Genetic Relatedness Among the Typhus Group of Rickettsiae

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The taxonomy of the typhus biogroup (Rickettsia prowazekii and R. mooseri [R. typhi]) of the genus Rickettsia and Rochalimaea quintana has been studied on the basis of the guanine plus cytosine content of the respective deoxyribonucleic acids (DNAs), the genome size, and the degree of DNA-DNA hybridization. The previously published guanine plus cytosine values for certain strains are confirmed, and the guanine plus cytosine values are described for some additional strains in each species. The genome sizes of four strains of R. prowazekii, three strains of R. mooseri, and two strains of R. quintana were very similar and averaged approximately 10^6 daltons. The various strains within the species boundaries hybridized at nearly the 100% level. The DNAs of various strains of R. prowazekii hybridized with the DNAs of various strains of R. mooseri rather consistently at the level of 70 to 77%. The DNAs of two strains of Rochalimaea quintana hybridized with those of R. prowazekii Breinl and R. mooseri Wilmington in the range of 25 to 33%.

The typhus group of rickettsiae is one of the biogroups that comprise the genus Rickettsia, the others being the spotted fever and scrub typhus groups. The typhus group contains three recognized species: R. prowazekii, R. mooseri (R. typhi), and R. canadensis. The classification is based on strong serological cross-reactions within the group as well as on other biological characteristics (41). A common guanine plus cytosine (G+C) content of deoxyribonucleic acid (DNA) has been demonstrated recently among strains of the typhus biogroup (37). Within the typhus group, recent studies of protein profiles have generally supported the established taxonomy based on serological and biological differences (13, 14, 19). This report provides additional information on the taxonomy of the group based on the G+C content of DNA, genome size, and DNA-DNA hybridization. Established strains and recent isolates of R. prowazekii and R. mooseri of the typhus biogroup are compared with each other as well as with two strains of Rochalimaea quintana.

MATERIALS AND METHODS

Rickettsial strains. The designations, sources, passage histories, and other pertinent information on the strains used in this study are presented in Table 1.

Reagents. Reagent grade liquefied phenol (Fisher Scientific Co., Pittsburgh, Pa.) was freshly distilled over metallic zinc into water. The phenol was stored in a brown bottle under nitrogen. Before use, the phenol was neutralized by adding 1/50 volume of 1.0 M sodium phosphate buffer, pH 6.8. Ribonuclease (Worthington Biochemical Corp., Freehold, N.J.) was dissolved in 0.05 M sodium phosphate buffer (pH 6.8) and heated at 80°C for 10 min to inactivate any possible deoxyribonuclease activity in the preparation.

Growth and purification of rickettsiae. The various rickettsial agents had been previously plaque purified in these laboratories (C. L. Wiseman, Jr., et al., unpublished data) to rid the cultures of extraneous agents. Seed pools of rickettsiae were then built up in Cofal/Marek-negative eggs (SPAFAS, Inc., Norwich, Conn.) and stored (−70°C) as 20% yolk sac homogenates in brain heart infusion medium. For rickettsial production, 6-day-old conventional eggs (Truslow Farms, Chestertown, Md.) were inoculated via the yolk sac route with seed pool inoculum diluted so that 30 to 50% of the eggs were killed in 7 to 8 days. Only live embryos were harvested. The yolk sacs were aseptically removed, blended, and diluted in brain heart infusion broth to a 20% (wt/vol) concentration. They were then shelf frozen in an alcohol-Dry Ice mixture and stored at −70°C until needed.

The rickettsiae were freed of host cell contaminants by methods developed and adapted in these laboratories and in use for several years (C. L. Wiseman, Jr., et al., unpublished data). This included an initial purification and concentration of the crude yolk sac homogenate by a sucrose batch method which removed about 90% of the contaminants, followed by a flocculation of the remaining host cell materials by a combination of albumin (2) and anti-yolk sac serum (21) precipitation, and finally two cycles of gradient separation, in this instance a modification of the Renografin gradient of Cahn and Fox (9) as adapted by Weiss et al. (14, 40) to rickettsiae. The purified rickettsiae were suspended finally in 1X saline-sodium citrate (SSC) (20), shell frozen in a Dry Ice-alcohol bath, and stored at −70°C until needed.

Growth and purification of R. quintana. The Fuller (ATCC VR-358) and Heliodoro (courtesy of J. W. Vinson, Harvard University School of Public Health, Boston, Mass.) strains of R. quintana were stored as seed pools in brain heart infusion broth at −70°C until needed. When cells were desired, an ampoule was quickly thawed, and several drops were placed onto each of a series of plates of Trypticase soy broth.
Rocha limaea quintana

Gimenez basic fuchsin stain but poorly with the Gram stain. After 10 to 12 h, and (iv) appropriate morphology and tinctorial properties, i.e., staining well with the Gram stain. 

The inoculum was spread on selected systems, Cockeysville, Md. The inoculum was spread on each plate by emulsifying the growth in 3 to 4 ml of 1x SSC. Any agar particles were first removed by low-speed centrifugation (21,000 lb/in² at a rate of 3 ml/min), followed by centrifugation at 12,000 x g for 15 min. The cell suspension was washed an additional two times in Ix SSC and finally resuspended in 1x SSC to an optical density (OD) of 2.00 at 440 nm and with a l-cm light path.

The DNA was then dialyzed against 2X SSC, and the resultant product was shell frozen in an alcohol-Dry Ice bath and stored at −70°C in flame-sealed ampoules.

**Determination of sheared DNA fragment length by electron microscopy.** The sheared DNA was spread as a film using the Kleinschmidt-formamide technique (23). DNA fragments were picked up on a grid and placed on a carbon film, which was burned at 5 cm from the lamp, and then rotary-shadowed in an Edwards evaporator. The shadowing material was 17 mm of platinum-palladium wire which was burned at 5 cm from the lamp and at a height of 4 mm. The photographic negative images of the DNA fragments were projected in a darkened room onto large sheets of paper (3 by 4 feet [ca. 0.91 by 1.22 m]) to give DNA fragments averaging 5 to 6 cm. The fragments were traced over with a felt-tipped marker pen, and the fragment lengths were then determined by using a map-measuring tool. The mean and standard deviation values so obtained were converted to true lengths by multiplying by the overall magnification factor. DNA fragment lengths were then rotary-shadowed in an Edwards evaporator and shadowed with copper grids coated with a colloidion carbon film, which was burned at 5 cm from the lamp and at a height of 4 mm.

**Purification and shearing of DNA.** The DNA extraction and purification procedure was basically that described by Thomas et al. (36). Pronase (1 mg/ml) was added to the rickettsial suspension (OD = 2.0 at 440 nm and with a 1-cm light path) followed by 0.2% sodium dodecyl sulfate. The mixture was incubated at 37°C for 7 h. The cell lysate was extracted with phenol for 30 min, and the DNA was precipitated from the aqueous phase with two volumes of chilled ethyl alcohol (95%). The partially purified DNA was dissolved in 0.1x SSC and then dialyzed against 0.05 M sodium phosphate buffer (pH 6.8). Ribonuclease (50 µg/ml) was added after dialysis, and the solution was incubated for 2 h at 37°C. The DNA was then adsorbed onto hydroxyapatite (DNA grade, Bio-Rad Laboratories, Richmond, Calif., 0.20 M sodium phosphate buffer, pH 6.8), followed later by elution (0.40 M sodium phosphate buffer, pH 6.8) according to the procedure suggested by Thomas et al. (36).

The DNA solution was then dialyzed overnight against 0.1X SSC. The DNA was sheared in a Ribi cell fractionator at 21,000 lb/in² at a rate of 3 ml/min (20), and the sheared DNA was concentrated under vacuum (rotating flask) to an OD in excess of 2.0 (260 nm, l-cm light path). The DNA was then dialyzed against 2X SSC, and the resultant product was shell frozen in an alcohol-Dry Ice bath and stored at −70°C in flame-sealed ampoules.

**TABLE 1. Source, passage history, and other pertinent information on the strains employed in this study.**

<table>
<thead>
<tr>
<th>Organism/strain</th>
<th>Source</th>
<th>Passage historya</th>
<th>Plaque purified</th>
<th>Reference</th>
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<tr>
<td><strong>Rickettsia prowazekii</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Breinl</td>
<td>Human</td>
<td>E155/TC3/E3</td>
<td>Yes</td>
<td>44</td>
</tr>
<tr>
<td>Madrid E</td>
<td>Human</td>
<td>CRD-3*/TC3/E4</td>
<td>Yes</td>
<td>11</td>
</tr>
<tr>
<td>Bur X-16</td>
<td>Human</td>
<td>TC5/E2</td>
<td>Yes</td>
<td>—9</td>
</tr>
<tr>
<td>GV-P12f</td>
<td>Flying squirrel</td>
<td>E7</td>
<td>No</td>
<td>4</td>
</tr>
<tr>
<td><strong>Rickettsia mooseri (R. typhi)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wilmington</td>
<td>Rat</td>
<td>E42/TC3/E3</td>
<td>Yes</td>
<td>28</td>
</tr>
<tr>
<td>Pak NA-18</td>
<td>Rat</td>
<td>GP/TC5/E3</td>
<td>Yes</td>
<td>—9</td>
</tr>
<tr>
<td>Ethio AZ-306</td>
<td>Rat</td>
<td>TC5/E3</td>
<td>Yes</td>
<td>—9</td>
</tr>
<tr>
<td><strong>Rochalimaea quintana</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fuller</td>
<td>Human</td>
<td>BA9</td>
<td>Yes</td>
<td>39</td>
</tr>
<tr>
<td>Heliodoro</td>
<td>Human</td>
<td>BA7</td>
<td>No</td>
<td>38</td>
</tr>
</tbody>
</table>

**a Abbreviations:** E, Yolk sac passage; TC, tissue culture plaque purification passage; GP, guinea pig passage; BA, blood agar passage.

**b Experimental vaccine lot CRD-3 (C. L. Wiseman, Jr., unpublished data).**

**c C. L. Wiseman, Jr., et al., unpublished data.**

**d Furnished by F. M. Bozeman, Bureau of Biologics, Food and Drug Administration, Bethesda, Md.**

agar with 10% sheep blood (BBL Microbiology Systems, Cockeysville, Md.). The inoculum was spread on each plate with a sterile glass rod. The plates were incubated at 35°C for 3 to 4 days in an atmosphere of 95% air–5% CO₂ and a relative humidity of 90 to 95% to obtain a confluent lawn. The cells were harvested from each plate by emulsifying the growth in 3 to 4 ml of 1X SSC. Any agar particles were first removed by low-speed centrifugation (200 x g, 5 min) followed by centrifugation at 12,000 x g for 15 min. The cell suspension was washed an additional two times in 1X SSC and finally resuspended in 1X SSC to an optical density (OD) of 2.00 at 440 nm and with a 1-cm light path. Bacteriological criteria for R. quintana included (i) the inability to grow on the blood agar medium included (i) the inability to grow on a Trypticase soy agar without blood enrichment, (iii) a generation time on the order of 10 to 12 h, and (iv) appropriate morphology and tinctorial properties, i.e., staining well with the Giménez basic fuchsins stained but poorly with the Gram stain.

The DNA extraction and purification procedure was basically that described by Thomas et al. (36). Pronase (1 mg/ml) was added to the rickettsial suspension (OD = 2.0 at 440 nm and with a 1-cm light path) followed by 0.2% sodium dodecyl sulfate. The mixture was incubated at 37°C for 7 h. The cell lysate was extracted with phenol for 30 min, and the DNA was precipitated from the aqueous phase with two volumes of chilled ethyl alcohol (95%). The partially purified DNA was dissolved in 0.1X SSC and then dialyzed against 0.05 M sodium phosphate buffer (pH 6.8). Ribonuclease (50 µg/ml) was added after dialysis, and the solution was incubated for 2 h at 37°C. The DNA was then adsorbed onto hydroxyapatite (DNA grade, Bio-Rad Laboratories, Richmond, Calif., 0.20 M sodium phosphate buffer, pH 6.8), followed later by elution (0.40 M sodium phosphate buffer, pH 6.8) according to the procedure suggested by Thomas et al. (36). The DNA solution was then dialyzed overnight against 0.1X SSC. The DNA was sheared in a Ribi cell fractionator at 21,000 lb/in² at a rate of 3 ml/min (20), and the sheared DNA was concentrated under vacuum (rotating flask) to an OD in excess of 2.0 (260 nm, 1-cm light path). The DNA was then dialyzed against 2X SSC, and the resultant product was shell frozen in an alcohol-Dry Ice bath and stored at −70°C in flame-sealed ampoules.

**Determination of sheared DNA fragment length by electron microscopy.** The sheared DNA was spread as a film using the Kleinschmidt-formamide technique (23). DNA fragments were picked up on copper grids coated with a colloidion carbon film, which were then rotary-shadowed in an Edwards evaporator. The shadowing material was 17 mm of platinum-palladium wire which was burned at 5 cm from the specimen and at a height of 4 mm. The photographic negative images of the DNA fragments were projected in a darkened room onto large sheets of paper (3 by 4 feet [ca. 0.91 by 1.22 m]) to give DNA fragments averaging 5 to 6 cm. The fragments were traced over with a felt-tipped marker pen, and the fragment lengths were then determined by using a map-measuring tool. The mean and standard deviation values so obtained were converted to true lengths by multiplying by the overall magnification factor. DNA fragment length was converted to daltons according to the Watson-Crick B configuration of double-stranded DNA (24). Assuming these conditions, the molar linear density is 1.91 x 10¹⁵ daltons/cm.

**Determination of DNA concentration and purity.** The DNA content was determined by the Burton (8) modification of the diphenylamine reaction. Ribonucleic acid and protein levels in the DNA samples were assayed by the methods of Meijbaum (30) and Lowry et al. (25), respectively.

**Determination of G+C content of DNA.** The G+C content (moles percent) of DNA was determined by the method of Marmur and Doty (27). All assays were performed in a Gilford spectrophotometer 2400.
hybridization experiments were performed between mixture, sample A, and sample B, respectively. When a decrease in absorbance per minute (260 nm) in the DNA rate, determination. The degree of hybridization, prepared. In all other respects, the analytical procedure was the same as that described under genome size. Four cuvettes. DNA molecular weight is related to equation: molecular weight \( \times \text{X} \) (98.37 - [0.91 \% (G+C)]) / k.

Measurement of DNA-DNA hybridization. The procedure followed (16) is, in principle, very similar to that involved in determining genome size. Four cuvettes are required. The adenine blank and the two DNA samples to be compared were set up as described in the section on genome size. In addition, a cuvette contained a 1:1 mixture of the two DNA samples to be compared. In all other respects, the analytical procedure was the same as that described under genome size determination. The degree of hybridization, D, is determined according to the equation: 

\[
D = \frac{4 V_m - (V_R + V_h)}{\sqrt{V_m} V_h}, \text{ where } V_m, V_R, \text{ and } V_h \text{ are the decrease in absorbance per minute (260 nm) in the DNA mixture, sample A, and sample B, respectively. When hybridization experiments were performed between R. prowazekii or R. mooseri and R. quintana, whose optimal renaturation temperatures are 62 and 66°C, respectively, an intermediate temperature of 64°C was chosen.}

RESULTS

Chemical purity and fragment length of DNA preparations. The levels of protein and ribonucleic acid in our DNA samples were below the levels of detectability with our assay procedures and were less than 1% of the DNA concentrations.

Two sheared DNA preparations of R. prowazekii Breinl and one of R. mooseri Wilmington were assayed by electron microscopy for DNA fragment size. The results were 2.1 \( \pm \) 1.0 \( \times \) 10^6 daltons (255 fragments), 2.9 \( \pm \) 1.2 \( \times \) 10^6 daltons (408 fragments), and 2.5 \( \pm \) 1.1 \( \times \) 10^6 daltons (206 fragments), respectively.

G+C content of DNA. Table 2 presents the G+C content of the DNA extracted from strains of several rickettsial species. The data shown for the Breinl and Madrid E strains of R. prowazekii and the Wilmington strain of R. mooseri confirm the previous results obtained by several investigators (37, 43); the results for the Burundi and flying squirrel strains of R. prowazekii and the Pakistan and Ethiopian strains of R. mooseri are new data.

Our value for the G+C content for the Fuller strain of R. quintana is consistent with the previously published value (37); the similar G+C value for the Heliodoro strain of R. quintana is new information. Escherichia coli K-12 DNA was analyzed for comparison. The G+C values of all DNA preparations, both rickettsial and E. coli, agreed well with the values previously published (27, 37, 43). The new data are consistent with groupings based on other data.

Genome size. The molecular weights of the DNAs of the four strains of R. prowazekii and the three strains of R. mooseri appear to be the same: approximately 1.1 \( \times \) 10^6. The genome size data on R. quintana Fuller and Heliodoro gave a value (1.0 \( \times \) 10^6 daltons) slightly lower than that obtained for the typhus group, but the differences may not be significant. The genome size obtained for the Fuller strain is in good agreement with a previously published value for this strain (22). E. coli K-12, as a reference organism, gave the appropriate value, 2.4 \( \times \) 10^9 daltons, consistent with published data (20). The genome in the rickettsiae tested, as a group, is thus seen to be about 45% of that of E. coli.

DNA-DNA hybridizations. Table 3 presents the degree of hybridization obtained under stringent renaturation conditions. When the DNAs of various strains recognized as belonging to the same species on the basis of serological criteria were hybridized, the values were invariably over 90% and were usually close to 100%. When the DNAs of various strains of R. prowazekii and R. mooseri were hybridized across the species line, the values rather consistently fell between 70 and 77%. The DNAs of R. quintana Fuller and Heliodoro hybridized against the DNAs of several strains of R. prowazekii and R. mooseri gave considerably lower values (25 to 33%).

DISCUSSION

Information on the molecular weight of a DNA can be obtained from the rate of reassociation of the complementary strands. Britten and Kohne (7) showed that the rate of reassociation of bacterial or viral DNAs is directly proportional to the size of the genome. They expressed this rate as the \( C_{0.65} \), the concentration of DNA...
TABLE 2. G+C content of the DNA and genome size of strains of R. prowazekii, R. mooseri, R. quintana, and E. coli

<table>
<thead>
<tr>
<th>Biogroup/strain</th>
<th>G+C content (mol%)</th>
<th>Genome size</th>
<th>DNA prepn</th>
<th>No. of runs</th>
<th>mol wt (×10^-1) (±SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Experimental^a</td>
<td>Published^a</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rickettsia prowazekii</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Breinl</td>
<td>29.0</td>
<td>29.0 (37)^d</td>
<td>1</td>
<td>5</td>
<td>106 ± 6.6</td>
</tr>
<tr>
<td>Madrid E</td>
<td>28.5</td>
<td>29.3 (37)</td>
<td>2</td>
<td>16</td>
<td>112 ± 9.2</td>
</tr>
<tr>
<td>Burundi (V-16)</td>
<td>29.7</td>
<td>1</td>
<td>1</td>
<td>6</td>
<td>108 ± 6.7</td>
</tr>
<tr>
<td>Flying squirrel (GV-F-12)</td>
<td>29.5</td>
<td>1</td>
<td>8</td>
<td></td>
<td>114 ± 5.8</td>
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<tr>
<td>Rickettsia mooseri (R. typhi)</td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>Wilmington</td>
<td>29.0</td>
<td>29.3 (37, 43)</td>
<td>1</td>
<td>17</td>
<td>109 ± 7.9</td>
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<tr>
<td>Pakistan (NA-18)</td>
<td>28.5</td>
<td>1</td>
<td>8</td>
<td></td>
<td>107 ± 6.1</td>
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<tr>
<td>Ethiopian (Eth-306)</td>
<td>29.0</td>
<td>1</td>
<td>3</td>
<td></td>
<td>107 ± 4.0</td>
</tr>
<tr>
<td>Rochalimaea quintana</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fuller</td>
<td>39.3</td>
<td>38.8 (37)</td>
<td>1</td>
<td>6</td>
<td>101 ± 3.5</td>
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<tr>
<td>Heliodoro</td>
<td>39.8</td>
<td>1</td>
<td>5</td>
<td></td>
<td>105 ± 2.4</td>
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<tr>
<td>Escherichia coli</td>
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<tr>
<td>K-12</td>
<td>51.7</td>
<td>1</td>
<td>6</td>
<td></td>
<td>241 ± 13</td>
</tr>
</tbody>
</table>

^a Our data.
^b Data from literature.
^c SD, Standard deviation.
^d Reference.

TABLE 3. DNA-DNA hybridizations between strains of R. prowazekii, R. mooseri, and R. quintana

in moles of nucleotide per liter times the time in seconds required for 50% reassociation. This relationship exists because the vast majority of the cistrons of a viral or bacterial DNA is composed of unique base sequences. As the result of these basic studies on the kinetics of renaturation by Britten and Kohne (7) and Wetmur and Davidson (42), a number of investigators have applied these concepts to determining the genome sizes of a variety of microorganisms (1, 10, 22, 34). This is readily accomplished with a recording spectrophotometer. The formation of double-
stranded complexes between complementary strands of the DNA is observed by the hypochromic shift. The rate of reassociation is determined by the rate of change in absorbance.

Gillis et al. (20) introduced a modified renaturation rate method for the determination of the molecular weight of genome DNA which was based on initial optical renaturation rate measurements of precisely known concentrations of fragmented DNA. The exact conditions and the effects of DNA concentration, renaturation time, size of DNA fragments, buffer concentration, optimal temperature, and G+C content on the renaturation rate were determined. Gillis et al. (20) applied the method to the determination of the genome size of 40 different bacteria. The method appeared to offer advantages in terms of simplicity, rapidity, and reproducibility over methods previously described (5, 18, 42), and for these reasons it was chosen for use in this study.

Renaturation rates may also be used to determine the degree of base sequence homology between DNAs. If two identical DNAs are allowed to reassociate under optimal conditions in the same incubation mixture, the reassociation rate of the mixture is expected to be the same as if they were incubated separately. If two DNAs with no base sequences in common are allowed to reassociate, the reassociation rate is expected to be equal to the sum of their rates when incubated separately. Thus, a mixture of two DNAs with partial base-sequence similarity will exhibit a rate of reassociation somewhere in the range between their rate alone and the sum of their rates. This spectrophotometric technique for determining the degree of DNA-DNA hybridization has been discussed both by Seidler and Mandel (34) and by De Ley et al. (16). Seidler and Mandel (34) based their hybridization procedure on the C_d technique developed by Britten and Kohne (7) and Wetmur and Davidson (42), whereas De Ley et al. (16) proceeded on the initial rate of reassociation concept, which was developed by Gillis et al. (20). We have applied the latter procedure to the hybridization studies presented in this paper because both genome size and degree of hybridization can be combined in the same experiment. In addition, there is no need to employ radiolabeled DNA, and the procedure is simple, rapid, and reproducible. Although the membrane filter (18) and hydroxyapatite hybridization procedures (5, 6) may permit a somewhat more detailed examination of the DNA relationships involved, the initial rate method employed here satisfied our immediate purposes of (i) distinguishing between species and groups within species, confirming relationships previously detected by serological means, and (ii) sorting new isolates into discrete groups, each closely related to an established species. The search for minor differences among strains of a given species is beyond the scope of this paper and would require somewhat different methods.

The typhus biogroup of the genus Rickettsia is currently recognized as containing three species: R. prowazekii, R. mooseri (R. typhi), and R. canada (41). In this study we have compared various strains of R. prowazekii, R. mooseri, and R. quintana, both within and without the genus and species boundaries, using the criteria of DNA base ratio, genome size, and degree of DNA-DNA hybridization. We did not include R. canada, a recent addition to the genus Rickettsia (3, 29), because plaque-purified cultures of this species were not yet available. Extraneous agents were present in the original seed obtained from the American Type Culture Collection, probably as a consequence of multiple egg passages. R. canada will, however, be included in a future study when an adequately purified culture is available.

The genome sizes found among the various strains of R. prowazekii, R. mooseri, and R. quintana are, in general, quite similar and are approximately 10^9 daltons. The validity of the genome size data is supported by the following. (i) The mean value of 2.4 x 10^9 daltons for the E. coli K-12 genome agrees well with the value obtained by Gillis et al. (20), who established the initial renaturation rate method for genome size determination. (ii) The value of 1.0 x 10^9 daltons obtained for the Fuller strain of R. quintana agrees reasonably well with the figure of 9.3 x 10^9 daltons given by Kingsbury (22). (iii) The molecular weight values obtained for the several strains of R. prowazekii and R. mooseri compare closely among themselves; no significant molecular weight differences would be expected between strains of the same species. (iv) Based on the studies of De Ley et al. (16) involving the effect of the molecular weight of the DNA fragments on the renaturation rate, our average value, from three samples, of 2.5 x 10^9 daltons compared to the value obtained by De Ley et al. (16), 4 x 10^9 daltons, is not sufficiently different to affect appreciably the renaturation rate.

Genome size data are now available on R. prowazekii, R. mooseri, and R. quintana (this paper) and on R. rickettsii and R. quintana (22), and all have approximately the same value (10^9 daltons). R. prowazekii, R. mooseri, and R. rickettsii are categorized as obligately intracellular parasites, whereas R. quintana grows extracellu-
ularly in the louse gut, in tissue culture, and on a blood agar medium (39, 41). Similarly, Kingsbury (22) obtained the same genome size for *R. rickettsii* as compared to *Neisseria gonorrhoeae* and *Neisseria meningitidis*. Thus, the genome sizes of a significant proportion of rickettsial species are well within the genome sizes of certain free-living bacteria and are, in fact, significantly larger than the genome size of members of the genus *Mycoplasma* (5 x 10^6 daltons) (1). If one accepts the dogma that essentially the entire bacterial genome codes for structural proteins, enzymes, and several types of ribonucleic acid and that there is little or no redundancy in the code (procaryotes), then one must assume that the failure of rickettsiae to grow extracellularly is not due to a genome size too small to be able to code for the essentials of extracellular growth but, rather, may be due to a relatively small number of evolutionary adaptations to a host-cell dependency.

The organisms of the typhus group included in this study had been identified as strains of one or the other reference organisms, *R. prowazekii* Breinl or *R. mooseri* (*R. typhi*) Wilmingtton, on the basis of conventional biological and serological criteria. The strains studied here had been isolated from a variety of sources, diverse geographical locations, and widely separated points in time and had had very different passage histories in the laboratory. Nonetheless, a remarkably high degree of hybridization (mean 97%, with a range of 92 to 104%) was found among the DNAs of strains identified on the basis of such conventional criteria as *R. prowazekii* and among DNAs of strains which had been identified as *R. mooseri*. Thus, as with conventional phenotypic characters, DNA hybridization values clustered tightly around the respective reference strains, with a range far less than that observed by others with certain different bacterial species (12, 17, 35). The DNA hybridization method employed here is not sensitive enough to detect minor differences among closely related strains. That some intraspecific differences may exist, however, is suggested by the studies of Dasch et al. (13), who recently studied the intraspecies differences of a number of strains classified as either *R. prowazekii* or *R. mooseri* by conventional criteria and who showed that the sodium dodecyl sulfate-polyacrylamide gel electrophoresis technique revealed no discernible differences between the *R. prowazekii* strains (Breinl [virulent], flying squirrel strains [four isolates], and Madrid E [avirulent, antibiotic resistant]) or between two *R. mooseri* isolates (Wilmington [human isolate] and a rat isolate). Dasch et al. (13), using the technique of isoelectric focusing of soluble rick-
groups (5, 33), the differences observed here appear to fall in an intermediate zone, sufficiently large to indicate species differences from some bacteria but not from others. In the case of the typhus rickettsiae studied here, the clear separation of strains into two DNA-DNA hybridization groups with no intermediate values might be taken as evidence that the 70 to 77% hybridization between the two groups does indeed constitute the basis for two species. Moreover, limited protein profile studies of purified rickettsiae have been performed in recent years by several investigators using the sodium dodecyl sulfate-polyacrylamide gel electrophoresis techniques. The consensus is that, although there is a general similarity in the protein profiles of strains of the species R. prowazekii and R. mooseri, there are too many differences observed to consider them as different strains of the same species (13, 19, 32).

It would appear, then, that the total available biological, serological, protein profile, and DNA-DNA hybridization information is consistent with the current classification of typhus rickettsiae into the two species R. prowazekii and R. mooseri. The relationship of R. canadensis to these two species will be the subject of a separate study.

We included strains of the agent of trench fever in this study primarily as a kind of negative control for the methodology. This organism was formerly included in the genus Rickettsia, but more recently it has been moved to a new genus on the basis of certain unique phenetic characteristics, and it is now designated Rochalimaea quintana (41). It does, however, display certain metabolic similarities to typhus group rickettsiae (41).

The accuracy of the initial rate method employed here suffers when the G+C content of the two DNAs differs by more than 8% (16). However, the degree of hybridization observed (25 to 33%) between the DNAs of R. quintana and R. prowazekii Breinl and R. mooseri Wilmington is within the maximum percent DNA homology between two DNA types which differ by 9 to 10% in G+C content but which have equal molecular weights and an average compositional nucleotide distribution (15). Thus, despite the limitations of the method, the results obtained are compatible with current views on the taxonomic relationship between R. quintana and the typhus rickettsiae (40). A more detailed study of R. quintana strains and Baker's vole agent appears elsewhere (31).

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