Human Ro Ribonucleoprotein Particles: Characterization of Native Structure and Stable Association with the La Polypeptide

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Abstract

Anti-Ro autoantibodies, found in sera of patients with systemic lupus erythematosus, Sjogren’s syndrome, and related diseases, target the Ro ribonucleoprotein particles (RNPs). Although the polypeptide and RNA components of the Ro RNPs have been characterized, much less is known about the native structure of these particles. We have now characterized by biochemical techniques intact Ro ribonucleoprotein particles from cultured HeLa cells. These particles segregated in three discrete subpopulations with characteristic physicochemical properties: one containing hY5 RNA (Ro\textsuperscript{hY5} particles), one containing only hY4 RNA (Ro\textsuperscript{hY4} particles) and one with hY1, hY3, and hY4 RNAs (Ro\textsuperscript{hY1-hY4} particles). The Ro\textsuperscript{hY1-hY4} particles were purified free of contaminating ribonucleoproteins; both the La and the 60-kD Ro polypeptides were stable components of this portion of the Ro RNPs. The La RNPs co-purified with the Ro\textsuperscript{hY1-hY4} particles and contaminated the Ro\textsuperscript{hY5} RNPs. The stable association between the La and the 60-kD Ro polypeptides provides a potential macromolecular target for the linked set of anti-Ro and anti-La antibodies, and suggests a possible functional association of these polypeptides. (J. Clin. Investig. 1990. 85:1182–1190) autoantibodies • anti-Ro antibodies • ribonucleoproteins • Ro protein • La protein

Introduction

Autoimmune diseases such as systemic lupus erythematosus (SLE) are characterized by autoantibodies in patient sera that bind highly conserved autoantigens (reviewed in 1–3). The spectrum of this autoimmune response is often restricted in any given patient (3, 4), with certain autoantibody specificities that correlate with clinical manifestations (reviewed in 1–3). Thus many efforts have been made to define the structural or biological features of autoantigens that might nominate them as targets of autoimmunity (for examples, see 5–8).

Anti-Ro antibodies are a prominent feature of the humoral autoimmune response in many patients with SLE and with Sjogren’s syndrome (9, 10). These antibodies target small ribonucleoprotein particles (RNPs)\textsuperscript{1} that consist of a 60-kD Ro polypeptide associated with 2–4 small RNAs (11, 12). Recent evidence also indicates that a 52-kD polypeptide, and a 54-kD polypeptide restricted to erythrocytes, are complexed with the Ro RNAs (13, 14). The number of RNAs differs among mammalian cells: for example, human cells contain four Ro RNAs, called hY1, hY3, hY4, and hY5, and murine cells contain two such RNAs, mY1 and mY2 (15). These RNAs are RNA polymerase III transcription products, and as such, are at least transiently associated with the La polypeptide (11). The 60-kD Ro polypeptide binds its Ro RNA at the base of a double-stranded stem formed by base-pairing of 5’ and 3’ ends (12); the binding site of the La polypeptide is believed to be at least partially constituted by a poly (U) stretch at the 3’ end of the RNA that is processed to form the mature Ro RNA transcript with release of this polypeptide (16–18). The RNA and/or protein binding sites for the 52- and 54-kD Ro polypeptides have yet to be defined.

Indirect evidence indicates that one 60-kD Ro polypeptide binds to one Ro RNA (12), implying the existence of distinct populations of Ro RNPs. We recently reported the frequent occurrence in patient sera of antibodies that targeted an antigenic epitope restricted to intact Ro particles containing the 60-kD Ro polypeptide and the hY5 RNA (Ro\textsuperscript{hY5} RNPs) (19). The production of antibodies to a specific Ro RNA suggested structural heterogeneity of these particles; moreover, the production of antibodies that only recognized intact Ro\textsuperscript{hY5} particles indicated that native Ro RNPs might serve as autoimmunogens, in addition to (or rather than) isolated Ro polypeptides.

Thus, to determine if Ro particles are indeed physically heterogeneous, and to identify structural features that could possibly designate a subset of these native Ro particles as targets of a specific autoimmune response, we have biochemically purified intact human Ro particles containing the 60-kD Ro polypeptide and the Ro RNAs. These particles segregated in three subpopulations with characteristic physicochemical properties. The La polypeptide, frequently targeted by the autoimmune response in conjunction with the Ro polypeptides, was stably associated with a portion of these purified particles.

Methods

Cells and sera. Human HeLa cells, initially obtained from American Type Culture Collection (Rockville, MD), were maintained at 37°C under 5% CO\textsubscript{2}, in RPMI 1640 (Gibco Laboratories, Grand Island, NY) supplemented with 10% heat-inactivated fetal calf serum, 60 μg of penicillin/ml and 100 μg of streptomycin/ml. Sera were obtained from healthy laboratory workers and from American and French Canadian patients with various connective tissue diseases. Control anti-Ro sera

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1. Abbreviations used in this paper: RNP, ribonucleoprotein particles; TBS, Tris-buffered saline; TSE, Tris-Cl, NaCl, and EDTA.
were defined as those which immunoprecipitated the 60-kD Ro polypeptide and all four hY RNAs from radiolabeled HeLa cell extracts (19), while anti-Ro sera immunoprecipitated only the 60-kD polypeptide and the hY5 RNA from identical extracts (19). The anti-Ro sera used did not immunoprecipitate the La RNAs from HeLa cell extracts nor recognize the 50-kD La polypeptide in immunoblots. As previously reported (12), a protein that comigrated with the La polypeptide was faintly visible in [35S]methionine labeled anti-Ro immunoprecipitates. None of the sera used in these studies immunoprecipitated the 52-kD Ro polypeptide from HeLa cell extracts, nor recognized this protein in immunoblots of these extracts (13).

Preparation and immunoprecipitation of cell extracts. For analysis of in vivo labeled RNAs, HeLa cells were radiolabeled with [32P]orthophosphate (10 μCi/ml cells; Amersham Corp., Arlington Heights, IL) for 14 h as previously described (20). Cells were collected by centrifugation, washed in TBS (10 mM Tris-Cl [7.5], 150 mM NaCl), and sonicated in NET-2 buffer (50 mM Tris-Cl [7.5], 150 mM NaCl, 0.05% Nonidet P-40). Immunoprecipitation of radiolabeled and unlabeled cell extracts was performed as described (19). Immunoprecipitated 3P-labeled nucleic acids were visualized by autoradiography on XRP film (Eastman Kodak, Rochester, NY) while the unlabeled RNAs were visualized by silver staining (21). For analysis of proteins, HeLa cells were labeled with [35S]methionine (10 μCi/ml cells; Amersham Corp.) and immunoprecipitations performed as described (19).

In experiments to assay the stability of the association of hY RNAs with the 60-kD Ro polypeptide, 4 M NaCl was added to parallel samples of the HeLa cell sonicates to increase their salt concentration to 0.25, 0.35, 0.50, 0.75, and 1.0 M, respectively. After rotation for 1 h at 4°C, these samples with high salt concentrations were added to antibody-coated Sepharose beads resuspended in NET-2 buffer that had been previously modified to contain the same NaCl concentrations as the sonicates.

Biochemical purification of the Ro particles. The first steps of the purification of the Ro particles were performed as described (19), with modifications. In brief, 1–2 × 10⁹ HeLa cells were collected by centrifugation and washed twice in chilled TBS. Subsequently all procedures were performed at 4°C and all buffers were supplemented with 1 mM diithiothreitol and 1 mM phenylmethylsulfonyl fluoride. A cytoplasmic fraction was prepared without detergent: cells were allowed to swell in six pellet-volumes of buffer A (10 mM Tris-Cl [7.5], 1.5 mM MgCl₂, 10 mM KCl) for 10 min on ice, spun down, resuspended in two pellet-volumes of buffer A and disrupted by 10 strokes of an all-glass Dounce homogenizer; 100 U of RNAse (Promega Biotec, Madison, WI) were added after disruption of the cells. Extracts were then clarified by centrifugation at 13,000 g for 20 min. The salt concentration of the supernatant was increased to 140 mM by adding 0.1 vol of buffer B (0.3 M Tris-Cl [7.5], 1.4 M NaCl, 1.5 mM MgCl₂). The extract was then layered over a 7.5-mL sucrose cushion (800 mM sucrose, 10 mM Tris-Cl [7.5], 5 mM MgCl₂) in 30-mL tubes and centrifuged at 26,000 rpm in a SW28 rotor (Beckman Instruments, Inc., Palo Alto, CA) for 90 min. The supernatant was collected and diluted 1:1 with TSE (50 mM Tris-Cl [7.5], 150 mM NaCl, 1 mM EDTA) and applied to a 40-mL column of DE52 (Whatman, Maidstone, Kent, England) pre-equilibrated in TSE. After extensive washings with TSE, step elutions of the column using TSE containing 175, 190, 210, 250, and 300 mM NaCl were performed. After addition of 40 U of RNAse, each eluate was separately concentrated by vacuum dialysis (Pro-Di-Con; Bio-Molecular Dynamics, Beaverton, OR) against TSE. Elution profiles of the Ro RNPs in each of the eluted fractions were determined by immunoprecipitation with control anti-Ro sera.

The concentrated DE52 eluates (0.5 ml) containing the different Ro particles were each layered on a 10–30% (wt/wt) sucrose gradient (10.5 ml) and centrifuged at 35,000 rpm in an SW41 rotor (Beckman Instruments, Inc.) for 18 h. After centrifugation, serial 0.5-ml fractions were collected from the bottom of the tube and assayed by immunoprecipitation with the sera of Ro particles. The fractions containing the peaks of the Ro RNPs were further fractionated by HPLC in TSE at room temperature on a gel permeation column (TSK G3000 SW 600 × 7.5 mm; Phenomenex, Rancho Palos Verdes, CA) at a flow rate of 0.5 ml/min (260 ps). Molecular weight standards included thyroglobulin (669 kD), ferritin (440 kD), catalase (232 kD), aldolase (158 kD), bovine serum albumin (67 kD), chymotrypsinogen A (25 kD), and uridine (0.244 kD). The eluates were collected in 250-μl fractions and assayed by immunoprecipitation for the presence of Ro particles.

SDS-polyacrylamide gel electrophoresis and immunoblotting. Proteins present in extracts at each step of the purification were either concentrated in Centricon-30 cones (Amicon Corp., Danvers, MA), or precipitated in 25% TCA, spun down by centrifugation at 12,000 g for 15 min, the pellet washed twice in cold acetone, and dried. Proteins were then dissolved in SDS-sample buffer (63 mM Tris-Cl [6.8], 2% SDS, 2% mercaptoethanol, 20% glycerol, 0.004% bromophenol blue) before fractionation in SDS-7.5% polyacrylamide gels (22) (acrylamide/bis, 30:0.8). 4 μg per lane of 60-kD Ro polypeptides affinity-purified from human placenta (kindly provided by Dr. M. Mamula, Yale University School of Medicine) were fractionated as a positive control on each gel. After transfer to nitrocellulose (Schleicher and Schuell, Keene, NH) in 25% methanol, Tris-glycine [8.3] (23), the nitrocellulose sheets were blocked with 3% BSA in TBS overnight and subsequently incubated with human antisera diluted 1:50 in TBS containing 0.1% Tween 20 and 1.0% BSA. Bound antibodies were detected with 32P-labeled protein A (1 × 10⁶ cpm/ml; ICN, Irvine, CA) followed by autoradiography on XRP film (Kodak).

Affinity purification of antibodies. Anti-La antibodies were affinity-purified (24) from a serum containing high titers of anti-La antibodies and a paucity of anti-Ro antibodies, as determined by immunoblotting assays. The La polypeptide from a sonicate of total HeLa cells was localized on immunoblots; the corresponding area was then cut out, incubated with the anti-La serum, and extensively washed. The bound antibodies were eluted with 0.2 M glycine-HCl [2.5], neutralized with 2 M Tris-Cl [7.5], and assayed by immunoprecipitation. Control antibodies eluted from an unrelated strip of the nitrocellulose sheet lacked anti-La activity in the 32P immunoprecipitation assay (19, 20).

Results

The Ro RNAs dissociate from the Ro particles at high ionic strength. To define the conditions of purification compatible with isolation of native Ro particles, we verified the stability of the Ro particles in buffers of increasing ionic strength. As shown in Fig. 1 A, relatively low salt concentrations dissociated the Ro RNAs from the 60-kD Ro polypeptides. Anti-Ro sera specific for the 60-kD polypeptide could not immunoprecipitate Ro RNAs from extracts containing 500 mM NaCl (Fig. 1 A, compare lanes 3 and 6). This did not result from the loss of binding of anti-Ro antibodies to the polypeptide itself, since the intensity of the 60-kD band immunoprecipitated from [35S]methionine labeled cell extracts did not decrease in buffers containing up to 1 M NaCl (Fig. 1 B; compare lane 2 to lane 3). The sensitivity to dissociation in buffers of high ionic strength varied among the Ro RNPs: hY3 and hY4 were totally dissociated from Ro polypeptides in 350 mM NaCl, while hY1 and hY5 were only partially dissociated in the same salt concentration (Fig. 1 A, lane 4). The presence of low millimolar concentrations of EDTA in the buffers appeared to slightly stabilize the Ro RNA-protein bond (data not shown), thus all buffers used in the purification procedure contained 1 mM EDTA.

Dissociation in these salt concentrations is a characteristic of Ro particles, since the immunoprecipitation of the La RNAs by anti-La sera was not affected in up to 750 mM NaCl (Fig. 1 A; compare lane 7 to lane 8). It is interesting that under conditions where the Ro RNAs have dissociated from the
60-kD Ro polypeptide, an anti-La serum still immunoprecipitated an RNA co-migrating with hY5; however, bands corresponding to hY1 and hY2 (a breakdown product of hY1) have disappeared (Fig. 1A; compare lane 7 to lane 8).

Three subpopulations of Ro particles can be biochemically purified from HeLa cells. As reported by Kato et al. (25), more than 90% of the Ro particles were recovered in the cytoplasmic fraction of HeLa cells prepared by gentle cell disruption, even without detergent (data not shown). The cytoplasmic fraction was then loaded on a DE52 anion exchange column, washed extensively, and the bound material eluted stepwise with buffers of increasing ionic strength. As illustrated in Fig. 2, and as previously reported (19), the Ro hY3 particles were present in both the 175 mM and 190 mM NaCl eluates (lanes 3 and 4, lanes 10 and 11), a group of particles containing all four Ro RNAs (Ro hY5 particles) eluted between 210 mM and 250 mM NaCl (lanes 6 and 13), and Ro particles containing hY1, hY3, and hY4 RNAs (Ro hY1-hY4 particles), but nearly devoid of any hY5 RNA, eluted between 250 and 300 mM NaCl (lanes 7 and 14). The Ro RNAs in each of these fractions were immunoprecipitated with an anti-Ro serum that bound the 60-kD Ro polypeptide (lanes 10–14), indicating that this polypeptide was complexed with its cognate RNAs. Contaminating tRNA and 5S RNA were present in each of the three fractions; however, it was noteworthy that the hY5 RNA and the La RNAs constituted major RNA species in the 175 and 190 mM and in the 210 to 250 mM NaCl eluates, respectively (Fig. 2, lanes 3 and 4, and lane 6). From these elution experiments, it was evident that Ro particles existed as at least two populations: the Ro hY3 RNPs, and the Ro hY1, hY3, hY4 RNPs. The existence of a second population of particles, one that contained all four Ro RNPs (Ro hY1-hY4) and that eluted between 210 and 250 mM NaCl could simply result from a contamination with each of the other two populations; alternatively, it could represent a group of Ro particles containing the Ro RNAs in approximately equimolar ratios.

After concentration to ~ 500 μl, the three eluates were loaded separately onto 10–30% (w:w) sucrose gradients and fractionated by rate zonal ultracentrifugation. The Ro hY5 RNPs, eluted at 175 mM NaCl from DE52, sedimented as a single peak where the major contaminating RNA was SS RNA (Fig. 3A, lane 6); in the peak fraction, SS RNA was not immunoprecipitable with anti-La antibodies (data not shown). In the 250 mM NaCl DE52 eluate, the Ro RNPs appeared to sediment slightly ahead of the La RNPs (Fig. 3B, compare lanes 5 and 6 to lanes 7 and 8). This observation was confirmed by immunoprecipitations of individual gradient fractions with an anti-Ro serum and with an anti-La serum as
shown in Fig. 3 C (note that the peak of immunoprecipitable Ro RNPs was in fractions 14 and 15, whereas the peak of immunoprecipitable La RNPs was in fractions 16 and 17). A population of RoY4 RNPs did sediment with the peak of the La RNPs (Fig. 3 C, lanes 4 and 9), however, indicating that RoY4 particles could represent a third discrete subpopulation of Ro RNPs, in addition to RoY5 RNPs and a population that appeared to contain the hY1-Y4 RNAs (RoY1-Y4 RNPs). The displacement of the RoY4 particles from the other Ro RNPs was not secondary to in vitro dissociation of the hY4 RNA from the antigenic 60-kD Ro polypeptide since these RNPs could be immunoprecipitated by a serum that targeted the 60-kD protein.

The peaks of Ro particles from the sucrose gradients were further fractionated by gel filtration HPLC. Again, RoY4 particles behaved differently from the other Ro RNPs (Fig. 4). While the Stokes' radius of the RoY5 particles and the RoY1-Y4 particles appeared compatible with the elution profile of a globular protein of 300-350 kD, the RoY4 particles eluted at ~230 kD (Fig. 4) with the bulk of the La particles (data not shown), consistent with their cosedimentation in sucrose gradients. After HPLC purification of the RoY5 RNP fraction from sucrose gradients, hY5 RNA represented the predominant RNA visible in silver-stained gels of phenol/chloroform extracts of the peak eluates (Fig. 5 A, lanes 4 and 5), although several polypeptides were present in these fractions (Fig. 5 B, lanes 5 and 6). The peak fractions of the RoY1-Y4 particles and of the RoY5 particles were still heavily contaminated with La RNPs; in these fractions, the Ro and La RNAs were not degraded (data not shown).

Characterization of the purified Ro particles. After each purification step, the Ro RNAs were immunoprecipitated with anti-Ro sera specific for the 60-kD Ro polypeptide. This implied that this protein remained bound to the immunoprecipitated Ro RNAs through the final stages of purification. To investigate this possibility, we performed immunoblots using anti-Ro, as well as anti-La sera, of the peaks of the HPLC eluates containing the RoY5, the RoY1-Y4, and the RoY4 RNPs, respectively. Both the 60-kD Ro polypeptide (Fig. 6, top, lane 4) and the 50 kilodalton La polypeptide (Fig. 6, bottom, lane 4) were present in the peak HPLC fractions containing the RoY4 RNPs, with the quantities of these proteins paralleling the peak of the RoY4 particles (compare Fig. 5 A, lanes 5 and 7 to Fig. 6, lanes 4 and 5). These fractions lacked contaminating La RNAs, suggesting that the La polypeptide was stably associated with the RoY4 particles. As expected, both the La and the 60-kD Ro polypeptides in the HPLC eluates containing the peak fractions of the RoY1-Y4 and the RoY4 RNPs (data not shown), since these fractions contained both Ro and La RNAs.

Of note, the La polypeptide in the HPLC eluates was predominantly composed of the intact 50 kilodalton molecule, with minimal degradation (Fig. 6, bottom, lanes 1, 2, 4, and 5), compared to degradation of contaminating La protein in the immunoaffinity-purified Ro polypeptide preparation (Fig. 6, bottom, lane 3). Similar results were obtained with immunoblots of the HPLC eluates containing the peak fractions of the RoY1-Y4 and the RoY4 RNPs (data not shown). Since the La polypeptide is highly sensitive to protease (26), this indicated that the Ro particles purified under our conditions were not substantially degraded.

To investigate more precisely the protein constituents of the HPLC-purified, intact RoY5 particles, we immunoprecipitated the peak fraction using antibodies specific for the 60-kD Ro...
Figure 3. Sucrose gradient rate zonal ultracentrifugation of Ro RNPs. (A) Total RNAs present in successive fractions of the gradient of the 175 mM DE52 eluate. Lane 1 shows total RNAs present in the 175 mM eluate from DE52, and lane 2 shows an immunoprecipitate of a HeLa cytoplasmic extract using a control anti-Ro serum to help identify the Ro RNAs. Successive fractions (1–22) were collected from the bottom of the tube; total RNAs present in representative fractions are shown. (B) Total RNAs present in successive fractions of the gradient of the 250 mM DE52 eluate. Lane 1 shows the immunoprecipitated control La RNAs and lane 2 shows immunoprecipitated control Ro RNAs, both from a cytoplasmic extract. Successive fractions were collected from the bottom of the tube. (C) Immunoprecipitates of the fractions of a gradient of the 250 mM DE52 eluate using control anti-Ro (lanes 2–6) and anti-La (lanes 7–11) sera. The fractions correspond to those in B.

Figure 4. HPLC gel filtration of the fractions from the sucrose gradients containing the peaks of Ro particles. The elution profile of molecular weight standards is shown (BSA, bovine serum albumin). The abscissa shows fraction number (see Methods for details). The elution profiles of the RoY5, RoY4, and the RoY1–Y4 particles are shown on the ordinate as – to ++, where – is undetectable by immunoprecipitation, + is barely detectable, ++ is clearly detectable, and +++ is the maximum intensity. Blank spaces indicate that a particular fraction was not tested by immunoprecipitation.

Ro polypeptide and the La polypeptide, respectively; antibodies that selectively immunoprecipitated the RoY5 particle from crude cell extracts (19) were also used in these experiments. As shown in Fig. 7, lanes 2, 3, and 5, these three antibodies immunoprecipitated the RoY5 particle. Since the RoY1–Y4 and the RoY4 fractions were contaminated with La RNPs, a similar immunoprecipitation of these purified particles was not attempted.

Discussion

We have partially purified Ro particles from HeLa cells that appear immunologically and biochemically intact, without significant degradation of the RNA or protein components. These particles likely exist as distinct populations of RNPs, since they partitioned into three subgroups, RoY1–Y4, RoY4, and RoY5, with at least the latter fraction being stably associated with the La polypeptide.

Several lines of evidence indicate that the Ro RNPs purified during these studies were intact. First, the RNA components of the purified RNPs did not degrade during the purification, since their electrophoretic mobility remained constant and identical to that of freshly immunoprecipitated Ro RNAs. Second, immunoblots of the final HPLC eluates indicated that the 60-kD Ro polypeptide, as well as the protease-sensitive La polypeptide, were largely intact. Thirdly, the autoantigenic
Figure 5. HPLC purified RoVY5 particles. (A) Total RNAs present in successive fractions from the HPLC gel filtration column containing the purified RoVY5 particles. Lane 1 shows total RNAs present in cytoplasmic extracts and lane 2 shows the control Ro RNAs immunoprecipitated from these extracts. The fractions were numbered from the first fraction eluted after the dead volume of the column. Lane 10 shows the RNAs immunoprecipitated from fraction 10 using a control anti-Ro serum. (B) Silver-stained SDS-polyacrylamide gel showing the proteins present in the 175 mM NaCl eluate from DE52 column (lane 2), the peak RoVY5 fraction in the sucrose gradient (lane 4), and the peak RoVY5 fraction in HPLC gel filtration (lanes 5–7). Extracts loaded in each lane were derived from approximately equal numbers of cells. Lane 3 illustrates 60 kD Ro polypeptides affinity purified from human placenta.

60-kD Ro polypeptide remained complexed with its RNAs throughout the steps of purification, since Ro RNAs could be immunoprecipitated with anti-Ro sera; similarly, the La polypeptide remained stably associated with the RoVY5 particle since it was immunoprecipitable with an affinity-purified anti-La serum. Finally, the conformation-dependent antigenic determinant specific to the RoVY5 particle was still intact since these particles could be immunoprecipitated with an anti-RoVY5 serum.

Previous studies have concentrated upon purification of the protein component of the Ro RNPs, either biochemically (27–29), or by affinity purification (30); in both cases, the conditions of purification made it unlikely that intact ribonucleoprotein particles could be purified. The recent identification of anti-Ro antibodies, that are apparently restricted to intact RoVY5 particles (19), as well as the discovery of other autoantibodies that target conformational epitopes (31, 32), indicates that isolated RNPs containing their cognate RNAs will be useful in further analysis of the humoral autoimmune response.

In extension of our previous immunological and biochemical observations suggesting that the Ro particles are heterogeneous (19), sensitivity to salt disruption was uneven among these RNPs: the RoVY4 and RoVY5 RNAs began to dissociate from their autoantigenic polypeptide at 250 mM NaCl, with the RoVY1 and RoVY5 RNPs being somewhat more resistant to disruption in high salt. Binding of all four Ro RNAs to the 60-kD Ro polypeptide was totally disrupted in buffers of rela-
During purification, since three populations of these RNPs were obtained that differed in discrete ionic strength, the binding of the La RNAs to the La protein did not seem restricted to the particles from the peak of the HPLC gel filtration column (19) (lane 2), or the 60-kD Ro polypeptide (lane 3); immunoprecipitates were also formed with a normal human serum (lane 4) and with affinity-purified anti-La antibodies (lane 5) (24). Lane 1 shows the control Ro RNAs immunoprecipitated from a HeLa cytoplasmic extract using the same anti-Ro serum shown in lane 3.

The first population containing the hY5 RNA, clearly hY5, was enriched in RNAs (RoY5) particles (19) (lane 2), or the 60-kD Ro poly-peptide (lane 3); immunoprecipitates were also formed with a normal human serum (lane 4) and with affinity-purified anti-La antibodies (lane 5) (24). Lane 1 shows the control Ro RNAs immunoprecipitated from a HeLa cytoplasmic extract using the same anti-Ro serum shown in lane 3.

Heterogeneity of Ro particles was further substantiated during purification, since three populations of these RNPs were obtained that differed in discrete biochemical properties. The first population containing the hY5 RNA (RoY5 particles) eluted from an anion-exchanger at low salt, sedimented in sucrose gradients somewhat faster than the La RNPs, and had a Stokes' radius compatible with a globular protein of 300 to 350 kD. A second one enriched in RoY4 RNPs eluted from DE52 at higher salt, sedimented in sucrose gradients with the La RNPs, and had an apparent molecular weight of ~230 kD. Finally, a group of particles containing the hY1, hY3 and hY4 RNAs (RoY1-3Y4 RNPs) eluted at an intermediate salt concentration, sedimented in sucrose gradients ahead of the La RNPs, and had an apparent molecular weight of 300 to 350 kD. At present we are unsure if this latter group of particles consist of individual Ro RNPs with similar physical properties, or if they exist as a complex.

Although others have previously reported that Ro particles fractionated with an apparent molecular weight of 100–150 kD in gel filtration (30, 35), the purification procedures used in these earlier studies made it unlikely that the sized particles were intact. In the current study, Ro particles had unexpectedly high Stokes' radii corresponding to globular proteins of 230 and 300–350 kD. Molecular weight determinations of RNPs by gel filtration are imprecise, however, at least in part because these particles in solution may not behave as globular proteins (36). For example, the 7S RNP, comprised of one copy each of the 5S RNA and a 34-kD polypeptide with a combined molecular weight of 73 kD, has a Stokes' radius by gel filtration HPLC corresponding to a globular protein of 120 kD. On the other hand, the actual molecular weight of one 60-kD Ro polypeptide plus one Ro RNA (27 kD for hY5, and 37 kD for hY1) is similar to that of the 7S RNP, yet the Stokes' radius is much greater for Ro RNPs; this observation at least suggests the possibility that Ro particles or their individual components exist as multimers, or that they contain polypeptide components in addition to the 60-kD protein.

Although we did not address all the possibilities that could account for the observed size of Ro RNPs, we did show that the RoY5 RNPs contained the La and the 60-kD Ro polypeptides since antibodies specific for these proteins immunoprecipitated the purified RoY5 particles. Moreover, in the HPLC purified RoY5 fractions, immunoblots demonstrated the presence of both the La and the 60-kD Ro polypeptides; since immunoprecipitates of these fractions lacked the La RNAs, it did not seem likely that contamination with La RNPs accounted for the presence of the La polypeptide. Further, since the HPLC fraction that was enriched in RoY5 particles eluted at 300–350 kD, it is unlikely that free La polypeptide specifically contaminated this fraction and accounted for the immunoblot results (Fig. 6, bottom). Thus, the immunoprecipitation and immunoblot experiments indicate that both polypeptides are capable of associating with the hY5 RNA and are likely stably complexed with this RNA, at least under the conditions utilized in this study. To account for the 300-kD molecular weight of the RoY5 RNPs, however, these particles may exist as dimers. At present, we do not know if the newly described 52 kD Ro polypeptide (13) is also a constituent of these purified RNPs, as the sera we used as probes in these studies only targeted the La and the 60-kD Ro proteins.

It is unclear if the other populations of Ro RNPs (RoY1-4Y4 and RoY4 RNPs) are stably associated with the La polypeptide. These fractions were contaminated with the La RNPs and immunoprecipitation and immunoblotting experiments with anti-La antibodies were obscured by this contamination; however, the size of these particles (> 220 kD) suggest that they are composed of more than one copy of the 60-kD Ro polypeptide and more than one 30-kD RNA.

Certain observations suggest that the La polypeptide may have a unique relationship with RoY5 RNPs. For example, under conditions where all Ro RNAs were dissociated from the 60-kD Ro polypeptides (Fig. 1 A, lane 2), anti-La antibodies still immunoprecipitated a fraction of the hY5 RNAs, but none of the hY1 and hY2 RNAs (Fig. 1 A, lane 7). Similarly, we also have observed that immunoprecipitates of total HeLa cell extracts using affinity-purified anti-La antibodies show a clear hY5 band but no hY1 or hY2 RNAs (hY3 and hY4 are obscured by the very abundant La RNAs).
It is significant that among all mature, processed RNA polymerase III transcripts, Ro RNAs (or a portion of these RNAs such as hY5) are the only ones stably associated with the La polypeptide, and that these RNAs are the RNA polymerase III transcripts still produced in cells grown under adverse conditions (11). It thus follows that the function of Ro RNPs may be related to that of the La polypeptide, which is thought to serve as a terminator of RNA polymerase III transcription (17, 18). Ro RNPs could serve as a cofactor that modulates La activity, or as a “sink” to help recycle used La polypeptides for another round of transcription termination. Alternatively, Ro RNPs bearing the La polypeptide could contain incompletely processed Ro RNAs (17, 18); however, this latter possibility appears less likely since no difference in mobility of the hY5 RNA can be detected when RoY5 RNPs are immunoprecipitated with either anti-Ro or anti-La antibodies.

Antibodies specific for the 60-kD Ro and the La polypeptides frequently occur in tandem, or as a linked set (4, 10). The close proximity and the stable association of these two polypeptides on at least a subset of the Ro RNAs provide a potential macromolecular target for this linked autoantibody set, and thus favors the hypothesis that autoantigens drive immune responses in SLE and related illnesses (4).

In conclusion, both protein and RNA components have been shown to be essential for complete function of certain other RNP autoantigens (37-39). Thus, intact Ro particles such as those obtained in these experiments may represent reagents to study the activity of this RNP, either alone or in conjunction with the La polypeptide, perhaps in the regulation of transcription by RNA polymerase III.

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