins, hemagglutinin (HA1 subunit) and neuraminidase, or to separate the faster-moving hemagglutinin (HA2 subunit) and membrane protein (see 11, 18). However, it is clear from the data presented in Fig. 5 that neither of the latter two proteins was present in those gradient fractions (fractions 3 and 4, Fig. 4 and 5) which contained the RNA-ribonucleoprotein complexes. It can also be concluded that the majority of the HA1 subunit and neuraminidase were also absent from those fractions and that they were recovered in the top gradient fractions which also contained the HA2 and membrane protein.

In unpublished similar experiments involving 3H-glucosamine-labeled WS virus (grown in de-embryonated eggs), we have observed that 99.7% of the viral 3H label is recovered in the top gradient fractions after dissociation by the procedure we have described. Less than 0.3% of the 3H label was recovered in the region which contained the RNA-ribonucleoprotein complexes. We are therefore led to conclude that neither the glycoproteins (NA, HA1, and HA2) nor the virion membrane protein (M) are part of the RNA-ribonucleoprotein complexes. The virion minor protein P, which can sometimes be seen as two barely separated bands (P1 and P2), was found in the same gradient fractions which contained the nucleoprotein NP, the RNA, and polymerase enzyme activity (gradient pools 3 and 4, Fig. 4 and 5). Not all of the P protein was recovered in these fractions; some was apparently also recovered in the top fractions of the gradient and also on the 70% sucrose pad (see Fig. 5 legend). The relationship of the P proteins to the NP protein and polymerase is currently under investigation. Methods to dissociate the polymerase from the virion RNA and obtain active enzymes are also being investigated.

Other investigations into the protein composition of the ribonucleoprotein complexes of influenza and other viruses have not been conducted in relation to the retention of transcriptase enzyme activity (2, 12, 13, 17). Since in those reports only one protein was identified (NP), this could be due either to the different conditions employed for virion dissociation (e.g., high DOC concentration) or to the small amount of P proteins associated with these complexes.

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