Comparison of the Proteins and Polypeptides of the Eight Serotypes of *Ureaplasma urealyticum* by Isoelectric Focusing and Sodium Dodecyl Sulfate-Polyacrylamide Gel Electrophoresis

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Analysis of representative strains of the eight human serotypes of *Ureaplasma urealyticum* by polyacrylamide gel electrophoresis identified 36 to 40 polypeptides for each strain. At least 80% of the peptides were common among strains, but unique major peptides were identifiable in ureaplasmic types. Type 1 had a polypeptide of 85,000 daltons, type 3 had a polypeptide of 72,000 daltons, type 5 had a polypeptide of 64,000 daltons, and type 8 had a polypeptide of 95,000 daltons. The unique polypeptides in types 1 and 8 were identified as membrane components. Two common major components of 44,000 and 70,000 daltons were observed. Several components were common to some, but not all, serotypes. Patterns obtained from *U. urealyticum* strains were strikingly different from the patterns of *Acholeplasma laidlawii*, *Mycoplasma gallisepticum*, *Mycoplasma arginini*, and *Mycoplasma hominis*. Isoelectric focusing demonstrated a unique membrane protein for type 1 at pH 6.4, whereas type 8 possessed an assembly of five unique proteins at pH 7.0. Ureaplasmata were strongly similar to each other by isoelectric focusing, but strikingly different from members of the other genera studied. Although a filtered, strongly buffered dialysate medium with 1% serum and 30 mM urea was used both to maximize yields and to minimize contamination, minor contaminants were detected, which comigrated with horse transferrin (pH 6.0) and cytochrome c (molecular weight, 14,000). The similarities of the polypeptide patterns of *U. urealyticum* strains affirm their close relationships to each other, in contrast to the diversity shown in the genus *Mycoplasma*, and our recognition of type-specific membrane peptides will enhance the identification of serotypes and the classification of strains.

The "T" *Mycoplasma* strains, isolated from humans, have been classified as a single species, *Ureaplasma urealyticum*, with eight serotypes (23). The basis for differentiation of the serotypes was the fact that strains of *U. urealyticum* can be typed reasonably specifically by the metabolic inhibition test or by growth inhibition on agar. However, the metabolic inhibition test shows much greater species specificity than type specificity in the differentiation of *Mycoplasma* species. Thus, we really do not know whether types of *U. urealyticum* are as similar to each other as strains of a classic *Mycoplasma* species or whether the relationship between types is more like that observed between *Mycoplasma* species which are serologically related (7).

The identification of *Mycoplasma* species and the determination of the relationships among strains of a species and of related species have been carried out by polyacrylamide gel electrophoresis; patterns from related organisms are quite similar, and those from unrelated organisms are markedly dissimilar (17). The purpose of the present study was to compare the polypeptides and proteins of representative strains of the eight serotypes of *U. urealyticum* by examining both molecular weights (sodium dodecyl sulfate [SDS]-polyacrylamide gel electrophoresis) and isoelectric points (isoelectric focusing). Comparisons were also made with strains of *Mycoplasma* and *Acholeplasma* species, and extensive control measures for monitoring contamination with medium components were carried out to verify the results.

MATERIALS AND METHODS

Mollicutes. The strains obtained from the American Type Culture Collection, Rockville, Md., for use in this study were *Acholeplasma laidlawii* ATCC
14192, Mycoplasma gallisepticum ATCC 15302, Mycoplasma hominis ATCC 14027, U. urealyticum type 1 ATCC 27813, type 2 ATCC 27814, type 3 ATCC 27815, type 4 ATCC 27816, type 5 ATCC 27817, type 6 ATCC 27818, and type 7 ATCC 27819. U. urealyticum type 8 as 960 (cl × 8) and Mycoplasma arginini strain G-230 were kindly supplied by M. C. Shepard and M. F. Barile, respectively.

**Stock solutions and medium supplements.** The following solutions were sterilized by filtration through a membrane filter (pore diameter, 0.22 μm): 1 M 2-(N-morpholino)ethanesulfonic acid (MES), pH 6.0; 1 M N-tris(hydroxymethyl)methyl-2-aminoethanesulfonic acid (TES) pH 7.3; 1 M sodium sulfite; 0.5 M dithioerythritol; and 1 M urea. Agama horse serum (Flow Laboratories, Rockville, Md.) was refiltered to ensure removal of insoluble serum precipitates. Phenol red (sodium salt; 1%); phosphate-buffered saline (pH 7.1; lacking Mg<sup>2+</sup> and Ca<sup>2+</sup> [3]), and TES-saline (5 mM TES plus 150 mM NaCl [6]) were sterilized by autoclaving.

**Culture media.** Soy peptone-fresh yeast dialsyate medium (5, 8) was sterilized by filtration rather than by autoclaving. The routine culture medium used contained dialysate broth supplemented with 5% agama horse serum, 30 mM MES buffer, 10 mM urea, 100 U of penicillin per ml, and 0.001% phenol red (final pH, 6.1). Culture methods were as described previously (8, 20).

The mass cultivation medium for U. urealyticum strains contained dialysate broth supplemented with 1% (vol/vol) agama horse serum, 90 mM MES, 30 mM urea, 1 mM fresh sodium sulfite (reducing agent [21]), and 100 U of penicillin per ml. An actively growing culture (which had just shown color change on the indicator) was diluted 1:1,000 into 10 to 12 liters of growth medium (approximately 2 × 10<sup>9</sup> colony-forming units per ml) and incubated aerobically at 37°C for 28 to 36 h (final yield, ~6 × 10<sup>9</sup> colony-forming units or 2.25 μg of protein per ml). Acholeplasma and Mycoplasma strains were grown in dialysate broth containing 10% agama horse serum, 100 U of penicillin per ml, and 20 mM TES at pH 7.3. Cultures in late log phase were centrifuged at 12,000 × g for 40 min. The sediment was washed three times with a total of 2 liters of TES-saline. Cells were suspended in 5 ml of sterile distilled water, freeze-dried, and stored at −70°C. For analysis, cells were reconstituted with sterile water (140 μg of protein per ml) and sonicated at 4°C for a total of 2 min per ml of suspension. Bovine serum albumin was used as the standard for protein determination (10).

**Optimal conditions for lysis of organisms for membrane preparation.** Two methods of lysis were tested, alkaline (carbonate lysis [4]) and saponin (digitonin lysis [16]). Tissue-culture monolayers were suspended in 0.15 M NaCl at 37°C, and 0.1 ml of the suspension was mixed with 2.5 ml of an appropriate buffer or digitonin at a concentration sufficient to give an optical density of 0.3 to 0.35 at 500 nm. For alkaline lysis, the following buffer solutions were prepared at 50 mM concentrations in 50 mM NaCl: citric acid-sodium citrate, pH 3.0, 4.0, and 5.0; MES, pH 6.0, TES, pH 7.0 and 8.0; and carbonate-bicarbonate at a range of pH 9 to 11. Digitonin (98% pure; BDH Chemicals, Ltd.; obtained from Gallard Schlessinger Corp., Carle Place, N.Y.) was dissolved in 0.25 M NaCl, and serial twofold dilutions were prepared in 0.25 M NaCl from an 80 μM stock solution. The lytic abilities of the various buffers and digitonin were tested by adding 0.1 ml of cell suspension at 37°C to 2.5 ml of agent also at 37°C. The optical density of the cell suspension was measured at intervals at 500 nM. Percent lysis was calculated by dividing the observed density by the control value.

**Preparation of membrane and cytoplasmic fractions.** Cells were lysed by using either carbonate-bicarbonate buffer or digitonin. Washed cells (5 ml containing a total of 30 mg of cell protein) were suspended in a 200-ml volume of 50 mM carbonate-bicarbonate buffer in 50 mM NaCl at pH 10.2 and 37°C and incubated for 10 min; the pH was adjusted to 7.0 by adding 10 ml of 1 M MES buffer, pH 5.5. The viscosity of the suspension was reduced by the addition of 2 mg of deoxyribonuclease I (bovine pancreas) and incubation at 37°C for 15 min. Membranes were separated from the cytoplasmic fraction by centrifugation. Sediment 1 was obtained by centrifuging the lysed cell suspension at 4,000 × g for 10 min. Sediment 2 was the pellet obtained by centrifuging the supernatant at 40,000 × g for 1.5 h, and the supernatant was the cytoplasmic fraction. Sediments 1 and 2 were each washed three times with TES-saline. The cytoplasmic fraction was centrifuged again at 40,000 × g for 2 h to remove sedimentable membrane components. The supernatant was dialyzed exhaustively against TES-saline for 48 h, concentrated 40- to 50-fold by placing the casings in Aquacide II-A (a hydrophilic polyethylene glycol; molecular weight, 250,000; Calbiochem, La Jolla, Calif.), and stored frozen at −70°C. Digitonin lysis was carried out in the same manner by using 200 ml of 40 μM digitonin in 0.25 M NaCl with incubation for 15 min at 37°C. Adjustment of the pH was not required before the addition of 2 mg of deoxyribonuclease I. The remaining procedure was identical to that used for alkaline lysis of cells.

**Polyacrylamide gel electrophoresis.** Electrophoresis was carried out in a model SE-500 (Hoeffer Scientific Instruments, San Francisco, Calif.) electrophoresis apparatus with 1.5-mm-thick polyacrylamide gels. The procedure was adapted from that of Laemmli (9); however, there was 3.5% cross-linkage in the gel and dithioerythritol was used instead of mercaptoethanol. The test sample contained 140 μg of total cell protein. Gels were stained with 0.2% Coomassie brilliant blue R-250 in methanol-water-acetic acid (4:5:4.5:1) at 56°C for 1 h. Slabs were destained with water-ethanol-alcohol-acetic acid (8:3:1) at room temperature. Destained gels were stored in aqueous 6% (vol/vol) acetic acid-2% (vol/vol) glycerol.

Isoelectric focusing live organisms. Isoelectric focusing in polyacrylamide gels in the pH range 3 to 10 was carried out in Multiphor 2117 apparatus (LKB Instruments, Inc., Rockville, Md.), as described elsewhere (19, 22). The samples were dissolved in equal volume of a sample-solubilizing solution (6 mM dithioerythritol, 500 mM glycerol, and 20 mM Tween-20 in water). The sample (80 μg of protein) was applied directly onto the surface of the gel. Polycrylamide gels were prepared in a manner similar to that described previously (19); however, urea was replaced with 500 mM glycerol, and...
Triton X-100 was replaced with Tween-20. The "anolyte" was 0.5 M phosphoric acid, and the "catholyte" was 70 mM ethylenediamine. Samples were focused at a constant current of 35 mA for 2.25 to 2.5 h or until voltage had increased to 1,000 V. The pH gradient, formed in the flat bed, was measured (at points 0.5 cm apart) on the surface of the gel by using an MI-410 micro-glass electrode and a 1.2 mm pH probe (Micro-electrodes, Inc. Londonderry, N.H.).

Serum protein fractions. The following commercially available equine serum fractions (U.S. Biochemicals, Cleveland, Ohio) were used as controls for comparative analysis: albumin, gamma globulins, and transferrin. Bovine serum proteins included albumin, alpha globulins, beta globulins, gamma globulins, transferrin, glycoprotein, Cohn fraction VI, and lipoprotein Cohn fraction 111-0; 10 to 12 μg of each protein was used per sample. Agamma horse serum and whole horse serum were titrated with 1 N HCl to a pH of 6.0. The precipitate was collected by centrifugation and washed three times with a 150 mM NaCl solution (pH 6.0).

RESULTS

Comparison of polypeptides by molecular weight. From 36 to 40 polypeptides (protein bands) could be distinguished in samples which contained 140 μg of protein (Fig. 1). Peptides of similar molecular weight were observed for the U. urealyticum serotypes at both the high-molecular-weight end of the gel (greater than 90,000 daltons) and at the low end of the gel (less than 30,000 daltons), whereas in the middle range of the gel striking differences were observed among serotypes (Fig. 1). Serotypes 1 and 8 possessed distinct major peptides of 85,000 and 95,000 daltons, respectively, which were not found in the other types. Types 3 and 5 contained polypeptides of 72,000 and 64,000 daltons, respectively, which also were not found in other types. Some polypeptides were located in only certain types; e.g., polypeptide 59,000 was found in types 6, 7, and 8. A polypeptide of 51,000 daltons was common in types 2, 4, 5, and 8, whereas a 46,000-dalton polypeptide was observed in types 2, 4, 5, 7, and 8. However, certain prominent polypeptides (e.g., polypeptides of 44,000 and 70,000 daltons) were common to all types (Fig. 1, upper two arrowheads). The polypeptide profiles of the U. urealyticum strains were strikingly different from those of strains of Mycoplasma and Acholeplasma (Fig. 2). M. hominis ATCC 14027 and M. arginini G-230, known to be serologically related (7, 21), showed only slight similarities in patterns, and the level of similarity was far less than the similarities seen among the U. urealyticum strains. As expected from their unique serological and biochemical features (7), A. laidlawii ATCC 14192 and M. gallisepticum ATCC 15302 were not only strikingly different from each other but also markedly different from the other organisms.

Comparison of organisms by pH value. The patterns from U. urealyticum strains showed strong similarities to each other but little resemblance to strains of Mycoplasma and Acholeplasma when compared by isoelectric focusing (Fig. 3). Serotype 1 contained unique proteins with pH values of 6.4, and the strains of types 2, 4, 5, and 7 contained proteins which focused at pH 5.3 and which were not found in the other strains tested. A. laidlawii ATCC 14192 showed a number of proteins in the acidic range (pH 3.7 to 4.8) not seen in any of the other
strains examined. Although *M. arginini* G-230 had shown some similarities to *M. hominis* ATCC 14027 by polyacrylamide gel electrophoresis (Fig. 2), less similarity was observed by isoelectric focusing; *M. arginini* G-230 showed a number of acidic proteins (pH 5.0 to 5.5) not seen in *M. hominis* ATCC 14027.

**Comparison of membrane and cytoplasmic fractions.** Lytic conditions for the *U. urealyticum* strains were studied to determine the optimum method for separation of membranes and cytoplasm. The lysis of *U. urealyticum* was a function of pH (Fig. 4); there was 50% lysis at pH 9.8, 70% lysis at pH 9.8, 70% lysis at pH 10.0, and greater than 80% lysis at pH 10.4. For further study, pH 10.2 was used to prepare membranes and cytoplasm to minimize destruction of membrane proteins while still providing effective lysis of cells. *M. gallisepticum* ATCC 15302 was the most sensitive organism in the experiment, showing 86% lysis at pH 9.4 and significant dissolution (96% cell lysis) at pH 10.0. In contrast, *A. laidlawii* ATCC 14192 was the most resistant organism, showing only 50% lysis at pH 10. The *U. urealyticum* strains and *M. hominis* ATCC 14027 were markedly sensitive to digitonin lysis, whereas *M. gallisepticum* and *A. laidlawii* were less sensitive (Fig. 5). For further experiments, 40 µM digitonin was used because with it 70% lysis could be achieved for the four organisms.

Membrane profiles obtained by SDS-polyacrylamide gel electrophoresis did not differ greatly from the whole-cell profiles of strains of serotypes 1 and 8; however, the cytoplasmic fractions were strikingly different from both the membrane fractions and the whole-cell fractions (Fig. 6). A 95,000-dalton polypeptide, which was clearly unique to type 8 (Fig. 1), was found to be a membrane component (Fig. 6). An 85,000-dalton polypeptide, which was unique to type 1, was also a membrane component. Nearly all of the peptides with molecular weights greater than 70,000 were found in the membrane fraction. A component of 70,000 daltons, which was observed in all of the ureaplasmic serotypes, was found to be clearly cytoplasmic (Fig. 6, tracks 3, 7, 10, and 14). Both methods of lysis appeared to give effective separation of cytoplasm from membrane, but certain differences were evident. The common component of 70,000 daltons was recovered more poorly by digitonin lysis than by alkaline lysis of type 8 cells (Fig. 6), whereas no such differentiation was observed in lysis of type 1 cells. On the other hand, the membrane component of 95,000 daltons contaminated the cytoplasm of type 8 cells when the cytoplasmic fraction was prepared by alkaline lysis (Fig. 6). We found that the cytoplasmic fraction was more effectively separated from the whole organism than the membrane fraction was, most likely because the membrane fraction contained some whole cells or because cytoplasmic components adhered to membrane fragments. At least five to seven polypeptides could be specifically assigned to the cytoplasmic fraction and excluded from the membrane fraction.

Testing of the fractions by isoelectric focusing also permitted differentiation of cytoplasmic and membrane components (Fig. 7). Membrane fractions of type 1 showed two unique proteins fo-
Fig. 3. Polyacrylamide gel isoelectric focusing of whole-cell lysates of strains of the eight serotypes of U. urealyticum, three species of Mycoplasma, and a representative of Acholeplasma; horse serum proteins were used as controls. From left to right the gels contained strains of U. urealyticum serotypes 1 through 8; M. argunii G-230, M. gallisepticum ATCC 15302, M. hominis ATCC 14027, A. laidlawii ATCC 14192, whole horse serum, horse transferrin, horse serum precipitate obtained at pH 6.0, and horse gamma globulins. Slanted arrows indicate major specific and common components.

Fig. 4. Lysis of ureaplasmata, acholeplasmata, and mycoplasmata according to pH. The lysis buffers used were 50 mM MES (pH 6.0), 50 mM TES (pH 7.0 and 8.0), and carbonate-bicarbonate buffer (pH 9.0, 9.4, 9.8, 10.0, 10.4 and 10.8). Symbols: ●, strain of U. urealyticum serotype 8; ▲, M. gallisepticum ATCC 15302; Δ, M. hominis ATCC 14027; □, A. laidlawii ATCC 14192. Percent lysis = (A - B)/(B - C) x 100, where A is the optical density of a suspension in phosphate-buffered saline (no lysis), B is the optical density of a suspension in the test system (variable degree of lysis), and C is the optical density of a suspension in an aqueous solution of 75 μM SDS (complete lysis).

Fig. 5. Lysis at various concentrations of digitonin. Symbols: ●, strain of U. urealyticum type 8; ▲, M. gallisepticum ATCC 15302; Δ, M. hominis ATCC 14027; □, A. laidlawii ATCC 14192. Percent lysis was measured as described for carbonate lysis (see legend to Fig. 4).

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Censing at pH values of 6.5 (Fig. 7). These components were clearly absent in the cytoplasmic fraction and also in the fractions of the type 8 strain (Fig. 7). However, at pH 7, type 8 mem-

branes were greatly enriched for a component (Fig. 7) which appeared to be an assembly of five proteins and which was specific to this type. Prominent cytoplasmic components focusing at pH 5.6 and 7.2 were found in both strains.

Contamination of U. urealyticum sediments with medium components. Washed cells of U. urealyticum and other members of the Mycoplasmatales are frequently, if not usually, contaminated with medium components, particularly insoluble peptone, yeast extract, and serum components, which cosedimented with the
organisms and cannot be removed by washing of the organism pellet (2, 6, 7, 13, 28). This problem has greatly frustrated attempts at analyzing cell fractions either biochemically (13) or immunologically (7). This problem can be evaded to an extent by the use of a dialysate broth base supplemented with agamma serum, which has been selected for its ability to remain clear during cultivation of the organism (5-8). In addition, in this study we filtered the dialysate broth base and used 1% serum (which we found to give cellular yields equivalent to the more usual higher serum concentrations) with a higher urea concentration in a strongly buffered medium to maximize yields for U. urealyticum (8). The degree of contamination with serum components was best shown by isoelectric focusing. We showed, for the first time, that a component migrating in the same location as a component of serum transferrin is found not only in ureaplasmata but also in mycoplasmata and acholeplasmata (Fig. 3, lower arrowhead). Other serum components are found in the acidic region of the gel (Fig. 3, upper arrowhead). Mycoplasmata are not known to possess cytochromes (12); however, a component common to all organisms was observed which comigrated with cytochrome c in SDS-polyacrylamide gels in the 14,000-dalton region (Fig. 1 and 2, dense line which extends across bottom of all gels; another possible serum contaminant is indicated by the arrowheads in Fig. 2). Thus, contaminating components having molecular weights similar to those of transferrin, cytochrome c, and immunoglobulin fragments were identified in the organisms studied. The fact that ureaplasmata can be grown in dialysate broth with 1% serum, in contrast to the crude medium with 5 to 10% serum used in other studies (11, 15), probably accounts for the greatly reduced contamination observed in our study. An important ancillary control is the fact that the profiles of U. urealyticum strains are strikingly different from those of mycoplasmata grown in a similar medium, indicating that the major differences and similarities shown are specific to U. urealyticum. In fact, the failure in previous studies (19) to distinguish specific peptides in various U. urealyticum types must be
protein composition of U. urealyticum

**FIG. 7.** Comparison of isoelectric focusing of membrane and cytoplasmic fractions with isoelectric focusing of the whole-cell lysates (prepared by the digitonin lysis procedure) of strains of U. urealyticum serotypes 1 and 8. Slanted arrows indicate specific components.

due in large part to contamination of the minute pellets with medium components.

**DISCUSSION**

The polypeptide patterns of the *U. urealyticum* strains were strongly similar when tested by SDS-polyacrylamide slab gel electrophoresis. However, most strains had at least one unique polypeptide present in a major amount. This association of specific polypeptides with given serotypes has not been demonstrated previously by polyacrylamide gel electrophoresis (15) using different techniques, namely, solubilization with phenol-acetic acid-urea (1) and employing tube-gels (15, 17, 29), in contrast to the SDS used for solubilization and estimation of molecular weights (22, 26) used in our study. The *U. urealyticum* patterns overall were sharply distinct from the patterns produced by three highly dissimilar organisms in the *Mycoplasma*ales: *A. laidlawii*, which does not require cholesterol (13) and whose genome is twice as large as that of *M. gallisepticum*, an organism which has a unique structure and utilizes glucose (13); and two serologically related arginine-utilizing organisms, *M. arginini* and *M. hominis* (7, 25). However, no specific common polypeptide could be used to identify *U. urealyticum* because at least one of the heterologous organisms showed peptides of similar molecular weights in each case, an event which would occur by chance because of the large number of polypeptides present (Fig. 2). However, it may be possible to identify *U. urealyticum* by using a combination of peptides which are in common to all strains. For example, the pattern produced by peptides of 70,000 and 44,000 daltons (Fig. 1 and 2) appears to permit identification of the species *U. urealyticum*. The "gaps" between the polypeptide bands at 33,000, 29,000, and 25,000 daltons (Fig. 1, three lowest arrowheads) are particularly important markers since the organisms in the other genera had peptides observable in these areas, whereas the gaps were consistently present in all human ureaplasmic strains (Fig. 1 and 2). The close similarities shown among the strains of the *U. urealyticum* serotypes indicate strongly that these organisms are very closely related and that the original decision to divide these organisms into types rather than species (23) was probably correct. Although further comparative studies will be required to encompass the diversity observed in the *Mycoplasma*ales (6, 7, 24), recognition of these specific markers or patterns or both may prove useful for identification of this species. Our study was focused at determining differences and similarities among the type strains of the eight serotypes of *U. urealyticum*. It remains to be determined whether the unique peptides demonstrated for the single representative strain of each serotype tested in our study will correlate with the antigenic determinant responsible for serotype differences or whether the unique peptides reflect biovar differences other than serotype among strains. Although few strains of a given serotype have been identified, considerable progress at identifying clinical isolates is now being made (23); this should permit comparisons of a number of strains of a given serotype.

The lytic methods used for ureaplasmata gave effective separation of cytoplasm from membranes, striking enrichment of the cytoplasmic fractions for specific components, and elimination of the major specific peptides of the types tested (85,000 daltons for type 1 and 95,000 daltons for type 8). However, the membrane fractions were not so clearly distinguishable from the whole lysed organisms by SDS-polyacrylamide gel electrophoresis; the major cytoplasmic peptides were greatly reduced in quantity, but a number of the peptides in the lower-molecular-weight region were found in both the membrane fractions and lysed whole organisms.
Nonetheless, the present data suggest strongly that the major specific peptides are membrane components. The fact that ureaplasmata are susceptible to digitonin (11) is not surprising in view of their requirement for cholesterol, nor is the lysis of *A. laidlawii* by digitonin remarkable since Razin and Argaman (14) have shown that acholeplasmata only resist digitonin lysis when the organisms are grown in cholesterol-free medium (which was not done in the present experiments). In a recent study, Romano and LaLicata (16), using the procedure of Masover et al. (11) and ultrasonic disruption, claimed that ultrasonic treatment provided membranes of superior quality compared with those obtained by digitonin lysis. However, it is now evident from our studies that effective lysis of ureaplasmic cells requires both a higher concentration of digitonin and a lower amount of protein (150 µg of protein per ml of 40 µM digitonin) than used in those studies (11, 16).

The degree of separation of the fractions separated much greater when they were tested by isoelectric focusing (Fig. 7). The cytoplasmic fractions were clearly greatly enriched for the major specific membrane proteins could be resolved for type 1 at pH 7.0, whereas the membrane fractions yielded mechanisms, p. 40

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**REPRINT REQUESTS**

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