Transcription of naked double-stranded RNA molecules in a fraction containing large vesicles plus mitochondria from the basidiomycete Agrocybe aegerita

GÉRARD BARROSO and JACQUES LABARÈRE

Laboratoire de Génétique Moléculaire et d'Amélioration des Champignons Cultivés, Université de Bordeaux II–INRA, CRA de Bordeaux, BP 81, 33883 Villenave d’Ornon Cedex, France

(Received 10 June 1992; revised 4 September 1992; accepted 23 September 1992)

Actinomycin D-resistant RNA synthesis was detected in the cellular (VM) fraction which contains large vesicles plus mitochondria from Agrocybe aegerita. It involved the RNAase-resistant complex of naked (unencapsidated) double-stranded RNAs (dsRNAs). The RNA polymerase assay showed [γ-32P]UTP incorporation in dsRNAs of a size corresponding to the previously described naked M-dsRNAs and in single-stranded RNAs of related size, which are assumed to act as intermediaries in the replicative cycle of the dsRNA molecules. Moreover, the incorporation of ribonucleotides into large mitochondrial dsDNA fragments was detected in the dsRNA-containing strain. This result favours a mitochondrial location for the naked dsRNAs and their associated polymerizing enzyme(s). No replication of the encapsidated large L-dsRNA viral genome was detected in the VM fraction, supporting the hypothesis that the two types of dsRNAs observed in this A. aegerita strain are independent entities.

Introduction

In lower eukaryotes, such as fungi or protozoa, most of the viruses store their genetic information in double-stranded RNA (dsRNA) molecules, whose transcription and replication are DNA-independent. These dsRNA segments are generally encapsidated in isometric virus particles (for reviews, see Buck, 1986 and Patterson, 1990). However, besides these typical viral genomes, naked (unencapsidated) dsRNAs have been described which are located in membranous structures (vesicles or mitochondria) of parasitic or saprophytic fungi (Rogers et al., 1987; Barroso & Labarère, 1990).

The best studied of the naked dsRNAs are those associated with hypovirulence of the chestnut blight fungus Cryphonectria (Endothia) parasitica (Anagnostakis, 1982). By sequencing the largest (12712 bp) dsRNA (L-dsRNA) present in a hypovirulent strain of this parasitic fungus, Shapira et al. (1991a) have shown that, although dsRNAs are not packaged within particles and do not exhibit an extracellular phase, their organization and expression strategies are closely related to viral genetic information. In this context, they propose the introduction of hypovirulence-associated virus (HAV) as a descriptive term for these dsRNAs. Moreover, they demonstrate that the smaller dsRNA molecules (from 10 to 0.6 kbp) of the hypovirulent strains are the result of the generation and maintenance of internally deleted forms of the L-dsRNA (Shapira et al., 1991b).

In another pathogenic fungus, Ophiostoma ulmi, Rogers et al. (1987) established that the dsRNA segments of the fungal diseased isolates copurified with mitochondria and were associated with a cytochrome aa₃ deficiency.

In the saprophytic edible basidiomycete Agrocybe aegerita, two types of RNAase-resistant complexes have been found. The first complex was the encapsidated genome (L-dsRNA of 6200 bp) of an isometric mycovirus; the second one comprised naked M-dsRNAs (three molecules of closely related size from 1900 to 1700 bp) associated with the large vesicles and mitochondria (VM) cellular fraction (Barroso & Labarère, 1990). In contrast to the discrete viral L-dsRNA molecule, M-dsRNAs were recovered in large quantities from the infected strain (10 μg M-dsRNAs from 20 g dried vegetative mycelium). However, the two types of molecules were, at that time, cryptic, i.e. their presence was not related to a phenotype or a modification of the hyphal growth or physiology (Barroso & Labarère, 1990).

To study the replication and the relationship between
the dsRNAs of the two RNA:se-resistant complexes of the A. aegerita strain, we have investigated the location and characterization of the dsRNA synthesizing enzyme of each complex.

Methods

Strains and cultures. The tetrapolar basidiomycete Agrocybe aegerita belongs to the class Agaricales. The wild dikaryotic strains used, SM47 (mating-type alleles: A7A8 B4B7) and SM51 (A7A4 B3B4), were derived from in vitro cultures of fragments of fruiting bodies collected from nature. The dikaryotic strain SM47 was cultivated in vitro to obtain basidiospores from which homokaryotic progeny A1 was obtained (Noël et al., 1991). The two strains SM51 and A1 were shown to lack dsRNA molecules and, accordingly, were used as controls in the RNA polymerase assays. Both homokaryotic and dikaryotic vegetative mycelia were cultivated on solid or liquid complete CYM medium (Raper et al., 1972). Growth of the vegetative mycelium was obtained in Roux culture flasks, at 28 °C in the dark. After incubation, mycelium was harvested by filtration on sterilized gauze, washed with distilled water, and large membrane and cell wall fragments. DNAase I (100 pg ml⁻¹), human placenta ribonuclease inhibitor (HPRI; Appligene) was omitted. The nucleic acid hydrolases were eliminated by treatment. The nucleic acid hydrolases were eliminated by

Cellular fractionation. Twenty grams of dried vegetative mycelium was disrupted in 30 ml cold 15% sucrose suspension buffer (SSB: sucrose 15%, w/v, 10 mM-Tris/HC1 pH 6.8, BSA 0.1%, w/v, 5 mM-β-mercaptoethanol) using a homogenizer (model HO2-1 from Edmund Bühler: 30 s at 20000 r.p.m., 5 s at 30000 r.p.m. and 5 s off, twice), then filtered through four layers of sterile gauze without squeezing the hyphae, to eliminate remaining large hyphal fragments. The filtrate was centrifuged once at low speed (28000 g, 10 min, 4 °C) to remove nuclei and large membrane and cell wall fragments. DNAase A (100 μg ml⁻¹) and RNase A (100 μg ml⁻¹) were added to the supernatant and incubated for 30 min at 37 °C. Then, a second low speed centrifugation was carried out. One volume of 15% sucrose containing 100 mM-EDTA was added to the supernatant and the whole was subjected to ultracentrifugation (150000 g, 150 min, 4 °C; Kontron T-2070 centrifuge with a T70-38 rotor).

For sucrose gradient fractionation, the C80000 pellet was resuspended in 1 ml SSB, then layered on the top of a discontinuous gradient (200000 g, 150 min, 4 °C; TST 55-5 rotor). The discontinuous sucrose gradient was preformed by adding successive 1 ml layers of 60, 45, 30 and 20% sucrose. After centrifugation, the gradient was fractionated using the Helirac 2212 (LKB). The mitochondria were obtained as a broad band in the 45% sucrose layer and shown by electron microscopy to be contaminated by large vesicles of related size (Barroso & Labarère, 1990). This large vesicles plus mitochondria sub-cellular fraction was called VM fraction.

Purification of virus particles. Viral nucleocapsids were recovered from the C80000 pellet by a 5 ml discontinuous sucrose gradient centrifugation (60-45-30-20-15% sucrose; 250000 g, 150 min, 4 °C; TST 55-5 rotor) and further purified by an isopycnic centrifugation (250000 g, 150 mins, 4 °C; TST 55-5 rotor) through a preformed CsSO₄ (15-20-30-45-60%, w/v) density gradient. The nucleocapsid band was recovered, diluted in 5 ml sterile distilled water, pelleted by centrifugation (250000 g, 150 min, 4 °C; TST 55-5 rotor) and resuspended in 100 μl sterile distilled water.

RNA polymerase assay. The standard reaction mixture contained, in a final volume of 200 μl: 100 mM-Tris/HC1 pH 7.5, 10 mM-magnesium acetate, 1 mM each of ATP, CTP, GTP and UTP, 10 μCi of [α-3²P]UTP (Amersham; 3000 Ci mmol⁻¹; 1 Ci = 37 GBq) and 100 μl of the VM fraction or viral nucleocapsids. pH 7.5 was previously established as the optimum pH for the RNA-dependent RNA polymerase activity associated with the Saccharomyces cerevisiae killer dsRNA virion (Welsh et al., 1980) or with a virus-enriched fraction of the basidiomycete Agaricus bisporus (Sriskantha et al., 1986). In the same way, the magnesium acetate concentration was equal to 10 mM, an intermediate concentration between the optimal concentrations of 5 and 15 mM for the RNA replicase activity of the S. cerevisiae and Ustilago maydis dsRNA virus (Welsh et al., 1980; Ben-Tzvi et al., 1984), and of the ‘La France’ disease virus partially purified from A. bisporus basidiospores (Sriskantha et al., 1986), respectively. Forty units of human placenta ribonuclease inhibitor and actinomycin D (100 μg ml⁻¹) were added. The RNA polymerase reaction was allowed to proceed at 37 °C, for 16 h. The reaction was stopped by lysing the cellular or viral fraction by addition of SDS to a final concentration of 2% (w/v). Each RNA polymerase assay reported in the text was performed at least in triplicate with, in each case, a cellular or viral fraction arising from independent mycelium cultures.

In some experiments, the RNA polymerase assay was carried out in the presence of DNAase I (Sigma) or RNA:se A at a final concentration of 25 μg ml⁻¹, at low (standard conditions) or high (2 × SSC; 0.3 M-NaCl, 0.03 m-sodium citrate) salt concentration. In these cases, the human placenta ribonuclease inhibitor (HPRI; Appligene) was omitted. The nucleic acid hydrolyses were eliminated by 30 min incubation at 37 °C with 50 μg proteinase K ml⁻¹, then 15 min at 56 °C in the presence of 2% SDS, followed by a phenol treatment for 15 min at 56 °C.

Extraction and analysis of nucleic acids. Nucleic acids were extracted from C80000 from VM fraction or from purified nucleocapsids, after lysis in SDS at a final concentration of 2% for 120 min at 56 °C. Then, a volume of phenol (saturated with TE buffer) was added. The resulting emulsion was centrifuged (8000 g, 10 min, 4 °C). The aqueous phase was taken and added to an equal volume of chloroform/isoamyl alcohol (24:1, v/v). The emulsion was treated as described above for the phenol emulsion. The nucleic acids were precipitated with 2 vols 95% (w/v) ethanol in the presence of 0.3 M-sodium acetate for 120 min at -20 °C, then pelleted by centrifugation and resuspended in sterile distilled water. The radioactivity of the pellet was determined using a liquid scintillation counter (Kontron).

Agarose gel electrophoresis and autoradiography. The nucleic acids were analysed in a horizontal 0.8% (w/v) agarose gel buffered in TBE (0.09 M-Tris/HCl, 0.09 m-boric acid, 0.002 M-EDTA). Nucleic acids were visualized by ethidium bromide staining (Maniatis et al., 1982). For autoradiography, the gel was incubated twice for 15 min in 10% (w/v) acetic acid then dried between sheets of Whatman 3MM chromatography paper. The dried gel was placed in a plastic bag and exposed to Kodak XAR-5 film at -80 °C with an intensifying screen (Cronex).

In vitro manipulations of nucleic acids. DNA:se A (Sigma) digestions were carried out in the presence of 10 mM-MgCl₂. RNA:se A (Sigma) digestion tests were performed in sterile distilled water or in 2 × SSC for 30 min at 37 °C. Then, to eliminate the enzyme, proteinase K was added in 2 × SSC at a final concentration of 50 μg ml⁻¹, incubated at 37 °C for another 30 min and a volume of phenol was added. The resulting emulsion was centrifuged. Nucleic acids of the aqueous phase were precipitated with 2 vols 95% ethanol, pelleted by centrifugation, resuspended in sterile distilled water and analysed by agarose gel electrophoresis.

Estimation of the size of the nucleic acids. The nucleic acid fragment sizes were estimated from the relative rate of migration in 0.8% agarose gel electrophoresis in TBE buffer by comparison with HindIII-cleaved λ DNA for DNA fragments, dsRNA segments of the maize rough dwarf fijivirus, as described previously, for dsRNAs (Barroso &
Transcription of dsRNAs in A. aegerita

Labarkre, 1990), and the 0.24-9.5 kb RNA ladder from Gibco-BRL for ssRNAs.

Results

The vegetative mycelia of the dikaryotic A. aegerita SM47, SM51 wild-type strains and of the homokaryotic progeny A1 were disrupted in 15% SSB to avoid osmotic lysis of the cellular organelles. These subcellular organelles were then separated by differential centrifugation (2500 g twice followed by ultracentrifugation at 85000 g) in the presence of DNAase I and RNAase A, as described in Methods. As reported previously (Barroso & Labarbre, 1990), an electrophoretic analysis of nucleic acids from the C5000 pellet of SM47 revealed three types of molecules (Fig. 1, lane B): (i) the high Mr dsDNA fragments resulting from mechanical breakage of the mitochondrial genome during nucleic acid extraction (Moulinier et al., 1992), (ii) the quantitatively minor encapsidated L-dsRNA (6200 bp) and (iii) the major naked M-dsRNA molecules resolved into two bands (M1 and M2) of related size (1900-1700 bp) by agarose gel electrophoresis.

Only the mtDNA fragments were seen in the C5000 pellets of the SM47 homokaryotic progeny A1 (Fig. 1, lane A), or from the dikaryotic strain SM51 (Barroso & Labarbre, 1990). No dsRNA molecule was evident in these two strains. The same negative result was obtained with both these strains using the ‘miniprep’ dsRNAs extraction procedure of Seroussi et al. (1989).

The C5000 pellets from the three strains were re-suspended in SSB and fractionated on a discontinuous sucrose gradient. Each of the three pellets was resolved as a broad band in the 45% sucrose layer which did not enter the 60% sucrose cushion. This band (VM fraction) was shown, by electron microscopy, to consist of large vesicles and mitochondria (Barroso & Labarbre, 1990). mtDNA fragments and the naked M-dsRNAs from SM47 were recovered mainly in this VM fraction. Indeed, this fraction was demonstrated previously to package naked dsRNAs in a RNAase-resistant membranous complex. In the corresponding fraction purified from the homokaryotic progeny A1, as well as from SM51, only mtDNA fragments were seen.

In addition, from extracts of SM47, the sucrose gradient fractionation allowed recovery of a second thin light-scattering band at the 20–30% sucrose interface. This band was previously shown to consist of nucleocapsids including a 6200 bp L-dsRNA genome in a RNAase-resistant complex (Barroso & Labarbre, 1990). This band was not seen in the extracts of the other two strains.

The presence of a RNA polymerizing system was investigated in the two types of RNAase-resistant complexes from the SM47 strain: the L-dsRNA including nucleocapsid and the M-dsRNAs including VM fraction. This subcellular fraction was compared with the corresponding ones, purified from the dsRNA-free SM51 and A1 strains, used in control experiments.

RNA synthesis by the M-dsRNA-packaging VM fraction

RNA synthesis activity of the A. aegerita SM47 VM fraction was investigated by electrophoretic analysis of the products incorporating [α-32P]UTP. For this, the fraction was incubated in the presence of the four nucleoside triphosphates (ATP, CTP, UTP and GTP; 1 mM each) at pH 7.5, with 10 mM-magnesium acetate and 10 μCi [α-32P]UTP. Forty units of human placenta ribonuclease inhibitor and actinomycin D at a final concentration of 100 μg ml−1 were added to inhibit cellular RNAase and transcriptional activity, respectively.

The membranes of the VM fraction were submitted to SDS (2%) treatment before and after 1, 2, 4 and 16 h incubation at 28 °C (i.e. the optimal mycelial growth temperature) under the conditions of the RNA synthesis assay. The nucleic acids were extracted and ethanol-precipitated. After 16 h incubation, the nucleic acid pellet had a specific radioactivity higher than 107 c.p.m. The nucleic acids were separated by agarose gel electrophoresis and revealed by ethidium bromide staining as well as by autoradiography of the fixed and dried
agarose gel (Fig. 2a and b). As shown in the autoradiograms, after 16 h incubation under the conditions of the RNA synthesis assay, all the nucleic acids of the fraction which were visible by ethidium bromide staining had incorporated [γ-32P]UTP. More precisely, three types of nucleic acid molecules were detected on the gels and autoradiograms: (i) a high M₃ band (H), with an apparent size > 20 kbp, (ii) two bands, M₂ and M₃, with the same electrophoretic mobility as the naked M-dsRNAs molecules, and (iii) two additional bands, S₁ and S₂, of smaller size.

The kinetics of labelling of the three types of molecules differed: a slight incorporation of [γ-32P]UTP was evident in the M and S bands after 1 h of the RNA polymerase assay. In contrast, the H band was detected on the autoradiograms only after a longer time, i.e. between 4 and 16 h incubation.

It should be noted that, during gel fixing and drying, the faint nucleic acid bands tended to spread out and, accordingly, the similarly sized M₁ and M₂, and S₁ and S₂ bands were resolved in two unique broader bands on the autoradiograms.

Identical electrophoretic patterns and autoradiograms were obtained when, not only the cellular fractionation leading to the VM fraction, but also the replicative assay was carried out in the presence of 25 μg DNAase I ml⁻¹ and 25 μg RNAase A ml⁻¹. A1 represents the autoradiogram of the nucleic acids extracted from the VM fraction purified from the control (dsRNA-free) strain A1, and subjected to a 16 h RNA polymerase assay in the absence of actinomycin D. For experimental details, see Methods.

The 32P-labelled nucleic acids extracted from the cellular fraction, after a 16 h RNA synthesis assay, were submitted to various enzymic treatments, then analysed by agarose gel electrophoresis and autoradiography (Fig. 3).

The M₁ and M₂ bands were degraded by RNAase A (5 μg ml⁻¹) in sterile distilled water, i.e. in low salt concentration (Fig. 3, lane Rl), but were unaffected by the same concentration of RNAase A in the high salt concentration buffer (2 × SSC; Fig. 3, lane Rh). Hence, the M₁ and M₂ bands corresponded to the previously described naked M-dsRNAs, of size 1900–1700 bp.

The small bands S₁ and S₂ were degraded by RNAase A in low and high salt concentrations, suggesting that they corresponded to single-stranded RNA (ssRNA) molecules (Fig. 3, lanes Rl and Rh). Accordingly, their sizes were established equal to 1900–1700 pb.

As shown in Fig. 3, the high M₃ band had the following characteristics: (i) it incorporates [γ-32P]UTP, (ii) it had the same electrophoretic mobility as the large within an RNAase-impermeable compartment, i.e. within structurally intact cellular organelles.

The 32P-labelled nucleic acids extracted from the SM47 VM fraction after 16 h incubation under RNA polymerase assay conditions (T), then submitted to various hydrolytic treatments: digestion by RNAase A (5 μg ml⁻¹) in sterile distilled water (Rl) or 2 × SSC (Rh), or digestion by DNAase I (5 μg ml⁻¹) in the presence of 10 mM-MgCl₂ (D). Nucleic acids were separated by agarose (0.8 %) gel electrophoresis before autoradiography. T' represents the autoradiogram of the nucleic acids from the SM47 VM fraction, after a 16 h RNA polymerase assay carried out in the presence of 25 μg DNAase I and RNAase A ml⁻¹. For experimental details, see Methods.
DNA fragments previously assessed as the fragments generated by mechanical breakage of the mtDNA molecule (Barroso & Labarère, 1990; Moulinier et al., 1992), and (iii) it was degraded by a DNAase I (5 µg ml⁻¹) treatment and not by RNAase A or RNAase H treatment (Fig. 3, lanes D and RI).

Despite the high sensitivity of detection of the ³²P-labelled nucleic acids, no incorporation of [α-³²P]UTP could be detected in a dsRNA molecule of a size corresponding to the viral genome H1 (6200 bp) in the VM fraction of the SM47 strain.

The RNA polymerase assay was performed also over 16 h, on the dsRNA-free VM fraction from the homokaryotic strain A1 or from the dikaryotic strain SM51, used as control strains. No incorporation of [α-³²P]UTP in nucleic acids was seen in the presence of 100 µg actinomycin D ml⁻¹. In contrast, when actinomycin D was omitted [α-³²P]UTP was incorporated in the high M, band corresponding to the mtDNA fragments and mainly in ssRNA molecules leading to a smearable signal from 5000 bp to the bottom of the gel (Fig. 3, lane A1). These results confirm the transcriptional inactivation of the cellular, more precisely mitochondrial, DNA-dependent RNA polymerase by a concentration of actinomycin D equal to 100 µg ml⁻¹. Consequently, all the dsRNA molecules seen in the VM fraction of the SM47 strain in presence of actinomycin D, seem to be imputed to naked M-dsRNA-associated RNA polymerase activity.

Search for a replicative activity associated with the isometric nucleocapsid

The isometric nucleocapsids of the virus present in A. aegerita SM47 were purified by fractionation in a discontinuous sucrose gradient in which they were resolved as a faint light-scattering band at the interface of the 20–30% sucrose layers. The band was recovered and subjected to a centrifugation in a preformed 20–60% CsSO₄ density gradient. A faint light-scattering band of nucleocapsids was obtained at the 20–30% CsSO₄ interface.

The nucleocapsids were used in a RNA synthesis assay carried out under the conditions used for the VM fraction: i.e. in the presence of the four nucleoside triphosphates, at pH 7.5, with 10 pCi [α-³²P]UTP, for 1, 2, 4 or 16 h. No incorporation of [α-³²P]UTP in nucleic acids could be detected on the autoradiograms. The same negative result was obtained when RNA synthesis activity was tested at 5, 10, 15 or 20 mM magnesium acetate. No incorporation of [α-³²P]UTP was found when the entire polymerase assay mixture was spotted on a 1 cm² Whatman 3MM filter paper and the radioactivity of the trichloroacetic acid precipitate determined, even when nucleocapsids were disrupted by sonication (5 min, 50 W; Vibra-cell, Sonics Materials) before the replicase assay.

Discussion

Our results have demonstrated that A. aegerita contains a fraction in which ribonucleotides are incorporated in dsRNA molecules, in a DNA-independent manner, i.e. in an actinomycin D-resistant mode of synthesis, at the vesicular or mitochondrial level. As full-length ssRNAs were produced, this is the first report of the presence of an effective dsRNA transcriptional system in fungal vesicles or mitochondria, although a polymerase activity was detected by [α-³²P]UTP incorporation in the vesicular RNAase-resistant complex of dsRNAs from the hypovirulent strains of the chestnut blight fungus Cryphonectria parasitica (Hansen et al., 1985). In the same way, naked dsRNA ‘plasmids’ were shown to replicate in the presence of actinomycin D in mitochondria of maize plants with S-type cytoplasm (Finnegan & Brown, 1986).

In the A. aegerita RNAase-resistant complex packaging the naked dsRNAs, analysis of the [³²P]UTP incorporation products revealed three types of molecules in the VM fraction: (i) the previously described naked dsRNA molecules, resolved in two bands (M₁ and M₂) of related size (around 2000 bp), (ii) two bands of ssRNAs (S₁ and S₂) of about 2000 bases, and (iii) ³²P-labelled high M₃ molecules.

Accordingly, our data are consistent with the presence in the membranous complex of a RNA-dependent RNA polymerase able to catalyse the transcription of the M₁ and M₂ dsRNA molecules and leading to ribonucleotide incorporation in dsRNA molecules, as described previously for the virion-associated replicase of the Penicillium stoloniferum virus S (Buck, 1975) or of the virus partially purified from ‘La France’ disease-affected A. bisporus basidiocarps (Sriskantha et al., 1986). The presence of this enzyme in the VM fraction strengthens the idea that these naked dsRNAs represent the segmented genome of a mycovirus which has co-evolved with its fungal host (Barroso & Labarère, 1990). This evolution would have maintained the replicative function in the packaging compartment of the fungal hyphae and allowed a concomitant loss of the capsid and of the viral extracellular phase.

The S₁ and S₂ ssRNAs appear as full-length transcripts of the dsRNA molecules or might be considered as replicative intermediates, i.e. they would reflect one part of the replication cycle, as described in the case of the polymerase enzymes associated with U. maydis (Ben-Tzvi et al., 1984) or S. cerevisiae virus (Welsh et al., 1980).
The incorporation of [\textsuperscript{32}P]UTP into the third type of molecules, those with high $M$, ones, is more difficult to explain, especially in the light of the fact that a DNAase I treatment is necessary to obtain degradation of the radioactively labelled product. In other basidiomycetes like *Coprinus cinereus* or *A. bisporus*, a unique DNA polymerase b has been detected which is able to polymerise dNTP on activated DNA templates or ss rRNA templates in the presence of an oligo(dG)$_{13-18}$ primer, or on dsRNAs without primer (Lu & Sakaguchi, 1984). By analogy, the [$\alpha$-\textsuperscript{32}P]UTP-labelled high $M$, molecules might be the result of polymerization of NTP on the mitochondrial dsDNA templates by the dsRNA-associated transcriptional complex. This idea is strengthened by the absence of radioactive labelling of the mtDNA fragments from the dsRNA-free strains SM51 and the homokaryotic progeny A1. In our conditions, i.e. a long time course (16 h) in presence of actinomycin D, the M-dsRNAs associated RNA polymerase would be able to use mtDNA as template and the newly synthesized ssRNAs would be used as primers for the mtDNA replication system. It is noticeable that this mtDNA replication occurred in *organello*, when the RNA polymerase activity (DNA dependent) of the VM fraction from the dsRNA-free strains SM51 or A1 was tested in the absence of actinomycin D. However, further work is needed to demonstrate that the incorporation of ribonucleotides in dsRNAs and dsDNAs is catalysed by the same enzyme. If this was the case, the actinomycin D-resistant [$\alpha$-\textsuperscript{32}P]UTP labelling of the dsRNAs and dsDNAs in presence of DNAase and RNAase, would suggest that all the templates (M-dsRNAs and large dsDNA fragments) are present with RNA polymerizing enzyme in the same intact cellular compartment. Accordingly, data would be in favour of a mitochondrial location of the M-dsRNAs molecules and of the dsRNAs-associated transcriptional complexes, as described previously for dsRNA molecules of male-sterile maize (Finnegan & Brown, 1986) or of the fungus *Ophiostoma* (*Ceratocytis*) *ulmi* (Rogers et al., 1987). It is of interest that the major mitochondrial naked dsRNAs found in maize cells or in the two fungi *O. ulmi* and *A. aegerita* show a narrow size range (between 1700 and 2900 pb).

In contrast to the results obtained with the VM fraction, no [$\alpha$-\textsuperscript{32}P]UTP incorporation was detected with partially or extensively (CsSO$_4$ gradient-) purified nucleocapsids from the SM47 strain. The absence of replicative or even transcriptional activity associated with the nucleocapsids, can be explained by two hypotheses. First, the two types of dsRNAs molecules (naked and encapsidated) are independent entities replicated by different enzymic systems and our polymerase assay, which allows the labelling of naked dsRNA molecules by the enzyme located in VM fraction, does not allow the activity of the nucleocapsid-associated replicase. Second, the naked and encapsidated dsRNAs are replicated by the same polymerase enzyme but, contrary to the results obtained with various dsRNA virions, the replicase activity requires a cellular environment or factor, available in the VM fraction but absent from the purified nucleocapsid.

The absence of a \textsuperscript{32}P-labelled dsRNA molecule of a size corresponding to the encapsidated viral genome (6200 bp) in the VM fraction, favours the first hypothesis.

Finally, the results confirm that, although the two types of RNAase-resistant complex seem to have a viral origin, the large (6200 bp) encapsidated dsRNA (possible totivirus; Francki et al., 1991) and the naked M-dsRNAs fragments constitute different viral entities which have been maintained in two different forms in the hyphae. In accordance, the nature of the different *A. aegerita* dsRNAs molecules varies greatly from that of the hypovirulent strains of the chestnut blight fungus, *C. parasitica*, where the complexity of the dsRNA population was shown to be due to defective dsRNAs from the L (12700 bp) HAV dsRNA genome (Shapira et al., 1991b). In *A. aegerita*, a primary infection would have led to the appearance and maintenance of the naked M-dsRNAs and of an associated RNA polymerase activity, in a RNAase-protecting fungal compartment (mitochondria or vesicles). However, the role of these naked M-dsRNAs is still unknown. Additionally, the presence in the strain of the isometric cryptic particle would be the result of an independent additional infection by a second mycovirus.

This work was supported by grants from the Conseil Scientifique de l'Université de Bordeaux II and the Institut National de la Recherche Agronomique.

References


Transcription of dsRNAs in A. aegerita


