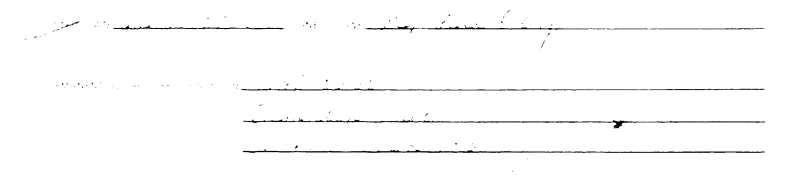
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FOR UNIVERSITY OF ALBEITA

ENTEROPACTE FACIATION OF GROUND MEATI

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I THESIS

SUBMITTED TO THE TAGU. TY OF GRADUATE STUDY S AND RESEARCH
IN ITTLE FULFILMENT OF THE REQUIFEMENT. A THE DEGREE

OF MASTER OF SCIENCE

PACULTY OF HOME ECONOMICS

PMON' ALBERTA

CPFING, 1977

THE UNIVERSITY OF ALBERTA FACULTY OF GRADUATE STUDIES AND RESEARCH

The undersigned certify that they have read, and recommend to the Faculty of Graduate Studies and Research, for acceptance, a thesis intimate inumeration and Identification of Enterotacteriassae in Ground Meats, submatted by Ng, Lai-King in partial fulfilment of the requirements for the degree of Master of Science.

Supervisor

Supervisor

November 1, 1976

ABSTSACT

The Health Protession Branch of Canada Health and wellfare, is proposing amendmence to the Canadian Food and Drug Act that would introduce pacterial standards for round meats. The proposed standards include limits for <u>Pscherichia</u> coll, determined by the Most Probable Number technique and the elevated temperature test. This is a time consuming method, which is not suitable for routine quality control purposes. This study compared different methods of enumerating Enterobacteriacore, coliform bacteria and E. coli in ground meats, using Violet Red Bile agar (VRFA), Mossel's <u>Enterobacteriaceae</u> medium (VRFA + 1% qlucose, VRBG) and Most Probable Number techniques ing Lauryl Tryptose broth (LST), Brilliant Green Bile 2% broth (BGB), Levine's EMB agar and EC medium. The counts obtained by these different methods and the types of organisms included in the counts were compared.

of the 169 samples of ground by included in the study, 62% fell within the proposed <u>E. coli</u> standard of less than 100 per gram, 27% had counts between 100 and 500 <u>E. coli</u> per gram, and 11% exceeded 500 per gram. The relationship between <u>E. coli</u>, coliform and <u>Enterobacteriaceae</u> counts was such that direct plating techniques (VRBA and VRBG) could not be used to estimate <u>E. coli</u> concentrations. However, based on the results of this study, VRBA and EC medium at elevated temperature could be used to give more

tipld results suitable for une in loutine quality control.

The bile precipitating of onlos isolated from VRRA vero primarily E. coli, E. agglowerans and S. liquefacions, whereas the typical "coliform" colonges growing on EMB vero primarily E. coli, C. freundii, K. pneumoniae and E. cloacae. When the isolates from all selective media were replated onto VRBA plates, over 50% of E. coli, C. freundii, K. pneumoniae, K. ozaenae and E. cloacae isolates grew on VRBA as bile precipitating (coliform) colonies. On EMB, over 50% of E. coli, C. freundii, K. pneumoniae and E. cloacae grew as typical "coliforms". In addition, other Enterobalt-eriaceae were frequently present in the ground meats, including E. agglomerans and S. liquefaciens.

In EC medium at 44.5°C and 24h incubation, the majority (92.2%) of organisms producing gas were \underline{E} . \underline{coli} type I (IMViC ++--). Increasing the temperature to 45.5°C, decreased the number of false positive \underline{E} . \underline{coli} results, but created the need to incubate the samples for 48h instead of 24h.

The FBL Minitek technique, using the biochemical tests selected for this study, proved to be satisfactory for identifying Enterobacterlaceae isolates from ground meats.

ACENUALEDGEMENTS.

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The author wishes to acknowledge the assistance of her chesis a Fisor, Dr. Michael Stiles, for the guidance and encouragement is the research and preparation of this thesis.

Thanks are also extended to Mr. Clare Shier of the Department of Educal Economy and Mr. Ray Weingardt of Educal Computers Services, inspectity of Alberta, for advice and quidance on situ analysis.

Bost of the research was parried out in the Department of Microbiology at the University of Alberta, and the technical assistance provided by the Department is gratefully acknowledged.

The research was carried out in association with a study to determine the microbiological quality of meat, sponsored by Agriculture Canada.

The advice of Dr. M. Finlayson, Alberta Provincial Laboratory of Public Health, and members of the Department of Medical Bacteriology, University of Alberta, is gratefully acknowledged.

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frquie	Flow diagram illustrating the tests used to differentiate the phenylulanine deaminase lacking <u>Enteropacteriaceae</u> which produce H ₂ S
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The significance of E. <u>Coll</u> is foods to econown.

different. B. <u>Coll</u> can survive on equipment and in food, and their presence does not accessifily indicate entire direct of recent faecal contamination. Herever, if <u>B. Coll</u> of faecal origin is present, enteric pathogens might survive with them. Although unbygien—andling of foods is a possible source of contamination, inadequate sanitation of equipment represents their most protable source. Organisms on equipment are from various sources, therefore other colliforms, lesides <u>B. <u>Coll</u>, are also emphasized to give a wider and more accurate indication of sanitation and hygiene.</u>

Collitorns are usually present to raw milk, therefore their presence in pasteurized milk and daily products would

And rester in a comparison of the entry of the entry of particular and contamination. The most of contaminations to indicate potential nealth hazards in all food on a feet with a red, which some forms of the menting anticopacter particular contamination of the entry of the entr

mealth Association (ABSA, 1965, 1971) recommended the Most Probable Sumber (MPN) technique, hackone or Lauryl Sultate Tryptose broths in used as a presumptive test, Brilliant Green File L% broth as a confirmed test and Bosin Methylene blue again or Endo again for the completed test. Subsequently, faecal and non-faecal strains must be differentiated. The Lauryl can be differentiated. The

enumeration of collisions (lactose-ferment <u>ferchact</u> enumeration of collisions (lactose-ferment <u>ferchact</u> enumeration of collisions and Desoxycholate Lactose agar have been d. The use of these solid media has been employed for ellucration of collisons in other foods. Modification of Violet Red Bile with addition of 1% plucose gives a medium on which <u>Enterobacteriaceae</u> can be enumerated (Massel, 1957; Mossel et al., 1962; Mossel and Ratto, 1970; Thatcher and Clark,

1 (6:21) .

Atypical colonies have been observed on Violet Red Bile war. There itypical colonies include coliform and non-objection factoria. When VSPA was used to enumerate coliforms in meats, large numbers of atypical, non-bile precipitating colonies were observed (Etiles, 1973). Whether these colonies should be included in the counts is questionable, because the identity of the colonies was not known.

objectives of Study

The objectives of this study were to determine the types of <u>Enterobacteriaceae</u> present in ground meats, and the influence of the method of enumeration on the types of <u>Enterobacteriaceae</u> included in the counts.

The study was also planned to evaluate the use of IMViC typing and the elevated temperature test for classification of colitorms in teneral, and faecal $\underline{\varepsilon}$. \underline{coli} , in particular.

LITERATURE REVIEW

The <u>Enterobacteriaceae</u> are facultative, anaerobic, Gram negative, nonspore forming rods that can ferment glue e usually with, but sometimes without, gas production. They are exidase necative, can reduce nitrate to nitrite, and, if motile, they are peritrichously flagellated (Bergey's Manual, 1974). They are commonly found as intestinal organisms, and as a result they are frequently referred to as the enteric bacteria. Within the <u>Enteropacteriaceae</u> are the "coliform" bacteria. These organisms are of considerable importance in water and food microbiology, so they are considered in this review as a separate entity.

I. The family ENTEROBACTERIACEAE

The <u>Enterobacteriaceae</u> are often referred to as the "Coli-Typhoid" group, but not all members of this family are intestinal parasites or human pathogens. Typical sources and habitats of the members of this family, as well as their pathogenicity are given in Bergey's Manual and are summarized in Table 1. <u>E. coli</u> is generally regarded as a non-pathogenic, commensal intestinal organism. However, some strains are enteropathogenic (E.E.C.) causing enteritis in inf. s and adults.

The most common intestinal pathogens among the <u>Entero-</u>, <u>bacteriaceae</u> are <u>Salmonella</u> and <u>Shiqella</u> spp., which cause salmonellosis and shigellosis, respectively. <u>Salmonella</u> are

widespread in Nature and can be found in foods and water. The ubiquitous nature of Salmonella has developed from their endless ecocycle (Taylor and McCoy, 1969). Shigella, on the other hand, are obligate parasites of humans and monkeys and are seldom associated with foods. Shigelloses generally result from drinking contaminated water or contact with infected persons.

In addition to these intestinal pathogens, other general and species of the Enterobacteriaceae have been implicated as pathogens, but they are not generally considered to be "food poisoning" organisms. For example, Proteus spp., Citrobaci T spp., Klebsiella spp. and Yersinia spp. are considered to be pathogenic. Edwarisiella tarda, although not usually considered a food poisoning microorganism, has been associated with outbreaks of diarrhoea (Bryan, 1969). The Erwinia spp. are plant pathogens and have not been implicated in animal diseases.

The organisms in Table 1 are listed according to the 8th edition of Bergey's Manual. There are marked differences in taxonomy and nomenclature of the Enterobacteriaceae between the 8th and the previous editions of Bergey's Manual. As a result, there could be difficulties in interpreting the literature on Enterobacteriaceae, for example £. coli used to describe lactose-fermenting, aerogenic strains; now non-lactose fermenting, anaerogenic strains such as the Alkalescens-Dispar group and

Table 1. The habitat, and sources of members of the family Enterobacteriaceae, based on Bergey's Manual (1974)

Qrqanisms	<u> Habitat and Sources</u>
<u>E. coli</u>	Intestine of warm blooded animals
<u>E. tarda</u>	Intestinal tract of snakes, Times faeces and water
Citrobacter spp.	Water, pod, taeces and urine
<u>Salmonella</u> spp.	Intestinal tract of warm-blooded animals and reptiles, and food
<u>Shigella</u> spp.	Intestine of man and higher monkeys
K. pneumoniae	Soil, water, grain and intestinal tract of man and animals
<u>K. Saenae</u>	Intections of respiratory tract
<u>loscleromatis</u>	Rhinoscleroma disease
<u>b</u>	Faeces of man and animals, sewage, soil and water
E. <u>aerogenes</u>	Faeces of man and animals, sewage, soil, water and dairy products
S. marcescens	Water, soil and food
P. vulgaris P. mirabilis	Paeces of animals, sewage, soil, especially with protein decompostion
P. morganii	Paeces
P. rettgeri	Chicken faeces, clinical specimens
P. inconstans	Urine and faeces
Y. pestis	Buboes, blood, sputum and lung exudate
Y. pseudotuberculosis	Pseudotuberculosis lesions, intestinal tract
Y. enterocolitica	Faeces, lympn nodes, milk and ice cream
<u>Erwinia</u> spp.	Plant pathogens, saprophytes or epiphytic flora

Paracolobactium op. have been included. The purpose of this section is to indicate the relationship between the Tribos, Genera and species of Enterobacteriaceae in the 8th edition of 1 gey's Manual and those in the 7th, 6th and 5th editions (Argey's Manual, 1939, 1948, 1957, 1974). The Tribe and Genus classifications used in these editions of the Manual are shown in Table 2.

Tribe I. <u>Escherichieae</u>. Consists of 5 genera in the 8th edition. The relationship between these genera in the 8th edition and those of the previous 3 editions are shown in Table 3.

The genus <u>Escherichia</u> in the 8th edition of Bergey's Manual has only one species: <u>E. coli</u>. In previous editions there were other species of this genus. Of these, only the pigmented <u>E. aurescens</u> has been incorporated into <u>E. coli</u>, the other species have been assigned to the genus <u>Citro-bacter</u>. However, it appears that some <u>Shigella</u> species, <u>S. alkalescens</u> and <u>S. dispar</u> of the Alkalescens-Dispar group, have now been incorporated with <u>E. coli</u>. In addition, slow and non-lactose fermenting <u>Paracolobactrum coliforme</u> of the 7th edition, are also included in <u>E. coli</u>. These changes have broadened the biochemical characteristics of <u>E. coli</u> to include anaerogenic and non-lactose fermenting strains.

The other <u>Escherichia</u> species, <u>E. freundii</u> and <u>E. intermedia (um)</u>, have been incorporated into the genus <u>Citro-bacter</u>, as <u>C. freundii</u> and <u>C. intermedius</u>, respectively.

Table 2. Tribes and genera of the family <u>Enterobacteriaceae</u> from the 5th to 8th editions of Bergey's Manual

<u>Genera</u>

Trabe	oth edition	6th edition	7th edition
Eschell_heae (lae)	Eschelichia Aerobacter Klebsiella	<u>Aerobacter Klebsiella</u>	Escherichia Aerobacter Klebsiella Paracolobactrum Alginobacter
Erwineae	<u>Erwinia</u>	<u>Erwinia</u>	<u>Erwinia</u>
Serrateao	<u>Serratia</u>	<u>Serratia</u>	<u>Serratia</u>
<u>Proteas</u>	Proteus	Proteus	<u>Proteus</u>
<u>Salmonelleae</u>	Salmonella Eberthella Shiqella	<u>Salmonella</u> <u>Shiqella</u>	<u>Salmonella</u> <u>Shigella</u>
		8th_edition	,
Escherichieae		Escherichia Edwardsiella Citrobacter Salmonella Snigella	
<u>Klebsiellede</u>		<u>Klebsiella</u> Enterobacter <u>Hafnia</u> Serratia	
<u>Proteae</u>		<u>Proteus</u>	·
<u>Yersinieae</u>		<u>Yelsinia</u>	
<u>Erwineae</u>			

Table 3.	01 801	gey'n Nanual w un elitions	: t h	Eschetichiede a	nd	
		th edition	6 t	h •ditton	٠, t	h waatten
Genus I: <u>Escherich</u>	<u>T</u>					
E- CO12	<u>E</u> .	 coli alkalescens aurescens coliforms 	<u> </u>	<u>coli</u> alkalescens	E •	<u>coli</u> alkalescens
Genus II: E <u>dwardsie</u>	<u>11a</u>					
<u>E. tarda</u>	- -		**		-	
Genus III <u>Citrobact</u>						
C. freund C. interm	<u>ii </u>	<u>treundiı</u> <u>intermedia</u>	<u>E</u> •	<u>freundii</u> intermedium	<u>E</u> -	<u>freundii</u>
Genus IV: Salmonell	<u>a</u>					
No signif	icant ch	nanges between	ed:	itions		
Genus V: <u>Shiqella</u>						
<u>s. sonnei</u>	<u>s</u> .	sonnei	<u>s</u> .	<u>sonnei</u> cey <u>lonensis</u>	<u>s</u> .	<u>sonnei</u> <u>ceylonensis</u>
S. flexne.	<u>ti S</u> .	<u>flexneri</u> <u>boydii</u> <u>dysenteriae</u>	<u>s</u> .	<u>paradysenteriae</u>	<u>s</u> .	paradysenteriae
S. dysent	5.	dysenteriae schmitzii arabinotarda	<u>s</u> .	<u>lysenteriae</u> <u>ambigua</u>	<u>s</u> .	<u>dysenteriae</u> <u>ambiqua</u>
				·		

⁻ No cross reference of this nomenclature to organisms in the 7th, 6th or 5th editions.

Other genera added to the Strie Epcherichiese in the 9th clitical include: Edward in the Strie Epcherichiese in the 9th ends Edward in the only one openies, no targat. The association of Edward in Edward with organisms in previous editions is not clear. Esimonely and shipplic were previously in the Strie Salmonellese, but this tribe has been eliminates from the classification in the 6th edition.

The main employers in classifying both <u>Salmonella</u> and <u>Shigella</u> in the 8th eartich is their antiquate structure. However, the brochemical characteristics have not been completely eliminated at the species level. Most of the species that are still described according to their blochemical reactions in the 8th edition have been carried forward from previous editions. The genus <u>Eberthella</u> ment oned in the 6th edition has been included with <u>Salmonella</u> in the 7th and 8th editions.

Tribe II. <u>Klebsielleav</u>. This tribe is new to the classification, and consists of 4 chera. The relationship between these genera and those in revious editions is shown in Table 4. The tribe <u>Klebsielleav</u> consists of some organisms from the <u>Escherichieav</u> in previous editions, as well as from the tribe <u>Serrateav</u>.

The genus <u>Klebsiella</u> has 3 species: <u>K. pneumoniae</u>, <u>K. ozaenae</u> and <u>K. rhinoscleromatis</u>. In the 6th edition, the latter two species were classified as strains of <u>K. pneumoniae</u>, nowever in the other editions these organisms

re rainy Cartae	to Manual with 85	or <u>hen ny hanga</u> er anolo <i>n</i>	n the "th eqition of grightege in previou
			Sth • 11tion
	~ <u> </u>		
GENUD 1: Klubbielka			
K. Susamonias	K. EnermoniaA. derogenus(non-motile)E. derogen∋ides	A. a <u>erogenes</u>	 F. Pheumoniae A. aprogene: (non-motile) E. granulomati E. paralytica K. capsulata
K. okaenae k. thinosclero- matis	h. <u>Ozaenae</u> E. <u>rhinosoler</u> e- <u>matis</u>	K. Fuenmoniae	K. ozaenae K. rhinosclero- matis
GENUS II: Enterolacter			
F. derodenes	A. <u>gloagae</u> A. <u>aerogenes</u> (motile)	A. <u>cloacae</u> A. <u>aerogenes</u> (motile)	<pre>A. cloacae A. aerogenes (motile)</pre>
GENUS 111: <u>Hafnia</u>			
<u>H. alvel</u>	-	-	-
GENUS IV: <u>Seirat</u>	<u>‡</u> <u>d</u>		
<u>S. marçeuvens</u>	 marçescens plymuthica kiliensis piscatorum indica 	S. marcescens S. plymuthicum S. kiliensis S. piscatorum S. indica	S. marcescens S. plymouthensis S. kilensis S. piscatora S. indica S. anolium

⁻ No cross reference of this nomenclature to organisms in the 7th, 6th or 5th esitions.

Pere separate species. K. poguagorae has been expanded to include the act motive strains of Agropacter actor actor (7th, eth and oth edition), Paracolopacitium acrogenorades (7th edition).

the genus <u>Enteropagies</u> was primarily letived from the revious <u>Acrologics</u>, and constats of two species: i. closed: and <u>E. acrogenes</u>. S. <u>acrogenes</u> constate of entry the motale strains of the previous <u>A. acrogenes</u>. <u>Enteropagies</u> agglomerans, escribed by Eving and Fife (1977) is placed in the Herbrecka group as <u>Ervinia herbreola</u> in the current edition of Bergey's Manual, and is fiscussed later.

The genus <u>Haipia</u> consists of one species: <u>H. alvel</u>.

Furthermore, there are some references to <u>E. alvel</u> (Pergey's Manual, 1474) and <u>E. naipiae</u> (Roche Encise III) in the literature, but these names are not user within the family <u>Enterohacteriaceae</u> is any of the editions of Bergey's Manual. The association between this organism in the 8th edition and those in the other editions is not specifie:

The genus <u>Serratia</u> also consists of only one species:

<u>S. marcescens</u>, and includes all of the <u>Serratia</u> spp. in previous editions. Some species are recommended for addition to this genus, and will be discussed later.

The Encise II (<u>Enterobacteriaceae</u> Numerical Coding and Identification System for Enterotube) was deviloped by Foche Diagnostic, Professional Services Development, 340 Kingsland Street, Nutley, N.J., N.S.A.

Tribedial Protector for tribe consists of one denus, with 6 species. There has been little change in this group between elitions, as shown in Table 5.

which was derived from Pasteurella in previous entrone and now consists of a species, as shown in Table 5.

True V. signed. Thus has two one denum Figure. Some electer have been dides of eleminated between eartions of the Manual (. e Table e).

Former organisms firsted in the previous elitions of beingey's Manual were not included in Tables 3-e, because there is insufficient information to show how they were related to organisms in the n w taxonomic scheme of the 3th edition. These organisms include Alginobacter sp. (7th oi.): several Erwinia spp. (eth ed.); Proteus hydrophilus, Proteus ichthyosmus, Erwinia flavida and Erwinia erivanesis (5th ed.). According to the 3th elition, Erwinia mimipressualis and Erwinia dissolvans should be claced under the genera Enterobacter and Klebsiella, respectively, however the exact nomenclature or these organisms under these genera is not indicated.

In 1972, Ewing and Fife described an organism named $\underline{\underline{\varepsilon}}$. $\underline{\underline{aqqlomerans}}$. This organism includes the Herbicola-Lathyri pacteria which have been retained in the genus Erwinia in

Eth edition	7th etitien	oth edition	Sth odrtion
anier Proteen			
GENUS I: Protous			
P- vulgatis	P. vulgarits P. mitabilis	P. vulgario P. mirabilio	P. americanus P. pseudovalerie
E. morganii P. rettgeri P. inconstans	P. morganii P. Lettyeii P. inconstans	P. morganii P. rettgeri	P. ammoniae P. mo inii G. rettyeri
TRIPE: Yeleinieae			
GENUS I: Yersinia			
Y. pestis Y. pseudo-	- -	- -	- -
<u>tuberculosis</u> <u>Y. enterocolitica</u>	-	-	-

Table (r. 8) staon hap of the tribe liwiness is the office critical or the office the following with liwipess of the previous critical

eth entien	7th whithon	eth (diffon	oth edition
<u> </u>			
- →ENUS I: <u>Er</u> yiti <u>a</u>			

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amylorova ballorb tarcherraria higrifluens	amyiciov <u>balici</u> tarcheipailla	amylorova salicis tarche iphila	amylorova salicis tarches; sila
<u>quercina</u>		-	-
<u>rubrifaciens</u>	·-		_
<u>merbicola</u>	<u>vitivold</u> milletiae	<u>vitivola</u> milletiae	milletia-
	<u>cassayae</u>	<u>Cassavae</u>	<u>cassavas</u>
	<u>ananas</u>	<u>ananas</u>	<u>ananas</u>
	<u>citrimaculans</u>	<u>citrimaculans</u>	
	mgudfisfaa	<u>mangiferae</u> <u>lathyri</u>	<u>manganitelae</u> <u>lathyri</u>
stewartii	~	- Identer	<u> </u>
uredovora	-	-	_
<u>carotovora</u>	<u>carotovora</u>	<u>carotovora</u>	<u>carotovara</u>
	<u>carnegleana</u>	<u>caruedreaua</u>	<u>solanisarri</u>
	atrospetica aroideae arsideae	atrosertica cytolytica	<u>phytophthal</u> aroideae
<u>chrysanthemi</u>	<u>chrysanthemi</u>	-	-
<u>cypritedil</u>	-	-	-
rhapontici ·	rnarontici	<u>rhapontici</u>	-
		(

⁻ A proper reference of this emenciature to ordanisms in the 7th, 6th or 5th editions.

the Streintion of bendey's Manual. i. adjlomejans was shown not be seen astrict plant pathoden, in that it has been assoluted from clinical openimens of humans and lower animals. This etganism forms chromopenic (yellow) colonies and one into it acrospenic and anaeropenic strains. The authors places this organism under the genus <u>EnteroLacter</u> instead of in a new genus because of its similarity to other <u>EnteroLacter</u> organism in this text, because it is more readily associated with this name in identification, rocedures.

<u>En_obacteriaced</u>. Your: et al. (1971) ecause of its unique blochemical and serological proper was. Two species are proposed, which have been isolated from numan clinical and faecal samples. Although these organisms appear to be

closely related to <u>Citrobacter</u> and <u>Enterobacter</u>, they can be distinguished by prochem: if differences.

II. The Colitorm Bacteria

The term "collitorms" is widely used by bacteriologists, sometimes without strict regard for the definition.

Furthermore, there are many definitions of colliforms, depending on the method and purpose of their detection and enumeration. According to the Standard Methods for Water and Sewage analysis (APHA, 1971), colliforms are defined (using liquid media) as "the aerobic and facultative anaerobic, Gram negative, nonsporeforming rod-shaped bacteria which terment lactose with gas production within 48h at 35°C." The colliform group, thus derined, is equivalent to previously used descriptions, such as: B. coli, coll-aerogenes and the colon group (APHA, 1965).

An alternate definition used by water bacteriologists is hased on the membrane filter technique, in which coliforms are defined as " e organisms that produce dark colonies with metallic en within 24h on an Endo-type medium containing lactose" (APHA, 1971). In dairy bacteriology, when coliform counts are high, they are usually enumerated on Violet Red Bile agar (VRBA) or Desoxycholate agar (APHA, 1967). On VRBA, the coliforms grow as typical dark red colonies, surrounded by a zone of bile precipitation, within 24h incubation at 32°C (APHA, 1967).

Although there are many definitions for the colliforms, they are all based on the fact that these organisms ferment lactose and can grow in the presence of bile salts at 32-37°C. Since there have been extensive taxonomic changes in the family <u>Enterobacteriaceae</u>, the names, of the organisms included within the colliforms have varied.

A. Classification and Characterization of Coliform Bacteria

Lactose was originally selected as the basis of differentiating between the commensal intestinal organisms and those with pathogenic implications (Winslow et al., 1919). The lactose fermenting bacteria in this group were turther divided into those that were typical intestinal parasites (B. coli) and those that were most frequently isolated from grass, grains and soil (A. terogenes) (MacConkey, 1905; Rogers, 1918). The classification and characterization of the coliforms was greatly influenced by the practical considerations of water bacteriologists, who wanted to identify intestinal coliforms as a sanitary index (Parr, 1939; Rogers et al., 1918; Winslow et al., 1919).

There were many attempts to relate types of coliforms and their biochemical characteristics to the ecological source of the organisms (Bardsley, 1934; kogers et al., 1914, Rogers et al., 1918; Winslow et al., 1919).

Carbohydrate fermentation was first used by MacConkey (1905) as a means of differentiating lactose fermenters from

various ecological sources. He used sucrose and dulcitol to divide the lactose fermenters into 4 groups. These groups were further subdivided using gelatin liquefaction and the Voges-Proskauer reaction as the basis of differentiation. Other studies failed to give a system for classifying the source of coliforms using carbohydrates alone and Parr (1939), established a series of tests for classifying coliforms:

Indole production
Methyl Red test
Voges-Proskauer reaction
Citrate utilization
Uric acid utilization
Gelatin liquefaction
Eijkman elevated temperature test
H₂S production
Sucrose and inositol fermentation
Alpha-methyl-d-glucoside fermentation.

It had been reported (Bardsley, 1934; Levine, 1918), that the lactose fermenting bacilli could be divided into two ecological groups based on their Methyl Red (MR) and Voges-Proskauer (VP) reactions. The coli group (E. coli) of intestinal origin were MR+, VP-, while the aerogenes-cloacae group from the soil were MR-, VP+. Using these two criteria alone was found to be inadequate because other combinations of MR-VP reactions were observed. As a result Koser's citrate reaction and the indole test were added to the classification.

Parr noted that Indole, Methyl Red, Voges-Proskauer and the Citrate tests (IMViC tests) were those most frequently

used to differentiate the "faecal" and "non-faecal" colltorms (Parr, 1939). E. coll (IMViC ++--) was shown to be the
most typical, numerous and constant colliform organism found
in faeces, and therefore most useful as a sanitary index for
water (Bardsley, 1934).

The ability of <u>B. coli</u> to produce gas at elevated temperatures (46°C) was established by Eijkman (1904). In 1949, the Coliform Sub-Committee of the Society of Applied Bacteriologists (Report, 1949) proposed the classification of the lactose fermenting coli-aerogenes bacteria hased on the IMViC tests and ability to produce gas from lactose in MacConkey broth at 44°C. This classification was revised by the Coliform Sub-Committee in 1956, using gelatin liquefaction as an additional characteristic, as follows:

Coli-aerogenes Organism	I M ViC	44°C	Gelatin
(1956 Classification)			•
<u>E. coli</u> I	+ +	+	-
E. coli III	+ +	-	<u>-</u> .
E. coli II	- +	-	-
<u>C. freundii</u> I	- + - +	<u>-</u>	-
<u>C. treundți</u> II	+ + - +	-	-
K. <u>aerogenes</u> I	+ +	-	-
K. <u>cloaçae</u> 1	+ ; +	-	+
E. carotovora	+ +		•
K. aerogenes II	+ - + +	_	-

Subsequently, the American Public Health Association (12th edition, 1965) classified coliform bacteria into three groups, based solely on the IMViC tests.

Organism		M	۷.	i C
<u>E. coli</u> I	+	+	-	-
<u>E. coli</u> II		+	-	-
<u>E. freundii</u> I	-	+	_	±
E. freundii II.	+	+	-	-
A. <u>aerogenes</u> I	-	-	+	±
A. aerogenes II	±	-	÷	+

B. Taxonomy and Nomenclature of Coliform Bacteria

The taxonomy and nomenclature of the <u>Enterobacteriaceae</u> has been greatly influenced by changes in classification of the coliforms, and vice versa, as illustrated by Cowan (1956) when he reviewed the report of the Coli-aerogenes Sub-Committee. Besides the coliforms, the non-lactose fermenters which are of clinical significance have also undergone changes in classification. Both have affected the taxonomy of the Family <u>Enterobacteriaceae</u> as a whole.

C. Atypical Coliforms

Typical coliforms can produce acid and gas from lactose broth within 24h. Atypical coliforms are frequently encountered (Parr, 1939). Some are slow lactose fermenters, that

is, they do not ferment lactore for a considerable number of days. Some atypical forms ferment lactore but fail to produce gas (anaerogenic). Some strains give all of the reactions of <u>Escherichia</u> except the fermentation of lactore, and have been referred to as the "paracoli" (Parr, 1939).

Some variants ferment lactore at room temperature but not at 37°c.

Sometimes, co iforms become atypical when grown in adverse environments. For example, sodium benzoate in weak glucose broth retarded the fermentative activities of <u>F</u>. <u>coli</u>, whereas other characteristics were only slightly affected. No gas producers could be detected if foods were preserved with sodium benzoate (Smirnow, 1910). Sodium acetate also inhibits coliforms from producing gas from sugars, though they can ferment alcohol derivatives of the same sugars.

Some variants are unstable, with a variable colony type as well as biochemical activity (Jones et al., 1966; Parr, 1939). There have been many reports of atypical colonies on VRBA (Hartman, 1960; Jones et al., 1966; Kereluk and Gunderson, 1959; Morris and Cerny, 1954; Ross and Thatcher, 1958). In some cases the atypical colonies have been identified as E. coli (Hartman, 1960; Jones et al., 1966) and should be included in the VRBA "coliform" count. In other cases, the atypical, non-bile precipitating colonies have been shown to be other organisms such as "Proteus,

Aerobacter, Pseudomonas, paracolon organisms, and Flavobacterium" (Ross and Thatcher, 1958). Although some of these organisms might be included in the "coliform" count, the pseudomonads and flavobacteria would give spur;ous results.

D. Significance of Coliform Bacteria in Water and Foods

In water bacteriology, <u>E. coli</u> I was selected as an indicator among the coliform bacteria, because of the specificity of <u>E. coli</u> i as an indicator of faecal contamination. In food bacteriology, the presence of coliforms indicates either unhygienic conditions by the presence of <u>E. coli</u> I or unsanitary conditions by the presence of other coliform bacteria. According to Buttiaux and Mossel (1961), a suitable indicator of faecal contamication should have the tollowing characteristics:

- 1. The bacteria selected should occur only in the intestinal environment.
- 2. They should occur in very high numbers in faeces.
- 3. They should have a high resistance to the extraenteral environment.
- 4. They should permit easy and reliable detection even when present in very low numbers.

Besides E. coli I, Buttiaux and Mossel (1961) considered that the <u>Klebsiella</u> group (non-motile, capsulated, VP+ and urease+) are always of faecal origin,

that the <u>Citiological</u> group are not only found in faeces but also in [cil, and that the <u>Enterobacter</u> proups are rarely found in the human intestine. In roods that have been dehydrated, prozen or retrigerated, Buttiaux and Mossel (1901) suggested that <u>E. goli</u> may not survive as well as some enteric pathogens that might also be present. As a result, <u>E. goli</u> might not be the best indicator organism for unnygienic handling of foods. They suggested <u>Klebsiella</u> as an alternate indicator to <u>E. goli</u> and stressed that non-lactose fermenting <u>E. goli</u> (<u>Paracolobactrum</u> coliforme).

Where routine tests are being conducted on foods for non-lactose fermenting enteric pathogens e.g. <u>Salmonella</u> spp., indicator organisms are of little concern. However, when safety is based on indicator organisms, Buttiaux and Mossel (1961) considered that the <u>Enterobacteriaceae</u> as a whole should be used, in place of the coliform bacteria.

III. Development of Media

The development of solid and liquid media for the enumeration or estimation of Enterobacteriaceae occurred simultaneously, MacConkey (1908), summarized the development of bile salts as a selective agent for "colon bacilli" (Bacillus coli communis). The medium was intended to be versatile by changing the carbohydrates.

In the case of differentiating B. typhosus and B. coli, lactose was the carbohydrate selected because these organisms grew as bile and non-bile precipitating colonies (MacConkey, 1908). In water and dairy product analysis, lactose also became the carbohydrate of choice, because emphasis was placed on detection of E. coli and lactose fermenters (Coliform bacteria) (Hall and Ellefson, 1918a,b).

Winslow and Dolloff (1922) studied the Θ t of bile added to Brilliant Green Lactose and Gentian Violet Lactose proths and they recommended the use of bile in the media to reduce the inhibitory effect of the dyes on the coliforms. The use of dyes was introduced by Churchman (1913) and Hall and Ellefson (191da, b), to eliminate the false results in the presumptive lactose broth tests, due to sporogenous anaerobic and aerobic, Gram positive bacteria growing in the broth (Hall and Ellefson, 1918a,b). However, it was shown that selective media, such as Brilliant Green Bile broth inhibited some coliform organisms (Mallmann and Darby, 1941; Winslow and Polloff, 1922). The Standard Method Committee of the American Public Health Association suggested the use of Lactose broth as a presumptive test and Brilliant Green Bile broth as a confirmed test for coliform organisms (Mallmann and Darby, 1941).

Mallmann and Darby (1941) observed that tryptose added to Lactose broth caused many "slow lactose fermenters" to produce gas in greater quantities in a shorter period of

time. Adding lawryl sulfate also gave selection of collforms. This resulted in the development of Lauryl Sulfate
Tryptose froth (Mailmann and Darby, 1941), which was shown
to be more reliable for isolation of colliforms to a Lactose
broth (Hajna and Perry, 1943; Mailmann and Darby, 1941;
Perry and Hajna, 1944), and to be a good presumptive medium
for use with foods (Hall, 1964).

other selective broths were also developed for the isolation of coliforms. These included: Formate bicinpleate broth (Stark and England, 1935), buttered behoxycholate Glucose broth (Hajna and Damon, 1955), Ec medium (Eijkmin, 1904) and its modifications (Hajna, 1937; Perry and Hajna, 1933,1944) and Boric Acid Lactose broth (Clark et al., 1957; Njoku-Obi and Skinner, 1957; Vaughn et al., 1951). The latter two broths, Ec medium and Boric Acid Lactose, became involved in the development of elevated temperature tests for the detection of faecal E. coli, which is discussed below (see p. 27).

Salle (1930), summarize: the development of selective media for the confirmatory test of $\underline{E} \cdot \underline{col} \cdot \underline{l}$ as follows:

"Very few of the media now extensively in use were compounded solely for water examinations. This accounts for the large number of modifications of Endo and other differential media. This is a point that should be kept in mind.

Drigalski and Conradi (1902) used crystal violet to eliminate other organisms interfering with the isolation of <u>B. typhosus</u> from stools. Litaus was added to the medium to identify the presence of the acid producing colon organisms. The well known medium of Endo (1903) is composed of basic fuchsin decolorized by the use of sodium sulfite. Modifications of the original formula have been reported by Kendall

and Walker (1910), Kendall and Day (1911), Kinyoun and Deiter (1912), Harding and Ostencerg (1912), Robinson and Rottger (1916), Levine (1918), and other apting the medium to special investigation. Holt: and Teague (1916), incorporated the dyess methylen: and eosin in again hase containing lactose and sucrose were able to isolate the typhoid and dysentery organisms on this preparation. This medium has been modified and simplified by Levine (1918 and 1921), adapting its use to water works laboratories."

the isolation of coliform bacteria. They recommended Violet hed Bile age, as well as Neutral field file agar and Brilliant Green Bile broth as giving reliable confirmatory tests. Violet hed Bile agar was shown to be satisfactory for direct plating and enumeration of <u>E. coli</u> in dairy products (Miller and Prickett, 1939). Leitson (1935) developed selective media containing sodium desoxycholate as the inhibitory agent. Selectivity was modified by changing the energy source, for example, lactose was used in the medium to isolate coliform bacteria, whereas citrate was used to isolate non-lactose fermenting, intestinal pathogens.

Eijkman (1904) developed the elevated temperature technique to differentiate <u>E. coli</u> from other coliform bacteria. Since then, there have been many modifications of both the medium and incubation temperature. Perry and Hajna (1933) added phosphate buffer, reduced the amount of glucose to 0.3%, and used a pH of 5.6 as opposed to 4.5. Other carbohydrates were also suggested to replace glucose, such as lactose and mannitol (Hajna 1937). In a comparison of the modified Eijkman lactose broth (EC medium) with

MacConkey broth, the EC medium was shown to see superior (Majna and Perry, 1939). Another medium used for the elevated temperature test was Boric Acri Lactose broth (Clark et al., 1957) pokurobi and Skinner, 1957; Vaughn et al., 1955; Njokurobi and Skinner, 1957; Walford, 1954). However, Boric Acid Lactose broth is considered less desirable than EC med am (Mail, 1964; Skinner and Njokurobi, 1957.

compensature requirements are critical for the differentiation of E. coli (Hall, 1964). Eijkman (1904) originally used 40°C. This temperature was found to be too high (Hajna and Perry, 1939; Levine et al., 1934; Perry and Hajna, 1944; Skinner and Brown, 1934), inhibiting the growth of some strains of E. coli. Other workers suggested 43 and 44°C as the incubation temperature (Hajna and Perry, 193°; Levine \underline{e}_{\pm} \underline{al}_{\pm} , 1934; Waltord, 1954). Most strains of Aerobacter aerogenes (E. derogenes) produce gas from glucose, lactose and mannitol at 42° C, many at 44° C, but few at 46°C. The same temperature selection applies to $\underline{\mathtt{A}}$. cloacae (E. cloacae) and Citrobacter spp., however they lo not produce gas from glucose or lactose at 46°C (Hajna and Perry, 1939). Hajna and Perry (1939) recommended the reduction of the incubation temperature to 45.5°C, but indicate; that this could be lowered to 44°C for some foods (e.g. Shellfish). As a result, a range of elevated incubCommittee on M solutions has come into asse. The International Committee on M solutional appearance into the French (That her and clark, 1908) cited two conditions into (1), the use of aC broth at 45.50 ± 0.70, and (11) the edominantly ameropean method, after Mackennia et al. (1948), using brilliant Green Lactore file. A broth at 460 ± 0.10%, other elevated temperatures have been proposed such as EC medium at 44.50 ± 0.00 for oyster: (Kelly, 1969), and trozen mods (Pishbein and Surkiewicz, 1964). More recently, Morsol (1962) reported on the usefulness of two modifications of Eigkman's test for raceal contamination of toods, using the European method of Mackenzie et al. (1948) and Chapman's Lactore-Tergitol-Triphe epitetrazelium Chlorice-Bromothymol blue against 440°C, de reported that both methods were equally reliable.

Fishedin (1962) studed the effect of elevated temperature (44.5 to 46.5°C) on gas production by <u>Aerobacter</u> (<u>Enterobacter</u>) and <u>E. col</u>: strains. At 44.5°C, it was found that as many as 72% of <u>Aerobacter</u> cultures produced gas.

Above 45.1°C there was a rapid drop in gas forming cultures by <u>Aerobacter</u>. At 45.5°C, only 2% of <u>Aerobacter</u> strains were gas positive, whereas 42% of the <u>E. coll</u> strains while gas production up to 40°C. Above 46°C the incidence of gas production declined rapidly. The most resistant gas producers at elevated temperature were the <u>E. coll</u> type II (IMVIC -+--) rather than the typical <u>E. coll</u> type I (++--).

elevated temperature test. Subsequently, Fishbein and Surkiewicz (1904) studied the effect 44.5 and 45.5% on the reliability of detecting faecal <u>E. goli</u> in for roods and nutmeats. They reported that 88% of 6,472 LST (Laury) Sulfate Tryptose broth) tubes live identical response at both temperatures. For those showing a difference in result at the two incubation temperatures, results at *5.5% gave fewer talse positives (threefold effect) but a 4% loss in recovery of <u>E. goll</u> over the lower temperature. Depending on the intention of the test, 44.5 or 45.5% would be selected as the elevated incubation temperature.

effect on jas production by <u>E. coli</u>, but that gas production by <u>Merobacter</u> (<u>Enterobacter</u>) and <u>E. treundii</u> (<u>C. freundii</u>) was inhibited. However, they also showed that spore-forming pacteria capable or termenting lactose 42.5 to 43.5°C were not significantly inhibited by boric acid.

Clark et al. (1957) studied the specificity of the boric acid medium and observed that coliforms of IMViC type ++-- gave a reasonably good percentage of positive reactions with foric Acid Lactose broth at 43°C, but that a large number of positive reactions were also observed with IMViC types: --++, ++-+, and -+--. As a result, they did not recommend Boric Acid Lactose broth for rapid classification of the ++-- IMViC type.

The Health Protection Branch (HPB) of Canada Health and Welfare (Health Protection Branch 1974) recommends the use of a 5-tube most probable number technique, using Lauryl Tryptose (LST) broth as presumptive test, Brilliant Green bile 2% broth as confirmed test, and Levine EMB agar as completed test (Harrevijn et al. 1973). Proposed standards for ground meat specify limits of E. coli per gram. The HPB methods also recommend transfer of LST positive tubes to EC media incubated at 45° ± 0.2°C. To confirm faecal E. coli, the Laure test for E. coli type I (++--) must be carried out. This is a laborious procedure, compared to the direct plating techniques recommended for dairy products, requiring 5 days for initial results, and a further 3 days to carry

out the IMViC tests.

Several rapid methods for the enumeration of coliforms or faccal <u>E. coli</u> in foods have been suggested, but they have not been thoroughly investigated, and their application to ground meats is not known. Abshire and Guthrie (1973) described a fluorescent antibody technique for the detection of taccal pollution (<u>E. coli</u>). Anderson and Baird-Parker (1975) used a direct plate method for enumeration of <u>E. coli</u> biotype I in roods. The method is based on the ability of <u>E. coli</u> to produce indole when growing at 44°C, on a cellulose acetate membrane, placed on a bile medium in a petri dish.

In a study of the significance of coliform bacteria meats, therefore, it is important to consider not only the selective counts, but also the typical and atypical organisms that are included or excluded from the counts. In addition, Mossel and co-workers (1957; 1961; 1962) suggested the enumeration of <u>Enterobacteriaceae</u> instead of coliforms as an indicator of hygiene and sanitation. Enumeration of <u>Enterobacteriaceae</u> can be achieved by using glucose as the carbohydrate in a selective medium for enteric bacteria, or in the case of Violet Red Bile agar, by adding 1% glucose to the medium.

METHODOLOGY

I. Colitorm and ENTEROBACTERIACEAE Counts in Ground Meats

Three qualities of ground beef and frozen and thaved pork sausage samples were purchased at retail stores and analyzed for bacteriological quality. An 11g sample of ground meat was homogenized with 99ml of 0.1% peptone water in a Waring Blendor jar, for 2 minutes at high speed as described by Al-Delaimy and Stiles (1975). Coliforms and Enterobacteriaceae were enumerated using different solid and liquid media.

A. Enumeration using the Direct Plating Technique

- 1. <u>Violet Red Bile Agar</u> (<u>VRBA</u>). Appropriate dilutions of the blended samples in 0.1% peptone water blanks were plated using the pour plate technique, with 15ml VRBA per plate and a 3-4ml overlayer (Difco Manual). The plates were incubated at 37°C for 24h. Differential counts of typical, bile precipitating colonies (Difco Manual) and atypical, non-bile precipitating colonies were recorded.
- 2. Mossel's Modified VRBA (VRBG). Difco VRBA with 1% glucose added was prepared and used as specified for VRFA above (Mossel et al., 1962).

B. Most Probable Number Technique

- 1. <u>Presumptive Test for Coliforms</u>. Appropriate dilutions were pipetted in triplicate into Difco Lauryl Tryptose broth (LST). Each tube contained an inverted Durham tube for reading gas production. The tubes were incubated at 37°C and read for gas production at 24h. Tubes that were gas negative at 24h were re-incubated and read at 48±2h.
- used to transfer cultures from all gas positive LST tubes to Difco Brilliant Green Bile 2% broth (BGB), containing an inverted Durham tupe. Transfers were only made from LST tubes after 48h incubation. These BGB tubes were incubated at 37°C and examined for gas production at 24±2h. Gas negative tubes at 24h were re-incubated at 37°C and examined for gas production at 48±2h.
- 3. Completed Test for Coliforms. All BGB tubes were streaked onto Levine EMB agar (Difco) after 24h incubation. Gas negative BGB tubes were noted, by marking the EMB plates accordingly. Plates were incubated at 37°C and inspected after 24h for black colonies with a metallic sheen, or black centered (nucleated) colonies with transparent peripheries, or nucleated, mucoid colonies.
- 4. <u>Elevated Temperature Test</u>. LST positive tubes were inoculated into Difco EC broth, containing an inverted Durham tube and incubated at 45.5°C for 24h. Gas positive

tubes were noted at the end of each incubation period. This test was also carried out using direct inoculation of the sample dilutions into EC broth and similarly examined for gas production, as specified above.

The MPN was computed using the MPN index and 95% confidence limits for the combination of positive and negative tubes, using the 3-tube table reproduced from the American Public Health Association by Thatcher and Clark (1908).

II. <u>Determination of the Types of ENTEROEACTERIACEAE</u> <u>Enumerated by the Different Techniques</u>

A. Selection and Purification of Cultures

Cultures were selected from VRBA, VRBG and EMB plates. The subsurface colonies on VRBA and VRBG were selected and picked according to the amount of bile precipitation and other visual differences in colony morphology. Colonies with a large, dark zones of bile precipitation (B++), faint zones of bile precipitation (B+) and those with no visible bile precipitation (B-) were selected, wherever possible, from plates representing each sample. Colonies were generally picked from plates with low densities to reduce chances of contamination by invisible or adjacent colonies in the medium.

Representatives of all colony types were picked from

EMB plates. These included typical colonies with a metallic sneen (MS+), and black-centered, "nucleated" colonies (NU+) and atypical mucoid colonies (MU+). In addition, non-colitorm, pink to transparent, non-nucleated colonies (NU-) were selected.

MacConkey agar (MA) to obtain isolated, surface colonies. A single, isolated colony on MA was selected and streaked onto a Nutrient agar (NA) plate to obtain a pure culture. The pure culture on NA was saved for biochemical tests. At the time that biochemical tests were carried out, purified cultures were transferred to Tryptic Soy broth (TSB) and frozen at -40°C for possible use in supplementary identification tests.

B. Identification of Cultures

1. Identification Tests

The methods selected to determine the identity of the selected cultures were based on the results presented in Method Development (p. 39).

- (i) Screening tests: Gram stain for Gram negative rods; Oxidase test (Steel, 1961) for oxidase negative organisms; and acid from glucose.
- (ii) Biochemical tests: Using the Minitek system, the following discs were included: arabinose, dulcitol,

mositol, raffinose and thammose for acid production; expositor segalactosidase production; lysine and ornithine decarboxylation; phenylalanine deamination; H₂S and indole production; citrate and malonate as sole source of carbon.

other tests using conventional tube techniques, included: gas from glucose, acid and gas from lactose using Phenol Red broth base (Difco) with 1% carbohydrate added (glucose broth was sterilized at 121°C for 15 min; a 10% lactose solution was filter ster 'ized using a 0.2µ millipore filter, and added aseptically to the sterile Phenol Red broth base); MR-VP (Difco) broth; and urea broth (Edwards and Ewing, 1972).

- (iii) Supplementary and additional tests: LST (Lauryl Tryptose broth, Difco) for gas production; EC medium (Difco) incubated at 44.5 and 45.5°C for gas production; TSI (Triple Sugar Ir agar, Dirco); Motility test using motility test medium (Difco). Supplemental tests necessary for further identification of the organisms were carried out using the conventional tube technique. The tests used were those recommended by "Roche Encise II".
- (iv) Morphological studies: The selected cultures were re-plated onto VRBA, Mossel (VRBG) and IMB agars to confirm their morphology, and to check that the selected cultures had been tested, as opposed to adventitious "contaminants". In addition, the comparative morphology of the cultures on the selective media was determined.

2. Bacteriology

The Minitek plate: re-inoculated and incubated at 37°C as directed for the Minitek method. All carbohydrates and lysine, ornithine, and H_DS/indole discs were o erlayed with 0.1ml sterile paraftin oil prior to inculation. Results were read after 18 to 24h incubation.

The other tests, except the Mk-VP and E3 broths were also incubated at 37°C. Mk-VP was incubated at 30°C for 48h for the Voges-Proskauer reaction and for 5 days for the Methyl ked to st. EC proths were incubated at 44.5 and 45.5°C t 0.05°C in circulating, constant temperature water baths for 24 and 48h. Glucose, lactose, LST and TSI were incubated and read at 24 and 48h. TSI was only checked for delayed H₂S production at 48h. Motility test agar was read at 24h. Urea Production was read at 24h, 48h, 4 days and 7 days.

3. Tests used to identify Cultures

Cultures were identified using the Roche Encise II characteristics: acid and gas from glucose, Lysine ornithine, H₂S, indole, lactose, dulcitol, phenylalanine, urease, and citrate. The supplementary tests recommended by Roche Encise II were carried out, if necessary.

METHOD_DEVELOPMENT

Lacteriagege. Apart from the conventional methods of Cowan and Steel (1968) and Edwards and Ewing (1972) there are commercially prepared identification kits, e.g.

Auxotab (Wilson Diagnostics, Inc., & Science Rd., Glenwood, III., U.S.A.);

Enterotuse (Roche Diagnostics, Division of Hoffmann-La Roche Inc., Nutley, New Jersey, U.S.A.);

the MyB system (Diagnostic Besearch Inc., Roslyn, N.Y., U.S.A.); and
the Minitek system (RBL, Division of Becton, Dickinson and Co., Mississauga, ont., Canada).

The userulness and accuracy of these commercial systems has been extensively studied, by comparison to the conventional methods (Blazevic et al., 1973; Hansen et al., 1974; Leers and Arthurs, 1973; Nord et al., 1974; Rhoden et al., 1973a,b; Robertson and MacLowry, 1974; Rosner, 1973; Tomfohrde et al., 1973).

Both the R/B and Enterotube systems are priented to medical bacteriology for identification of human enteric isolates. They are based on a limited range of biochemical criteria, and lack versatility for use with the broader spectrum of <u>Enterobacteriaceae</u> that might be present in foods. The Minitek system, on the other hand, allows greater versatility in selection of biochemical criteria to be

tested and it is cheaper.

.. Comparison of Identification Systems

An initial study of 17 cultures (10 known and 7 unknown) was carried out to compare the performance of the conventional, Enterotupe, R/B and Minitek systems. The 10 known cultures were coded and treated as "unknowns" to eliminate researcher bias in the identification of the cultures. The criteria used in the Minitek system were lased on the system used by Hansen et al. (1974). The objective of this preliminary study was to determine the convenience, ease of interpretation and the accuracy of the results using each system.

The results of the R/B system were difficult to interpret, and with the results obtained, the cultures could not be identified with any degree of reliability. The identification of the cultures by the other systems is shown in Appendix A.

The known cultures could be identified more reliably than the unknown cultures. In general, positive identification was obtained for known cultures, but not for the unknown cultures, even with the complete conventional method. In some cases, the different lystems resulted different identities being assigned to the same cultures. This resulted from discrepancies in the results of the biochemical is between systems. The principal

discrepancies with the Enterotupe technique resulted from false positive indole tests, false negative citrate and unease tests. In the Minitek system, the principal discrepancies occurred in H₃S production, gas from glucose, unreliable urease, and false negative citrate tests.

especially with limited media preparation facilities.

Furthermore, the conventional technique did not result in the definite identification of all cultures. Enterotube and Minitex were both convenient, but some difficulty was experienced in interpreting the tests which lead in part, to the disc spancies between the systems. Supplementary tests were required for both the Enterotube and Minitek systems to complete the identification. It was not known, whether these discrepancies were due to deficiencies in the system, or incorrect interpretation of the results. Furthermore, the "known" cultures used in this study were not "type cultures, so the need for supplementary tests might be due to the "known" cultures peing atypical.

II. Comparison of Enterotube and Modified Minitek System to Identify Known Cultures

A second preliminary test was carried out using ATCC type cultures, referred to as designated strains in the 8th

American Type Culture Collection, 12301, Parklawn Drive, Rockvil Caryland 20852, U.S.A.

edition of Sergey's Me. II (1974). The objectives of this test were to determine the accuracy of interpreting the interotube tests; the influence of incubation temperature on the Minitek results (for those cultures with recommended growth temperature of 30°C); and the ability to reduce the number of supplementary tests ising the Minitek system.

bystem were based on the flow diagrams in Figures 1-4. The minitek discs included: arabinose, citrate, H₂S/indole, inositol, lysine, maionate, ONPG, ornithine, phenylalanine, raffinose, rhamnose and urea. It was intended that the modified Minitex system should include all of the Enterotube tests, except dulcitol. In addition to the Minitek discs, glucose and lactose fermentation in phenol rea broth (Difco Manual), H₂S on Triple Sugar tion (TSI) agar (Edwards and Ewing, 1972; Difco Manual), urease activity in 3 separate media (Christensen, 1940; Vuye and Pijck, 1973), and motility in Motility Test medium (Difco), were tested.

The identity of the cultures is shown in Appendix B.

Considerable discrepancy occurred in the reading of the

Enterotube by the two laboratories, and between the Minitek

as well as Enterotube systems. The temperation of the

Minitek results was based on a special developed computer

program. The combination of Minitek technique and the

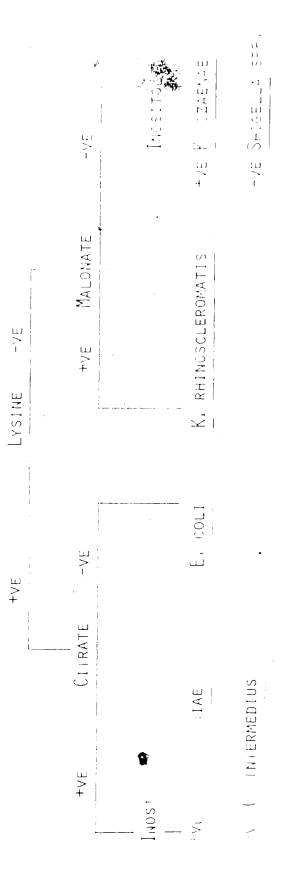
computer program gave an accurate identification of the

cultures without additional supplementary tests, especially

UREA I tve P. RE ISER! ZE P. TORGANII TOSITO +<E -VE P. MIRABILIS +VE P. VULGARIS ±∨E INDOLE

1 111				
ARABINOSE		 CITRATE	+VE C, FREURDII	-VE Y, PESTIS
		đ	+VE C	
	38182			
	+ \ E	 	SALMONELLA SPP.	

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∃>.	INOSITOL	+VE -VE	* +VE -VE	*VE K, RHINOSCLEROMATIS K, OZAENAE	-ve E, cloacae
+VE LYSI		- V E	>-	S, MARCESCENS "VE	S ∧ -
	ARABINOSE		- VE	K, OZAENAE	,
		TALONATE	+VE MOTILITY	+VE E, AEROGENES	-VE K, PNEUMONIAE

CITRATE, UREA/E	GAS FROM GLUCOSE	SHIGELLA SPE	C
+ VE		H, ALVEI	

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required supplementary tests to identify the cultures. Occasionally, the Enterotube resulted in incorrect identification for the known ATCC cultures. It was therefore decided to proceed with the modified Minitek and the computer program for identification of the unknown organisms. In the previous preliminary test, difficulty was reported in identifying the unknown cultures. The computer program will therefore be tested on the first series of unknown cultures (approximately 230) to test its ability to give a definite identification.

For the organisms grown and tested at 30 and 37°C, there were some minor differences in the test result, but not significant enough to affect the final identification.

As a result 7°C was considered an appropriate incubation temperature of 30°C.

III. Evaluation of Computer Program to Identify Unknown Cultures

The computer program developed to identify the unknown cultures consisted of a matrix of biochemical characteristics of known cultures (based on 8th edition of Bergey's Manual and the results for the ATCC type cultures). The program matched the unknown cultures with the organisms in the matrix, according to their biochemical characteristics.

The program determined the b highest matching organisms and the percentage match. No allowance was made for weighting of the importance of the different tests in the analysis.

The first 230 organisms tested by the modified Minitek technique were subjected to computer analysis. Although some cultures could be identify I with a high degree of confidence, many of the cultures could not be given a positive identification, because more than one organism had the same percentage match. Furthermore, over 20% of the unknown cultures were identified as most probably being Shigella spp. However, Shigella spp. are not expected to be present in food (Bergey's Manual, 1974), and many of these suspected Shigella were motile. The difficulty with this program appeared to be that the various identifying characteristics were neither weighted nor exclusive.

The same known and unknown cultures were checked against the Roche Encise II identification scoring system. Because dulcitol had not been—cluded in the modified Minitek system at this time, all dulcitol results were scored as negative, since this was the most likely test result (Bergey's Manual, 1974). The Roche Encise II system gave accurate identification of the ATCC type cultures, and reduced the number of doubtful or unidentifiable cultures for the unknowns. For example, most of the suspected Shiqella spp. were identified as other organisms.

The result of this analysis indicated that the modified

Minitek method with Roche Encise II would be the best method tor identifying the unknown cultures. The additional results obtained could then be used as supplementary tests. However, even with this technique, additional supplementary tests might be necessary, but the incidence of these tests was not sufficient to add any tests besides dulcitol to the modified Minitek technique.

IV. Selection of Biochemical Methods

The results with the Minitek discs were compared to the alts of the conventional tube methods. In general, good agreement was observed between these results, except for urea and $\mathrm{H}_2\mathrm{S/indole}$. The main difficulty with the urea disc was the indefinite colour of the positive disc reaction. The Christensen's urea agar slant technique (Christensen, 1946) was first compared with the disc method. However, this agar slant method also gave intermediate colour reactions and was more difficult to prepare and inoculate than a broth. Therefore, Christensen's urea broth (Edwards and Ewing, 1974) was compared with the disc method for a large number of cultures. This method was more satisfactory and enabled delayed urease production, notably by some <u>Klebsiella</u> spp. (Edwards and Ewing, 1972), to be detected. In comparison, the Minitek urea discs were totally unreliable, resulting in numerous false positive and negative results.

RESULTS

I. <u>Distribution of E. coll. Coliform and ENTEROBACTERIACE) E</u>
Counts

A total of 188 samples were analyzed using direct plating (VRBA and VRBG) and the MPN techniques (LST, BGB, EMB and EC). The distribution of coliform, Enterobact-eriaceae and E. coli counts observed for the ground beef-samples is shown in Table 7. Of the 169 ground beef samples, 104 (62.3%) samples had counts less than 10 <u>r</u> coli per gram and 18 (10.8%) samples had counts exceeding 500 per gram. Similarly a total of 19 (79.2%) frozen pork sausage samples had counts below 100 E. coli per gram, and some of them exceeded 500 per gram. The only pork sausage samples that exceeded 500 E. coli per gram were those that had been thawed prior to sale.

II. Comparison of Counting Techniques

A. <u>Differences_between_Sample_Types</u>

A log₁₀ transformation of the data was performed and an analysis of variance¹ of the log transformed data was carried out. The analysis of variance results, shown in Table 8, indicate that although there is a statistically

¹ ANOVAR program from SPSS. Computing Services, University of Alberta, Edmonton.

Table 7. Distribution of <u>Fiterobacteriaceae</u> counts of ground beef

Order of Counts per gram* 100 101 102 103 104 105 MPN_TECHNIQUE Number of Samples in each category: Presumpti oliforms 0 . 20 6.8 55 18 Confirmed coliforms 56 1 28 66 1.3 Completed coliforms 13 47 65 35 8 Faecal E. coli 28 76 44 15 1 DIRECT PLATING VRBA - Total 17 62 65 20 - Bile ppt 51 65 u_3 VRBG - Total 10 24 61 61 - Bile ppt 14 64 58 19

^{* 10° --} count < 10 per gram

^{101 --} count ≥ 10 < 100 per gram

Table 8. Statistical analysis of the <u>E. coli</u>, coliform, and <u>Enterobacteriaceae</u> counts on raw ground meats:

1. Analysis of variance to test for significant differences between bacterial counts from meats and plating media.

Source ¹	Degraes of treedom	Mean <u>Squares</u>	r <u>ratio</u>	Probability
A	4	18.954	4.524	0.002 **
Error (A)	179	4.189		00.1
В	/	21.148	75.340	0.001 ***
A x B	23	0.284	1. 0 1 3	0.447
Error (B)	1253	0.28 1		

2. Analysis of variance to test for differences between the 8 counts on pooled data.

<u>Source</u>	Degrees of <u>treedom</u>	Mean <u>squares</u>	r ratio	Probability
Sample Error (S(within)) B B x S(within)	183 7 1281	4.614 39.465 0.284	138.99	0.001 ***

A: All samples (i.e. high, low and economy ground beef, frozen and thawed pork sausages).

B: Differential media counts (i.e. VRBA-T, VREA-P, VRBG-T, VRBG-P, LST, BGB, EMB and EC).

3. Duncan's Multiple Range test.

Means in increasing order

EC EMB VRBA-P BGB LST VRBG-P VRBA-T VRBG-T

1.704 2.342 2.370 2.754 2.926 2.932 2.949 3.076

AT 5% level of confidence?

EC EMB VRBA-P BGB LST VRBG-P VRBA-T VRBG-T

The counts under the common line are not significantly different

interaction effect between product and method of enumeration (plating media). Since the objective of this part of the study is to compare the counts obtained on the different selective media, the differences between product types are not considered. Because of the lack of interaction effects between products and media, differences between media can be studied from these data (see below).

B. <u>Differences between Media</u>

The analysis shown in Table 8 indicates a significant difference between media (p=0.001). Duncan's Multiple Range test¹ based on the description of the test by Steel and Torric (1900), was used to study media differences. At the 5% level of significance, the media could be separated into 5 separate groups, as shown in Table 8 (part 3): (i) The total count on Mossel's <u>Enterobacteriaceae</u> medium (VRBG-T), which was significantly higher than any of the other 7 counts. (ii) The presumptive coliform (LST) count, the bile precipitating colony count on Mossel's medium (VRBG-P) and the total count on Violet Red Bile agar (VRFA-T), which were similar. (iii) The confirmed coliform (BGB) count, which was significantly greater than the bile precipitating coliform count on Violet Red Bile agar (VRBA-P) and the completed coliform (EMB) count. (iv) The VRBA-P and EMB counts, which

¹ APL-DUNLN program. Computing Services, University of Alberta, Edmonton.

were similar, but significantly greater than (v) the E_{\star} colinger (EC) count.

C. Correlation between Different Counts

results are presented in Table 9. All correlations between counts were highly significant (p=0.001). For ease of presentation, the results are considered as 3 groups.

- (a) Direct plating metia. The correlations between these counts were high, ranging fit.

 to 0.97. The lowest correlation coefficients and lower obtained for the Violet Red Bile agar, he witating count (VRBA-P) with counts on the other media. On the other hand, VRBA-T, VRBG-T and VREG-P were highly correlated, with greater than 80% predictability be seen the counts.
- between the MEN coliform counts ranged from 0.72 to 0.92.

 The predictability of the BGB (confirmed) count from the LST (presumptive) count was 85%, however for the LST and BGB coliform tests the predictability of the EMB (completed) test was only 52 and 64%, respectively. The correlation coefficients between the MPN coliform counts (LST, BGB and EMB) and the elevated temperature <u>E. coli</u> count were low,

Pearson's correlation SPSS program. Computing Services University of Alberta, Edmonton.

Table 9. Pearson's correlation coefficients between 8 different <u>Enterobacteriaceae</u>, colliform and <u>E. collicounts</u>*

r – A el si v	L ABHY-i.	V RBG-T	V KBG-P	LST	BGB	EMB

				į	•		
EC	0.27	0.46	0.31 %	0.36	0.44	(52	0.68
EMB	0.51	0.63	0.48	0.51	0.72	0.80	
BGB	0.67	0.70	0.69	0.72	0.92		
LS,T	0.71	0.70	0.73	0.72			
VRBG-P	0.90	0.80	0.97				
VRBG-T	0.93	0.77					
VRPA-P	0.83						

^{*} VPBA Violet Red Bile agar
VRBG Mossel's <u>Enterobacteriaceae</u> medium
(VRBA+1% glucose)

LST Lauryl Tryptose broth

BGB Brilliant Green Bile 2% broth

EMB Levine FMB agar

EMB Levine EMB agar EC EC broth at 44.5°C

⁻T Total count of bile precipitating and non-bile precipitating colonies at 24h

⁻P Count of bile precipitating colonies at 24h

resulting in poor prodictability of the E. coli count.

relationships exist between the LST and BGB counts and the direct plate counts, but only 40 to 50% of the count on the direct plating media could be predicted from the MPN counts. The relationship between the EMB (completed) coliform and the EC (E. coli) counts and the direct plating media was much less.

D. Effect of Temperature and Incubation Time on VRBA and VRBG Counts.

It was observed that both during refrigeration or on plates held at room temperature, more non-bile precipitating (B-) colonies grew on the VRBA and VRBG plates. Extended incubation at 37°C also resulted in higher numbers of B-colonies appearing on the plates. A fixed time-temperature of incubation, researched 37°C for 24±2h, was necessary for consistent results to be obtained.

E. EC Test

1. Comparison of the Direct and Indirect EC Tests

A total of 54 samples were analyzed by the direct and indirect methods in EC medium at 44.5°C for 24 and 48h.

Log₁₀ transformed data were compared using a t-test* which

gave a highly significant difference between the two counts (p<0.01). The mean of the counts for the direct test was 7.4 x $10^{3}/3$ compared to 1.4 x $10^{2}/9$ for the indirect test.

2. Comparison of EC Tests at 44.5 and 45.5°C.

The 1,030 <u>Enterobacteriaceae</u> isolates from ground meats were inoculated into duplicate sets of EC broth and incubated at 44.5 and 45.5°C. The results for gas production are shown in Table 10, and may be summarized as follows:

- (a) Of the 734 isolates that were gas negative after 48h incubation at 4 °C, only one isolate produced gas at 45.5°C, resulting in 99.9% agreement in the tests.
- (b) Of the 295 isolates that were gis positive after 24h incubation at 44.5°C, 7.8% (in isolates) exhibited delayed gas production at 45.5°C, and 1.4% (4 isolates) railed to produce gas after incubation at 45.5°C for 48h. The 23 isolates giving delayed gas production at 45.5°C included 22 E. coli I and one E. agglomerans. The isolates that were gas negative with extended inculation at 45.5°C, included two E. coli I and one E. pneumoniae and an E. cloace.
- (c) There were only 7 delayed gas producers at 44.5°C, and three of these failed to produce gas after 48h at 45.5°C. None of these isolates were \underline{E} . \underline{coli} I.

tetest program of the SPSS Computing Services, University of Alberta, Edmonton

ible 10 Relationship between elevated temperature tests in EC broth at 44.50c and 45.50c

		luced in EC		 	
Gas produced in EC at 44.5°C			Negative 48h	dumber of cultures	
l Posit re 24h	1 +0.8	7 8	1 1.4	295	
Pos tive 48h	28.0 j	28 • b	1 42.4	7	
Negative 48h 	0.0	0.1	99.9 - 99.9 -	/ } 4 	

III. Identification of ENTEROBACTERIACEAE Isolates

A. Confirmation of Colony Morphology

The colonies isolated from VRRA and VRBG plates for identification were described on the basis of the presence and amount of visible bile precipitation at the time of the ection. The colonies on EMB were selected according to their ociphology (see methodology p.35). Because of the changes of contamination in selecting the isolates, especially whe picking sub-surface colonies from the direct plating media (VRBA and VRBG), it was necessary to check the colony types of the puritied cultures on the selective media.

were re-plated onto VRPA (see Table 11). For the original B++ isolates, a few (4.4%) were b+ on re-plating, but others (17.0%) formed B- colonies. Similar manges in morphology of original B+ and B- colonies occurred when they were replated onto VRBA. There was little concern for changes when a contaminant or a marked change in the culture during purification and storage. For B+ isolates, it was difficult to interpret the changes from B+ to B++, and B+ to B- for the re-plated cultures, because the conditions on the original plates (e.g. meat particles in the inoculum) could have influenced the appearance of the colonies. For B-

Table 11. Comparison of morphology of colonies originally isolated from Violet Red Bile agar (VRBA) with their morphology when resplated on VRBA after purification*

	Morphology	on VRBA	(r plated)	Number -
Morphology on VAHA	 H++ 	H +) Lturess
3 + +	 	4.4% (3)		[68
B +	 32.7% (36)	33.5% (39)	31。8第 (35)	1 110 1 0
8 - ((6)	14.4% (19)	81.1% (107)	1 132
Totals	 95	31	154	310

Table 12. Comparison of morphology or colonies originally isolated from Violet Red Bile agar + 1% glucose (VRBG) with their morphology when re-plated on VRBG after purification*

Original morphology on VEBG	Morpholog	gy on VRBA	(re-plated) B-	Number of cultures
B++	1 80.2x (109)	15.4% (21)	4.4% (6)	130
1 · B+ 	 38.8% (40)	50.5% (52)	10.7%	
B-	1 18,7% (*)	50.0% (40)	31.3% (25)	80 9
Totals	164	113	52	319

^{* 3++ --} Large dark zone or bile precipitation

B+ -- faint zone of bile precipitation

B- -- no bile precipitation

replating were considered similar, but change to B++ suggested either a contaminant or marked change in the culture during laboratory handling and storage.

Data for VRBG isolates re-plated onto VRBG are shown in Table 12, and the same considerations applied.

The data for the isolates from EMB, re-plated to EMB agar, are shown in Table 13. Considerable changes occurred in colony morphology, nowever the chance of contamination, compared to VRBA and VRFG, was much less because the isolates were surface colonies. Considerable changes could be observed between original MS+ and NU+ isolates, in fact as many as 60.8% NU+ isolates became MS+ (nucleated, with metallic sheen) on re-plating. Such changes in morphology were most likely due to pleomorphism, which in the case of the non-coliform NU- colonies, could be of considerable importance to the interpretation of the coliform count on EMB.

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B. Association of Colony Morphology with Types of 2rganisms

The distribution of organisms between colony types is shown in Tables 14-15. The description of the colony morphology in these tables refers to the morphology both when the isolates were originally selected and after replating (confirmed colony morphology). This was necessary

Table 13. Comparison of morphology of colonies originally isolated from EMB agar with their morphology when re-plated on EMB after purification*

Original morphology	Morpho	Morphology on EMB (re-plated)					
on EMB	MS+	NU+	N U -	MU+	ot cultures		
MS+	85.9% (128)	9.4% (14)	2.0% (3)	2.6% (4)	140		
N U +	60.8% (76)	24.0% (30)	10.4% (13)	4.8% (6)	125		
NU - 1	17.1% (14)	12.2% (10)	67.1% (55)	3.7%	82		
MU+ 1 1	27.6 % (16)	15.5%	15.4% (7)	41.4% (24) 美国	58		
Totals	234	63	80	37	414		

^{*} MS+ -- metallic sheen, nucleated colony NU+ -- nucleated (pink + black centre) colony

NU- -- non-nucleated colony

MU+ -- mucous with or without a nucleus

for a reliable association of organisms with colony morphology to be obtained.

1. Violet Red Bile agar: Coliform Medium

categories, based on the degree of bile precipitation. The identity of the organisms associated with the different colony types (B++, B+, B-) is shown in Table 14.

(a) B++ colonies

-

The main organisms were <u>E. coli</u> I (46%), aerogenic <u>E. aqqlomerans</u> (15%) and <u>E. clóacae</u> (15%). On re-plating, the <u>E. coli</u> I accounted for an even greater proportion of the organisms associated with this colony type (60%). <u>E. cloacae</u> remained important, but <u>E. aqqlomerans</u> decreased in relative importance. Changes on re-plating could be attributed primarily to <u>E. coli</u> I colonies that were originally B+ becoming B++.

(b) B+ colonies

The main organism associated with this group was E.

Coli I (30% of the original isolates) but this decreased to

14% on re-plating because so many of the E. coli I gave

greater zones of bile precipitation when re-plated in pure

culture on VRBA. The principal organisms in this group

became S. liquetaciens (30%) and aerogenic E. agglomerans

(21%).

Table 14. Identity and number of organisms associated with different colony types on Violet Red Bile agar (VRBA) from original isolates and for purified (re-plated) cultures*

l 	Orig 0rig B++	ginal (Type B+	Colony B-	Conf:	irmed Type B+	Colony B-
<u>F</u> . <u>col</u> l I	1د ا	34	4	57	g	.3
E. coli II	1	2	2	2	1	2
ther <u>E. coli</u>	0	1 '	1	0	1	1
C- <u>fr∈undii</u>	3	6	В	8	4	5
K. <u>pneumoniae</u>	5	4	3	6.	5	1
K. <u>ozaenae</u>	1	2.	0	0	,0	3
E. <u>agglomerans</u> aerogenic anaerogenic	10	20	30 19	1 3 1	13 . 1	44 22
<u>E. aerogenes</u>	0	3	2	0	1	4
<u>E. cloacae</u>	10	5	6	11	2	8
<u>E. hafniae</u>	3	3	5 .	4	3	. 4
<u>S. liquefaciens</u>	2	23	3 વ	2	18	44
S. marcescens	0	. 0	1	0	0	1
<u>S. rubidaea</u>	Û	2	۷	0	. 0	4
Y. enterocolitica	0	0	1 .	0	0	1
Uridentifiable	1	2	1	1	1	2
Oxidase positive	0.	1	16	0	2	1 5
Total number	68	112	14 Č	95 	61	1 64

* See Table 11

(c) B- colonies

The main organisms associated with this group were S. liquefaciens, aerogenic and anaerogenic E. agglomerans and the non-Enterobacteriaceae. Of the Enterobacteriaceae isolated as B- colonies, S. liquefaciens and E. agglomerans accounted for 63% of the isolates.

2. Mossel's Violet Red Bile Glucose agar: ENTEROBACT-ERIACEAE medium

The organisms growing on VRBG were handled and analyzed as described for VRBA, and the data are shown in Table 15.

(a) B++ colonies

The main organisms isolated originally and confirmed by re-plating as B++ were <u>E. coli</u> I (24% and 31%) and <u>S. liquefaciens</u> (26% and 31%). Other B++ isolates included <u>C. freundii</u>, <u>K. pneumoniae</u>, <u>E. aqqlomerans</u>, <u>E. cloacae</u> and <u>E. hafniae</u>, these organisms on re-plating, accounted for 30% of the B++ colonies. The same trend observed on VRBA, where B+ colonies became B++ on re-plating was observed on VRBG.

(b) B+ colonies

The main organisms in this group were aerogenic and anaerogenic \underline{E} . asglomerans. The relative importance of \underline{E} . coli I and \underline{S} . liquefaciens decreased as a result of the replating, and the change of B+ colonies to B++.

Table 15. Identity and number of organisms associated with different colony types on Mossel's <u>Enterohact-epiaceae</u> medium (VRBG) for original isolates and for purified (re-plated) cultures*

	Original Colony Type			 Confi	 Confirmed Colony Type		
Organism	B++	B+	В -	B++	B+	B-	
<u>E. coli</u> I	34	14	5	51	2	0	
E. coli II	4	1	2	4	2	1	
C. <u>freundii</u>	! 8	3	2	10	3	()	
K. pneumoniae	9	1	- 2	111	1	0	
K. <u>ozaenae</u>		3	1	1 3	2	1	
E. <u>aqqlomerans</u> aerogenic anaerogenic	15 - 15	11	20 20	8	38 25	9 16	
E. <u>aerogenes</u>	0	5	3	1 0	5	3	
<u>E. cloacae</u>	12	4	2	1 12	6	0	
E. <u>hafniae</u>	5	6	3	7	5	2	
<u>S. liquefaciens</u>	36	26	5	51	15	1	
S. marcescens	1	0	1	1	1	0	
<u>S. rubidaea</u>	0	0	2	0	0	2	
Proteus s	0	0	2	0	0	2	
Y. enterocolitica	0	0	2	0	0	2	
Unidentifiable	1	1	3	2	3	0	
Oxidase positive	3	7	14	1 1	2	20	
Total number	139	106	89	1 164 1	112	58	

^{*} See Table 12

(c) by colone a

The main organisms among the <u>Enterobacteriaceae</u> in this croup were aerogenic and anaerogenic <u>E. agglomerans</u>. After to-plating, they accounted for 66% of the <u>Enterobacte aceae</u> growing as B- colonies on VRBG. Of the re-plated organisms, 50% of the B- colonies were not <u>Enterobacteriaceae</u>.

3. Eosin Methylene Blue agar: Confirmed Coliforms by MPN Technique

The organisms growing on EMB were identified by 4 different colony types: metallic sheen (MS+), pink with black centre (nucleated, NU+), non-nucleated (NU-) and mucous (MU+). The data are shown in Table 16. MS+ and NU+ are considered typical coliform colonies. MU+ are atypical coliform colonies but NU- are considered non-coliform bacteria.

(a) MS+ colonies

The principal organism in this group, from original isolates and after re-plating, was E. coli (67 and 66%, respectively). C. freundii and K. pneumoniae accounted for 24% of the isolates producing metallic sheen on re-plating. Some E. agglomerans also produced a metal sheen, but the colonies did not have typical dark centres and the metallic sheen was faint. The increased number of MS+ colonies on replating could be primarily attributed to NU+ colonies

Table 16. Identity and number of organisms associated with different colony types on Levine EMB aq.C from original isolates and for purified (re-placed) cultures*

	- T		. =			2-1-2	, 	
1					Conf	 		
Organism	MS+		N11 -	M () +	i MS+		ИΩ-	M (1 +
<u>E. coli</u> I	100	5.8	9	4	1 154	14	1	2
E. coli II	1 3	3	1	()	1 6	O	1	ο,
C. freundii	1 13	17	17	ή.	1 27	16	8	1
K. Pneumoniae	1 19	16	6	20	1 1 29	17	4	11
K. ozaenae	1 1	3	3	5	1 5	1		2
E. agglomerans ae ogenic an erogenic	5 0) 1	3 1	2 0	1 1 1 5 1 0	0	~ 2	5 0 1
I E. <u>aerogenes</u>	1 1 1	0	1 -	Ö	0	0	1	1
E. cloacae	i i 1	13	11	15	}	11	14	12
I <u>E. hafniae</u>	1 1	1	5	0	3	0		0
S. <u>liquefaciens</u>	1 1	6 3#	12	5	. 1	3		1 1
l <u>S. rubidaea</u>	1 0	ा : 0	1	0	0	0	1	0 1
 Unidentifiable	! 1	1	0	2	2	О	0	2
Oxidas positive	1 1 3 1	2	12	0 1	O	1	16	0
Tot . number	149	126	82	58	235	63	80	37

* See Table 13

producing metallic sheen. This sup and pleomorphism in colony types on EMB agar.

(b) NU+ colonies

Because of the above-mentioned change from NU+ to MS+ on re-plating, the number of NU+ colonies decreased from 126 to 63. Of the re-plated NU+ colonies, E. coli I (22%), C. treundii (25%), K. pneumoniae (27%) and E. cloacae (17%) were the main organisms.

(c) NU- colonies

NU- colonies are considered non-coliforms in the MPN technique. However, some pleomorphism was observed between NU- (non-coliform colonies) and typical coliform type colonies. For the re-plated colonies, the principal organ types included S. <u>liquefaciens</u> (24%), <u>E. cloacae</u> (18%) an 20% non-<u>Enteropacteriace</u>ae organisms.

(d) MU+ coloni.

The principal organisms in this group were \underline{K} . $\underline{Pneumoniae}$ and \underline{E} . $\underline{cloacae}$.

C. Interrelationships of Colony Morphology between the Selective Media

All of the isolates obtained in this study were plated onto all 3 of the selective media (YRBA, VRBG and EMB) to determine the interrelationships of their colony morphology.

Dilutions that gave between 30 and 50 colonies per plate were inoculated onto each medium. The plates were incubated at 37°C and observed after 24%? The results are shown in Tables 17-19.

On Mossel's VRBG medium (Table 18) most of the Enterobacteriaceae produced a zone of bile precipitation around the colonies, only 49 (4.8%) of the Enterobacteriaceae failed to produce a visible zone of bile precipitation. Of these, 71.4% were E. agglomerans, representing 17.2% of the E. agglomerans isolated in this study. Of the total B-colonies on VRBG, 31.2% were non-Enterobacteriaceae. In contrast on VRBA (Table 17) a total of 404 (39.7%) Enteroba eriaceae isolates grew as non-bile precipitating colonies. The principal organisms producing B- colonies on VRBA were E. agglomerans (41.3%), S. liquefaciens (28.0%), E. cloacae (7.4%), C. freundii (5.7%) and E. hafniae (4.4%). Of the non-Enterobacteriaceae B- isolates, 95.1% grew as B-colonies on VRBA, representing 8.8% of the total B- isolates on VABA.

All of the recognised coliform type organisms (E. coli, C. freundii, K. pneumoniae, E. aerogenes and E. cloacae) formed B++ or B+ colonies on VRBA. However, 27.4% of C. freundii isolates, 71.4% of E. aerogenes and 38.0% of E. cloacae failed to produce a visible zone of bile precipitation. Some non-coliform type organisms produced visible zones of bile precipitation on VRBA, including 28.9% of S.

Table 17. 6: bwth characteristics of Enterobact Flaceae isolates on Violet Red Bile agar (VRRA) *

I I	l I gion	Tot '		
Organism	1 0++	8+	B+	Numi
i <u>E. coli</u> I	85.7	9.3	0.7	288
E. gy 5 41	(to 0	. () . ()	25.0	1 20
l C. <u>freun</u> dii	i i : 0	41.7	27.4	1 84
K. <u>Ingumoniae</u>	1 1 7	21.2	5.9	85
K. <u>Ozaenae</u>	! ⊿3.3	19.0	47.6	1 21
E. agglomerans aerogenic anaerogenic	8.4 1.4	17.6	74.0 95.4	
l E. a.e. Menes	7.:	21.4	71.4	 14
l <u>E. cloacae</u>	3 . 1	13.9	₹ 38.0	79
E. <u>bafnias</u>	25,8	16.1	58 .1	31
<u>S. liquefaciens</u>	3.8	25.1	71.1	159
<u>S. rutidaea</u>	14.3	0.0	85.7	7 7
; 				

^{*} B** -- large zone of bile precipitation .
B+ -- small zone of bile precipitation

B- -- no bile precipitation

Table 18. Growth characteristics of Enterohacteriaceae isolates on Mossel's Violet Red Bile agar + 1% glucose (VRBG) *

İ	' - 	Perc nt growing on VRBG as:				
Organism	B++	B+	B-	Number		
E. coli i	95.1	4.5	0.3	238		
E. coli II	70.0	25.0.	5.0	1 20		
C- freundii	40.5	9.5	0.0	84		
K. pneumoniae	98.8	0.2	0.0	85		
K. ozaenae	76	19.0	4.8	21		
E. auglomerans aerogenic anaerogenic	25.4 2.7	63.8 68.5	10.8 28.8	1 1 131 1 73		
E. aerogenes	42.9	57.1	0.0	14		
E. cloacae	87.3	12.7	0.0	79		
E. hafniae	74.2	19.4	6 - 4	31		
S. liquefaciens	74.2	25.2	0.6	1 159		
S. rubidaea	28.6	28.6	42.8	7		

^{*} B++ -- large zone of bile precipitation & -- B+ -- small zone of bile precipitation

⁻⁻ no bile precipitation

l quefact to and 18.1% of E. andlowerans.

Type a colliform colonies on EMB are nucleated, with or without metallic sheen (MS+ and NU+), atypical mucous colonies (MU+) also occur, and non-colliform colonies are non-nucleated (NS-). The results of lating the Enterobact-eriaceae is plates from this grady onto EMB are shown in Table 19.

The principal organitims growing as MS+ and NU+ olonies on EME included \underline{E} . \underline{coli} , \underline{C} . $\underline{freundii}$, \underline{K} . $\underline{pneumoniae}$ and \underline{E} . auglomerans. Of these, E. coli, C. freundii and K. * ? <u>pneumoniae</u> represented typical coliforms. Some <u>K. pneumoniae</u> grew as atypical coliforms (MU+), however 3.8% of \dot{E} . coli, 21.4% of C. freundii, and 10.6% of K. pneumoniae grew as non-coliform organisms (NU-) on EMB. The majority of $\underline{\mathbf{E}}$. agglomerans 3.8%) were NU-, but those that were MS+ (12.2%) could be distinguished from the coliform organisms producing metallic sheen. The principal NU-sorgantsms were S. liquefaciens and E. agglomerans. The non-Enterobacteriaceae organisms generally grew as NU- colonies on EMB. Some of the cultures failed to grow within 24h when replated onto selective media. However, some grew on VRBG and VRBA as B- colonies and on EMB as NU- colonies when incubated for a further 24h at room temperature.

الرثيق



Table 19. Growth characteristics of <u>Enterobacteriaceae</u> isolates on Levine EMB agar

l (Orjanism	Percent Percent growing on EMB as: Originism MS+ NU+ NU- MU+					
 <u>E- coli</u>	84.1	13.1	1.7	1.0	290	
E. coli II	1 65.0	10.0	25.0	0.0	20	
C- <u>treundri</u>	1 50.0	27.4	21.4	1.2	1 84 1	
<u> K- Fuenmoniae</u>	1 44.7	28.2	10.6	16.5	1 185 I	
i <u>K. ozaenae</u>	1 48.0	4.8	57.1	9.5	1 21 1	
E. <u>agglomerans</u> aerogenic anaerogenic	1 10.8	16.8 12.2	60.3 83.8	6. 1	1 1310	
i <u>E. derogenes</u>	0.0	14.3	78.6	7.1	1" 8 1	
D cloacae	 ₋ 6•3	26.6	39.2	27.8	79	
E. <u>natnia</u> .	21.9	15.6	62.5	0.0		
5. <u>liquetaciens</u>	1.9	5.7	91.1	1.3	1 158 1	
S. <u>rubidaea</u>	0.0	14.3	85.7	0.0		
	<u> </u>			*		

♦

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D. Growth Characteristics of ENTEROBACTERIACIAE

Loclates in Lauryl Sulf. to Tryptose broth (LST, and

Lactose broth

i.

زنز

were inoculated into Lo and Lactose broths. The results indicate which of the <u>Hobacteriaceae</u> isolated in this study would be scree of by these presumptive coliform tests. The sults for LST broth are shown in Table 20. The organisms that were able to produce gas from LST were E.

Coli I (93.6%), E. coli II (75.0%), C. freundii (67.9%), K.

Pheumoniae (90.6%, aerogenic E. agglomerans (46.3%), and E. cloacae (79.7%). Only 15.7% of S. liquefaciens isolates produced gas in LST tubes.

The results for growth in Lactose broth are shown in Table. The majority of E. coli I. C. freundii and K. pneumoniae produced acid and gas within 24h. Additional insubation for 48h only increased the lactose positive E. coli I by 6.0% and K. pneumoniae by 4.7%. Anaer remic E. coli were observed, but their numbers were not high, and only 1. or E. coli I were lactose negative E. coacae isolates were generally aerogenic, la tose fermenters (86.1%): Of the 15 E. aerogenes isolates, 7 were non-lactose fermenters, the lactose fermenters were either aerogenic or anaerogenic.

The sijority of S. <u>liquefaciens</u> were either non-lactose fermenters (71.1%) or argerocent for lactose (5.0%). The

Table 20. Growth characteristics of <u>Enterobacteriaceae</u> isolates in Laury Sulphate Tryptose broth (LST)

	Perceit	rowing or	LST as:	
Organis	Positive 24h	Posit.ve 48h	Negative 48h	Total Number
E- COLL I	93.6	0.7	5.7	296
E. coli II)	0.0	25.0	20
C. freundii	67.9	16.7	15.5	84
K. pneumoniae	! !	3.5	. 5.9	l 85 l
I <u>K. ozaenae</u>	1. 71.4	9.5	19.0	21
E- agglometans aerogenic anwerogenic	18 1 46.2 1 5.4	28.0 9.5	25.8 85.1	132 74
l <u>E. aerogenes</u>	13.3	13. 3	73.3	15 1
! ! E. <u>cloaçae</u>	79.7	6.3	13.9	7 9
E. hafniac	40.6	12.5	46.9	32
S. <u>liquefaciens</u>	15.7	4. 8	60.5	159
l <u>S- rubidaea</u> l	14.3	0.0.	85.7	7 , 1
L			<u>`</u>	

Table 21. Growth characteristics of Enterobacteriacede isolates in Lactose broth*

1	Perce	Percent growing in Lactose hroth as:				
Organism	A+G+ 1 24h	A+G+ 48h	A+G- 48h	A-G- 48h	Total Number	
E- coli T	94.3	0.7	4 - 1	1.0	296	
<u>E. coll</u> II	70.0	5.0	15.0	10.0	1 20 1	
C. <u>freundii</u>	83.3	6.0	4.8	6.0	84	
l <u>K. pneumoniae</u>	 88.2	4.7	5.9	1.2	1 85	
l <u>K</u> . <u>ozaenae</u>	1 61.9	14.3	4.8	19.0	1 21 1	
E. agglomerans aerogenic anaerogenic	 47.7 8.1	16.7 8.1	7.6 32.4	28.0 51.4	1 132 1 74	
l <u>E. aerogenes</u>	1 1933.3	* <i>₽</i> > 1 3.3	6.7	46.7	l , 15	
l <u>E. cloacae</u>	1 74.7	11.4	0.40	13.9		
l <u>E. hafniae</u>	40.6	12.5	್ಷ್ಯ₃ 3.1	43.8	 3^	
l <u>S. liquefaçiens</u>	1 12.6	11:3	5.0	71.1	1 159	
· · · · · · · · · · · · · · · · · · ·				Sur	ķ € "	

^{*} A+G+ -- Acid and Gas produced

A+G- -- Acid produced, but no gas A-G- -- No Acid or Gas produced

data in Table 21 indicate that aerogenic E. agglomerans were primarily lactose termenters (64.4%), whereas the anaerogenic E. agglomerans were lactose negative (51.4%), and a further 32.4% were anaerogenic for lactose.

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The comparison of LST and Lacto: Table 22. Anaerogenic lactose fermen tose broth were also negative in LST. In LST, 18% of tose negative isolates were gas positive. The majority of the organisms accounting for this difference were E. agglomerans and S. liquefaciens. Simularly, 23.6% of anaerogenic lactose termenters gave LST positive results. There were only 17 of these isolates, and of these, 5 were E. agg merans and 4 were C. freundii. The delayed lactose and LST negative cinarily S. liquefaciens and Enterobacter isolate The delayed lactose and LST cultures were mainly E Aglomerans; and delayed lactose and delayed LST positive isolates were principally C. freundii and E. agglomer Lactose positive cultures consisted of 4.3% delayed LST and 5.2% LST negative cultures. The delayed LST cultures were primarily C. freundii and E. agglomerans, whereas the LST negative cultures were principally C. freundii, S. liquefaciens, E. agglomerans and E. hafniae.

Table 72. Comparison of growth characteristics of <u>Entero-bacteriaceae</u> isolates in Lauryl Tryptose (LST) and Lactose broths

	Percent	growing	in LST as:	Total
Lactose !	Gastat 20h	Gaş at 48h	no Gas at 48h	
Acid+gas at 24h	<u>90.5</u>	<i>L</i> , 3	5. 2	 629
Acid+gas at 48h	33.3	<u>=6.9</u>	39.7	l [` €
Acid only at 48h	9.7	13.9	<u>76.4</u>	72
Negative at 48h 	7.2	10.8	<u>82.1</u>	l 251

Underlined results indicate tests which have compatible results $\inf_{\mathbb{R}^n}$ both tests

Table 23. Occurrence of IMViC types within Enterobact-eriaceae isolates with different Lauryl Tryptose broth (LST) and Lactose Lauryl Tryptose

IMViC types	LST+	LST-	Lactose+	Lactose-
++	294	24	295	19
-+	1 50	26	59	25
 +-++	37	2	38	: 0
 -+-+	67	15	73	8
}	11	15	25	14
1 1+ 1	30	30	33	21
++	164	17∠	208	130
[· ·	L	·		

E. Relationship of ENTEROBACTERIACEAE Isolates with IMViC Types

The <u>Enterobacteriaceae</u> isolates, have been considered on the basis of their lactose fermentation and production of gas in LST. The IMVic types associated with these Lactose and LST results are shown in Table 23. IMViC types that were represented by less than 10 cultures have been eliminated from the results. Of the 7 remaining IMViC types. LST and Lactose positive cultures were mainly IMViC ++-- and --++, while for LST and Lactose negative cultures the principal IMViC type was --++.

Enterobacteriaceae (coliforms). For these isolates, the association between IMViC type and type of organism is shown in Table 24. The ++-- types were generally E. coli I, but if few other lactose organisms were also ++--. The -+-- organisms were E. coli II, but some C. freundii and other organisms including K. ozaenae, Enterobacter spp. and S. liquefaciens were also observed as -+--. For the lactose fermenting E. agglomerans d S. liquefaciens, the principal IMViC group was --++, but these organisms were also distributed across most of the other IMViC groups presented in Table 24.

The results of IMViC tests on lactose negative isolates are shown in Table 25. The 5 lactose negative <u>E. coli</u> gave typical [MViC ______s. Lactose negative <u>E. aqqlomerans</u>

Table 24. Relationship of lactose positive organisms and LMViC type::

	¢;à·						
(IMVic types of lactose positive Enterobacteriaceae						
Organisms	1 ++	+ + +	·· • · •	-+	+-	1,	++
E- coli 1 A.	290		_	_	-	*	
E. coli II	i -	-	-	18	· _	-	-
<u>C. freundii</u>	i –	-	51	1 8	-	7	1
K. pneumoniae	1 3	35	-	-	<u>-</u>	2	31
<u>K. ozaenae</u>	i -	_	3	, }	-	Q	1
E. <u>agglomerans</u> aerogenic anaerogenic	i 1 5 1 -	2 2	8 1	5 - 4	13	6, 6	51 17
E. <u>aerogenes</u>	<u> </u>	-	-		-	-	6
E. cloacae	<u> </u> -	-	2	1	3	-	62
E. <u>hafniae</u>	1 1	-	3	2	3	4	5
S. <u>liquefaciens</u>		-	ч 4	1	3	9	2 7
	1			•		i	



roble 25. Relationship of lactose negative organisms and IMViC types

	IMViC types of lactose negative Enterobacteriaceae					
Organisms	++	-+-+	-+	+-	<u>-</u> 	++
E. coli I	3	_	_	-	-	
b. soli II	-	-	2	-	_	-
C. freundii 1	-	1	4		_	-
l K. Puenwonfae	-	-	-	-	-	1
<u>K. ozaenae</u>	-	-	3	-	-	_
E. agglomeros aeroge anaerogenic	1.5 1	2 2	8 8	1 6	3 5	2 5
E aerogenes	, -	-		-	_ -	7
E. cloacae	-	1	-	2	-	7
E. <u>hafnigs</u>	-	1		3	1	7
S. <u>liquefaciens</u>	-	1	_	1	12	99
S. marcescens	-	-	-	1	-	
	·.					<u> </u>

included 15 that were ++-- and 16 -+--. Most of the lackness were --++-

Results for LST positive isolates were similar to t lactose positive results, shown in Table 24. For LST negative cultures the pattern was similar to that for Lactose negative isolates, but there was a greater number of E. coli that were ++-- (16 isolates) or -+-- (5 isolates).

The use of the elevated temperature test at 44.5°C in SEC broth, in association with the IMViC tests, eliminated the organisms that were not \underline{E} . \underline{coli} in the type \underline{I} (++--) and type II (-+--) groups, and will be discussed in the following section.

F. Association of ENTEROBACTERIACEAE with Elevated Temperature Test Results

The elevated temperature tests in EC broth were carried out at 44.5 and 45.5°C. The association of different organisms with the growth characteristics in EC broth at 44.5°C are shown in Table 26. The EC positive organisms at 44.5°C were primarily E. coli, however a small percentage.

(7.1%) of E. coli I were EC negative which would result in false negative results for 21 out of 296 isolates. There were only 20 -+-- isolates, but of these, 55% were EC positive and 45% EC negative. If elevated temperature (44.5°C) EC tests were used to identify faecal E. coli, false positives would result from E. coli II (-+--), K.

Table 26. Distribution of or janisms with different growth characteristics in EC broth at 44.5%

	EC at 44.5°C Percent growing as:				
l Organism		Positive 48h	Negative 48h		
E. coli I	92.2	54.1	2.9		
E. coli II	3.7	_	1.2		
C. <u>freundii</u>	-	-	11.4		
K. pneumoniae	(1.0)	14.3	11.0		
l <u>K. ozaenae</u>	<u> </u>	<u>14.3</u>	2.7		
E. auglomerans aerogenic anaerogenic	(1.0)	<u>14.3</u>	17.4 10.1		
i i <u>·Ē· āerodeņēs</u>	<u>.</u> -	4 -	2.0		
E. clogcae	(0.3)	-	10.6		
E. hafniae	.0	-	3.9		
S. liquetaciens	-		21.6		
Unidentifiable	0.7	-			
Others.		- -	3.5		
Totals	295	7	734		

Numbers in brackets have one (0.3%) of the cultures negative at 45.5°C

Numbers underlined were negative at 45.5°C



presumerror, is apploantian, proclosed and by harmony of the operator temperature test, come talsed proclesses with the elevated temperature test, come talsed proclesses with the elevated, but talse negatives work as perfect material temperature.

ENTERORA TERIACRAS GROW Most semiliar for Ident semiliar of

In the fire nemical terms to identify the includes in this country, whiteous and lactors formentation were determined in Prenct Restrict base. Triple Sugar Ston (1881) again plant: were also incompated to detect His grade comp However, acreant pas production from discose and acre from lactor, goald also by reactin the ESI plants. Examination of the glucose and lactors results by both methods revealed some differences, as shown in Table 27. Using the tube :lecose and lactose results as references, 4.3% of the gas positive isolates tailed to produce gas from glucose on CT slants, whereas 42.1% or cas negative asolates produced gas M. TSI. Inis anvolved H4 is lites but of 1, 🖎, tested. Similarly, for isolates projucing gas and/or acid from lactore by the tube method, 13.4% failed to produce acid from lactose on the TSI slants. Of the lactose negative isolates, 24.7% produced acid on TSI (false positives).

From the solution Minitek method (see Methodology 1.00) during out on the isolates in this study, the rasid data necessary for identifying these cultures using the Roche

Calle 27. Comparison of glusome and lastome fermentation in Stiple Sugar Bron (Tal) slant and Prenol med broth tures

		TEI	I slants.	•	T I	
I Phyenolikodi I Phyenolikodi	1	Tropis	1.A	LACTOMY		
		astra en l	y	I 1 ·)	I Total	
GLUCOSE acid+gar. (at .4h)	(1965)) • 15 K () 4)	T ' 		4	
acid+qa; (at 48h) (តែកា≟្ក ុ (11)	51 <u>,</u> 7% (5)	i i		1 16	
idad only (at ush)	42.1% (45)	57.9 % (62)	1		1 1 1)7	
LACTOSE acid+gas (24/48h)	· · · · • • · · · · · · · · · · · · · ·	*** **********************************	1 1 87.4% 1 (618)	12.6% (39)	 	
acid only (at 48h) (73.6% (53)	26.4% (19)	1 1 72 1	
no act: (at 4ch)			1 1 24.7% 1 (62)	75.3% (194)	1 1 25 1 1	

Encire if system could be obtaine. Into resulted in the socialistication of swe out of the social included additional feets which enables a further occupation of the identification for the foch starts and is system, i.e. any .07 cultures to be identified identified as in a supplementary test.

of the SC equations identified with tests from the modified Minitek to Snique, the tests were used with the following frequenties:

Vogen-Pronkager 47.
ONPG 11
Motility 234
Ararinoso 181
Phamnon 237
Inosito; 104
Refrinoso 102
Methyl Fed 5
Malonats 5

First the supplementary tells recommended by the Boche Encise II system but not include in the modified Minitok technique used in this body, a function 194 isolates were identified. The supplementary tests used and their frequencies are given below:

Cellibiose 9:
Salicin of
Archine 44
KC: 1
Jorin's recate ...
Aic:
DNa
Muc: 4
Sucrose 3
Capsule stain 3

DISCUSSION and CONCLUSIONS

1. Coliform and ENTERORACTERIACEAE Counts

The distribution of the colliform and Enterobacteriaceae counts in these products were such, that they included samples that fell within the proposed Canadian E. colistandard (Pivnick, (1., 1975), exceeded the lower limit of 100 per gram, an exceeded the upper limit of 500 per gram. Larked differences were noted between product types however, the absence of interaction effects between product and plating media allowed the significant difference between media to be examined from this analysis.

According to Mossel et al. (1901), and based on the definition of Enterobacterlageag as glucose fermenting organisms (Bergey's Manual, 1974), the bile precipitating colonies on Mossel's VRBG medium should represent the Enterobacterlageag count. However, some of the non-bile precipitating colonies on VRBG were Enterobacterlageag. This was confirmed by the re-plated isolates. Although there was a significant difference at the 5% level, but not at the 1% level, between the total and bile precipitating counts on VRBG, there was less than a 2-fold difference between the means of these counts. Many of the 3- isolates (68.8% of 78 isolates) were Enterobacterlageag; as a result, using the solates) were Enterobacterlageag; as a result, using the precipitating VRBG count to represent the Enterobacterlageag; as a result, whereas using

the total VRBG count would give some false positive results. However, the differences are not great and the total count might be the easiest, since differentiation between low levels of bile precipitation and non-bile precipitating colonies is sometimes questionable, and this difficulty would be avoided.

In comparison, the total count on VRFA was statistically similar to the bile precipitating count on VRBG. The total count on VEBA, therefore, also estimates Enterobac eriaceae. The be colonies on VRBA were principally E. agglomerans and S. liquetaciens, nowever some were atypical coliform-type organisms: The bile precipitating isolates were lactose fermenting organisms that included not only the generally recognised collitors organisms (E. coli, E. aerogenes, E. cloacae, K. pneumon de, and Citrobacter spp.), but also other <u>Enteropacteriaceae</u> not normally included with the "coliform" bacteria, such as E. agglomerans, E. hafniae, $\underline{\kappa}$. ozaenae and \underline{s} . lique giens. As a result, the classification of the organisms typically expected in the coliform count (i.e. lactose fermenting <u>Enterobacteriaceae</u>) needs to be expanded to include the lactose fermenting strains of the organisms listed above. On the other hand, the bile precipitating count on VRBA does not represent all of the "coliforms" because of false negative (atypical) coliforms.

Since the tal count on VRFA estimates Enterobact-

erraceae and the bile precipitating count estimates lactore fermenting Enterobacteriaceae, the use of the differential counts on VRBA might be preferred because they give more information than Mossel's VRBG medium.

In the MPN technique, the statistically significant differences between the counts did not represent a large, practical difference. There was only a 4-fold difference between the means of the completed (EMB) and the presumptive (LST) counts. None-the-less, the LST count estimates total Enterobacteriaceae, whereas the EMB count estimates "coliform" bacteria, and is the equivalent of the bile precipitating count on VRBA. Goepfert 1976) in his study of the microbiclogy of raw ground beef follows Hajna and Perry's (1943) proposal of using LST counts to estimate coliforms. Many studies have indicated that false possives are include: in the LST count and the results of this study corroborate this. The fact that the LST count was statistically similar to the <u>Enterobacteriaceae</u> (total VFFA and bile precipitating VRFG) counts, and statistically different from the completed coliform (EMB) count, suggests either the occurrence of false positives in LST or inhibition causing false negatives in BGB, and subsequently on EMB.

In this study, 18% of lactose negative organisms in Phenol Red Lactose broth produced gas in LST. This might be expected because LST promotes lactose fermentation by slow stose fermenters and anaerogenic coliforms (Mallman and

Darby, 1941). However, the organisms growing in 45T included a large number of non-typical "coliforms", including some strains of E. agglomerans and 3. liquetaging, yet the majority of these organisms do not produce das in LST. E. igglomerans was softom identified to an MS+ of NU+ isolate from EMB, despite the fact that some E. agglomerans isolates on re-pliting, onto EMB were shown to produce light coloured colonies with a faint metallic shoes. E. liquetaging and S. liquetaging were observed primarily as NU- colonies on EMB, we as a meconiform Enterological actually of the types of organism producing the in-terminal producing the in-the broth would be necessary.

The elevated temperature test in EC broth is used to indicate faecal <u>E. coli</u> or <u>E. coli</u> type I. Some workers recommend direct inoculation of the meat homogenate into EC broth instead of inoculation (indirect), from the presumptive coliform test. The results of this study indicated that there was a significant difference between the direct and indirect methods, in which the indirect method gave results 2-fold greater than the direct method. In relation to the proposed <u>E. coli</u> standard for ground beef (Pivnick <u>et al.</u>, 1975), this represents a major difference in the result, and the recommendation of the Health Protection Branch for this test must be clearly specified.

The elevated temperature test may be carried out at different temperatures. The most common temperatures in use

The MEN technique outlined by the Health Protection

Branch, used in conjunction with the IMViC tests, is

reliable for detecting faecal <u>E. coli</u> in meats, with only a

limited number of false negative results. But the MPN

technique is time consuming, and while suitable for

regulatory purposes, it is totally unsatisfactory for

quality control purposes. Based on the correlation

coefficients between the EC counts and the other enumerating

or estimating techniques used in this study, none of the

other tests were capable of giving a reliable estimate of

terms of this standard.

faccal <u>F. goli</u>. It seems unlikely that a reliable, quick test could be obtained because the brochemical characteristics of β_{+} <u>coli</u> are very closely related to <u>Qther Enterobacteriaceae</u> (Bergey's Annual, 1974).

As a result, the use of VRIA and direct or indirect EC counts at 45.50c may be considered as routine methods to estimate <u>Enterobacteriaceae</u>, coliform bacteri and faecal <u>E</u>. <u>Coli</u>. The deficiencies in this proposal should be apparent from the results of this study, none-the-loss this recommendation is made because result; would generally be available in .4h (except for gas negative EC tubes which should be incupated for a further 24h to include delayed gas producers).

Using VERA and the direct or indirect EC test gives the following information:

- 1. It the hile precipitating count (coliform bacteria) is less than 1000 per gram, then the sample will mose likely fit within the proposed standards of faecal \underline{E} . \underline{coli} .
- 2. If the gas positive tubes by direct EC at 45.5°C are below 50 per gram and not greater than 250 per gram the sample will probably fall within the lower and upper faecal E. CC21 limits of the proposed standard, respectively.

For 113 samples with VRFA counts <1,000 per gram, 72.3% were <100 faecal <u>E. coli</u> per gram, 93.3% were <500 per "am, and only 6.7% were >500 per gram. Por 44 samples with VRFA counts >1000 per gram, 52.3% were <100 faecal <u>E. coli</u> per gram, 77.3% were <500 per gram and 22.7% were >500 per gram.

are below 100 per gram and not greater than 500 per gram the sample will probably fall within the lower and up, I limits of the proposed standard, respectively.

4. The total VRBA count could be used as an indicator of sanitary handling and proper temperature control of the meats.

temperature for 24:2h, to limit growth of non-<u>Enterobact</u>eriaceae, non-bile precipitating colonies (8-) on the
plates./If 45.5°C is selected for the EC tests, fewer false
positive results would occur, but false negatives would be
observed at 24h. Extended incubation, for an additional 24h,
reduces the false negative results. Conversely, incubation
at 44.5°C reduces the need for extended incubation but false
positive results will increase.

The use of <u>E</u>. <u>coll</u> as an indicator of faecal contamination of ground beef has been criticized by several workers (Hill, 1975; Goepfert, 1976). They suggested that contaminating <u>E</u>. <u>coll</u> could come from the intestine or hide of animals at slaughter. Furthermore, it is considered that <u>E</u>. <u>coll</u> could enter the food indirectly, from equipment. Goepfert (1976) claimed that <u>E</u>. <u>coll</u> thrive both in and out of the intestine. This contradicts the established premise of water tacteriologists that <u>E</u>. <u>coll</u> do not survive well in

an unfavourable environment. however, there can be little doubt that <u>E</u>. <u>coll</u> murvive and grow, even at relatively low temperatures in ground meats (Ai-Delaimy and Stiles, 1975). The purpose of identifying <u>E</u>. <u>coll</u> in foods is to indicate the possibility that intestinal pathogens might also be present. We such, the presence of <u>E</u>. <u>coll</u> of faecal origin in a food, by direct or indirect contamination, could indicate that intestinal pathogens are also present.

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Mossel and Co-workers (Buttraux and Mossel, 1961; Drion and Mossel, 1971; Mossel, 1957; Mossel et al., 1962; Mossel et al., 1963) proposed he use of Enterobacteriaceae as indicators or sanitation and hygiene in foods. Not many of the other faecal Enterobacteriaceae referred to by Mossel, such as Proteus spp. and Klebsiella spp. (IMVIC type ++++), were identified among the isolates of this study. However, a large number of E. agglomerans and S. liquefaciens were detected in this study. E. agglomerans was previously Erwinia sip. of the Herbicola-Lathyri bacteria. With its description by Ewing and Fife (1972), the origin of this organism is described as "from a wide variety of environmental sources and from various kinds of specimens from man and lower animals." It is described as an "opportunist" organism that may cause human infections

¹ Suspected <u>Salmonellae</u> obtained from Selenite Cystine enrichment, streaked onto Bismuth Sulfite and Brilliant Green agars, that were H S posit /e and lactose negative on TSI slants, were frequently shown to be <u>Proteus</u> spp., indicating that they were present in low numbers.

especially in the debilitated or young. The presence of E. agglometable in a food, therefore, indicates light of sanitation and/or hygiene. On the other hand, S. Lingefaciens is reported as being found primarily in food, and it implications in cols are not indicated (Ewing et al., 1973).

other <u>Enteropacteriaceae</u>, should be further studied to ascertain the importance of the total VRBA count in assessing sanitation and hygienic handling of media. This is particularly important because the coliforms were significantly less than <u>Enteropacteriaceae</u> in meats, and the total VRBA count might be a better parameter for use in quality control. For the reliable interpretation of the significance of other <u>Enteropacteriaceae</u> in quality control.

more information on the growth characteristics of these organisms is required.

II. Classification of Colifor Bacteria

Lactose fermentation is used as a screening test for coliform bacteria. However, atyp: 1. liform (non-lactose fermenters or anaerogenic) bacter. The observed in this study. In the case of E. coli I, there were not many atypical strains (approximately 5% of 296 isolates). In contrast, the Enteropacter spp. (cloacae and aerogenes) usually considered as coliform bacteria had much larger

numbers of atypical strains (180% of 90 isolates). Last one and the sole basis of classifying colliform bacteria results in many organisms expected to be included in the colliform count, being omitted. The use of lactore fermentation as the criterion for colliform bacteria became further complicated by the fact that 5. liquidiaciens and E. anglomerans are variable lactose fermenters (Ewing and F.fe, 1972; Ewing it

IMVIC tests were established for subdividing coliform bacteria into groups. This study revealed that the IMVIC grouping was specific for <u>E</u>. <u>coli</u>, but not for other coliform bacteria. The non-lactose fermenting <u>Enterobact-Eriaceae</u>, typed by IMVIC tests, were also found to be distributed across all IMVIC types. Because of this, atypical coliforms cannot be classified by their IMVIC reactions. Furthermore, the elevated temperature test data used in conjunction with the IMVIC test data, refers only to the IMVIC types ++-- or -+--, and most, or all, of these organisms are <u>E</u>. <u>coli</u>, especially if 45.5%C was used as the incubating temperature, instead of 44.5°C. As a result, the classification scheme proposed by the Coli-Aerogenes Subcommittee (Report, 1956) is only useful for screening out <u>E</u>.

The Ccli-Aerogenes Sub-committee (Report, 1956) also proposed the use of gelatin liquefaction in the classification of coliform bacteria. Gelatin liquefaction

could and not et value and it tinguishing reteem to compagning and R. Close is, theret is, it was not need in this istudy. The uses of an affitional took such is geliting. In quetaction expands the tests used to classify the collisions to the point where the reolites within as well to identified by enterplaceing agentation but a

ili. Minitek liğeniztication (Meghnigo)

Enterotate and k/b system, without supplementary tests.

The modified Minitor System used in this study enabled a further 50c cultures to be identified from the book. Encise II system, indicating that the modified Minitok technique was more increase than Enterotule. Based on the overall identification data it is recommended that for identification of <u>Enterotacteriaceae</u> from means the following tests should be carried out:

- . Acid and gas from glucose and lactose
- 2. Acid from arabinose, cellibiose, dulc.tol. inositol, raftinose, rhamnose, and callein.

. .

- or Utilization of citiate as a sole source of carbon
- s 4. From the or sectyl methyl carbinel (Venice = ${\rm From} {\sf Kauer}({\bf 1}, {\bf action})$
 - or Production of Typine and ornithing decarboxylands.
 - to liberation of promylatinance deaminars
 - I. Production of "Galactomaga" (ONEG to, t)
 - H. Production of undam
 - A. Production of a board indole
- 16. Growth on Priggs Sugar from (281) clants.
- 11. Motility fout.

All of these tests, except the 1st slints and tetrlity test and heddines out using Minitox discs. However, the firmitations experienced with detection of gas production from glucese and lactose, the indefinite colour changes on uses discs; and the lack of agreement between results on the Hyd Minitox disc? compared to HyS production on TSI slants, caused conventional ture media to be used for these tests in this study. Although the 1st slant is capable of giving acid and gas from glucose, and acid from lactose, in addition to HyS production, the lack of agr — petween TSI glucose and lactose results and the re in the conventional tube techniques prompted the use of — chal tube methods as the reference for this study.

I Subsequent to this study, a new urea disc was developed by FBL Minitex to give a more clearly defined colour change in $\dot{}$ this test.

[?] Subsequent to this study, a new $\rm H_2S$ disc was developed by BED Minitek to give improved performance in this test.

For the non-results, the difference between Minitok and a I methods might be subspected state most tescriptions of the production are based on TSI clants (Bergey's Manual, 1970), and Vausan and bevine (1900) reported that again had a marked influence on his production. The ures a test in Christensen's urea profit (Edwards and Evine, 1977) revealed many slow means producers that required up to 7 days for a positive reaction. Slow urease production could account for the difficulty experienced with the Minitek urea test.

Encise II system. The Roche Encise II system uses serological typing or Salmonella, however this was not used in this study and a NPG positive tests were used to screen out <u>Salmonella</u>. The proposed tests for identification of <u>Enterobacteriaceae</u> from meats are outlined below:

Miniter discs

Phenylalanine CNPG
Citrate
Malchate
Indole
Lysine
Crnithine
Arabinose
Collibiose
Dulcitol
Inositol
Raffinose
Rhamnose
Salicin

Conventional Media ,

Glucose Lactose MP-VP TSI Ur•a Motility

if other studies indicate a different range of <u>Enterobact</u>—
<u>eliadeae</u>, other brochemical tests might is necessary.

Minitek allows this flexibility. In this study, the Roche
Encirce II system for identification of <u>Enterobacteriaceae</u>
was used, however it would appear that a computerized system
appropriate or the selected modified Minitek system should
be developed.

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1. Statistically significant differences were observed between the different $\underline{Enterobacteriaceae}$, coliform and \underline{E} . \underline{col}_1 counts obtained on the ground meats.

- Enterobacteriaceae, however nome atypical Enterobacteriaceae that failed to prepicipitate the bile salts were observed on this medium.
- 3. The total count on VERA and the bile precipitating count on VRBG were statistically similar, and estimate the total Enterobacteriaceae count.
- 4. The bile precipitating count on VRBA represents typical coliform bacteria, hence using VRBA total and bile precipitating counts gives more information than enumerating Enterobacteriaceae on VRBG.
- 5. Although there was no statistical difference between coliforms enumerated on VRBA and EMB, the organisms comprising the counts were different. Hence the lack of statistical difference might be fortuitous.
- 6. The direct plating techniques could not be used to estimate \underline{E} . \underline{coli} counts.

7. To confirm \underline{E} , \underline{coli} I, EC at elevated temperature (45.50C) should be used to screen \underline{E} , \underline{Coli} , \underline{VRBA} and \underline{EC} tests which be used to give more rapid, less laborious with type accurance information.

(

8. <u>E. coli</u> I was isolated from most samples in this study and represented 29% of the isolates. <u>E. addlomerans</u> and <u>S. liquetaciens</u> were frequently isolated and represented 21 and 16% of the isolate, respectively. However, the significance of these organisms in ground meats is not known.

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APPENDIX A. Identification of 10 Known and 7 Unknown <u>Enterobacteriaceae</u> Cultures using Different Identification Systems

Cultures	Conventional Enterotube technique & Encise III		Minitek technique
<u>S. flexneri</u>	<u>Shiqella</u>	#2040	<u>Shiqella</u>
r- pneunoniae	K. <u>Pneumoniae</u>	#3423	<u>Klebsiella</u>
E. coli	<u>Escherichia</u>	#2060	Escherichia
Unknown	S. <u>liquefaciens</u>	#3200	<u>Serratia</u>
E. <u>hafniae</u>	E. <u>hafniae</u>	#3600	<u>E. hafniae</u>
P. vulgaris	P. mirabilis	P. milabilis	P. bilis
Unknown	<u>Escherichia</u>	#2060	Escherichia
C. freundii	C. <u>freundii</u>	#3240	Escherichia
S. typhimurium	<u>Salmonella</u>	#3311	E. <u>cloacae</u>
Unknown	<u>Escherichia</u> Shiqella	# 3060 .	<u>Shigella</u>
Unknown	E. cloacae	#3020	<u>Shiqella</u>
Unknown	Unidentifiable	#2070	Shiqella
K. Pneumoniae	<u>Klebsiella</u>	#2000	<u>Klebsiella</u>
Unknown	Unidentifiable	#2070	Citrobacter
Inknown	K. pneumoniae	K. <u>Pneumoniae</u>	<u>Klebsiellä</u>
S. marcescens S.	<u>marcescens</u>	#3600	Serratia
	E. aerogenes	E. aerogenes	Arizona
		·	

¹ See addendum for probable identity of organisms, associated with these I.D. numbers in Encise II.

APPENDIX B. Results for Identification of ATCC Type Cultures

C u	ltures	Identity of	t cultures	Modified Minitek
<u>K</u> .	pneumoniae	#2425	#3427	K. pneumoniae
<u>K</u> .	<u>ozaenae</u>	#2007	#2403	<u>K. ozaenae</u>
<u>K</u> .	rhinoscleromatis	#2000	#2000	K. rhinoscleromatis
<u>E</u> .	<u>çolı</u>	#2620	#3630	E. coli
<u>c</u> .	<u>freundii</u>	#2021	#2122	C. freundii
<u>c</u> .	<u>intermedius</u>	#3021	#3101	C. <u>intermedius</u>
<u>s</u> .	<u>cholerae-suis</u>	# 250 1	#32 1 3	<u>S. cholerae-suis</u>
<u>s</u> .	typhimurium	#3601	# 3 7 0 1	S. typhimurium
<u>s</u> .	<u>flexmeri</u>	#2000	#2000	Unidentifiable
<u>P</u> .	<u>vulgaris</u>	#2105	#2147	P. vulgaris
<u>p</u> .	<u>morganii</u>	#2201	#2207	P. mirabilis
<u>E</u> .	<u>tarda</u>	#2600	#3740	E. tarda
<u>s</u> .	marçescens	#2601	#2603	S. marcescens
<u>E</u> .	cloacae	#3 22 1	#3221	E. cloacae
<u>E</u> .	<u>aerogenes</u>	#3601	#3621	E. <u>aerogenes</u>
∄. 3 7° 30°	•	#2600 #260 1	#3600 #3602	E. <u>aerogenes</u> E. <u>aerogenes</u>
P. 37° 30°	=	#2005 #2005	#2006 #2006	P. <u>inconstans</u> A P. <u>inconstans</u>
<u>Y</u> .	<u>enterocolitica</u>	# 220 1		<u>Shiqella</u> or <u>Y. enterocolitica</u>

I A = Enterotybe result from Enterobacteriaceae laboratory, University of Alberta, Edmonton. The numbers are the identity number from the Roche Encise II (see Addendum).

B = Enterotube result from researcher's laboratory. The numbers are the identity number from the Roche Encise II (see Addendum).

Encise II Identity #	Organism (s)	Reaction score
2000	identity not well defined	
2005	Anaerogenic <u>E. agglomerans</u> <u>P. vugaris</u>	0.9999 0.0001
2006	Anaerogenic <u>E. agglomerans</u> <u>P. rettgeri</u> <u>P. vulgaris</u>	0.8860 0.0822 0.0318
2007	Anaeroyenic <u>E. agglomerans</u> <u>P. rettgeri</u> <u>P. vulgaris</u>	0.5103 0.4887 0.0010
∠0 20	identity not well defined	
2021	S. <u>rubidaea</u> Anaerogenic <u>E</u> . <u>aqqlomerans</u> <u>K</u> . <u>ozaenae</u> <u>C</u> . <u>freundii</u>	0.6699 0.2729 0.0555 0.0016
2040	identity not well defined	
2060	identity not well defined	
2070	Alkalescens Dispar <u>Escherichia</u> <u>C. freundii</u>	0.9861 0.0139 0.0001
:105	P. vulgaris	1.0000
122	C. freundii	1.0000
147	P. vulgaris	1.0000
201	identity not well defined	
20 7	P. mirabilis	1.0000
343	C. freundii	1.0000
403	K. ozaenaeE. aerogenes	0.9989 0.0011
125	unidentifiable	
600	identity not well defined	

2603	S. marcescensS. liquefaciensK. ozaenaeE. aerogenes	0.9451 0.0516 0.0020 0.0013
2620	E. <u>aerogenes</u> S. <u>liquefaciens</u> K. <u>ozaenae</u> Escherickia	0.4166 0.2902 0.1817 0.1114
3020	identity not well defined	
3021	identity not well defined	1
3060	Escherichia Aerogenic E. agglomerans C. freundii	0.5436 0.4558 0.0006
3 10 1	<u>C. freundii</u> <u>Salmonella</u> spp. H ₂ S +v e <u>Escherichia</u>	0.9906 0.0094 0.0001
3200	identity not well defined .	
3213	<pre>E. cloacae C. freundii</pre>	0.6660 0.3340
3221	E. cloacae S. liquefaciens C. freundii K. ozaenae	0.6708 0.3116 0.0099 0.0078
3240	Escherichia C. freundii	0.9970 0.0030
3311	<u>Salmonella</u> <u>C. freundii</u> H ₂ S +ve <u>Escherichia</u>	0.7140 0.2859 0.0001
3423	 K. pneumoniae S. rubidaea K. ozaenae E. aerogenes 	0.9615 0.0258 0.0072 0.0055
3427	unidentifiable	
3600	identity not well defined	
3601	identity not well defined	
3602	identity not well defined	

3621	E. delogenes S. liquefaciens E. hafniae K. ozaenae	0.8564 0.0813 0.0317 0.0006
3630	Escherichia E. aerogenes	0.7494 0.2506
3701	identity not well defined	
3740	<u>Edwardsiella</u> H ₂ S +v e <u>Escherichia</u>	0.9821 0.0179
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