Deoxyribonucleic Acid Base Composition of Acetic Acid Bacteria

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SUMMARY

The base composition of purified DNA from 28 strains of acetic acid bacteria was determined. Most strains of the genus Gluconobacter clustered closely together at 60.6-63.4% (guanine + cytosine) of total base. All strains of the Acetobacter aceti biotype lay within the range 55.4-64.0% (guanine + cytosine). The close relationship and possible common phylogenetic origin of the genera Gluconobacter and Acetobacter is again stressed by these results. The base composition of DNA from acetic acid bacteria and from species of Pseudomonas was very similar, confirming the suspected close relationship between these groups. There is a noticeable agreement between the sequences of Acetobacter strains, arranged according to increasing % (guanine + cytosine) and arranged according to increasing enzymic equipment: strains with greater biochemical activity have on the whole also a higher % (guanine + cytosine) in DNA. The range of the compositional distribution of DNA molecules is on the whole broader in Acetobacter than in Gluconobacter. The results corroborate previous conclusions that both biotypes contain clusters of strains without species differentiation.

INTRODUCTION

DNA base composition appears to be a promising new tool in bacterial taxonomy (Lee, Wahl & Barbu, 1956; Belozersky & Spirin, 1958; Lanni, 1960; Sueoka, 1961; Schildkraut, Marmur & Doty, 1962; Marmur & Doty, 1962; Marmur, Seaman & Levine, 1963). The present knowledge in this field can be summarized as follows.

1. Different 'strains' of the same well-established species have almost identical DNA base compositions. Thus, for seven strains of Escherichia coli, the molar content of guanine + cytosine (G + C) expressed as a percentage of the total base in the DNA was 50% ± 0.4 (Marmur & Doty, 1962). Different strains of E. coli had 50% (G + C), even when determined by different authors.

2. Different 'species' which are known for morphological, physiological and other reasons to belong to the same genus, have identical or closely related % (G + C) values. This is well illustrated, for example, in the genera Escherichia, Salmonella, Clostridium and Hemophilus (Lee et al. 1956; Marmur & Doty, 1962). Exceptions are believed to be indicative of wrong classification. For example,
Catlin & Cunningham (1961) found all the strains of Neisseria they examined to have 49.5-51.5% (G + C), except for N. catarrhalis, which had 40.0-41.3%. They conclude that 'the inclusion of “catarrhalis” strains in the genus Neisseria appears illogical from the evolutionary point of view.' The suspected divergency within Proteus was confirmed by the results of Falkow, Ryman & Washington (1962), who found that P. morganii had a 50% (G + C), whereas the other species had 39%. On physiological and biochemical grounds P. morganii had previously been thought to be only remotely related to the other species ‘vulgaris’ and ‘mirabilis’ and a new genus Morganella had even been proposed for it (Fulton, 1943). The genus Bacillus offers another example where base composition tends to confirm the interpretation which was reached by quite different arguments. Base composition in Bacillus varies between 33 and 50% (G + C); this is an unusually wide range for one genus. The explanation might be (De Ley, 1962) that Bacillus should not be considered a single genus, but a collection of genera or biotypes, which would form the bridge between the Enterobacteriaceae (about 50% G + C) and the saccharolytic clostridia (about 27% G + C). Bisset (1962) suggested that ‘there is room for several genera in the family Bacillaceae’.

(3) Different genera, which are known or suspected for morphological, physiological, biochemical and other reasons to be closely related, also have base compositions which are very similar such as Escherichia, Salmonella and Shigella (Lee et al. 1956; Marmur & Doty, 1962), Proteus and Providence (Falkow et al. 1962). Unrelated genera have widely different % (G + C) values, the range being from 25 to 75% (G + C).

(4) If the mean % (G + C) of the DNA of two strains is different by 10%, there are few DNA molecules of the same (G + C) content common to both of them (Sueoka, 1961).

(5) Organisms with a closely related base composition are believed to be phylogenetically related (Sueoka, 1961) when they also display related morphological, physiological, biochemical and other characteristics.

However, up to now, the DNA base composition method has not been applied systematically to a wide set of strains of a group of bacteria other properties of which have been thoroughly investigated. The acetic acid bacteria seemed to be suitable objects for an application and test of the above conclusions. In view of the finding that Gluconobacter and Acetobacter each consist of a series of strains with a gradation of properties, it was proposed to regard all previous ‘species’ as variants within the biotypes Gluconobacter oxydans and Acetobacter aceti, which might phylogenetically be derived from the same common pool of ancestors (De Ley, 1961a). It was thus expected that the DNA base composition of Gluconobacter would be similar to that of Acetobacter, that the DNA base composition of all strains within each biotype would cluster in the same region, and that the base composition of acetic acid bacteria would be in the neighbourhood of that of Pseudomonas, namely 60-67% (G + C), in view of their suspected close taxonomic relationship.

Two methods were used: (1) direct estimation of purine and pyrimidine bases after hydrolysis and paper chromatographic separation; (2) thermal denaturation ('melting point'). It was also the aim of the present work to compare the practicability of both methods for routine analysis.
DNA compositions of acetic acid bacteria

METHODS

Organisms. We used the same strains as described previously (De Ley, 1961a) and will adhere to the nomenclature proposed in that paper. In addition, we also used Acetobacter aceti (strains paradoxus P1 and P2) and A. aceti (strains peroxydans 3 and 4) obtained through the courtesy of Dr J. W. M. la Rivière (Laboratory of Microbiology, Delft, the Netherlands), Gluconobacter oxydans strain melanogenus 116 (courtesy of Dr A. H. Stouthamer, Laboratory of Microbiology, Utrecht, the Netherlands) and G. oxydans strain melanogenus 49 (courtesy of Dr J. W. M. la Rivière, Laboratory of Microbiology, Delft, the Netherlands). Repeated plating and re-determination of the properties of the strains according to the criteria of Frateur (1950) and De Ley (1961a) assured that only pure cultures were used. Pseudomonas fluorescens CCEB 488 was obtained through the courtesy of Dr O. Lysenko (Institute of Biology, Prague); Agrobacterium tumefaciens through the courtesy of Dr M. Bernaerts (Ministry of Economic Affairs, Brussels) and the Escherichia coli was a National Collection of Type Cultures (Colindale, London, Great Britain; NCTC) strain.

Mass cultures. All strains of acetic acid bacteria were incubated at 30° for 2 to 3 days on solid media in Roux flasks, except for the strains of ‘peroxydans’ and ‘paradoxus’, which needed 1 week. Strains of Gluconobacter and the ‘mesoxydans’ group of Acetobacter were grown on a medium containing (% w/v): 10, glucose; 3, CaCO₃; 1, Yeastex (Ned. Gist en Spiritusfabriek, Brugge, Belgium); 2.5, agar. Most strains of the ‘oxydans’ and ‘peroxydans’ groups of Acetobacter were grown on 1.5% malt extract (Difco), 0.5% Yeastex, 3% (v/v) ethanol, 2.5% agar. A. aceti (paradoxus) was grown on 50% (v/v) red wine, 0.5% Yeastex, 1% DL-lactate Ca salt, 2% CaCO₃, 2.5% agar. Sometimes other media were used, e.g. with sorbitol instead of glucose. No differences in DNA base composition were then observed.

Pseudomonas fluorescens was incubated for one day at 30° in a liquid medium with 0.5% peptone (Difco) + 0.25% Yeastex and with passage of sterile air. Agrobacterium was grown in Roux flasks on a medium with 1% Yeastex, 1% glucose, 2% CaCO₃ and 2.5% agar for two days at 30°. Escherichia coli was grown for two days at 30° in Roux flasks on 0.5% peptone, 0.25% Yeastex and 2.5% agar.

Estimation of DNA base composition by paper chromatography. Organisms were harvested and washed as previously described (De Ley, 1961a). The pellet was extracted according to the method of Smith & Wyatt (1951). Two to three g. wet wt. living bacteria were suspended in 10 ml. n-NaOH and incubated at 87° for 20 hr. In some experiments the organisms were treated with 1% Na lauryl sulphate before NaOH extraction. No significant difference in the final yield was observed. After the extraction the residue was centrifuged down in a Servall SS-1 at 13,000g for 15 min. The DNA was precipitated by adjusting to pH 4 with acetic acid and adding one vol. of 95% (v/v) ethanol in water. The precipitate was centrifuged down and dissolved in 10 ml. dilute NaOH. Protein was removed by gel-formation with chloroform according to the method of Sevag, Lackman & Smolens (1938). The solution was shaken with 0.25 vol. chloroform and 0.1 vol. octanol for 30 min. and the chloroform and aqueous layers then separated by centrifugation at 2700g for 15 min. The upper aqueous layer was repeatedly treated with chloroform for 15 min. and centrifuged until almost no protein was observed at the liquid/liquid interface. DNA was again precipitated with ethanol at pH 4; the precipitate collected by
centrifugation, redissolved in 10 ml. dilute NaOH and the amount of nucleic acid
and protein determined by measuring the extinction at 260 and 280 m\(\mu\) according
to Warburg & Christian (1942). DNA was again precipitated with ethanol at pH 4
and dried in a vacuum desiccator. It was hydrolysed with 72% (v/v) perchloric
acid and the purines and pyrimidines of 10 pl. spots separated by paper chromatog-
raphy with a mixture of 65% (v/v) isopropanol 2N with respect to HCl, as the
solvent (Wyatt, 1951). After separation the bases were detected on the chromato-
grams by the photographic method of Markham & Smith (1949). The chromato-
grams were first dried overnight at room temperature. The source of ultraviolet
radiation was a germicidal G.E. lamp; Gevaert Reflex Document paper was used
for contact prints. The spots on the chromatograms were copied from the prints,
cut out, eluted with 4 ml. 0·1 n-HCl (Wyatt, 1951) by standing for 15–20 hr. at
37\(^\circ\). Blanks of equal size were cut out at distances corresponding to the \(R_f\) value
of the bases, from a strip of the same chromatogram, on which 10 pl. 2N-\(\text{HCl}_4\) had
been spotted between triplicate sample spots. The eluates of the bases were read
against the corresponding blanks in a Beckman spectrophotometer (model DU) at
the wavelengths indicated by Vischer & Chargaff (1948), except for cytosine, for
which 274 m\(\mu\) was used since this gave more reproducible results. The amount
of each of the bases was calculated from the difference \(\Delta\) between the extinction at the
absorption peak and at 290 m\(\mu\). For test solutions containing 10 \(\mu\)g. bases/ml.
0·1 n-HCl \(\Delta\) is: \(\Delta\) adenine = \(E_{262.5} - E_{290} = 0·900\); \(\Delta\) guanine = \(E_{249} - E_{290} = 0·475\); \(\Delta\) thymine
= \(E_{264·5} - E_{290} = 0·545\); \(\Delta\) cytosine = \(E_{274} - E_{290} = 0·458\). \(\Delta\) for cytosine was deter-
mined with a pure sample from Calbiochem (3625 Medford Street, Los Angeles 63,
California, U.S.A.)

**Estimation of DNA base composition by thermal denaturation.** DNA was isolated
according to the method of Marmur (1961). Thermal denaturation was followed at
260 m\(\mu\) in a Beckman spectrophotometer (model DU) as described by Marmur &
Doty (1962). To protect the photocell and the slit of the monochromator against
excessive heating, which would result in inaccurate readings, it was found advan-
tageous to use the following arrangement. The cell holder compartment was flanked
at each side by the following set up: first two thermospacers for the circulating hot
water from a Haake Ultra-Thermostat NB, followed by a 5 mm layer of cork
and finally another thermospacer through which passed dropwise a current of tap
water at 12\(^\circ\). To prevent traces of water vapour reaching the photocell (which would
again result in erratic readings) a suitable well was machined into the side of the
cell-holder compartment, which faced the photocell, and a circular quartz window
was glued into the opening. The cell-holder compartment itself was insulated on all
sides at the outside by cork. Tubing between the circulating water bath and the
thermospacers has to be as short as possible and insulated by either plastic or
asbestos. In this way the difference in temperature between the liquid in the
cuvettes and the bath did not exceed 1·2\(^\circ\). The circulating water contained either
25% ethyleneglycol or 30% glycerol. The temperature in the cell-holder compart-
ment was measured in a cuvette, containing water, tightly covered with a plastic
cover, through which passed a calibrated thermometer. A suitable hole was drilled
through the compartment cover. The thermostated cell holder compartment is
schematically represented in Fig. 1. Blanks and the samples were measured in
quartz cuvettes with ground glass stoppers. The \(T_m\) values were determined graphi-
DNA compositions of acetic acid bacteria

Estimation of the range of base composition distribution of DNA. Assuming that the distribution of the compositional distribution of DNA molecules around $T_m$ was Gaussian, the standard deviation $\sigma$ of the distribution, expressed as $\% (G + C)$ and corrected for the natural transition width of adenine-thymine DNA, was determined for each strain from the absorbance-temperature curves according to Doty, Marmur & Sueoka (1959). The mean $\% (G + C)$ was calculated with Marmur & Doty’s (1962) formula: $\% (G + C) = (T_m - 69.3)/0.41$. The Gaussian distribution around the mean $\% (G + C)$ was calculated with the equation $y = 1/(\sigma \sqrt{2\pi}) \exp(-x^2/2\sigma^2)$, in which $x$ was expressed as $\% (G + C)$ around the mean. The results are represented graphically in Figs. 4 and 5. The ratio of the area under the $y$ curve in any $\% (G + C)$ interval to the total area represents the fraction of the DNA molecules within this $\% (G + C)$ range of the total number of DNA molecules.

RESULTS

The results are summarized in Table 1. The $T_m$ value is an average of at least two determinations; $T_m$ values were in most cases reproducible to within $0.1^\circ$, exceptionally to $0.15^\circ$. $\sigma$ was usually reproducible to about $0.25 \% (G + C)$. The values given for the paper chromatographic method represent the average values of at least six estimations from two different extractions. The reproducibility of this method can be seen in Table 2, which illustrates the actual results obtained with three strains. The mean deviation on the ratio $(A + T)/(G + C)$ was 2.8, 2.9 and
3.7% for these three examples, which is the common range of deviation in quantitative paper chromatographic extractions. When the base composition, expressed as % (G+C), as determined by the paper chromatographic method, was plotted against the $T_m$ values of the same strains (see Fig. 2), the result was in good agreement with Marmur & Doty's (1962) formula $\% (G+C) = (T_m - 69.3)/0.41$. Two examples of thermal denaturation curves are given in Fig. 3. The compositional distribution of DNA molecules is represented in Figs. 4 and 5.

Table 1. Base composition, expressed as % (G+C), 'melting points' $T_m$ values and standard deviation $\sigma$ of the compositional distribution about the mean % (G+C) of purified DNA of various acetic acid bacteria

(Methods: see text)

<table>
<thead>
<tr>
<th>Biotype</th>
<th>Strain</th>
<th>Paper chromatographic method</th>
<th>Thermal denaturation method</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>% (G+C)</td>
<td>$T_m$</td>
</tr>
<tr>
<td>Acetobacter aceti</td>
<td>liquefaciens 20</td>
<td>65.4</td>
<td>95.6°</td>
</tr>
<tr>
<td></td>
<td>xylinum 25</td>
<td>—</td>
<td>95.1°</td>
</tr>
<tr>
<td></td>
<td>xylinoides 4940</td>
<td>—</td>
<td>94.95°</td>
</tr>
<tr>
<td></td>
<td>estunensis E</td>
<td>—</td>
<td>94.85°</td>
</tr>
<tr>
<td></td>
<td>xylinum 87-47</td>
<td>61.1</td>
<td>94.25°</td>
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<tr>
<td></td>
<td>mesoxydans var. saccharovorans 4</td>
<td>61.0</td>
<td>93.8°</td>
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<tr>
<td></td>
<td>pasteurianus 11</td>
<td>—</td>
<td>93.85°</td>
</tr>
<tr>
<td></td>
<td>acet. Ch31</td>
<td>59.5</td>
<td>92.8°</td>
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<tr>
<td></td>
<td>aceti var. muciparus 5</td>
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<td>93.75°</td>
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<td></td>
<td>mobilis 6428</td>
<td>58.8</td>
<td>93.5°</td>
</tr>
<tr>
<td></td>
<td>vini aceti 4989</td>
<td>—</td>
<td>92.9°</td>
</tr>
<tr>
<td></td>
<td>rancens 15</td>
<td>—</td>
<td>92.4°</td>
</tr>
<tr>
<td></td>
<td>cerinum var. rosiensis 22</td>
<td>56.5</td>
<td>92.2°</td>
</tr>
<tr>
<td></td>
<td>paradoxus P1</td>
<td>—</td>
<td>92.2°</td>
</tr>
<tr>
<td></td>
<td>rancens 23kl+</td>
<td>—</td>
<td>92.1°</td>
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<tr>
<td></td>
<td>paradoxus P2</td>
<td>—</td>
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</tr>
<tr>
<td></td>
<td>peroxydans 3</td>
<td>—</td>
<td>95.35°</td>
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<tr>
<td></td>
<td>peroxydans 4</td>
<td>—</td>
<td>95.35°</td>
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<tr>
<td></td>
<td>peroxydans 8618</td>
<td>61.0</td>
<td>95.0°</td>
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<tr>
<td>Gluconobacter oxydans</td>
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<td>62.1</td>
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<tr>
<td></td>
<td>suboxydans 26</td>
<td>61.0</td>
<td>94.85°</td>
</tr>
<tr>
<td></td>
<td>capsulatus 4943</td>
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</tr>
<tr>
<td></td>
<td>suboxydans 3734</td>
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</tr>
<tr>
<td></td>
<td>gluconicum 4739</td>
<td>—</td>
<td>94.55°</td>
</tr>
<tr>
<td></td>
<td>melanogenus 49</td>
<td>—</td>
<td>94.4°</td>
</tr>
<tr>
<td></td>
<td>melanogenus 8086</td>
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<td>94.2°</td>
</tr>
<tr>
<td></td>
<td>melanogenus 116</td>
<td>—</td>
<td>94.2°</td>
</tr>
<tr>
<td></td>
<td>suboxydans su</td>
<td>58.1</td>
<td>92.8°</td>
</tr>
</tbody>
</table>

Three other bacteria were included as controls of the methods. Pseudomonas fluorescens 488 had a DNA base composition of 59.5% (G+C), determined by paper chromatography, which is in good agreement with reported values for Pseudomonas in the literature (Lee et al. 1956; Marmur & Doty, 1962). Agrobacterium tumefaciens DNA had 58.8% (G+C), also determined by paper chromatography. For other strains 58.2 and 58.8% have been reported (Lee et al. 1956). For Escherichia coli a (G+C) % value of 50.0 was calculated from the $T_m$ value.
Table 2. Reproducibility of the chromatographic method for molar base composition determination

(All the values given represent the average of three spots from one chromatogram. Methods; see text. Abbreviations: A = adenine; T = thymine; G = guanine; C = cytosine. As an evaluation of the method, the values \( \frac{A}{T}, \frac{G}{C} \) and \( \frac{A+G}{T+C} \) (which ought to be equal to 1), as well as \( \frac{A}{G} \) and \( \frac{T}{C} \) (which ought to be equal), are presented.)

<table>
<thead>
<tr>
<th>Strain</th>
<th>( \frac{A}{T} )</th>
<th>( \frac{G}{C} )</th>
<th>( \frac{A+G}{T+C} )</th>
<th>( \frac{A}{G} )</th>
<th>( \frac{T}{C} )</th>
<th>( \frac{A+T}{G+C} )</th>
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<tbody>
<tr>
<td><strong>Gluconobacter oxydans</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Expt. 1</td>
<td>0.93</td>
<td>0.93</td>
<td>0.93</td>
<td>0.73</td>
<td>0.73</td>
<td>0.73</td>
</tr>
<tr>
<td>Expt. 2</td>
<td>0.96</td>
<td>0.93</td>
<td>0.94</td>
<td>0.75</td>
<td>0.73</td>
<td>0.74</td>
</tr>
<tr>
<td>Expt. 3</td>
<td>0.97</td>
<td>0.97</td>
<td>0.97</td>
<td>0.70</td>
<td>0.71</td>
<td>0.70</td>
</tr>
<tr>
<td><strong>Acetobacter aceti</strong></td>
<td></td>
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<tr>
<td>(liquefaciens 20)</td>
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<td></td>
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</tr>
<tr>
<td>Expt. 1</td>
<td>1.05</td>
<td>1.00</td>
<td>1.02</td>
<td>0.53</td>
<td>0.50</td>
<td>0.52</td>
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<tr>
<td>Expt. 2</td>
<td>1.05</td>
<td>1.02</td>
<td>1.03</td>
<td>0.53</td>
<td>0.51</td>
<td>0.52</td>
</tr>
<tr>
<td>Expt. 3</td>
<td>1.06</td>
<td>0.99</td>
<td>1.02</td>
<td>0.53</td>
<td>0.50</td>
<td>0.52</td>
</tr>
<tr>
<td><strong>Pseudomonas fluorescens</strong></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>(CCCEB 488)</td>
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<td></td>
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<tr>
<td>Expt. 1</td>
<td>0.93</td>
<td>1.08</td>
<td>1.02</td>
<td>0.65</td>
<td>0.75</td>
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<tr>
<td>Expt. 2</td>
<td>0.98</td>
<td>1.09</td>
<td>1.04</td>
<td>0.64</td>
<td>0.71</td>
<td>0.67</td>
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<tr>
<td>Expt. 3</td>
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<td>1.07</td>
<td>0.65</td>
<td>0.72</td>
<td>0.68</td>
</tr>
<tr>
<td>Expt. 3</td>
<td>1.05</td>
<td>1.03</td>
<td>1.04</td>
<td>0.64</td>
<td>0.66</td>
<td>0.66</td>
</tr>
</tbody>
</table>

Fig. 2. Relationship between the 'melting point' \( T_m \), determined by thermal denaturation, and the base composition \( \% (G+C) \) determined by paper chromatography, of purified DNA from several acetic acid bacteria. The numbers are taken from Table 1. The curve was drawn from Marmur & Doty's formula \( T_m = 0.41 (\% (G+C)) + 69.3 \).
Fig. 3. Thermal denaturation curves of purified DNA of two strains of acetic acid bacteria. The examples were selected to represent the strains with the lowest $T_m$ value (curve A, *Acetobacter aceti*, 'paradoxus' P2) and with the highest $T_m$ value (curve B, *A. aceti*, 'liquefaciens' 20). All other strains of Acetobacter and Gluconobacter had $T_m$ values within these limits. Each curve is the result of two different estimations. The results are expressed as relative extinction at 260 mp, being the ratio extinction at temp. $t$/extinction at 25°C.

Fig. 4. Approximation of the compositional distribution of the DNA molecules of strains of Acetobacter. The symbols pointing to each curve are the strain numbers (see Table 1). The position of the arrows indicates the mean % (G+C). The curves were calculated as described in the text.
**DISCUSSION**

All Acetobacter strains have base compositions of \((G+C)\) between 55.4\% (strain 'paradoxus' P2, 'rancens' 23 kl +) and 64.0\% (strain 'liquefaciens' 20). Nearly all Gluconobacter strains lay in a narrow cluster from 60.6\% (strain 'melanogenus' 116) to 63.4\% (strain 'viscosus' 8131). Strain 'suboxydans' SU with 57.2\% was the only one to be somewhat removed from the central cluster of Gluconobacter (Table 1; Fig. 5). These data are in agreement with the conclusion of De Ley (1961a) that each biotype *Acetobacter aceti* and *Gluconobacter oxydans* can be regarded as a cluster of strains, without inner sharp breaks, thus making species differentiation unnecessary. This is also clearly illustrated in Figs. 4 and 5 by the continuously overlapping curves of the compositional distribution. Incidentally, the range of the distribution of DNA molecules of each strain is broader in Acetobacter (average \(c\) 1.5) than in Gluconobacter (average \(c\) for the melanogenus strains: 1.06; for the other strains: 0.5).

On the basis of physiological, biochemical and enzymic data De Ley (1961a) noted that the strains of the *Acetobacter aceti* biotype could be arranged in a smooth gradation from the most complex one to the one with the poorest enzymic equipment. A comparison of the sequence of Acetobacter strains of Table 1 with the sequence of Fig. 5 of De Ley's (1961a) paper shows a noticeable agreement: strains with greater biochemical activities have on the whole also a higher \%(G+C)\. (De Ley & Schell, 1962, drew attention to the fact that the strain 'liquefaciens' 20 has to be removed...
from Gluconobacter and belongs in the biotype Acetobacter, where it is the most complex representative.) In Gluconobacter the base compositions are so close together that they cannot be arranged in a sequence of statistical significance. In Table 1 and Figs. 4 and 5, some strains are not where they would be expected according to their position in the biochemical sequence. This holds for ‘estunensis’, ‘pasteurianus’ 11, ‘suboxydans’ su and ‘peroxydans’ 3, 4 and 8618. However, it ought not to be expected that both sequences would agree completely. Indeed, the % (G+C) range encompasses the complete genotype and includes all properties, whereas the biochemical sequence was based mainly on carbohydrate metabolism. It might be that the strains ‘estunensis’, ‘pasteurianus’ 11 and ‘suboxydans’ su possess or lack some properties which have so far escaped examination. The peroxydans strains were expected to have very low base compositions in the vicinity of ‘paradoxa’ and ‘rancens’ in view of their biochemical similarities with these strains. The base composition of the ‘peroxydans’ strains is unexpectedly high. Fig. 4 shows that the ‘peroxydans’ strains have in fact only very few DNA molecules in common with the ‘paradoxa’ and ‘rancens’ strains. A similar situation holds for strain ‘suboxydans’ su, which is morphologically, physiologically and biochemically nearly indistinguishable from strain suboxydans 26, but nevertheless appears to have very few DNA molecules in common with it. Further work on the finer details of the DNA molecules in relation to the taxonomic position of these strains will be required.

As expected, Gluconobacter and Acetobacter strains have DNA with base compositions in the same range. These results support the hypothesis (De Ley, 1961a) which proposes that Gluconobacter and Acetobacter may have originated from a common pool of ancestors.

The DNA base compositions of the acetic acid bacteria are in the same range as those for Pseudomonas, which extend from 60 to 67 % (G+C) (Lee et al. 1956; Marmur & Doty, 1962). This stresses once more the suspected close relationship between the pseudomonads and the acetic acid bacteria. Phenotypically they have several features in common (Stanier, 1947; for a review see De Ley, 1961b). The similarity in the base compositions of the DNA molecules of both groups of bacteria also points to a possible genotypic relationship. It seems not unlikely that these bacteria derive from a common phylegetic origin. The comparison between the methods of paper chromatography and thermal denaturation for the estimation of DNA base compositions for routine analysis showed that the latter procedure was to be preferred; it is easier, faster, less elaborate and yields more reproducible results.

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DNA compositions of acetic acid bacteria

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