

# STUDIES ON THE CLASSIFICATION OF THE COLON-TYPHOID GROUP OF BACTERIA WITH SPECIAL REFERENCE TO THEIR FERMENTATIVE REACTIONS

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## INTRODUCTION

In recognition of the urgent need of progress along the lines of bacterial classification, the Society of American Bacteriologists at its meeting in Urbana appointed a special Committee on the Characterization and Classification of Bacterial Types. This Committee conceived its first duty to be a general revision of the families and genera of the Bacteriaceae, which was presented to the Society in preliminary form in 1917. In addition to this broad survey of the entire field the Committee projected a more intensive study of the colon-typhoid group, a group which, together with certain sharply defined species, includes many puzzling intermediate forms, difficult of classification, and yet of fundamental medical and sanitary importance. War work and other demands upon the time of the Committee have delayed the completion of this latter task, but certain members of the Committee, particularly Dr. Charles Krumwiede and Dr. L. A. Rogers have already published important individual contributions to the subject (Krumwiede, Pratt and Kohn, 1916a, 1916b, 1917; Krumwiede, Kohn and Valentine, 1918; Rogers, Clark and Evans, 1914, 1915; Rogers, Clark and Lubs, 1918). The present paper is the result of a detailed study of 160 cultures of colon-typhoid bacteria in the collection of the American

Museum of Natural History, conducted in large measure by the application of a series of standardized tests agreed upon by the Committee on Characterization and Classification of Bacterial Types. The laboratory work was for the most part carried out in 1915, 1916 and 1917 and preliminary reports upon its progress were made before the Society of American Bacteriologists in 1915 and 1916 (Winslow and Kligler, 1916; Winslow, Kligler and Rothberg, 1917). Final publication of the results has been delayed until time was available for a careful digestion of the data and a review of the literature concerning the classification of this group of bacteria.

#### THE MAJOR SUBDIVISIONS OF THE COLON-TYPHOID GROUP

*Primary division on the basis of lactose fermentation.* It seems reasonably certain that the typhoid bacillus was recognized in stained preparations by Eberth in 1880 and by Koch in the same year, and that Friedlaender saw *Bacterium pneumoniae* in 1882. General interest in this group of organisms dates however from the cultivation of the typhoid bacillus by Gaffky in 1884. This author described the organism as a motile rod, taking up anilin stains less readily than most bacteria, which failed to liquefy gelatin and gave a characteristic growth on potato and blood serum. Either from the presence of contaminating organisms, or a misinterpretation of irregular staining, he reported that it formed spores upon potato at 37°C.

In the following year Escherich (1885) described and figured *Bacterium coli* as a Gram-negative organism producing characteristic colonies and coagulating milk and forming gas in glucose broth. For the next decade attention was largely devoted to the establishment of distinctions between the types described respectively by Gaffky and by Escherich.

This differentiation was naturally based first of all on the production of gas in glucose media (facilitated by Smith's introduction of the fermentation tube in 1889) and on the formation of acid in lactose media. Petruschky (1889-1890) was probably the first investigator to introduce quantitative measurements of

acid production. He cultivated his bacteria in litmus milk and determined the amount of alkali necessary to produce neutrality. *B. suipestifer* gave an alkaline reaction, while the acidity produced by the other organisms studied was as follows: *B. typhosus* 0.2 to 0.3 per cent; *B. pneumoniae*, 0.3 to 0.4 per cent; *B. neapolitanus*, 0.7 to 0.8 per cent; *B. acidi-lactici*, 1.7 to 1.8 per cent. Chantemesse and Widal (1887) and Smith (1890) also pointed out that *B. typhosus* could be distinguished from *B. coli* by its less active fermentation of lactose. Chantemesse and Widal (1891), in controverting the claim of certain workers that *B. coli* and *B. typhosus* were identical, laid great stress on fermentation of sugars as a differential test and suggested the use of sugar broth containing calcium carbonate. They state that *B. coli* forms gas from glucose, sucrose, lactose, maltose, rhamnose, glycerol, erythritol and mannitol, but not from starch or glycogen; and they maintain that *B. typhosus* forms gas from none of these substances. Wurtz (1892) recommended the addition of an indicator to liquid or solid media for the easy separation of the two forms. Germano and Maurea (1893) considered the presence or absence of gas in 2 per cent glucose agar as the one absolutely diagnostic test.

In 1896 Capaldi and Proskauer suggested the use of two special media for the differentiation of the typhoid and colon groups. In medium I, containing 0.2 per cent asparagin and 0.2 per cent mannitol with the necessary salts, *B. typhosus* fails to grow and *B. coli* forms acid, while in medium II, containing 2.0 per cent peptone and 0.1 per cent mannitol, *B. typhosus* forms acid and *B. coli* does not. Medium I of course depends for its differential action on the inability of *B. typhosus* to utilize asparagin as a source of nitrogen, while the difference in the behavior of *B. coli* in the two solutions is presumably related to the balance of nitrogenous and carbohydrate foodstuffs provided. Durham (1901) urged the value of litmus whey as a test medium. Drigalski and Conradi (1902) reported that *B. coli* fermented arabinose and rhamnose while *B. typhosus* failed to do so; and they suggested a new medium for the differentiation of colon and typhoid colonies which, like the ordinary

lactose agar and the Endo (1904) medium, depends on the power of the former bacterium to produce acid from lactose. Meillère (1907) claimed that *B. typhosus* will attack inosite under aerobic conditions, while *B. coli* cannot ferment this substance.

Meanwhile many other distinctions between the typhoid and colon organisms had been described by various observers. Fremlin in 1893 differentiated *B. coli* from *B. typhosus* by its less active motility, fewer and less easily stained flagella, less tendency to occur in chains, more rapid and vigorous growth on various media (including the classical visible yellowish growth on potato), and the power to produce indol—as well as by the coagulation of milk and the formation of gas in glucose media. The active motility of the typhoid organism has since been made a basis for the selective media of Hiss (1902) and Hesse (Jackson and Melia, 1909), and for the isolation methods of Drigalski (1906) and Starkey (1906).

A new differential method of great importance was introduced in 1896 in the agglutination tests of Pfeiffer and Kolle and of Widal; and this test, too, quickly demonstrated the fundamental differences between the colon and typhoid organisms.

The more vigorous reducing power of the colon bacillus is another characteristic, which has been widely used in the form of the nitrate test. The greater ability of the colon bacillus to reduce dyes was suggested as a basis of differentiation by Dunbar (1892) using litmus, and by Rothberger (1898) using neutral red. Lead acetate on the other hand is turned brown by *B. typhosus* and not by *B. coli*, as pointed out by Orłowski, Saquépée and Chevrel (1905) and Burnet and Weissenbach (1915).

Finally the attempt to devise methods for the isolation of the typhoid bacillus has led to the recognition of a number of more or less specific differences in the tolerance of *B. typhosus* and *B. coli* toward a wide variety of antiseptic substances. The acid broths used for preliminary enrichment in the attempt to isolate the typhoid bacillus from water were designed merely to inhibit the ordinary water bacteria; but Hankin (1899) pointed out that within the colon-typhoid group *B. coli* could bear a

higher concentration of acid than *B. typhosus*. Winslow and Lochridge (1902) showed that the toxic action of the mineral acids was due to their free hydrogen ions and presented the following table of the comparative resistance of the colon and typhoid organisms.

*Disinfectant action of mineral acids*

	PARTS OF DISSOCIATED HYDROGEN PER MILLION NECESSARY TO EFFECT			
	99 per cent reduction		100 per cent reduction	
	HCl	H <sub>2</sub> SO <sub>4</sub>	HCl	H <sub>2</sub> SO <sub>4</sub>
<i>B. typhosus</i> .....	2.94	2.54	4.85	3.90
<i>B. coli</i> .....	7.49	7.68	12.80	12.60

Loeffler (1903, 1906) and Lentz and Tietz (1903, 1905) reported that *B. typhosus* was more resistant than *B. coli* to the action of malachite green. A whole group of green dyes seem to be favorable for the selective cultivation of the typhoid bacillus and its allies. Werbitzki employed China green for this purpose and Conradi, brilliant green. The whole group of the green dyes has been studied very carefully by Krumwiede and Pratt (1914). They found brilliant green most satisfactory for this purpose, the paratyphoid-enteritidis forms being most resistant, *B. typhosus* coming next, and the colon-aerogenes group being most readily inhibited. Roth (1903) and Hoffman and Ficker (1904) maintained that *B. typhosus* was more resistant to caffeine than the colon bacillus. Nowack (1905) thought that a slightly alkaline reaction exerted a selective influence on the typhoid organism. Jackson and Melia (1909) found *B. typhosus* more resistant to the action of bile salts than *B. coli*.

Davis (1914) has studied the resistance of the members of the colon-typhoid group to potassium tellurite, and finds that the colon-aerogenes forms are most resistant, followed in succession by the paratyphoid B and Gaertner group, *B. typhosus*, *B. paratyphosus* A and the dysenteries. Capsulated organisms vary widely in susceptibility, *B. capsulatus* appearing at the top and *B. pneumoniae* and *B. rhinoscleromatis* nearly at the bottom of

the list. *B. acidi-lactici* somewhat surprisingly proves one of the sensitive organisms, while *B. zopfi* is most susceptible of all.

Manfredi (1917) reports that a concentration of cholesterol of 0.5 per cent will check typhoid and the paratyphoids, while *B. coli* will grow in concentrations up to 1 per cent.

During the decade following the first cultivation of the typhoid and colon bacilli it was made clear that these organisms belong to a large group of Gram-negative non-spore-forming bacilli. They are plump straight rods with rounded ends which grow under both aerobic and anaerobic conditions on ordinary media producing colonies which vary from a thin translucent irregularly notched "grape-leaf" form to a more compact and opaque, very slightly yellowish, colony, and possessing more or less marked powers of decomposing carbohydrate materials.<sup>1</sup>

This main group may be sharply split into two primary subdivisions by the presence or absence of the power of fermenting lactose. The lactose fermenters are generally more vigorous in their attack on other carbohydrates as well. They are less actively motile. They grow more abundantly on media and exert a more powerful reducing action. They generally produce indol. They are as a rule of low pathogenic power. The non-lactose-fermenters differ in all of the characteristics cited; and each type has its specific agglutinative reactions and its more or less specific degree of tolerance of various antiseptic substances.

#### SECONDARY SUBDIVISION OF THE NON-LACTOSE FERMENTERS

Considering next the further subdivision of the typhoid or non-lactose-fermenting series, the first important step was the discovery that certain members of this series possessed the power of forming gas in glucose media. *B. suipestifer* discovered

<sup>1</sup> Our whole conception of the morphology of this group of organisms will be radically altered if the studies of Hort (1917) are confirmed. This investigator describes in acid broth (and to some extent in ordinary broth) cultures of typhoid, paratyphoid, dysentery and colon organisms extraordinary reproductive stages, characterized by budding, terminal, lateral, or superficial, with the production of cruciform and radial cell aggregates, and he believes that detached buds may constitute a filterable stage in the life cycle of the organisms.

by Salmon and Smith in 1885 and *B. enteritidis*, demonstrated as the cause of an outbreak of meat poisoning by Gaertner in 1888, were the first organisms of this type to attract attention. Smith (1893a) pointed out that *B. suispestifer* produced gas in glucose but not in lactose media, and Durham (1898) noted that the Gaertner bacillus differs from the typhoid bacillus, not only in its power to form gas but in the lower final acidity in lactose media. The table below presented by him is an exceedingly interesting one as a product of a period when quantitative methods were so rarely used in bacteriological investigations.

*Acidity, per cent normal acid*

MEDIUM	PERIOD (AT 37°C.)	B. TYPHOSUS	B. ENTERI- TIDIS	B. COLI
Neutral.....	24 hours	0.25-0.35	0.2-0.3	0.6-0.8
Litmus whey.....	4-5 days	Less than 0.6	-0.15	Over 1
Litmus medium.....	24 hours	0.15	0.3	0.5
Litmus medium containing 2 per cent peptone.....	Several days	0.6	0.35	-0.2
Litmus medium containing 1 per cent glucose.....	Eventually	Generally alkaline	Alkaline	Alkaline

In a later communication Durham (1901) shows that litmus whey is of special value for differentiating these forms, *B. enteritidis* producing like *B. typhosus* an initial acid reaction, but unlike *B. typhosus* causing a subsequent change to alkalinity.

Meanwhile other types of colon-typhoid intermediates were being reported from various sources. Gwyn in 1898 described a case of a disease clinically like typhoid fever from which he isolated an organism resembling *B. enteritidis*, which he called a "paracolon" form (using a term first introduced by Gilbert in 1895). Cushing (1900) described a similar organism, which he called Bacillus 0, and like Durham clearly recognized the existence of a series of intermediate forms, resembling *B. typhosus* in morphology and motility, in the possession of pathogenic power, in failing to ferment lactose or produce indol, and in

relatively meager growth, but resembling *B. coli* in producing gas in glucose and differing from both in producing a transient acid reaction in milk with later reversion to alkalinity.

In 1901 Schottmüller published his classical paper on the paratyphoids. Six different strains of intermediates were worked out in detail. It was shown that these forms like those studied by Durham, Gwyn and Cushing were characterized by active motility, growth intermediate in luxuriance between that of *B. coli* and that of *B. typhosus*, gas production in glucose media, and reduction of neutral red. From the typhoid bacillus they were also distinguished by colony structure on Piorkowski's alkaline urea-gelatin, by the decolorization of an indigo compound and by a peculiar transparency in milk due to alkali production. In litmus-whey he distinguished two types of the "intermediates," one maintaining an acid reaction, the other changing to a blue-violet color. These forms, which we now know as the paratyphoids A and B, were also differentiated by their growth on media and their agglutination reactions.

By the early nineties then the non-lactose-fermenters had been separated into two clearly defined subgroups which may most conveniently be distinguished by the presence or absence of the power to produce gas in glucose media.

#### FURTHER SUBDIVISION OF THE NON-LACTOSE FERMENTING ORGANISMS

Each of the two groups, to which reference has been made above, was further subdivided during the early years of the twentieth century.

The first step in the differentiation of types among the bacilli which fail to form gas in either glucose or lactose was the discovery of the bacillus of dysentery by Shiga in 1898. Similar organisms were isolated by Kruse in 1900 and by Flexner and by Strong and Musgrave in the same year. Flexner (1901) distinguished the dysentery from the typhoid organisms by less active motility, general tendency to produce indol, secondary alkaline reaction in milk, and characteristic agglutination reac-



tions. After a long controversy as to the specificity of the various dysentery organisms the investigations of Martine and Lentz (1902), Duval and Bassett (1902), Hiss and Russell (1903), Park, Collins and Goodwin (1903) and others made it clear that there were at least two quite distinct types of dysentery bacilli, the Shiga type which ferments only the monosaccharides and forms no indol and a second type including the Flexner, Strong and Hiss strains which produces indol and ferments both glucose and mannitol and in some cases other carbohydrates as well. The Flexner types are most nearly allied by their fermentative reactions to *B. typhosus*, while the Shiga strains represent a group of exceedingly limited metabolic powers. J. H. Smith (1915) is, we believe, correct in using the fermentation of mannitol for the primary classification of the forms which produce no gas in glucose, and we shall follow him in recognizing as the first two subdivisions of the colon-typhoid series:

*Group I.* Organisms fermenting no carbohydrates except the simple hexoses (including here for convenience *B. alcaligenes* as well as the Shiga type of dysentery).

*Group II.* Organisms fermenting the hexoses and mannitol and certain other carbohydrates; forming only acid but no gas (including *B. typhosus* and the Flexner and other mannitol fermenting types of dysentery bacilli).

Turning now to the paratyphoids themselves, the organisms which ferment glucose with gas production but fail to attack lactose, it will be remembered that Schottmüller even in 1900 recognized two distinct types of paratyphoid organisms according to their reaction in litmus milk. Kayser (1904) emphasized this distinction more clearly and showed that Paratyphoid B resembled most closely the organisms of the Gaertner food poisoning group. Morgan (1905) studied 21 B strains and 10 A strains, the former producing a permanent acid reaction in milk and forming indol, while the latter turned the milk first acid and then alkaline, failed to form indol and gave agglutinative reactions with sera of the *B. enteritidis* group. Boycott (1906) was perhaps the first investigator to point out that even with the A strains milk cultures eventually became alkaline. This

conclusion was confirmed by Bradley (1912), Krumwiede, Pratt and Kohn (1916b), Hadley (1917) and others, and Jordan (1918) shows that the difference between the two types is probably due merely to a difference in their rate of multiplication. Buxton (1902), Bainbridge (1909) and others emphasize the less vigorous action of the A strains in glucose media, and point out that with neutral red these organisms often fail to show reduction, which is always present with the B strains. Buxton noted the peculiar opalescence in milk cultures of the B organisms and attributed it to a solution of casein caused either by the alkalinity or by the presence of a casease.

Aside from the quantitative difference in their action upon milk, the A and B divisions of the paratyphoid series exhibit other distinct differences in fermentative power. Ford (1905) concluded that *B. suipestifer* fermented only rhamnose, manitol, and dulcitol besides the hexoses, while Schottmüller's paratyphoid A fermented arabinose also and *B. enteritidis* arabinose and xylose. Later investigations indicate that *B. suipestifer* does attack xylose and that dulcitol fermentation is generally absent in the A paratyphoids and *B. suipestifer*. The difference in dulcitol fermentation was noted particularly by Ditthorn (1913). The fact that *B. suipestifer* as well as paratyphoid B and *B. enteritidis* all ferment xylose, while paratyphoid A does not, was clearly shown by Harding and Ostenburg (1912) and Krumwiede, Pratt, and Kohn (1916a). Weiss and Rice (1917) and Hulton-Frankel (1918) point out that fermentation of inosite furnishes another differential test, paratyphoid B attacking this substance while paratyphoid A fails to do so. *B. suipestifer*, and curiously enough *B. enteritidis*, also resemble the A strains in this respect. Exhaustive studies by Krumwiede, Pratt and Kohn (1917), Jordan (1917), and Krumwiede, Kohn and Valentine (1918) have confirmed the fundamental fact that the paratyphoid group may be subdivided in the following manner on the basis of fermentative power.

	XYLOSE	ARABINOSE	DULCITOL	INOSITE
Para A.....	-	+	±	-
<i>B. suipestifer</i> .....	+	-	±	-
<i>B. enteritidis</i> .....	+	+	+	-
Para B.....	+	+	+	+

Another interesting line of differentiation between the two main subdivisions of the paratyphoid group is the color reaction in media containing lead or iron salts, first suggested by Orłowski. Sacquépée and Chevrel (1905) pointed out that on a gelatin medium containing the double tartrate of iron and potassium, or subacetate of lead, paratyphoid B and *B. typhosus* produce a black coloration (due presumably to precipitation of lead sulphide), while paratyphoid A and *B. coli* fail to do so. Recently Burnet and Weissenbach (1915), Weissenbach (1917), Kligler (1917), and Jordan and Victorson (1917) have confirmed these results, which occur with marked regularity, and the latter investigators show that, as in most respects, *B. suipestifer* behaves like paratyphoid A.

A similar color reaction, suggested by Orłowski, and used by Sacquépée and Chevrel (1905) and Weissenbach (1917), is the production of a green coloration in gelatin containing nitroprussiate of sodium. In such a medium paratyphoid B produces an intense green coloration in forty-eight hours and *B. typhosus* in three days; while the reaction fails entirely with *B. coli* and paratyphoid A.

The various investigations cited make it clear that we may recognize at least two more groups of the non-lactose-fermenters (in addition to those cited on p. 437).

*Group III.* Organisms fermenting glucose and the other hexoses with the formation of gas; fermenting mannitol and rhamnose and arabinose, in a similar way, but failing to ferment xylose and lactose and usually failing to ferment dulcitol; producing an acid reaction in milk which persists for from five days to six weeks; this group includes the A paratyphoids.

*Group IV.* Organisms differing from members of group III in fermenting xylose and generally dulcitol; sometimes failing to attack arabinose; producing a transient acid reaction in milk which turns to alkaline in two to five days; this group includes *B. suispestifer*, *B. enteritidis*, paratyphoid B and their allies.

#### SECONDARY SUBDIVISION OF THE LACTOSE-FERMENTERS

The non-lactose-fermenters may be separated, as indicated above, into four secondary groups which are rather clearly differentiated in each case by definitely correlated bio-chemical properties, as well as characteristic serological and pathogenic relations. The classification of the lactose-fermenting division of the colon-typhoid series has proved a much less simple task.

Escherich (1885) recognized two distinct types of organism in this group, *B. coli-commune* which he described as a Gram-negative rod producing characteristic colonies, gas in glucose media, and slow coagulation of milk, and *B. lactis-aerogenes* which he differentiated by the plumper form of the cells, lack of motility, and more rapid coagulation of milk. These differences seem somewhat slight but they have been shown by later observers to be correlated with much more fundamental characters. Smith (1893a) pointed out the heavier growth and tendency to capsule and zooglea formation on the part of *B. aerogenes*. In a later paper (Smith 1895a) he emphasizes the rapid gas production and relatively low acidity characteristic of this organism, and points out that the ratio of CO<sub>2</sub> to H<sub>2</sub> is higher in some strains of *B. aerogenes* (and in all the strains of *B. cloacae* studied) than is the case with *B. coli*. Russell and Bassett (1899) also laid stress on the difference in the CO<sub>2</sub> to H<sub>2</sub> ratio and suggested that the forms producing the larger proportion of CO<sub>2</sub> were not of intestinal origin, but were perhaps normal soil bacteria. Durham (1901) noted that the *B. aerogenes* types fermented starch and inulin and gave the peculiar reaction described by Voges and Proskauer in 1898. Grimbert and Legros in 1900 and Jordan (1903) pointed out that organisms of the *B. aerogenes* series frequently fail to form indol. A most important step in the

classification of this group was taken when Harden and Walpole (1905), and Rogers, Clark and Davis (1914) accurately measured the ratio of  $\text{CO}_2$  to  $\text{H}_2$  (which had been roughly estimated in an open fermentation tube by Smith and the other workers cited above); and showed that the colon forms actually produce these two gases in approximately equal proportions, while with the aerogenes type the true ratio of  $\text{CO}_2$  to  $\text{H}_2$  is about 2 to 1. Clark and Lubs (1915) discovered that the two types may be easily distinguished by a difference in the acid reaction produced in sugar broths of specified composition, and introduced the simple methyl-red test for measuring this difference. These investigators and others to be cited later have made it clear that in a number of respects the two types mentioned are sharply differentiated, so that we may recognize two more distinct groups of the colon-typhoid series, characterized as follows:

*Group V.* Organisms fermenting the hexoses, mannitol, xylose, rhamnose, arabinose and lactose, frequently salicin, dulcitol, sucrose, raffinose and glycerol; producing a strong acid reaction in media containing these substances and a moderate amount of gas, composed in equal parts of  $\text{CO}_2$  and  $\text{H}_2$ ; producing indol and failing to give the Voges-Proskauer reaction; generally motile and not capsulated; never liquefying gelatin.

*Group VI.* Organisms differing from Group V in producing a lower acidity in carbohydrate media and a larger proportion of gas, of which two thirds is  $\text{CO}_2$ ; in fermenting dulcitol and glycerol less frequently and adonitol, salicin, starch, sucrose, raffinose and inositol more frequently; in giving a generally negative indol and a constantly positive Voges-Proskauer reaction; in being generally non-motile and frequently capsulated; and in sometimes liquefying gelatin.

The distinction between these two groups is a very sharp and clean cut one, as indicated by the large number of correlated characters in which they differ. Lines of demarcation within these groups of the lactose-fermenting organisms are unfortunately much less clear. Smith as early as 1893 recognized that among the low ratio (*B. coli*) strains there are some which attack sucrose and some which do not, and that among the high ratio forms *B. cloacae* (as described by Jordan in 1890) has the

power of effecting a sluggish liquefaction of gelatin. Durham in 1901 limited the term *B. coli-communis* to the sucrose-negative form and named the sucrose-positive form *B. communior*. Winslow and Walker (1907) pointed out that the fermentation of raffinose varies together with that of sucrose; but this is perhaps less a correlated character than an independent measurement of the same character since these two sugars have a similar molecular configuration. As a matter of fact no very striking correlations are apparent between any of the characteristics of these organisms, and the attempts made by MacConkey (1905) Jackson (1911) and others to classify them according to their fermentative reactions are purely arbitrary and artificial. The same thing is true of classifications like those of Ford (1903) and Jordan (1903) in which the liquefaction of gelatin is given an important place. MacConkey (1905) laid special stress upon dulcitol for differential purposes, Kligler (1914) upon salicin and glycerol, while Levine (1917) has made perhaps the most important contribution to the whole problem. Certain types recognized by Levine will be discussed later on, but for the purposes of a general subdivision of the colon-typhoid series it is perhaps only necessary to point out that group V as defined above includes a sucrose-positive and a sucrose-negative subgroup, while group VI includes forms which liquefy gelatin and others which fail to do so.

#### THE VALUE OF CARBOHYDRATE FERMENTATION TESTS AS A BASIS FOR BACTERIAL CLASSIFICATION

It will be noted that almost all observers who have attempted to differentiate and classify the organisms of the colon-typhoid series have laid primary stress upon the fermentative reactions of these organisms in carbohydrate media, and it is important to know whether such reactions are of sufficiently fundamental biological importance to warrant such an emphasis.

The results obtained by certain investigators have tended to cast some doubt upon the constancy and reliability of fermentation tests as applied to the members of this group. In the first place intermediate forms have been described which differ from typical ones either in exhibiting a slow but ultimately distinct

action upon a particular carbohydrate, or in displaying a combination of fermentative characters the reverse of that which normally occurs. Thus Wilson (1910) describes a curious form isolated from the urine of a carrier which at 37° produced no acid in lactose media and but little gas in mannitol and maltose, while at 20° it formed an abundance of acid in lactose and an abundance of gas in mannitol, maltose and salicin media. Glucose and sucrose always showed acid but no gas. Oette (1913) notes a peculiar strain which possessed the general fermentative reactions of the paratyphoids and showed specific agglutination with paratyphoid B but produced no gas. Raubitschek and Natonek (1913) isolated 31 different strains of typhoid bacilli from the various organs of two patients and found marked quantitative differences in fermentative power between them. Tenbroeck (1916) describes a variant arising in an old culture of *B. suispestifer* which retained all its normal cultural and immunological characteristics except that it no longer produced gas in glucose and failed to reduce neutral red. Smirnow (1916) exposed a series of colon strains to the influence of strong glucose solutions (3 per cent), NaCl (4 per cent), Na<sub>2</sub>SO<sub>4</sub> (1.5 per cent) and phenol (0.25 to 0.75 per cent). After successive transfers, covering a period of one to three months, he found a marked suppression of biochemical activities, glucose and phenol exhibiting the most marked effects. Indol formation was first affected, then gas production in various carbohydrate media, then the characteristic growth on potato, then the coagulation of milk, and finally the production of acid in carbohydrate media. The cultures generally reverted to their normal characteristics on prolonged cultivation in ordinary media, but some of the induced modifications proved permanent. Hadley, Caldwell, Elkins and Lambert (1917) recognize two varieties of *B. pullorum*, one of which produces gas while the other fails to do so. Oliver and Perkins (1918) report the case of a dysentery-like organism which under ordinary conditions attacked glucose only but which when cultivated under diminished oxygen tension acidified galactose, maltose, levulose, mannitol, lactose and sucrose. It is very probable, as shown by recent work, that results were complicated by the buffer action of CO<sub>2</sub>. Bron-

fenbrenner and Davis (1918) have isolated a number of organisms which when first studied fermented lactose very slowly, but whose power to attack this sugar could be progressively increased by cultivation in lactose media.

Such differences as those described may be in large measure explained as the result of a direct response to different environmental conditions, or as the normal quantitative variations exhibited by all biological reactions. In using any biochemical characteristic for differential purposes it is necessary to specify a standard set of conditions and to use modal points in the curve of distribution of quantitative values, rather than arbitrary limits, in defining groups. In connection with some of the irregularities observed it must be remembered that the complex sugars used by bacteriologists frequently prove on careful analysis to contain hexose-impurities; and that if not originally present the hexoses may be produced by excessive autoclaving. Another type of investigation indicates however that the causes of observed variations may be of a more deep seated nature. Twort (1907) studied the action of 18 colon-typhoid organisms on 49 different glucosides, and on the basis of the results obtained concluded that no significant differences existed and that all the organisms studied were "varieties or hybrids of one or more species." He then tried to produce modifications of fermentative power by cultivating various strains for long periods in media containing carbohydrates which they would not at first utilize, and found that they slowly acquired new fermentative powers when treated in this way. All the members of the paratyphoid group acquired the power to ferment sucrose, *B. typhosus* began to ferment dulcitol and lactose, the dysentery bacilli of Kruse and Flexner were able to ferment sucrose within twenty-four hours. Neisser (1906), Massini (1907), Burk (1908), Mueller (1908, 1909), and Penfold (1910, 1911a, 1911b, 1912) have described what appear to be clear-cut and definite cases of mutation in this group of organisms. The phenomena recorded are in general as follows: an organism cultivated on a solid medium containing a carbohydrate, which it normally fails to attack, produces colonies which at first appear (as indicated by the reaction of the indicator contained in the medium) as of the



usual non-fermenting type. Later however papillary projections appear on the colony, which are red if litmus be present, and plates made from these papillae yield a certain proportion of colonies which from the first exhibit vigorous fermentative power. Penfold (1910) studied Twort's lactose-fermenting strain of *B. typhosus* and found that it readily threw off non-fermenting mutants. In general however he found the fermentation of lactose to be a fairly stable character but dulcitol and rhamnose exhibited the phenomenon of mutation in a large number of strains. According to Mueller (1908, 1909), of 120 strains of *B. typhosus* all produced rhamnose-fermenting mutants. Mueller and Penfold (1912) both observed raffinose-fermenting mutants in colonies of members of the paratyphoid group. Of 34 strains of non-lactose-fermenters isolated by Penfold from feces, 21 produced fermenting mutants of some sort. The relation between the phenomenon of papilla-formation and the gradual increase in fermentative power, often observed (as by Bronfenbrenner) in liquid media, is well brought out in one of Penfold's investigations (1911a). In a dulcitol broth culture of *B. typhosus* which became acid to litmus after ten days of incubation he found, by plate cultures made from day to day, that at first fermenting strains were rare but that their number suddenly increased after the first week. Successive transfers in dulcitol broth produced a mass culture which turned litmus broth red in one day, presumably as a result of progressive selection of the dulcitol-fermenting mutant. In a later paper Penfold (1911b) shows that cultivation of certain colon and paratyphoid strains on chloracetic-acid agar produces varieties which have lost all power of producing gas in sugar media, while the fermentation of the alcohols is unaffected. Revis (1911) notes a similar phenomenon, a colon organism grown in presence of 0.05 per cent malachite green losing the power of gas production and after more prolonged treatment failing to coagulate milk and to form acid in dulcitol. Ledingham (1918) reports two highly variable strains of Flexner-Y dysentery, one of which produced mutants fermenting rhamnose and arabinose, while the other normally acidified these sugars but produced a mutant which failed to do so.

A general survey of this evidence indicates that the fermentative characters of the colon-typhoid bacteria are not only influenced by environmental conditions but also exhibit inherent powers of spontaneous mutation; yet it does not materially weaken the general value of tests of fermentative ability for systematic purposes. It is significant that spontaneous mutations are most common with dulcitol and rhamnose, carbohydrates which have been shown in comparative studies to lack the correlations with other characters which are exhibited by lactose, sucrose, xylose, and other substances; and that other modifications, such as the suppression of gas production, have generally been associated with strikingly abnormal environmental conditions. Taking the great mass of colon-typhoid strains, as they are isolated from the bodies or intestines of men and animals, and cultivated under standard conditions, fermentative characters exhibit a high degree of constancy and what is even more important a high degree of correlation with other bio-chemical and serological and pathogenic properties. It is no accident that the disease-producing organisms of the colon-typhoid series are practically without exception organisms which fail to produce gas in lactose media, but a law, which can have its basis only in the principle of phylogenetic relationship; and the same principle holds for numerous correlations which have been cited in the description of the six principal subdivisions of the colon typhoid series which have been recognized in the preceding section. Even Twort's studies, when properly analyzed, exhibit clear evidence of the general relationships of the chief groups of organisms concerned and of their progressive increase in fermentative power in proceeding from one end to the other of the series. His *B. alcaligenes* strains attacked none of the glucosides studied. With dysentery and typhoid organisms 7 per cent of his glucoside tests were positive; with the paratyphoid and Gaertner forms, 15 per cent; with colon bacilli, 46 per cent; and with *B. aerogenes* types, 65 per cent. We are dealing here with a group in which the line of evolutionary development has been marked most clearly by the progressive loss or progressive acquisition of fermentative power. Intermediate types appear at times and mutants may occur; but on the whole,

the series of forms between *B. alcaligenes* at one extreme and *B. aerogenes* at the other may be broken up into six or more main groups on the basis of fermentative ability; and the groups so constituted are true biologic entities marked by a number of independent correlated characters which can only have their origin in phylogenetic relationship.

THE CHEMISTRY OF THE FERMENTATIVE REACTIONS OF THE COLON-TYPHOID GROUP

*The gaseous products of carbohydrate decomposition.* The production of gas is the most obvious and striking evidence of carbohydrate decomposition and ever since Smith (1890) introduced the fermentation tube into bacteriology the appearance of visible gas in the Smith or Durham tube has been used as one of the most common criteria of fermentative activity. It is obvious that such a test is but a rough and approximate one and that no sharp line can be drawn between gas-producers and non-gas-producers, since a certain amount of gas will be necessary to saturate the liquid medium in the tube before excess gas will collect and become visible in the closed arm. Clark (1913) criticised Penfold's emphasis on the non-gas-forming mutants produced by the influence of chloracetic acid on this ground, and has pointed out that since normal *B. coli* produces a smaller amount of gas in glucose and galactose media than in dulcitol and mannitol the suppression of gas formation in the former case and not in the latter is perhaps merely due to a general weakening of the organisms concerned. The method of studying carbohydrate fermentation by examining the gas which collects over a liquid, which is exposed freely to the air at another point, is still less suited for careful studies of the composition of the gases present. Smith in his classic investigations laid stress not only upon the proportion of CO<sub>2</sub> to H<sub>2</sub> (which he reported as 1:2 for *B. coli*—Smith 1890) but also upon the quantitative determination of the total amount of gas present (which he states as 40 to 70 per cent for *B. coli*, *B. proteus* giving less and *B. cloacae* more,—Smith 1893b). It seems obvious that the amount and composition of the gas collected in the Smith tube will be governed almost as much by the solubility of the gases produced, and the rate at

which they diffuse outward from the free surface of the liquid medium, as by the inherent biochemical activities of the organisms which produce them, and measurements made in the Smith or Durham tube will always be accompanied by the loss of a large and a variable proportion of the CO<sub>2</sub> actually produced.

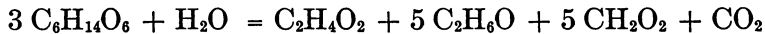
As a matter of fact, even before the introduction of the fermentation tube into bacteriology, the real composition of the gases produced in bacterial fermentation had been established by several observers who used the proper methods to secure a sample of all the gas actually produced, by cultivation in closed bulbs which were later evacuated by the air pump. As early as 1887 Hoppe-Seyler observed that calcium formate infected with river mud yielded a gas composed in equal volumes of CO<sub>2</sub> and H<sub>2</sub>. Frankland and Fox (1889) were perhaps the first investigators to work with a pure culture, using a form which they isolated from sheep dung and named *B. ethaceticus*. This organism was said to be a gelatin liquefier but curiously enough gave the fermentative reactions of *B. coli* since Frankland and Lumsden (1892) and Frankland and MacGregor (1892) report an equal volume of CO<sub>2</sub> and H<sub>2</sub> produced in glucose, mannitol, and arabinose. Macfadyen, Nencki and Sieber (1891) reported that the gas produced by *B. aerogenes* was 72 per cent CO<sub>2</sub> and 28 per cent H<sub>2</sub>. Frankland, Stanley and Frew (1891) estimated a ratio of CO<sub>2</sub> to H<sub>2</sub> of 10 to 8 for *B. pneumoniae*. Pakes and Jollyman (1901) also reported the 1:1 ratio in the decomposition of sodium formate by paratyphoid, colon and aerogenes types. By far the most exhaustive studies of this question were made, however, by Harden (1901, 1905) who placed the distinction between the 1:1 ratio for *B. coli* and the ratio of 2 or more parts of CO<sub>2</sub> to 1 of H<sub>2</sub> for *B. aerogenes* on a definite and solid basis.

In this country Bennett and Pammel (1896) showed that the gas formed in the Smith tube is at first only H<sub>2</sub> and that the ratio between H<sub>2</sub> and CO<sub>2</sub> depends mainly on the absorption of CO<sub>2</sub> by the liquid medium; and Longley and Baton (1907) made clear the slight value of measurements of the amount of gas and the ratio of its components in the open tube. Keyes (1909) and Keyes and Gillespie (1913) made the first exact studies in America of the actual gas production by exhaustion methods and

Rogers, Clark and Lubs extended and confirmed the work of Harden and broadened the basis of the radical distinction between the high and low ratio groups.

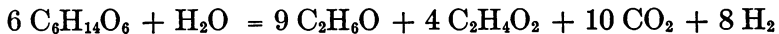
*The acid products of carbohydrate decomposition.* Studies of gas production have been given first place in this review of the subject because of the fact that gaseous products are most easily measured and have therefore attracted more general attention. A real comprehension of the processes involved demands however a knowledge of the soluble products as well.

In the classic studies of Frankland and his associates on *B. ethacetica* it was shown that this organism when decomposing mannitol and glycerol produces chiefly ethyl alcohol and acetic acid, with smaller amounts of formic and succinic acids, the ratio of alcohol to acetic acid being 2 molecules to 1 in the case of mannitol and 3 to 1 in the case of glycerol. (Frankland and Fox, 1889). In later papers (Frankland and Lumsden, 1892, Frankland and MacGregor, 1892) it is stated that the proportion of acetic acid is higher with glucose than with the alcohols mentioned above, still higher with arabinose and highest of all with glyceric acid. When the fermentation takes place in a closed space a considerable amount of formic acid accumulates and a formula for the decomposition of mannitol is suggested as follows:



In the open tube it is assumed that the formic acid is promptly decomposed into  $\text{H}_2$  and  $\text{CO}_2$ .

With *B. pneumoniae* under the conditions of ordinary cultivation Frankland, Stanley and Frew (1891) give a slightly different formula for the decomposition of mannitol



Frankland and his associates apparently failed to lay emphasis on the non-volatile acids, although they suggest the presence of traces of a fixed acid (probably succinic). Macfadyen, Nencki and Sieber (1891) however report ethyl alcohol and acetic and lactic acids in glucose broth cultures of colon-group organisms, the proportion of acetic acid being less in the case of *B.*

*aerogenes* than in cultures of *B. coli*. Grimbert (1896) gives the results tabulated below for the Friedlaender bacillus and *B. coli*.

*Products of permanent activity*

	GRAMS FORMED PER 100 GRAMS OF CARBOHYDRATE FERMENTED					
	B. pneumoniae				B. coli	
	Mannitol	Dulcitol	Arabinose	Xylose	Glucose	Lactose
Ethyl alcohol.....	11.4	29.3	0.0	6.9	Trace	6.8
Acetic acid.....	10.6	9.6	36.1	23.4	14.3	25.4
Laevo lactic acid.....	36.6	0.0	49.9	Trace	42.7	Trace
Succinic acid.....	0.0	21.6	0.0	19.9	Trace	29.8

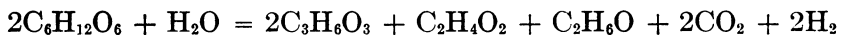
Duchacek (1904) found that in media exposed to air *B. coli* produced approximately equal amounts of lactic and acetic acids, but that under an atmosphere of hydrogen the proportion of lactic acid increased to 2:1. *B. typhosus* formed a considerably higher proportion of lactic acid than *B. coli*.

Harden's work constitutes by far the most extensive investigation into this question. Harden and Walpole (1906) give the following results of the fermentative activity of *B. coli* and *B. aerogenes*.

*Products of fermentative activity*

	GLUCOSE		MANNITOL	
	B. coli	B. aerogenes	B. coli	B. aerogenes
Per cent of carbohydrate fermented				
Alcohol.....	12.8	18.2	28.1	32.5
Acetic acid.....	18.8	8.6	9.5	2.1
Lactic acid.....	31.9	9.1	18.6	8.6
Succinic acid.....	5.2	4.5	8.9	2.8
Formic acid.....	0.0	1.7	3.0	1.6
Carbon dioxid.....	18.1	35.2	28.4	35.5
Cubic centimeters per gram				
CO <sub>2</sub> .....	91.8	178.5	143.0	180.3
H <sub>2</sub> .....	110.0	92.4	167.0	143.6
Ratio H—CO <sub>2</sub> .....	1.19	0.52	1.18	0.79

The results for *B. coli* with glucose would correspond roughly to the following formula:



Levulose behaves in the same way; and the general proportion of the end-products in the case of the hexoses and the alcohols is explained in the following scheme.

HEXOSE-GLUCOSE		PRODUCTS	ALCOHOL - MANNITOL		PRODUCTS
CH <sub>2</sub> OH		C <sub>2</sub> H <sub>6</sub> O +CO <sub>2</sub> +H <sub>2</sub>	CH <sub>2</sub> OH		C <sub>2</sub> H <sub>6</sub> O +CO <sub>2</sub> +H <sub>2</sub>
CHOH	CH <sub>2</sub> OH		CHOH	CH <sub>2</sub> OH	
CHOH	CHOH	2C <sub>3</sub> H <sub>6</sub> O <sub>2</sub>	CHOH	CHOH	2C <sub>3</sub> H <sub>6</sub> O <sub>2</sub>
CHOH	CHOH		CHOH	CHOH	
CHOH	CHOH		CHOH	CHOH	
CHOH	CHOH		CHOH	CHOH	
CHO	CHOH	C <sub>2</sub> H <sub>4</sub> O <sub>2</sub> +CO	CH <sub>2</sub> OH	CHOH	C <sub>2</sub> H <sub>6</sub> O +CO <sub>2</sub> +H <sub>2</sub>
	CHO			CH <sub>2</sub> OH	

For glucose this scheme would yield 2 molecules of lactic acid and 1 each of acetic acid and alcohol, for mannitol 2 each of lactic acid and alcohol; and to explain the observed results it is necessary to assume that in the latter case the lactic acid is largely destroyed by secondary reactions. Glycerol according to Harden is split directly into alcohol and formic acid.

In the case of *B. aerogenes* the proportion of lactic acid to alcohol formed is lower with glucose and very much lower with mannitol, while the acetic acid is even more reduced. It is evident that the portion of the carbohydrate molecule which furnishes the acids in the case of *B. coli* is here split in a different way and Harden and Walpole (1906) have shown that the end product in the *B. aerogenes* fermentation is chiefly 2:3-butylene glycol with some acetyl methyl-carbinol. The latter substance as shown by Harden (1906) and by Harden and Norris (1912) is the active agent in producing the Voges-Proskauer reaction. The acetyl methyl carbinol in the presence of potash and oxygen is oxidized to CH<sub>3</sub>.CO.CO.CH<sub>3</sub> (diacetyl) and the diacetyl reacts with some constituent of the peptone medium to produce the eosin-like fluorescence characteristic of this reaction.

Harden's studies were confined for the most part to the gas-producing organisms but he points out that with forms like *B. typhosus* formic acid must be produced from the fractions of the molecule which with *B. coli* yield CO<sub>2</sub> and H<sub>2</sub>. Sera (1910a, 1910b) has investigated this point and reports that typhoid and dysentery bacilli in decomposing glucose, glycerol or mannitol produce acetic and formic acids, with some propionic acid in the case of glycerol. In glucose and glycerol acetic acid is in excess, in mannitol formic acid. *B. typhosus* also forms a trace of alcohol.

An illuminating study was made by Grey (1914) of the fermentative products of Penfold's chloracetic acid mutants, in which the power of gas production had been reduced. He found that in each case the selected strains produced in glucose media more lactic acid and less alcohol, acetic acid and formic acid, and also decomposed less of the formic acid which they did produce. In mannitol however the change was limited to a decrease in decomposition of formic acid, the primary products formed being the same. Grey assumes that the glucose molecule is normally split, first into lactic acid and an Intermediate substance A, which in turn yields formic acid and an Intermediate substance B (possibly acetaldehyde). The Intermediate substance B finally yields alcohol and acetic acid. Grey believes that the decomposition of Intermediate substance B is due to a reduction, accomplished ordinarily by the excess of H<sub>2</sub> present in the case of mannitol but by a special reductase in the case of glucose. If the formation of this reductase were suppressed by chloracetic acid it would explain the fact that, while Intermediate substance B undergoes a normal splitting in the case of the selected strains when acting on mannitol, the process is fundamentally affected in the case of glucose. The decomposition of formic acid is of course equally affected in both instances.

It is evident that the changes which go on in the fermentation of carbohydrate media are complex and for the most part still obscure; but the broad facts appear to be established that the process leads to the production of alcohol and of acetic, lactic, formic and succinic acids (the latter in small amount); that the

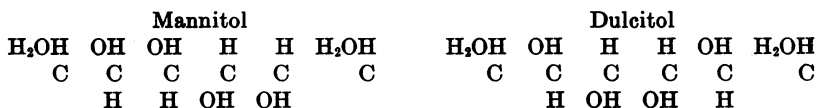


decomposition of the formic acid and perhaps of other constituents of the molecule produces a gas composed of carbon dioxide and hydrogen; that in the decomposition of mannitol alcohol is produced in much greater, and lactic acid in much smaller, amount than is the case with glucose; that in the case of *B. aerogenes* butyleneglycol, acetylmethylcarbinol and an excess of carbon dioxide are formed at the expense of the portion that with *B. coli* yields lactic and acetic acids.

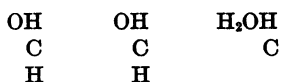
*Relation between the structural formula of the carbohydrates and their decomposition by colon-typhoid bacteria.* The question of the relation between molecular structure and fermentability is of the greatest interest to the biologist, since data in regard to such a relationship may reasonably be expected to throw important light upon the underlying bio-chemical processes involved. The studies made upon the colon-typhoid group justify a few fundamental generalizations in regard to this point.

First of all it is evident that the hexoses are more easily attacked than any of the other substances ordinarily investigated. All of the members of the colon-typhoid group, with the exception of *B. alcaligenes*, produce a marked increase of acidity in these carbohydrate media. In our own studies we find that fructose, mannose, and galactose behave essentially like glucose, all the strains which acidify glucose acidifying all the other hexoses as well.

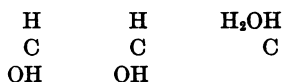
The hexahydric alcohol, mannitol, is the substance which comes next in order of availability, all of the groups studied except *B. alcaligenes* and the Shiga dysentery types acidifying this substance. Dulcitol however, which resembles mannitol so closely, is much more resistant. As a matter of fact dulcitol occupies a somewhat unique position among all the carbohydrates whose decomposition has been extensively studied. It is the only one whose fermentation seems wholly uncorrelated with that of other carbohydrate media. In all the principal groups of the paratyphoid, colon and aerogenes series we find forms which ferment this alcohol and others which fail to do so; and it has been pointed out that this particular fermentation is the one which exhibits the most general tendency toward mutative variability.



Sorbitol appears to behave like mannitol, being attacked by all the paratyphoid types according to Boycott (1906), Jordan (1917), and Krumwiede Kohn, and Valentine (1918). From the formulae of these alcohols it may be noted, as suggested by Revis (1910) that the configuration of mannitol and sorbitol resembles that of the alcohol half of the hexoses. In the hexoses the grouping is



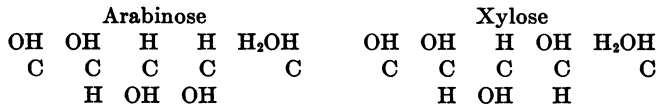
In mannitol and sorbitol it is



In dulcitol on the other hand there is no alcohol grouping with two adjacent carbon atoms having the hydroxyls on the same side; and in this difference may perhaps lie the difference in susceptibility to bacterial decomposition.

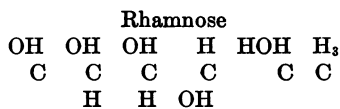
Next to these alcohols in availability for the colon-typhoid organisms comes the disaccharide maltose. We have not studied this carbohydrate ourselves but Hiss and Russell (1903), Hiss (1904), Smith (1915), and others show that the Shiga type of dysentery bacillus fails to attack maltose, while Morgan (1906, 1911) reports negative results for the peculiar organism which bears his name, as do Hadley, Elkins and Caldwell (1918) for *B. pullorum*. On the other hand Drigalski and Conradi (1902), Boycott (1906), Sacquépée and Chevrel (1905), Savage (1912), Jordan (1917), and Krumwiede, Kohn and Valentine (1918) find that the typhoid and paratyphoid organisms all attack maltose readily.

The next substances, in order of availability, appear to be xylose and arabinose with the structural formulae indicated below.



Here we may note the very interesting fact that among the less active fermenters of the typhoid and paratyphoid series (where the fermentation of these sugars first appears) action upon xylose and arabinose seems to be inversely correlated. Thus the Flexner dysentery and paratyphoid A types attack arabinose but not xylose, the typhoid and hog-cholera strains xylose but not arabinose. It would appear that the power to ferment the two different molecular groupings represented by these sugars appears or disappears independently and they are presumably attacked in a quite different way.

Rhamnose is the next carbohydrate to be attacked by the colon-typhoid bacteria.<sup>2</sup> This substance, which is a methylated pentose, is utilized by all the paratyphoid, colon and aerogenes series; and it may be assumed that the presence of the methyl group is the circumstance which prevents the Flexner dysentery and typhoid organisms from decomposing it.



From the five and six carbon carbohydrates to the disaccharide lactose there is a distinct and well marked step; and we may reasonably assume that the broad distinction between the pathogenic dysentery-typhoid-paratyphoid series on the one hand and the non-pathogenic colon-aerogenes series on the other is correlated with the absence or presence of an enzyme capable of breaking up this disaccharide into glucose and galactose. The

<sup>2</sup> In this discussion we have for convenience considered the carbohydrates in their order of availability, passing from those most easily to those least commonly fermented. It is not necessary however to assume that evolution has proceeded along the line of a progressive acquisition of fermentative properties. Since the organisms of low fermentative power are the pathogenic (and presumably more recent) forms it would be more reasonable to assume that development has been in the direction of a loss rather than a gain in fermentative power, in which case *B. aerogenes* might be the primitive ancestral type of the whole group.

difference between maltose and lactose in availability is a striking and significant one. It has been noted that all the members of the typhoid and paratyphoid series produce acid from maltose; but the power to decompose lactose appears only among the gas-producing forms of the colon and aerogenes groups.

It seems clear that the processes involved in the decomposition of the two disaccharides, maltose and lactose, must be fundamentally dissimilar; and it is of interest to note that while the difference in availability between mannitol and sorbitol and between the three pentoses are associated with differences in reactive organic groups, the even more fundamental difference between the availability of maltose and lactose can be explained only by a stereoisomeric difference in the molecules concerned. As in the case of animal enzymes the presence of maltase and lactase seems to vary quite independently.

Next to lactose in order of availability come sucrose and raffinose; and here we find another definite relation between fermentability and molecular structure. The correlation between the fermentability of these two sugars has been shown by Winslow and Walker (1907), Burk (1907), Howe (1912), Rogers, Clark and Davis (1914a), Kligler (1914), Levine (1916) Murray (1916), and Rogers, Clark and Lubs (1918) to be an almost perfect one. These two sugars, although one is a disaccharide and the other a trisaccharide, are alike in the absence of the aldehyde group; and this fact may be assumed to account for the similarity in their behavior.

The fermentation of the glucoside salicin is generally, but not always, correlated with that of lactose; being fermented by most, but not all, of the colon group and by all the *B. aerogenes* strains; while the cyclohexanhexol inosite appears to be fermented only by Paratyphoid B and certain *B. aerogenes* strains.

#### PROGRESSIVE CHANGES OF REACTION IN CARBOHYDRATE MEDIA AND THEIR SIGNIFICANCE

The estimation of the individual products of carbohydrate decomposition is of course a difficult and time consuming process; and even the determination of the gases evolved must be

carried out by exhaustion from a closed vessel if it is to have any serious value. The acidity produced in a given culture medium is on the other hand a characteristic which can be easily and accurately measured. To this point we have therefore devoted a considerable share of our attention.

When the work on the colon-typhoid group was begun we made our first studies of acid production by the old titration method, cultivating each strain in broth (containing 1 per cent Digestive Ferments Company peptone, 0.5 per cent  $K_2HPO_4$ , and 1 per cent of carbohydrate) at 30°C. After two days and seven days of incubation, tubes were withdrawn, held in steam or boiling water for ten minutes to drive off  $CO_2$ , cooled and titrated against N/20 NaOH with phenolphthalein as indicator. The general results of the observations made by this method on a series of colon-aerogenes strains have been already reported by one of us (Kligler, 1914). They indicated that in any carbohydrate medium there is a fairly sharp distinction between fermenting and non-fermenting forms, the line between the two groups being at about 1.5 per cent normal acid, (as ordinarily determined), the mode for the non-fermenters being near the neutral point and that for the fermenters varying between 2.5 and 3.5 per cent acid for the various carbohydrates.

In view of the rapidly accumulating evidence which indicated that the measurement of acidity by the titration method is inaccurate and often misleading, we determined the true acidity in the various fermentable media for our complete series by the use of the Clark and Lubs indicators. The media first used for this purpose contained 0.5 per cent Digestive Ferments Company peptone 0.5  $K_2HPO_4$  and 0.5 per cent of the fermentable substance to be studied, as recommended by Clark and Lubs. We found however that a somewhat sharper differentiation could be obtained by increasing the amount of sugar, and in our regular routine studies the medium was made up with 0.5 per cent peptone, 0.5 per cent  $K_2HPO_4$  and 1 per cent of the fermentable substance. The cultures were incubated at 30°C. and the acidity was determined after two, four and five days by the use of methyl red, dibromthymolsulphonphthalein and phenol sul-

phonphthalein, according to the methods suggested by Clark and Lubs (1917).

In general four distinct types of reaction were observed when the cultures were examined in this way, reactions which we have described as types I, II, III, and IV (Winslow, Kligler and Rothberg, 1917). In type I which is observed when the carbohydrate present is not attacked at all, the reaction remains throughout in the neighborhood of  $P_H$  7.0. Type II is the reaction characteristic of slow fermentation, the  $P_H$  value after two days being still at 6 or over, and only later reaching an acidity below 5.5. This type of reaction appears with substances which are decomposed with some difficulty, such as dulcitol, rhamnose, and salicin, and the principal results of this kind which we obtained are tabulated below.

*Type II reactions*

GROUP OF ORGANISMS	FERMENTABLE SUBSTANCE	AVERAGE PH VALUES AFTER		
		2 days	4 days	5 days
Flexner dysentery.....	Mannitol	6.5	5.5	5.7
Paratyphoid A.....	Rhamnose	6.4	5.0	5.1
	Arabinose	7.0	5.0	5.1
B. suipestifer.....	Rhamnose	7.0	6.4	5.8
Paratyphoid B.....	Rhamnose	6.7	5.0	5.1
	Dulcitol	6.7	5.3	5.2
B. communior.....	Dulcitol	7.0	5.6	5.5
	Salicin	5.9	5.1	4.8
B. communis.....	Sucrose	6.1	5.2	5.2
	Salicin	6.4	5.4	5.3
	Dulcitol	7.1	5.6	5.5

A type III reaction is the normal reaction for the colon-typhoid organisms in any carbohydrate which is readily fermented. In this case the acidity rises rapidly, usually within twenty-four hours and always within forty-eight hours, to a  $P_H$  value of 5.3 or less and remains practically constant at about that point.

We have observed this rapid rise and subsequent maintenance of high acidity with all the groups of organisms studied (except *B. aerogenes*) and in the case of all the substances fermented except those cited above as giving a somewhat delayed reaction. With these exceptions the  $P_{\text{H}}$  value after two days of incubation varied for all the groups studied and for all the substances fermented between the limits of 5.0 and 5.6, while the final values after five or six days varied between 4.8 and 5.3. This final reaction averaging about 5.0 is so constant that it may be fairly assumed to represent the limiting concentration of acid for the colon-typhoid organisms in the particular medium used for our studies. Clark (1915) has pointed out the same constancy in final reaction for *B. coli* in 30 different media, his range of variation (even with a series of variously buffered media) being only from  $P_{\text{H}}$  4.3 to  $P_{\text{H}}$  5.3. Michaelis and Marcora (1912) reported 4.8 as the limiting  $P_{\text{H}}$  value for *B. coli* in glucose peptone broth. Our results make it possible to extend these conclusions and to state that not only for *B. coli* but for the whole colon-typhoid group the final acidity in sugar broths is practically the same,—about  $P_{\text{H}}$  5.0—in the medium studied by us.

In a longer period of incubation than that used for our routine determinations (five days) there appears to be a slight reversion toward alkalinity even in the case of the type III reaction, as indicated by the results tabulated below for a series of 9 typical typhoid strains; and this slight reversion may probably be attributed to the formation of alkaline products of protein decomposition.

*Average acidity (PH) produced after various periods by nine typhoid strains*

	1 DAY	2 DAYS	4 DAYS	6 DAYS	8 DAYS	11 DAYS	13 DAYS
Glucose.....	5.3	4.9	5.0	5.2	5.3	5.3	5.4
Mannitol.....	5.5	5.1	5.4	5.6	5.8	5.9	5.8

It will be noted that the  $P_{\text{H}}$  values are slightly higher throughout in the mannitol as compared with the glucose broth. Our results indicate that there exist minor differences of this kind, the fermentation of carbohydrates which are less readily utilized being a trifle slower and producing a little lower acidity than the

decomposition of glucose. The line between a type II and a type III reaction is not therefore an absolutely sharp one.

The type IV reaction, in which the acidity rarely goes below a  $P_H$  value of 5.5 and quickly reverts to a value over  $P_H$  6 is characteristic of the *B. aerogenes* group and appears to be related to a fundamentally different action upon the fermentable substances. The differences between the type III and type IV reactions during the first forty-eight hours are indicated below for a series of *B. coli* and *B. aerogenes* strains.

*Average acidity (PH) produced by a series of strains of B. coli and B. aerogenes in glucose broth*

	2 HOURS	4 HOURS	8 HOURS	24 HOURS	48 HOURS
<i>B. coli</i> .....	7.3	7.2	6.0	4.9	4.7
<i>B. aerogenes</i> .....	7.2	6.8	5.8	5.7	6.2

The reactions cited below for *B. aerogenes* during a longer period of incubation in various carbohydrate media are for a selected series of 10 typical strains and may be compared with corresponding figures quoted above for *B. typhosus* in glucose and mannitol.

*Average acidity (PH) produced by a series of B. aerogenes strains*

	1 DAY	2 DAYS	4 DAYS	6 DAYS	8 DAYS	11 DAYS	13 DAYS
Glucose.....	5.3	5.3	5.8	6.2	6.3	6.6	6.3
Lactose.....	5.6	5.6	5.9	5.9	6.1	6.8	7.0
Sucrose.....	5.4	5.5	6.0	6.4	6.5	7.1	6.8
Rhamnose.....	5.5	5.4	5.7	5.1	5.9	6.3	6.3

In the type IV reaction, marked differences in rate of fermentation appear between individual strains and a high acidity (in some cases as high as  $P_H$  5.1) may be attained for brief periods. The reversion is very rapid however and  $P_H$  values lower than 5.5 are rarely observed.

The phenomenon of reversion from a preliminary acid reaction to a more alkaline one was explained by the earlier workers on this subject as due to exhaustion of the carbohydrate, followed by the utilization of nitrogenous foodstuffs with the release of alkaline products of protein decomposition.



Kendall, Day and Walker (1913) made a somewhat extensive study of this point, determining the reaction of various members of the colon-typhoid group in plain broth and glucose broth by the use of neutral red as an indicator, and simultaneously estimating the rate of protein decomposition by the increase in ammonia. Their results indicated a definite sparing action of the sugar. For example *B. dysenteriae* in plain broth produced 2.1 to 4.2 mgm. of NH<sub>3</sub> per 100 cc. broth with no material change in reaction, while in 1 per cent glucose broth the reaction reached an acidity of 2 to 2.8 per cent Normal to neutral red, while no ammonia was formed. *B. typhosus*, the paratyphoids, the Morgan bacillus, and *B. coli* all behaved in essentially the same way; while in the case of *B. cloacae* the carbohydrate was apparently destroyed so quickly that it made little difference in the protein metabolism whether glucose was present or not. In the case of *B. alcaligenes* there was no difference in reaction between the plain broth and the glucose broth, but growth was more luxuriant and more ammonia was produced in the latter case, suggesting some sort of utilization either of the glucose itself or of impurities contained therein. The range of values recorded for certain groups are of such interest as to warrant reproduction.

ORGANISM	BROTH	REACTION TO NEUTRAL RED	INCREASE IN NH <sub>3</sub>
<i>B. dysenteriae</i> .....	Plain	-0.3 to +0.2	2.1 to 4.2
	Glucose	2.0 to 2.8	None
<i>B. typhosus</i> .....	Plain	+0.1 to -0.9	4.2 to 10.5
	Glucose	2.9 to 4.9	None
Paratyphoids.....	Plain	-1.5 to +0.3	4.2 to 16.1
	Glucose	2.3 to 4.8	0.0 to 2.8
Morgan bacillus.....	Plain	-0.5 to -2.1	20.3 to 42.0
	Glucose	2.9 to 4.6	4.2 to 8.4
<i>B. coli</i> .....	Plain	-0.4 to 1.9	11.2 to 36.4
	Glucose	3.9 to 6.1	-1.4 to 2.1
<i>B. cloacae</i> .....	Plain	-0.9 to -1.4	20.3 to 21.7
	Glucose	-0.6 to +1.0	16.8 to 19.0

These results do indeed suggest that the difference in reaction in a carbohydrate medium may be partly due to the relative rate at which protein and carbohydrate, respectively, are being attacked. That in the case of certain organisms there is something much more fundamental involved is indicated by the peculiar behavior of *B. cloacae* (a form which belongs to the *B. aerogenes* group); and the same deep-seated difference is indicated by the progressive change of reaction in glucose broth recorded by Kendall, Day and Walker for the various groups of organisms studied. We have calculated below the average values for certain of these groups and in so doing have separated one of the groups presented by the authors under the name of *B. mucosus-capsulatus* into two subgroups since 9 strains called by Kendall, Day and Walker *B. lactis-aerogenes* are clearly different in their behavior from the rest of the *B. mucosus-capsulatus* groups.

*Average acidity (to neutral red) in glucose broth cultures of various colontyphoid organisms after various intervals*

DAY	B. ALCALIGENES	DYSENTERY	TYPHOID	PARATYPHOID	MORGAN	B. COLI	B. LACTIS AEROGENES	B. MUCO-SUS CAPSULATUS	B. CLOACAE
1	- 0.4	2.1	3.0	3.0	2.8	3.8	3.7	2.2	2.0
3	- 0.8	2.7	3.3	3.8	3.8	4.4	4.2	1.6	1.6
6	- 1.0	2.7	3.5	3.8	3.7	4.8	4.8	1.3	0.9
9	- 1.1	2.7	3.6	3.7	3.9	5.0	4.6	1.0	- 0.1

The results, as tabulated above, indicate a progressive increase in acidity (a type III reaction) for dysentery, paratyphoid, Morgan, typhoid and colon bacilli, although it is interesting to note that Kendall, Day and Walker's figures indicate a distinct difference in the final end-point, the dysentery bacilli producing a lower, and the colon bacilli a higher, acidity than the typhoid, paratyphoid and Morgan groups. The forms called by Kendall, Day and Walker *B. lactis-aerogenes* appear by their fermentative activity to be of the *B. coli* type but those designated as *B. mucosus-capsulatus*, as well as the *B. cloacae* forms, show the characteristic type IV reaction of *B. aerogenes*.

Ayers and Rupp (1918) have recently brought forward direct and convincing evidence that the reversion of reaction exhibited in *B. aerogenes* cultures is due rather to the secondary decomposition of organic acids with the formation of basic carbonates than to neutralization by basic products of protein decomposition. They show that the reversion takes place even in a synthetic medium containing sodium ammonium phosphate as a source of nitrogen (in which case ammoniacal products could not possibly be sufficient to account for the phenomenon); and that it is accompanied by a rapid destruction of formic, acetic and other acids. Even with *B. coli* they noted as might be expected that acid formation does not run parallel with the destruction of glucose, formic acid remaining constant or being slightly reduced during the later stages of fermentation. The distinction between *B. coli* and *B. aerogenes* may therefore be considered as lying chiefly in the difference in rate between the preliminary decomposition of the sugars into acids and the secondary decomposition of the acids themselves; yet it is evidently a distinction that is clearly marked and of important systematic significance.

The slower and much less marked reversion which is observed if organisms giving the type III reaction are cultivated for a prolonged period may perhaps be due to a different process, an exhaustion of the fermentable carbohydrate and a subsequent decomposition of nitrogenous foodstuffs with the formation of alkaline substances. The same explanation may probably account for the reversion shown in milk cultures by the members of the paratyphoid group, the transient acidity produced being due to the fermentation of the small amount of glucose present.

It must of course be remembered that the course of such reactions as those described will vary within wide limits with variations in the composition of the medium employed. Clark and Lubs (1915) studied the effect of varying the concentration of carbohydrate and showed that with 0.5 per cent peptone and 0.5 per cent  $K_2HPO_4$  and 0.1 per cent glucose *B. coli* shows a marked reversion, the curve of acidity by days resembling closely that given by *B. aerogenes* in the presence of 0.5 per cent glucose.

It has been shown by one of us (Kligler 1916) that the net result of the action of bacteria in a given medium not only depends on the relative amounts of peptone and glucose present but is influenced by the amount of phosphate as well.

CHARACTERISTICS OF COLON-TYPHOID BACTERIA OF GROUP I  
(FERMENTING NO CARBOHYDRATES MORE COMPLEX THAN  
THE HEXOSES)

The first well marked group of colon-typhoid organisms, as pointed out above, includes the forms of exceedingly low fermentative power which are capable of attacking only the simple hexoses if they ferment any carbohydrates at all, but are unable to produce acid in mannitol media.

The collection of cultures studied by us included 14 different strains belonging to this general group, which could be subdivided further by their action upon the hexoses and the formation of indol.

Five of the strains formed no acid in any carbohydrate media, the  $P_{\text{H}}$  value of glucose broth remaining over 7.0 and milk cultures turning gradually to a deeper and deeper blue.<sup>3</sup> One strain (no. 439) showed a decolorization in milk after four weeks incubation, the color changing from deep blue to a pale chalky tint. These cultures were sent in to the Museum collection with the following names: *B. pneumoniae*, *B. dysenteriae*, *B. alcaligenes*, *B. ozenae*, *B. bronchisepticus*; but all of them have lost the power of attacking carbohydrates if they ever possessed it. All are indol negative<sup>4</sup> and all fail to liquefy gelatin.<sup>5</sup>

<sup>3</sup> Observations of the behavior of our cultures in milk were made under the following conditions. Certified milk was steamed for forty-five minutes in the Arnold sterilizer and left over night in the ice box for the cream to rise. The milk was then siphoned off; tinted with Kahlbaum's azolitmin, tubed and sterilized in the Arnold for twenty minutes on three successive days. After inoculation the cultures were incubated at 37°C. and observed after twenty-four and forty-eight hours and one, two, four and six weeks.

<sup>4</sup> Indol production was studied by the following method which Marshall (1907) and others have shown to be an accurate and satisfactory one. A medium containing 0.3 gram tryptophane and 5 grams  $K_2HPO_4$  in 1000 cc. of water was inoculated from a fresh twenty-four hour culture and incubated at 30° for forty-

It is of some interest to note that the names borne by three of these five strains would indicate that they did originally possess fermentative powers. Strain 27 of our collection was obtained from The Rockefeller Institute, as a Kral culture of *B. pneumoniae*; Strain 543 was isolated at the University of Pennsylvania, from a case of atrophic rhinitis identified as *B. ozenae* and successfully used as a vaccine; strain 177 was described as *B. dysenteriae* in the Journal of Experimental Medicine, 6, 181.

We have made no tests of motility on our strains, but Kendall, Day and Walker (1913) and Stewart (1917) describe *B. alcaligenes* as motile. Kühnemann (1911) claims that it has polar flagella, which if correct would remove it from the colon-typhoid group entirely. Petruschky's original description includes the statement that the flagella are peritrichic, and stresses the brownish growth upon potato. Berghaus (1905) who studied *B. alcaligenes* with some care bases the characterization of this species on the strongly alkaline reaction in milk, a brown coloration on potato, specific agglutination reactions, and the fact that it is an obligate aerobe. The species may be described as follows, and strain 439 of the American Museum collection (isolated from feces at the University of Pennsylvania) may be taken as a type.

*B. alcaligenes* Petruschky. Gram-negative non-spore-forming motile rods, producing thin translucent irregular colonies on

eight hours. One cubic centimeter of a 2 per cent alcoholic solution of p-dimethylamidobenzaldehyd was added drop by drop so as to mix with the medium, followed by a few drops of concentrated HCl, a reddish-purple color indicating the presence of indol. A comparison of the results obtained over a year before with the same cultures in a pepton medium showed that the results with the two procedures were identical.

<sup>5</sup> Gelatin liquefaction was observed in small test tubes 1 cm. in diameter and 10 cm. long. Five cubic centimeters of standard gelatin made with Liebig's beef extract was placed in a tube and sterilized in the autoclave for five minutes under 20 pounds pressure. The tubes were inoculated by spreading a loopful of a twenty-four hour broth culture over the surface of the medium and they were incubated for twenty days at 20°C. in an atmosphere saturated with moisture. At the end of this time the depth of liquefied gelatin was estimated in centimeters.

gelatin and a brownish growth on potato. Reaction in carbohydrate media alkaline. Indol not produced. Gelatin not liquefied.

A second subdivision of the non-mannitol fermenting organisms, characterized by the decomposition of the hexoses with the production of acid but no gas, was represented in our collection by but two strains, the dysentery bacilli of Shiga and Kruse, respectively. They gave a type III reaction in glucose, mannose, fructose and galactose as indicated by the figures cited below.

*Reaction (PH) produced by non-mannite-fermenting dysentery bacilli in glucose broth*

	2 DAYS	4 DAYS	5 DAYS
Kruse.....	4.9	4.9	4.9
Shiga.....	5.0	5.1	4.9

No other carbohydrates were fermented; and in milk both strains produced a very faint initial reddening followed by a return to the original color of the medium. Indol was not formed, gelatin was not liquefied, and lead acetate media were not turned brown. Shiga originally described his dysentery organism as slowly motile but Kruse (1900), Vedder and Duval (1902) and all later observers agree that it is non-motile. All investigators who have studied the agglutination reactions of dysentery bacilli (among whom may be mentioned Pai and Krishnan, 1916 and Andrewes, 1918) report that the agglutinations of the Shiga type (including the Kruse and New Haven strains of non-mannite-fermenters) are sharply specific. Furthermore, as shown by Kruse, Flexner and Sweet (1906) and others the Shiga bacillus possesses the power, unique in the colon-typhoid group, of producing a soluble toxin. Nicolle, Debains and Loiseau (1916) note that this type produces in culture a characteristic odor of chestnut flowers. The Shiga type of dysentery appears to be an unusually definite and distinct one. Martin and Williams (1917) for example isolated 47 strains at

Cairo, all of which were true to type. While it is of course certain that both the mannit fermenting and non-mannit-fermenting types of dysentery bacilli are causally connected with the disease whose English name they bear, it is most desirable that they should be given definite Latin names in accord with scientific usage—particularly if, as we believe, they represent quite sharply differentiated bacterial types. It seems to us that the Flexner organism for instance is biologically more nearly related to *B. typhosus* than to the Shiga bacillus. Chester (1901) has given the Shiga type the name *B. shigae* and it may be characterized as follows.

*B. shigae* Chester. Gram-negative, non-spore-forming non-motile rods, producing thin translucent irregular colonies on gelatin. Form acid rapidly in media containing the hexoses, but not in other carbohydrates. Milk turned slightly acid and then neutral or slightly alkaline. Indol not produced. Gelatin not liquefied. Lead acetate media not browned. Produces a soluble toxin and exhibits characteristic agglutinative reactions. Found in human stools. The causative organism in one form of dysentery. Strain 197 of the American Museum collection is a type of this species. It was obtained by us from The Rockefeller Institute, labeled "Shiga, Japan, 293."

#### CHARACTERISTICS OF COLON-TYPHOID BACTERIA OF GROUP II (FERMENTING MANNITOL BUT NOT RHAMNOSE)

The second general group recognized above includes the forms which ferment mannitol and either xylose or arabinose but rarely both; and which produce acid but not gas in the sugars which are attacked. The arabinose-positive forms correspond to the Flexner group of dysentery bacilli and the arabinose-negative forms to *B. typhosus*.

Of the arabinose-fermenters we had only two strains in our series, both obtained from The Rockefeller Institute, strain 110 a Flexner-Harris strain, isolated by Flexner at Manilla in 1900 and strain 196, a strain also isolated in the Philippines by Strong. Both were alike in forming acid in glucose, mannose, fructose,

galactose, mannitol and arabinose, but not in the other carbohydrates studied. The acid production was however rather of our type II than our type III order, being slightly more sluggish than in the case of most of the organisms studied, as indicated below.

*Acidity (PH) produced by Flexner dysentery bacilli in glucose broth*

	2 DAYS	4 DAYS	5 DAYS
Strain 110.....	6.0	5.4	5.5
Strain 196.....	5.6	5.4	5.6

Milk cultures were practically unchanged, although strain 110 showed a very slight reddening after two days, with reversion to the original neutral tint. Indol was produced and gelatin was not liquefied. Lead acetate media were not turned brown.

While the Shiga dysentery organisms constitute a clear and well-defined type the mannitol-fermenting strains are much more variable. Lentz reported that the Flexner strains fermented maltose and dextrin, while his Strong strains (all mannitol-fermenters) failed to do so. Hiss and Russell (1903) described their Y strain as failing to attack maltose; while the Flexner strain they studied acidified both maltose and sucrose. Park, Collins and Goodwin (1904) reported the same reactions for the Flexner type, while their Seal Harbor strain failed to attack either maltose or sucrose. Hiss (1904) recognized three subgroups of mannitol-fermenting dysentery bacilli as follows:

TYPES	MALTOSE	SUCROSE	DEXTRIN
Y (Hiss), Seal Harbor (Park), etc.....	-	-	-
Flexner, Strong.....	-	+	-
Flexner, Harris, Wolstein.....	+	+	+

On prolonged cultivation however the Y strains did attack maltose and sucrose (and a non-mannitol-fermenting Kruse strain attacked maltose!). Gay (1904) gave a similar classification. Morgan (1911) examined a series of about 50 dysentery-



like organisms, half from England and half from abroad and reported a considerable proportion of positive results (acid-production) in media containing sorbitol, arabinose and raffinose, in addition to the carbohydrates included in the table above. He concludes that

In the mannite-fermenting dysentery group (excluding the "Strong" strains) must be incorporated a large and probably ever-increasing number of strains reacting with striking uniformity to one test (i.e. agglutination with "Y" or Flexner serum) but differing from one another markedly when their fermentation properties and receptor mechanisms are minutely investigated.

Smith (1913) gives an elaborate classification of this group based on reactions in dulcitol, sucrose, sorbitol, dextrin and maltose. He calls the Y type dextrin positive, and in a later paper (Smith, 1915) gives the following reactions for the three common types.

	DULCITOL	MALTOSE	SORBITOL	DEXTRIN	ARABI- NOSE	RAFFINOSE
Y (Hiss).....	-	-	-	+	+	+
Strong.....	+	+	+	-	+	+
Flexner-Harris.....	-	+	-	+	+	+

It will be noted that these results differ in some respects from those cited above from Hiss and Gay, and it is probable that there were errors in the work of the earlier observers. Hort (1915) gives the same characteristics as those tabulated by Smith.

Recent observers have emphasized particularly the variability of the members of the mannitol-fermenting dysentery group and the difficulty of drawing sharp lines of distinction between them. Thus Pai and Krishnan (1916) report that Flexner and Y types cannot be clearly distinguished by their agglutination reactions. Martin and Williams (1917) studied 76 mannitol-fermenting dysentery bacilli which were constantly positive (acid-producing) in glucose, galactose and mannitol and constantly negative in lactose, dulcitol, inulin and adonitol, but which varied widely in their action on maltose, sucrose, dextrin,

raffinose, arabinose, rhamnose, sorbitol, glycerol and indol,—the same strain giving variable results at different times. Andrews (1918) and Thøtta (1919) have recently described types of dysentery-like organisms which indicate a confusing series of varieties within this group. Strain 110 of the American Museum collection, as originally a Flexner strain, should ferment maltose but as a matter of fact does not. Strain 196 as a Strong strain should ferment both maltose and sucrose but does not, giving us further evidence of the instability of fermentative powers of these organisms.

On the whole we are inclined to think that it is wiser not to burden the literature of this highly unstable group with a multiplicity of specific names based on fermentative irregularities, and believe that it would probably be best to include all the mannitol fermenters under the species *B. dysenteriae* with the principal sub-types as varieties. It may be noted that the agglutinative characters and pathogenic properties of these organisms are much more stable and appear to warrant the recognition of the group as a distinct entity.

The species and varieties may be defined as follows:

*B. dysenteriae* (Flexner). Gram negative, non-spore-forming rods. Non-motile. Producing characteristic translucent irregular colonies. Gelatin not liquefied. Indol generally produced. Acid production in media containing the hexoses and mannitol, and usually in arabinose. Lead acetate media not browned. Found in stools, causal agents of one form of dysentery.

Variety Hiss-Y. Dextrin fermented, maltose not fermented, sucrose not fermented.

Variety Flexner. Dextrin and maltose both fermented. Sucrose not fermented.

Variety Strong. Maltose and sucrose fermented but not dextrin.

The biological relationships of *B. dysenteriae* (Flexner) are very hard to surmise. It is related to *B. shigae* by lack of motility and by the nature of the disease which it produces; it is related to *B. typhosus* by the fermentation of maltose and mannitol; it is allied to both these organisms by failure to produce gas. On

the other hand the fact that it forms indol and sometimes attacks such complex carbohydrates as sucrose and raffinose would suggest a transition to *B. coli* and the lactose fermenters. Andrewes' (1918) three species of non-pathogenic *B. dysenteriae*-like bacilli may be connecting links along such a line.

Turning now to the organisms which ferment xylose but not arabinose, (*B. typhosus*) we find a much more compact and homogeneous group.

The typhoid bacillus, as pointed out in an earlier section, is characterized by active motility, failure to produce indol, and failure to ferment the more complex carbohydrates such as arabinose and rhamnose. Of our series of cultures, 24 belonged to this group. The fact that all but one of these strains came in to the American Museum collection under the name *B. typhosus* (the single exception being called *B. paratyphosus*) is good evidence of the fixity of this type. All of our 24 strains were alike in producing a rapid and complete acid reaction (type III) in the hexoses, and in mannitol, and xylose (in xylose the reaction reverted later slightly to about  $P_x$  5.7), and in failing to attack lactose, sucrose, arabinose, rhamnose, dulcitol, or salicin. In litmus milk the medium was slightly reddened after two days but after two weeks turned neutral or slightly alkaline again. Indol was never formed, nor was gelatin liquefied. Lead acetate media were turned brown.

Sorbitol, which we did not study, is reported as acidified (Morgan, 1906, Smith, 1913, Robinson, 1915). So are maltose and dextrin (Smith, 1915, Harding and Ostenberg, 1912, Robinson, 1915). Smith (1915) states that raffinose is also attacked. This latter reaction has not been reported by other observers and seems improbable from the negative results in sucrose.

The typhoid bacillus is evidently a clearly marked and usually stable organism which may be defined as follows.

*B. typhosus* (Zopf). Gram-negative non-spore-forming rod. Actively motile. Forms translucent irregular colonies on gelatin media and faint nearly colorless growth on potato. Produces strong and prompt acid but no gas in media containing the hexoses maltose, mannitol, sorbitol, xylose and dextrin. Does

not attack arabinose, rhamnose or lactose. Produces a slight initial reddening of litmus milk, which after two weeks reverts to a neutral or slightly alkaline reaction. Fails to form indol or liquefy gelatin. Will not grow in asparagin-mannitol medium (Capaldi-Proskauer No. I). Does not reduce neutral red. Does cause browning of lead acetate media. Has low tolerance for acids but rather high tolerance for brilliant green dyes and alkaloids. Characteristic serum agglutination reactions. Found in human stools and urine as actual or potential cause of typhoid fever.

Our type of *B. typhosus* is strain 608, the Rawlings strain, obtained by Col. F. F. Russell from Colonel Leishman in 1908 and used in preparing the standard army vaccine.

There were two peculiar strains in our collection, both of which were sent in labeled *B. pyogenes-foetidus*. They were alike in giving the general fermentative reactions of *B. typhosus* but liquefied gelatin and casein rapidly.

CHARACTERISTICS OF COLON-TYPHOID BACILLI OF GROUP III  
(FERMENTING MANNITOL, ARABINOSE AND RHAMNOSE,  
GENERALLY FORMING GAS)

We may turn next to group III (see page 439) which includes the paratyphoid A types, fermenting the hexoses mannitol, maltose, rhamnose and arabinose (but not xylose) with the formation of gas, and producing like *B. typhosus* only a very slow reversion to an alkaline reaction in milk. Nine of the strains studied by us fell in this group. Six of them were sent in to the Museum collection labeled as Paratyphoid or specifically as Paratyphoid A; two as *B. pullorum*; and one as *B. alcaligenes*.

All fermented the hexoses, mannitol, rhamnose and arabinose. Two strains (nos. 294 and 322) attacked dulcitol. (Dulcitol is fermented by paratyphoid A according to Boycott, 1906, Morgan, 1906, Springer 1911, Smith 1915, and Jordan 1917). None of our strains produced acid in xylose, lactose, sucrose, salicin or inosite. Of the carbohydrates not studied by us, maltose is fermented by paratyphoid A according to Boycott 1906, Sac-

quépée and Chevrel 1906, Springer 1911, Smith 1915, Jordan 1917, and Krumwiede, Kohn and Valentine 1918; while dextrin is not attacked according to Jordan 1917, Weiss and Rice, 1917, Hulton-Frankel 1918 and Krumwiede, Kohn and Valentine 1918; and raffinose, inulin, erythritol and adonitol are not attacked according to Jordan 1917. Litmus milk was first turned slightly red but in the case of most of our strains by two weeks had reverted to a neutral or slightly alkaline reaction, very much as in the case of *B. typhosus*. Jordan (1917) states that his strains remained acid for two weeks, while Krumwiede, Pratt and Kohn (1916 b) report limits varying from five days to six weeks. Neutral red is said not to be reduced by Bainbridge 1919, but Sacquépée and Chevrel 1906, Hollande and Beauverie 1915, Nicolle, Raphael and Debains 1917 all report that this dye is decolorized. Lead acetate media were not browned by our cultures (as reported by Sacquépée and Chevrel 1906 and Hollande and Beauverie 1915). Gelatin was not liquefied and indol was not produced by any strain. Morgan (1906) stated that paratyphoid A produced indol but as shown by Marshall (1907) and Zipfel (1912) his methods of determining indol were unreliable. Among the minor characteristics of the paratyphoid A group may be mentioned the formation of agar colonies of a type intermediate between those characteristic of *B. typhosus* and *B. coli* respectively (Jordan 1917).

The designation of the paratyphoid bacilli as "A" and "B" has become firmly established in medical literature; but this terminology is quite clearly inadmissible from a systematic biological standpoint. The two forms are not varieties of one species but quite distinct forms, in spite of the fact that they produce a clinically similar disease. The A type is more nearly related to *B. pullorum* than to the B type; and the B type more nearly related to *B. enteritidis* and *B. suispestifer*, than to the A type of paratyphoid. It seems to us clear that definite specific names in a proper Latin form should be given to these organisms; and the name *B. paratyphosus* may properly be adopted for the A type to be defined as follows.

*B. paratyphosus*. Gram negative, non-spore-forming motile rods. Colonies on gelatin somewhat intermediate between the thin translucent irregular colonies of *B. typhosus* and the convex regular colonies of *B. coli*. Produces acid and gas in media containing the hexoses, mannitol, rhamnose, arabinose, maltose, sorbitol and sometimes dulcitol; but not in xylose, lactose, sucrose, salicin, inositol, dextrin, raffinose, inulin, or adonitol. Milk first turned slightly red and later (usually only after two weeks) neutral or slightly blue. Neutral red reduced, lead acetate media not browned. Gelatin not liquefied. Indol not produced. Characteristic serum agglutination reactions. Found in human stools and urine as actual or potential cause of one form of paratyphoid fever.

Culture 16 in our collection has been taken as a type of *B. paratyphosus*. It was obtained from the Rockefeller Institute in 1911 labelled Schottmüller A.

Clearly allied to *B. paratyphosus* is *B. pullorum*, described by Rettger in 1900 as the causative agent in bacillary white diarrhea of chicks. This organism, as shown by Rettger and Koser (1917), Hadley, Caldwell, Elkins and Lambert (1917) and Hadley, Elkins and Caldwell (1918) has all of the ordinary cultural and fermentative reactions of the A paratyphoids except that it is non-motile and fails to attack maltose or dulcitol, and that its agglutinative relations are closer with *B. typhosus* than with any of the paratyphoids. It may be defined as follows.

*B. pullorum* Rettger. Gram negative non-spore-forming non-motile rods. Colonies on gelatin somewhat intermediate between the thin translucent irregular colonies of *B. typhosus* and the convex regular colonies of *B. coli*. Produces acid and gas in media containing the hexoses, mannitol, rhamnose, arabinose and sorbitol; but not in maltose, dulcitol, xylose, lactose, sucrose, salicin, inositol, dextrin, raffinose, inulin or adonitol. Milk first turned slightly acid, later, but only slowly, neutral or slightly alkaline. Lead acetate not reduced. Gelatin not liquefied. Indol not produced. Exhibits group agglutination with *B. typhosus*. Causative agent of bacillary white diarrhea

in young chicks and found (without definite pathological symptoms) in ovaries of adult fowls.

Our strain 277 is a type of this species. It was isolated by Jones at Ithaca and sent to us by Parke Davis and Company in 1911 with the number 0233.

A special variety of this species has been described by Hadley (Hadley, Caldwell, Elkins and Lambert, 1917) as causing disease in adult fowls, differentiated from the typical *B. pullorum* by failure to form gas, but otherwise identical with it.

CHARACTERISTICS OF COLON-TYPHOID BACTERIA OF GROUP IV  
(FERMENTING MANNITOL, XYLOSE AND RHAMNOSE,  
GENERALLY PRODUCING GAS)

The fourth of the general groups into which we have divided the colon-typhoid series is that of which the B paratyphoids are the most typical examples. All of these forms ferment xylose and rhamnose and some of them arabinose as well, all produce a rather prompt alkalinity in milk, and all differ in agglutinative reactions from the A paratyphoid types. In general vigor of growth and fermentative power these forms stand nearer to *B. coli* than do any other of the non-lactose-fermenting organisms.

The first type of organism which we are inclined, with some doubt, to place in this group is the causative agent of fowl typhoid (*B. gallinarum*). This form, which was not represented in our own series, has the general fermentative reactions of the B. paratyphoids (see page 440) except that it attacks dextrin also, and that it fails to form gas. In view of the lack of correlation between gas production and power to attack various carbohydrates exhibited by the Morgan bacillus and the two varieties of *B. pullorum*, we are inclined to lay less stress on the former than on the latter characteristic. We cannot therefore agree with Hadley, Caldwell, Elkins and Lambert (1917) who believe that *B. gallinarum* is more closely allied to *B. typhosus* than is *B. pullorum*. All three organisms are however allied by their agglutinative reactions. According to Hadley, Elkins and Caldwell (1918) the milk reaction of *B. gallinarum* is of the B para-

typhoid type and the simultaneous fermentation of xylose and arabinose, a phenomenon which is so characteristic of paratyphoid B, seems to indicate that *B. gallinarum* may best be considered as a member of this group. It is certainly intermediate in its characteristics between *B. typhosus* and paratyphoid B; and it is of little consequence on which side of the arbitrary line between these types it may be placed.

The organism of fowl typhoid was described by Klein in 1889 as *B. gallinarum* and is probably identical with *B. sanguinarium* of Moore (1895). It may be characterized as follows.

*B. gallinarum* Klein. Gram-negative, non-spore-forming non-motile rods. Forming colonies on gelatin somewhat intermediate between the thin translucent irregular colonies of *B. typhosus* and the regular convex colonies of *B. coli*. Ferments the hexoses, mannitol, maltose, arabinose, xylose, dulcitol, dextrin, rhamnose, sorbitol, but not adonitol, salicin, lactose, sucrose or raffinose. Forms acid but no gas. Milk, first acid, quickly turning alkaline and developing the translucent appearance characteristic of paratyphoid B. Does not liquefy gelatin or produce indol. Exhibits group agglutination reaction with *B. typhosus*. Found as causative agent in fowl typhoid.

Hadley, Elkins and Caldwell (1918) describe two allied species, *B. pfaffi* which differs from *B. gallinarum* in failing to ferment dextrin or dulcitol and in fermenting salicin; and *B. jeffersoni* which fails to ferment dulcitol and produces no change at all in milk.

*B. avisepticus*, the causative organism in fowl cholera, is a member of the Pasteurella group and does not belong in the colon typhoid series at all.

The more familiar organisms of the paratyphoid B group include, as pointed out above (p. 439) three distinct types, *B. suipestifer* (xylose + arabinose - inosite -), *B. enteritidis* (xylose + arabinose + inosite -), and paratyphoid B, itself (xylose + arabinose + inosite +).

Of the first of these types we had 5 strains in our collection, 3 of which were sent to us as *B. cholerae-suis*, one as *B. paratyphi*, and one as *B. sternbergii*. All fermented the hexoses, mannitol,



xylose and rhamnose but not arabinose, lactose, sucrose, salicin or inosite. Two attacked dulcitol and three did not. Milk was in all cases turned blue in two days and deepened in color progressively. Gelatin was not liquefied, indol was not produced, and lead acetate was not blackened. One strain of *B. cholerae-suis*, otherwise the same as the five mentioned, failed to ferment rhamnose.

Of the paratyphoid B forms which ferment arabinose as well as xylose and rhamnose, we had 24 strains in our collection. Five were originally sent in as paratyphoids, five as *B. enteritidis*, six as mouse and rat viruses of various sorts (*B. danysz*, *B. murium*, *B. murisepticus*), three as *B. abortivus*, and five under other names (*B. typhi-suis*, *B. typhosus*, *B. pullorum*, *B. icteroides*, *B. paracoli*). All were alike in fermenting the hexoses, mannitol, xylose, arabinose, and rhamnose and in failing to ferment lactose, salicin and sucrose. Dulcitol was attacked by nineteen strains and inosite by six (strains 22, 30, 169, 235, 237, 589). None formed indol or liquefied gelatin. Lead acetate was browned by all but the three strains of *B. abortivus*. Litmus milk was slightly reddened in two days but by the sixth day was always distinctly alkaline and became progressively more blue with a characteristic translucency.

These results obtained by us are in accord with the findings of previous students of this group, of whom the most important have been Boycott (1906), Bainbridge (1909), Harding and Ostenberg (1912), Jordan (1917), Krumwiede, Pratt and Kohn (1917), and Krumwiede, Kohn and Valentine (1918).

All observers agree that the paratyphoid organisms are motile. Boycott (1906), Smith (1915) and Jordan (1917) point out that they ferment maltose, and sorbitol; Jordan (1917) and Krumwiede, Kohn and Valentine (1918) report failure to attack raffinose, adonitol, dextrin, inulin and erythritol. The reduction of neutral red is described by Sacquépée and Chevrel (1906), Bainbridge (1909), and Hollande and Beauverie (1915). High resistance to green dye is reported by various observers.

Of the three species included in this group the first, distinguished by failure to attack arabinose and generally negative

results in dulcitol, is the organism isolated by Salmon and Smith from cases of hog cholera in 1885. This form has been confused with an organism of the hemorrhagic septicemia group associated with swine plague from which it is quite distinct. The type found in hog cholera should bear the name *B. suipestifer* and the swine plague organism that of *B. suisepiticus*. *B. suipestifer* according to Jordan (1917) is a form quite constantly of porcine origin, while both *B. enteritidis* and paratyphoid B are commonly isolated from human sources. It is distinguished from the human forms by failure to brown lead acetate media and, according to Krumwiede, Kohn and Valentine (1918) by the power to reduce Andrade's indicator in glucose-serum-water. It may be defined as follows, culture 258 obtained from Parke Davis and Company in 1911 labelled "Boxmeyer, Belle Plain, 1903 No. 053" being taken as the type.

*B. suipestifer* Kruse. Gram negative, non-spore-forming, motile rods. Forming colonies on gelatin intermediate between the thin translucent irregular colonies of *B. typhosus* and the regular convex colonies of *B. coli*. Ferments the hexoses, mannitol, maltose, xylose, rhamnose and sorbitol with formation of gas. Does not ferment arabinose, lactose, sucrose, salicin, inosite, raffinose, adonitol, dextrin, inulin or erythritol. Generally fails to attack dulcitol. Turns milk first slightly acid, reverting in five days to an alkaline reaction, the color of the medium deepening with the development of a translucent appearance. Neutral red and malachite green reduced, lead acetate not blackened. Indol not produced. Gelatin not liquefied. Group agglutination with paratyphoid B. Found in intestines of hogs and as secondary invader of tissues in hog cholera.

*B. enteritidis* is distinguished from *B. suipestifer* by fermentation of arabinose, general fermentation of dulcitol, and blackening of lead acetate media, and from paratyphoid B by failure to ferment inosite. It exhibits distinct and characteristic serum agglutinative reactions. It may be defined as follows.

*B. enteritidis* Gaertner. Resembles *B. suipestifer* in all cultural characters except that arabinose is always, and dulcitol

generally fermented, and that lead acetate media are blackened. Exhibits characteristic serum agglutination reactions which distinguish it from either *B. suispestifer* or paratyphoid B. Found in human intestine and as causative factor in outbreaks of food poisoning. The type is our strain 18, a Gaertner strain obtained from the Rockefeller Institute in 1911.

Finally there remains for consideration the true paratyphoid B organism which is distinguished from *B. enteritidis* by the fermentation of inositol and by its serological reactions. So far as we are aware no one has ever given this very distinct type a specific name in proper Latin form; and since the identification of the paratyphoids merely as "A" and "B" is quite misleading as to their true biological relationships, we suggest the name *B. schottmulleri* for the paratyphoid B organism, to be defined as follows.

*B. schottmulleri*. Resembles *B. suispestifer* in all cultural characters except that arabinose and inositol are always fermented and dulcitol is generally fermented and that lead acetate media are browned. Its agglutinative reactions distinguish it from *B. enteritidis*. Found in human intestines and urine and as causative agent in paratyphoid fever and food poisoning outbreaks. The type is our strain 22, a Schottmüller strain obtained from the Rockefeller Institute in 1911.

The various forms of mouse and rat virus appear not to be distinct entities according to Bainbridge (1909), Savage (1912) and Krumwiede, Pratt and Kohn (1917); and our own results bear out their view that some of these viruses belong to each of the three species listed above, the Danyz virus being a variety of *B. enteritidis*.

The peculiar organism known as the Morgan bacillus may perhaps best be considered here, although its exact relationships are obscure, since it possesses the power of forming gas but can attack only a limited number of carbohydrates. Five of our cultures were of this type. Two of these strains came to the Museum as Morgan bacilli while the others were labeled respectively *B. communis*, *B. cuniculicida* and *B. pseudotuberculosis*, but all were alike in their failure to ferment the higher carbohydrates and in the production of a small but definite amount of

gas (about 5 per cent) in glucose, mannose, fructose and galactose. The acid production in the hexoses was vigorous in all these five strains when they were first tested in 1917, but when re-examined in 1919 one of them (strain 586) had lost its power to attack any of the sugars. This observation, together with the fact that one of the strains (strain 139) which attacked the hexoses, but only the hexoses, in both our tests was originally obtained from D. D. Jackson as *B. communis* A (fermenting mannitol, dulcitol, lactose and raffinose) again suggests a loss of fermentative power during prolonged cultivation in the laboratory. In milk cultures all of the strains produced an immediate alkaline reaction, later decolorizing the litmus to some extent so as to produce a chalky blue appearance. This fact is probably associated with the rapid decomposition of protein characteristic of this form. Kendall, Day and Walker (1913) note a production of 4.2 mg. of  $\text{NH}_3$  per 100 cc. of glucose broth for the Morgan bacillus in one day, as against less than 0.1 mgm. for *B. typhosus* and 0.2 for the paratyphoids. All of our strains formed indol and failed to liquefy gelatin.

Morgan (1906) describes this organism as motile and differentiates it from the hog cholera type by its reaction in litmus milk, active production of indol and failure to ferment maltose, arabinose and dextrin. Organisms of this type have been studied by Lewis (1912), Alexander (1912) and particularly by Graham-Smith (1912). The latter investigator prepared an elaborate system of classification of the non-lactose-fermenting bacteria, his type G including the forms which produce gas in glucose but not in mannitol media, being the most common of all the types in the lactose-negative group. Tribondeau and Fichet (1916), who isolated the Morgan bacillus from 13 cases of dysentery originating in the Dardanelles note, in addition to the characters described above, that it produces a fluorescence in neutral red broth and rapidly blackens lead acetate. They found several strains of related organisms which at first fermented maltose and sucrose but which later lost this power.

It is evident that the Morgan bacillus, wherever its closest relationships may lie, constitutes a fairly definite type, of com-

mon occurrence in the human intestinal canal. So far as we are aware it has nowhere been given a scientific name in proper form; and we have therefore called it *B. morgani*, characterizing the species as below and considering culture 692 of the American Museum collection as its type. This strain was isolated from the stool of an infant at the Providence City Hospital by H. E. Smiley.

*B. morgani*. Gram-negative, non-spore-forming motile rods, producing thin translucent irregular colonies on gelatin. Rapid formation of acid and slight gas production in media containing the hexoses. Milk turned gradually blue. Indol formation vigorous. Gelatin not liquefied. Produces fluorescence in neutral red broth and blackening in lead acetate media. Found in normal and diarrheal stools.

This organism would seem to be allied to *B. shigae* by its limited fermentative powers, to *B. dysenteriae* (or *B. coli*) by indol production, and to the paratyphoids by the formation of gas. It may perhaps represent an extreme variant of the variable *B. dysenteriae* group but we have considered it with the paratyphoids on account of its gas production.

One other type of non-lactose-fermenting organism often considered as a member of the colon-typhoid group is the *B. proteus*; but on account of the fundamental differences in morphology and metabolism (fermentation of sucrose but not lactose and very vigorous decomposition of proteins) we believe the *Proteus* forms should not form a part of this series at all.

CHARACTERISTICS OF COLON-TYPHOID BACTERIA OF GROUP V  
(FERMENTING LACTOSE AND THE SIMPLER CARBOHYDRATES WITH  
PRODUCTION OF EQUAL VOLUMES OF CO<sub>2</sub> AND H<sub>2</sub>)

This group of organisms, of which *B. coli* is the most familiar example, is distinguished from all the organisms of the dysentery-typhoid-paratyphoid series by the power of fermenting lactose and from the *B. aerogenes* types by the fact that in its fermentation it produces equal volumes of CO<sub>2</sub> and H<sub>2</sub> (low ratio fermentation of Clark and Lubs). Its members differ from the

typhoid and paratyphoid organisms in less active motility, in more vigorous growth on media (with more convex and regular colonies), in strong reducing action, in the formation of indol, and in characteristic differences in resistance to various antiseptics.

In the Museum collection we found 42 different strains belonging to this general group. All of them fermented the hexoses, mannitol, xylose, arabinose, rhamnose, and lactose, with the production of rapid and permanent acidity and gas. None attacked inosite while with salicin, dulcitol and sucrose results were variable. All acidified and coagulated milk promptly, a firm clot being generally formed in six days with marked decolorization of the litmus. Indol was formed by all but 10 strains. Gelatin was not liquefied.

In regard to other reactions which we did not study it may be noted that maltose is fermented, according to Levine and other workers; that adonitol is generally not attacked (Mac Conkey, 1909, Rogers, Clark and Lubs 1918); and that starch, glycogen, inulin and dextrin give negative results according to Chantemesse and Widal (1891), Drigalski and Conradi (1902), Burk (1907), MacConkey (1909), Rogers, Clark and Evans (1914, 1915), and Levine (1916). Murray (1916) however reports that a fair proportion of strains from bovine feces do attack inulin.

Twort (1907) in a study of the utilization of a large series of unusual glucosides found that *B. coli* generally attacked euonymin, iridin, senegin, coniferin, arbutin, salicin, syringin, quilajinic acid, populin, camellin, and globularin. The hydrolytic splitting of esculin with the production of sugar and a substance called esculetin which reacts with iron citrate to produce a brown color has been used by Harrison and van der Leek (1909) and others as a test for this group. The colon bacillus generally reduces nitrates and neutral red (Rothberger, 1898); but does not brown lead acetate media (Sacquépée and Chevrel, 1905, Burnet and Weissenbach, 1915). It produces an acid reaction in Capaldi and Proskauer's medium 1 but not in medium 2. It has a relatively high resistance to acid (Hankin, 1899, Winslow and Lochredge, 1902) potassium tellurite (Davis, 1914), and cholesterol (Manfredi, 1917); but a relatively low resistance to

the green dyes (Loeffler, 1903, 1906, Lentz and Tietz, 1903, 1905, Krumwiede and Pratt, 1914), caffeine (Hoffman and Ficher, 1904), and bile salts (Jackson and Melia, 1909).

The further subdivision of the colon group of bacteria was first attempted on the basis of the fermentation of sucrose. Germano and Maurea (1893) for example distinguished one type which fermented sucrose and decolorized jequirity solution and another which failed to give either of these reactions. Smith (1893, 1895a) in his classic investigations emphasized the importance of the sucrose-positive and sucrose-negative varieties. Durham (1901) gave the sucrose-positive form the name *B. communior*. Winslow and Walker (1907) and Howe (1912) pointed out that the forms which attack sucrose also attack raffinose and vice versa.

Very elaborate classifications of the colon group were developed by MacConkey (1905), Bergey and Deehan (1908), and Jackson (1911), based primarily on fermentation of sucrose and dulcitol and secondarily on fermentation of adonitol and inulin, the Voges-Proskauer reaction, motility, indol formation and liquefaction of gelatin. The principal types recognized by these authors were as follows.

- I. Sucrose - Dulcitol - *B. acidi-lactici*
- II. Sucrose - Dulcitol + *B. coli-communis*
- III. Sucrose + Dulcitol - *B. coscoroba* [*B. aerogenes* (V. - P. +), and *B. cloacae* (V. - P. +, Gelatin +) also belong here]
- IV. Sucrose + Dulcitol + *B. communior* [*B. neapolitanus* (non-motile) and *B. pneumoniae* also belong here]

These classifications were all defective in grouping the Voges-Proskauer positive, indol negative, gelatin positive organisms with the various fermentative types of the true colon group. We now know that these forms belong to the high ratio (*B. aerogenes*) group and should not be placed with *B. coli* and *B. communior*. *B. acidi-lactici*, *B. coli*, *B. coscoroba* and *B. communior* may however be considered to represent distinct types, if the fermentation of dulcitol is really a characteristic of specific importance. There seems grave doubt however whether this

alcohol should be given a prominent place in classification; for as pointed out above dulcitol throughout the colon-typhoid series shows less correlation with other properties than any other carbohydrate and under experimental conditions shows the most marked tendency to spontaneous variations.

Rogers, Clark and Evans (1914), in a study of colon organisms from bovine feces, found that sucrose and raffinose fermentations are directly and almost perfectly correlated, and that the sucrose positive forms are generally dulcitol positive and adonitol negative, while the sucrose negative forms vary in both dulcitol and adonitol. The sucrose negative forms may be divided (although the authors do not point this out) into a dulcitol + adonitol - group and a dulcitol - adonitol + group. Kligler (1914) emphasized the value of salicin as a differential test substance, finding dulcitol inversely correlated with salicin among the sucrose positive forms and positively correlated among the sucrose negative forms. Failure to attack glycerol was most common among the salicin-fermenters.

The most extensive study of this group of organisms is that conducted by Levine (1917). This investigator worked with 333 strains isolated from soil, sewage and the feces of men, horses, sheep, pigs and cows and devoted special attention to the mutual correlation of the various characters studied. One hundred and eighty-two of the strains were of the methyl-red-positive, Voges-Proskauer-negative type. Sucrose and raffinose were almost perfectly correlated and the author divides his methyl red positive organisms first on the sucrose fermentation. The sucrose negative forms are next subdivided according to their action on salicin, giving *B. coli* (sucrose - salicin +) and *B. acidi-lactici* (sucrose - salicin -) as species. Among the sucrose positive strains the most natural division, according to correlated characters, is between a motile type *B. communior* and a non-motile series which can be further subdivided into a salicin positive species *B. neapolitanus* and a salicin negative species, *B. coscoroba*. *B. communior*, *B. neapolitanus* and *B. coscoroba* were much more common in animal than in human feces, while *B. coli* and *B. acidi-lactici* were commonly isolated from human sources.



It is impossible to make a direct and comprehensive comparison of the results of these various investigators since each one used some tests not applied by the others. In our own investigations we did not study motility or the fermentation of adonitol and salicin. The following characteristics would appear to be indicated for the principal types recognized by Kligler and Levine.

	SUCROSE	SALICIN	DULCITOL	ADONITOL	MOTILITY
<i>B. neapolitanus</i> .....	+	+	-	-	-
<i>B. communior</i> .....	+	-	+	-	+
<i>B. coscoroba</i> .....	+	-	+	-	-
<i>B. coli-communis</i> .....	-	+	+	-	+
<i>B. immobilis</i> .....	-	+	+	-	-
<i>B. Grünthal</i> .....	-	-	-	+	+
<i>B. acidi-lactici</i> .....	-	-	-	+	-

All recent observers are agreed as to the superior value of salicin as compared with dulcitol for the primary subdivision of this group; and Levine's main classification, based on so large a series of strains can safely be accepted as in the main a correct one. We cannot feel certain however that the presence or absence of motility is a sufficient basis for the establishment of species, in view of the highly inconstant results obtained by other students of this property. We are therefore inclined to recognize four distinct species, as characterized below, with three varieties, based on motility.

*B. neapolitanus* Fraenkel. Gram-negative non-spore-forming rod. Non-motile. Vigorous growth on media, colonies more regular and convex than those of *B. typhosus*. Produces prompt and permanent acidity and gas composed of equal volumes of CO<sub>2</sub> and H<sub>2</sub> in media containing the following substances; the hexoses, maltose, mannitol, xylose, arabinose rhamnose, lactose, sucrose, salicin and esculin, but not as a rule dulcitol or adonitol, and never inosite, starch, glycogen, inulin or dextrin. Turns milk strongly acid and coagulates it in six days at 37°. Generally forms indol but does not liquefy gelatin. Produces an acid reaction in Capaldi-Proskauer medium 1 but

not in medium 2. Does not give Voges-Proskauer reaction. Reduces neutral red and generally nitrates. Does not brown lead acetate media. Exhibits high tolerance to acids, potassium tellurite and cholesterol, but relatively low tolerance to the green dyes, alkaloids and bile salts. Found in feces of higher animals and man, particularly the former.

Our type of this species is strain 126, isolated from a urinary fistula in 1911 and received by us from Dr. Jackson of the Mt. Prospect laboratory as *B. aerogenes* A2.

*B. communior* Durham. Differs from *B. neapolitanus* in failing to ferment salicin, in generally fermenting dulcitol and in being sluggishly motile. Our type is 137 isolated from feces by Dr. Frazer in 1911 and sent to us by Jackson as *B. communior*.

var. *coscoroba*, differs from *B. communior* in lacking motility.

*B. coli* Escherich. Differs from *B. neapolitanus* as described above in failing to ferment sucrose and raffinose, in generally fermenting dulcitol, and in exhibiting sluggish motility. Specially abundant in human feces. Type, 125 isolated from a case of cystitis in 1910 and sent to us by Jackson as *B. communis* B.

var. *immobilis* differs from *B. coli* in being non-motile.

*B. acidi-lactici* Grotenfelt. Differs from *B. neapolitanus* in failing to ferment sucrose, raffinose or salicin and in generally fermenting adonitol. Specially abundant in human feces. Type, 131 from Jackson as *B. acidi-lactici* B. Isolated from feces.

var. *Grünthal*. Differs from *B. acidi-lactici* in being motile.

All of the type specimens listed above are described by Jackson (1911). He states that type 137 does not ferment mannitol but we find that it does.

The organisms in our collection did not show by any means a clear cut correlation between the fermentation of sucrose, salicin and dulcitol. Classifying them according to their action upon sucrose and salicin we found 15 strains of *B. neapolitanus* (of which only 5 were negative in dulcitol), 4 of *B. communior* (of which 3 were negative in dulcitol), 16 of *B. coli* (of which 10 were positive in dulcitol), and 7 of *B. acidi-lactici* (of which 4 were negative in dulcitol). The names under which these organisms were sent to us corresponded fairly well with the species as

identified except that the *B. neapolitanus* series included two strains called *B. aerogenes*, one *B. pneumoniae*, and one *B. ozenae*, one *B. sternbergii* and one *B. bovisepiticus*; the *B. communior* series one *B. bronchicanis*; the *B. coli* series, a *B. diphtheriae-columbarum*, two *B. cholerae*, one *B. astheniae*, one *B. anaerogenes*, one *B. aerogenes*, and 3 *B. acidi-lactici*; and the *B. acidi-lactici* series two *B. coli*, one *B. voldagsen*, one *B. danysz*, and one *B. dysenteriae* (!).

CHARACTERISTICS OF COLON TYPHOID BACTERIA OF GROUP VI  
(FERMENTING LACTOSE AND THE SIMPLER CARBOHYDRATES  
WITH PRODUCTION OF TWO OR MORE VOLUMES OF  
CO<sub>2</sub> TO ONE OF H<sub>2</sub>)

The last group of the colon-typhoid series, of which *B. aerogenes* and *B. cloacae* are the principal types, was recognized by Escherich (1885) on the basis of greater plumpness of the cell form, lack of motility and more rapid coagulation of milk, by Smith (1893a) on the basis of heavier growth and tendency to capsulation, and later (Smith 1895a) on rapid gas production and higher ratio of CO<sub>2</sub> to H<sub>2</sub>. Durham (1901) notes the fermentation of starch and inulin and the Voges-Proskauer reaction as characteristic, Grimbert and Le Gros (1900) and Jordan (1903) the failure to form indol. Finally Harden and Walpole (1905) and Rogers, Clark and Davis (1914) demonstrated conclusively the fundamental difference in carbohydrate metabolism between these forms and those of the *B. coli* type. Rettger (1903) showed that the differences in protein metabolism are equally distinct, *B. aerogenes* being much less active in this respect forming mercaptans, skatol, phenols, aromatic oxy-acids and skatol-carbolic acid much more slowly. Ferreira, Horta and Paredes (1908) report that *B. aerogenes* and *B. cloacae* ferment sucrose but not dulcitol and give a rose color with no luster on the Endo medium. Rivas (1908) points out that *B. aerogenes* exhausts the carbohydrate constituent in glucose broth much more rapidly than does *B. coli*. Rogers, Clark and Davis (1914) described a series of high gas ratio cultures isolated

from milk, and pointed out that these forms attack sucrose, raffinose, and starch and liquefy gelatin more frequently than do the *B. coli* forms, and on the other hand attack dulcitol and glycerol less often. Rogers, Clark and Evans (1915) studied 166 high ratio cultures from grains and described six different types of which the most abundant had the following characteristics.

Gelatin-liquefying forms. Not capsulated, indol negative, sucrose positive, raffinose positive, mannitol positive, dulcitol positive, glycerol generally positive, starch negative, inulin negative, adonitol negative.

Gelatin-non-liquefying forms. Sometimes capsulated, sometimes fermenting starch, inulin and adonitol, rarely fermenting mannitol or dulcitol, otherwise as above.

Rogers, Clark and Lubs (1918) describe a series of high ratio strains from human feces as generally indol negative, sucrose, raffinose, mannitol and adonitol positive, inulin negative, and generally dulcitol negative. According to these results the high ratio type common in human feces differs from that normal on grains by the fermentation of mannitol and adonitol.

Levine (1916b) states that *B. aerogenes* usually ferments sucrose, raffinose, salicin, glycerol, dextrin and starch, but rarely dulcitol. In a later paper (Levine 1917) he confirms the conclusion of Kligler (1914) (opposed to that of Rogers, Clark and Evans) that gelatin liquefaction and glycerol fermentation are negatively correlated. Of 151 strains of the high ratio type (isolated from soil or in a few cases from sewage) all fermented mannitol, and almost all sucrose, raffinose, and salicin. He recognizes two species, *B. aerogenes*, which rarely liquefies gelatine, is non-motile and forms gas from glycerol and starch; and *B. cloacae* which is a liquefying organism failing to ferment either glycerol or starch.

In general it is evident that the organisms of this group may be subdivided into a type which liquefies gelatin and one which fails to do so. The former, *B. cloacae*, is not capsulated, and does not ferment starch, inulin or adonitol, while *B. aerogenes* is often capsulated and sometimes ferments starch and inulin

and adonitol. According to Rogers, Clark and Lubs the grain type of *B. aerogenes* differs from the fecal type in its failure to attack mannitol and dulcitol. Twort (1907) reports the frequent fermentation by *B. aerogenes* of a number of glucosides not attacked by *B. coli* such as cerberid, periplocin, cathartinic acid, amygdalin, sapotoxin, saponin, bryanin, convallamarin, digitalin, strophanthin, coronillin, gratiolin and phloridzin.

Our own series of cultures included 8 of the *B. cloacae* and 23 of the *B. aerogenes* type. All gave positive results in the hexoses, mannitol, xylose, arabinose, rhamnose, lactose, salicin and sucrose. All coagulated and decolorized milk, the clot being formed by 19 strains in six days at 37°, by the other 12 more slowly. Only 6 strains formed indol. All but 1 were alkaline to methyl red and all but 8 gave the Voges-Proskauer reaction. Only 9 fermented dulcitol and 16 fermented inosite. A special study showed that our gelatin-liquefying strains all failed to ferment glycerol, while the *B. aerogenes* did attack this substance.

We may therefore recognize at least two species in this group, as follows.

*B. cloacae* Jordan. Gram-negative non-spore-forming rod. Non-motile. Vigorous growth on media, colonies even more regular and convex than those of *B. coli*. Ferments the hexoses, maltose, mannitol, xylose, arabinose, lactose, rhamnose, sucrose, raffinose, and salicin, often inosite, but generally not dulcitol, glycerol, starch, inulin or adonitol. Produces vigorous and complete destruction of the carbohydrates which it attacks forming acetyl-methyl-carbinol (Voges-Proskauer reaction) and at least twice as much CO<sub>2</sub> as H<sub>2</sub>, the reaction of the medium becoming moderately acid and then reverting to a lower degree of acidity. Acidifies and coagulates milk but not quite so promptly as *B. coli*. Fails to form indol. Liquefies gelatin slowly. Found in human and animal feces, sewage and soil. Our type of this species is type 23 isolated by Jordan from the Chicago Drainage canal in 1899.

*B. aerogenes* Escherich. Gram-negative non-spore-forming rod. Generally non-motile. Frequently capsulated. Growth on media vigorous, colonies convex and often viscid in texture.

Ferments the hexoses, maltose, xylose, arabinose, rhamnose, lactose, sucrose, raffinose, salicin, usually glycerol, starch, and inulin, and sometimes inositol and adonitol, dulcitol and mannitol. Produces vigorous and complete destruction of the carbohydrates, forming acetyl-methyl-carbinol (Voges-Proskauer reaction), and at least twice as much  $\text{CO}_2$  as  $\text{H}_2$ , the reaction of the medium becoming moderately acid and then reverting to a lower degree of acidity. Acidifies and coagulates milk but not quite so promptly as *B. coli*. Fails to form indol or liquefy gelatin. Found in human and animal feces and sewage but particularly in soil and on grains.

Our type strain is type 240 received from Johns Hopkins in 1911 with following history—"probably a descendant of the original capsule bacillus of Pfeiffer . . . reaction identical with *B. aerogenes*."

A variety which fails to ferment mannitol and adonitol is the commonest form on grains.

*B. aerogenes* represents the extreme of fermentative power in the colon-typhoid series; and on the assumption that the course of evolution has been marked by progressive loss of fermentative power and acquisition of the parasitic habit it may be considered the most primitive type of the whole group. It has been suggested by one of us (Kligler, 1917) that three different lines of evolution may have started from this type, the first leading through capsulated streptococci to the diverse forms of streptococci and pneumococci of the present day, the second through *B. coli*, *B. typhosus* and the dysentery organisms to the hemorrhagic septicemia group, and the third through *B. cloacae* and *B. proteus* to the saprophytic spore-bearing and pigment bacteria.

There is one group of organisms usually considered to be related to *B. aerogenes* whose affiliations appear to be obscure. These are the capsulated forms frequently met with in association with certain pathological conditions. Capsulated bacilli were first described by Friedlander from pneumonia in 1883; and important studies of this group have been made by Fricke (1896), Strong (1899), Perkins (1904) and Coulter (1917).

All of these workers assumed that the pathogenic capsulated bacilli with which they worked (*B. pneumoniae*, *B. ozenae*, *B. rhinoscleromatis*) were related to *B. aerogenes*; but all agree that they differ from *B. aerogenes* in producing a more distinct capsule and colonies of a translucent syrupy consistency; in forming indol; and in fermenting sucrose but not lactose. The last property is a very rare one in the colon-typhoid group, although the fermentation of sucrose but not lactose is characteristic of the *Proteus* bacilli. In our Museum collection we had a number of organisms which were sent to us bearing the names of the capsulated pathogenic forms; but none of them exhibited the characteristics described by Perkins, Strong and Coulter. Of three cultures sent to us as *B. pneumoniae* one was finally classed as *B. alcaligenes*, one as *B. neapolitanus*, and one as *B. aerogenes*. Of two sent in as *B. ozenae*, one was *B. alcaligenes* and one *B. neapolitanus*. Two sent in as *B. capsulatus* and one as *B. rhinoscleromatis* all proved to be *B. aerogenes*. Thus of eight strains originally believed to be members of this group, four were of the methyl red negative, Voges-Proskauer positive type, while two belonged at the other extreme of the paratyphoid series, with no fermentative powers at all. It seems evident, either that we are dealing with an extraordinarily variable group or that forms which are not really related have been identified as of this type merely because of the possession of a capsule. (Fletcher, 1918, has recently described capsulated forms of paratyphoid and dysentery bacilli). The evidence brought forward by Strong (1899), Perkins (1904) and Coulter (1917) is so completely concordant as to make it clear that there is a distinct type of capsulated bacillus which ferments sucrose and not lactose and possesses considerable pathogenic powers; but in view of the incompleteness of our knowledge of its other reactions we are somewhat uncertain of its relations to the colon-typhoid group.

REVIEW OF THE GENERAL CHARACTERISTICS OF THE SPECIES OF  
THE COLON-TYPHOID GROUP

The principal characters of the species of colon-typhoid bacteria as defined above are presented for convenient comparison in the table below, and the fermentative relationships of certain of the more important forms are presented in figure I.

If *B. aerogenes* be taken as a representative of the most primitive type it is evident that there is a more or less steady decrease

SPECIES	HEXOSES	MALTOSE	MANNITOL	XYLOSE	ARABINOSE	RHAMNOSE	SORBITOL	DULCITOL	LACTOSE	SALICIN	SUCROSE	RAFFINOSE	INOSITE	DEXTRIN	GAS	VOGES-PROSKAUER	METHYL RED	MILK	GELATIN	INDOL	LEAD ACETATE	MOTILITY	PATHOGENIC
<i>B. aerogenes</i> .....	+	+	#	+	+	+	+	#	+	+	+	+	+	+	+	+	+	Coag.	-	-	-	-	-
<i>B. cloacae</i> .....	+	+	+	+	+	+	+	#	+	+	+	+	+	+	+	+	+	Coag.	+	-	-	-	-
<i>B. neapolitanus</i> ...	+	+	+	+	+	+	+	#	+	+	+	+	+	+	+	+	+	Coag.	-	+	-	-	-
<i>B. communior</i> ....	+	+	+	+	+	+	+	#	+	+	+	+	+	+	+	+	+	Coag.	-	+	-	-	-
<i>B. coli</i> .....	+	+	+	+	+	+	+	#	+	+	+	+	+	+	+	+	+	Coag.	-	+	-	-	-
<i>B. acidi-lactici</i> ...	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	+	Coag.	-	+	-	-	-
<i>B. morgani</i> .....	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	al.	-	+	+	+	+
<i>B. schottmulleri</i> ..	+	+	+	+	+	+	+	#	-	-	-	-	+	-	+	+	+	ac. al.	-	+	+	+	+
<i>B. enteritidis</i> ....	+	+	+	+	+	+	+	#	-	-	-	-	-	-	+	+	+	ac. al.	-	-	+	+	+
<i>B. suipestifer</i> ....	+	+	+	+	-	+	+	#	-	-	-	-	-	-	+	+	+	ac. al.	-	-	-	+	+
<i>B. gallinarum</i> ....	+	+	+	+	+	+	+	+	-	-	-	-	-	+	-	+	+	ac. al.	-	-	-	-	+
<i>B. pullorum</i> .....	+	-	-	+	+	+	+	+	-	-	-	-	-	-	-	+	+	ac. al.	-	-	-	-	+
<i>B. paratyphosus</i> ..	+	+	+	+	+	+	+	#	-	-	-	-	-	-	+	+	+	ac. al.	-	-	-	+	+
<i>B. typhosus</i> .....	+	+	+	+	+	+	+	-	-	-	-	-	-	+	-	+	+	ac. al.	-	-	+	+	+
<i>B. dysenteriae</i> ....	+	#	-	#	#	#	#	#	#	#	#	#	#	#	-	-	-	ac. al.	-	#	-	-	+
<i>B. shigae</i> .....	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	ac. al.	-	-	-	-	+
<i>B. alcaligenes</i> ....	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	al.	-	-	+	+	-

of fermentative power, down through the colon, paratyphoid and typhoid groups to *B. shigae* and *B. alcaligenes*. It cannot however be maintained that this progression necessarily or even probably represents the exact line of evolutionary development. In certain respects (as in its action on lead acetate media) *B. typhosus* seems more closely allied to *B. schottmulleri* than to *B. paratyphosus*. *B. dysenteriae* as noted exhibits highly variable reactions on maltose, xylose, arabinose, dextrin, rhamnose,



salicin and sucrose, which have therefore been left out of the chart entirely. *B. morgani* is another highly variable and unstable type, as is also *B. pullorum*; while the capsulated pathogenic forms allied to the Friedlander bacillus form an exceedingly puzzling complex. On the other hand *B. shigae* and *B. typhosus*

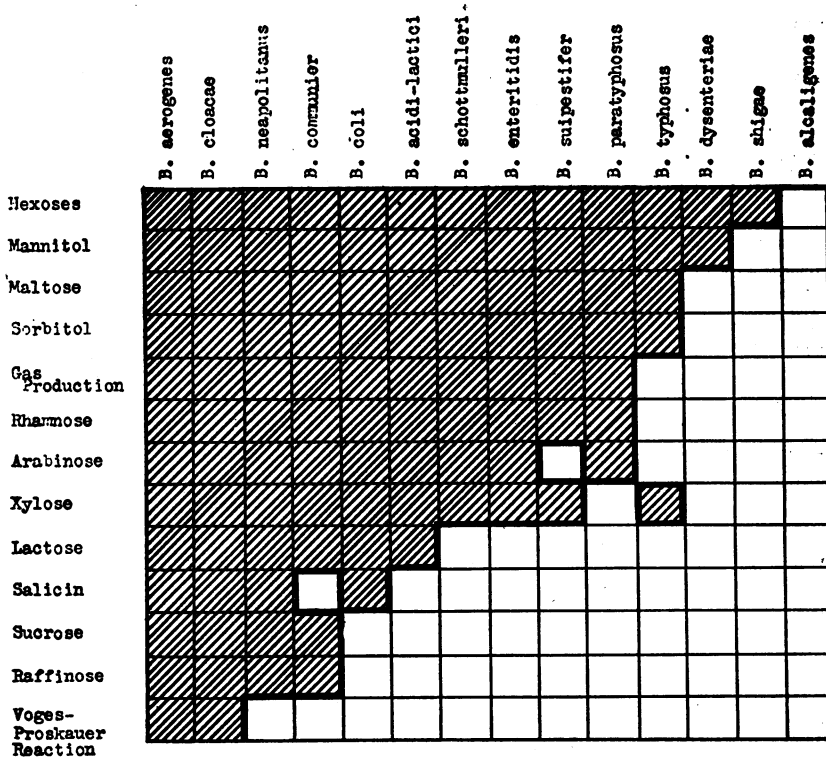


FIG. I

are species of very definite and constant characteristics; while *B. schottmulleri*, *B. enteritidis*, *B. suispestifer*, and *B. paratyphosus*, as well as the chief colon-aerogenes types, can be identified with reasonable ease.

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