

Biochemical identification of bacteria

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Outline

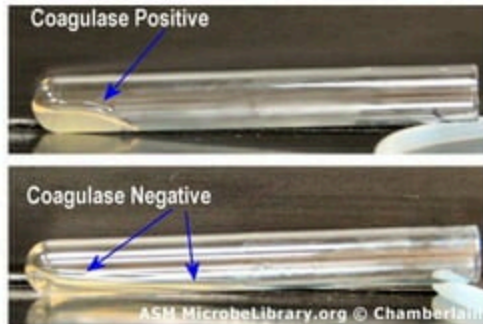
- Phenotypic vs genotypic tests
- Pros and cons of biochemical tests
- Basis of biochemical tests
- Examples of biochemical test
- Diagnostic algorithms
- The future of biochemical identification tests

Methods of bacterial ID

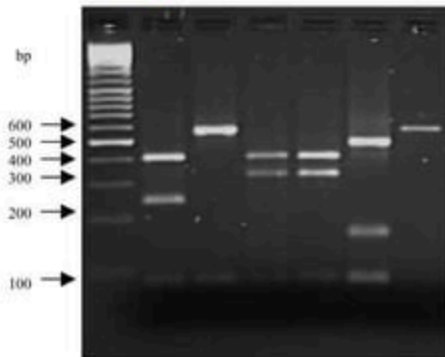
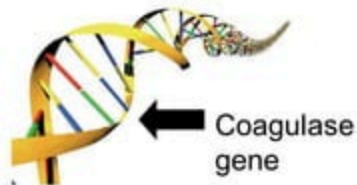
- Phenotypic
 - Detects the physical properties of bacteria
 - Influenced by gene expression
 - Includes biochemical tests
- Genotypic
 - Detects the genetic code of bacteria (DNA)
 - Not influenced by gene expression

Eg coagulase for staphylococcal ID

- Phenotypic test



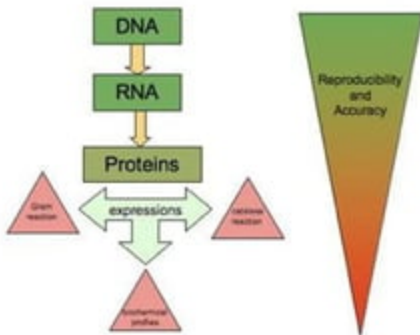
- Genotypic test



Biochemical ID:

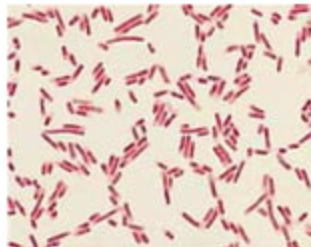
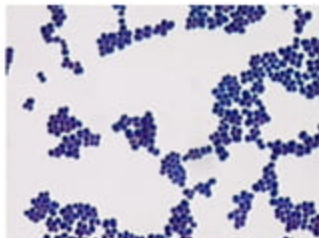
Pros and cons

- Pros
 - Cheap
 - Experience with use++
 - Does not require expertise
 - Potentially fast TAT (range: seconds to overnight)
- Cons
 - Biosafety risk (live organisms)
 - Less accurate, less discriminatory
 - Phenotype may be unstable
 - Eg inducible (ie influenced by gene expression)
 - Not possible if organism is slow growing or fastidious
 - Subjective interpretation (less reproducible)



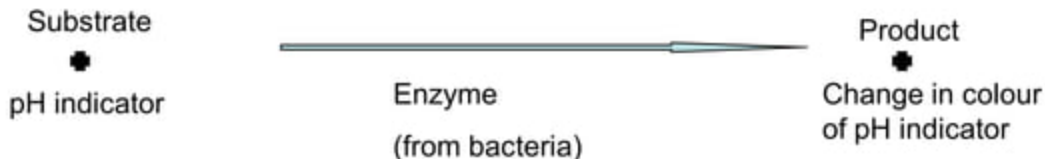
Type of phenotypic ID

- Appearance
 - Macroscopic
 - Microscopic (eg gram stain, rod vs coccus)
- Growth requirement/rate
 - Media
 - Atmospheric gases
 - Temperature
- Smell
- Motility
- Hemolysis on blood agar
- **Biochemical tests**



(See lecture on “Culture characteristics for bacterial identification”)

Basis of biochemical tests



- Important features
 - Standardisation of method
 - standardised amount of bacteria used for test (=inoculum)
 - +ve and -ve controls

pH indicators



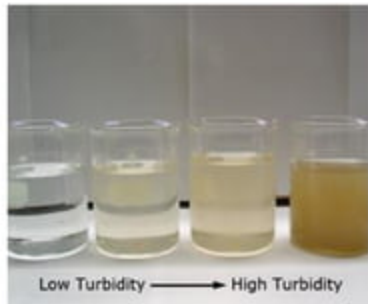
FIGURE 62.4 Glucose test. Tube on the left is positive (pink); tube on the right is negative (yellow). © The McGraw-Hill Companies/Robert H. Langlois/Photographic Services

- Colour changes occur at different pHs for different indicators

pH Indicator	pH range	Change from acid to alkaline
• Methyl red	4-6	red to yellow
• Andrades	5-8	pink to yellow
• Bromescol blue	5-6	yellow to purple
• Phenol red	6-8	yellow to red

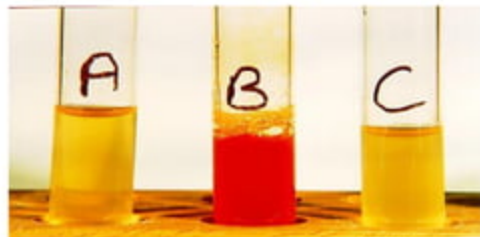
Standardisation of the inoculum

- Examples of solid phase:
 - Loop size (eg 1microL, 10microL)
- Examples of liquid phase
 - Turbidity of fluid
 - The ability of particles in suspension to refract and deflect light rays
 - Optical density
 - Nephelometry



Positive and Negative controls

- Positive control: bacteria with known +ve test result
- Negative control: bacteria with known -ve test result
- If either or both of the controls fail, then the test is not valid



-ve
control

+ve
control

test
isolate



Types of biochemical ID methods

- Manual vs automated
 - Automated systems have the advantage of automated reading which improves speed, consistency and removes subjective error.
- In house vs commercial



Examples of common biochemical tests used for ID of gram negative bacteria

- Urease
- Indole
- Oxidase
- Glucose fermentation
- Lactose fermentation
- Nitrate

Urease

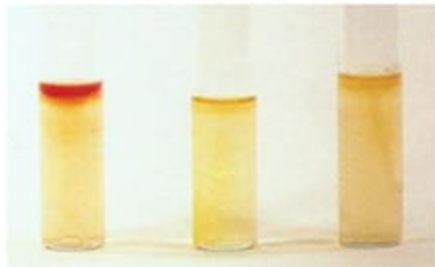
- Detects hydrolysis of urea to ammonia by urease enzyme
- Ammonia causes an increase in pH which is detected by the pH indicator (orange → pink)
- Urease +ve bacteria:
 - Proteus
 - Klebsiella



FIGURE 42.4 Urease test. Tube on the left is positive (Proteus); tube on the right is negative. © The McGraw-Hill Companies, Inc.

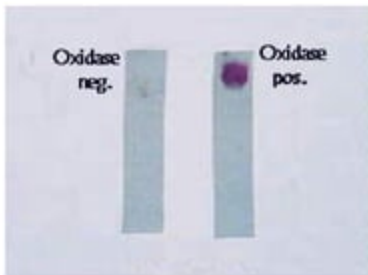
Indole

- Detects indole production from tryptophan, which produces a colour change in combination with dimethylaminobenzaldehyde (clear to red)
- Indole +ve bacteria:
 - E.coli
 - Citrobacter



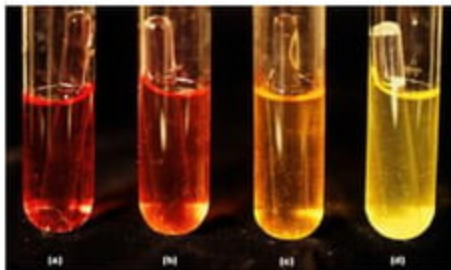
Oxidase

- Detects cytochrome oxidase enzyme that converts dimethylphenyldiamine to indophenol blue (clear to blue)
- Oxidase +ve bacteria:
 - Pseudomonas
 - Vibrio



Glucose fermentation

- Detects ability of bacteria to ferment glucose to pyruvic acid using the Embden Meyerhof pathway
- Detected by phenol red pH indicator (red/alkaline to yellow/acid)
- Bacteria that ferment glucose:
 - E.coli
 - Proteus



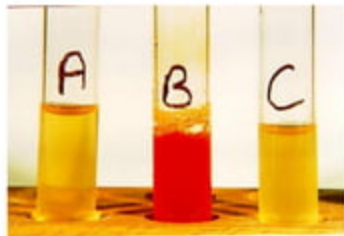
Lactose fermentation

- Detects ability of bacteria to ferment lactose to glucose then to pyruvic acid using the Embden Meyerhof pathway
- Detected by phenol red pH indicator (red/alkaline to yellow/acid)
- Bacteria that ferment glucose:
 - E.coli
 - Klebsiella



Nitrate

- Detects nitrate reductase enzyme which converts nitrate to nitrite.
- Nitrite then revealed by addition of naphthylamine and sulfinic acid to form diazonium dye (clear to red)
- Nitrate +ve bacteria:
 - E.coli
 - Klebsiella



TSI slope

- Incorporates multiple substrates and pH indicators into 1 tube
- By streaking bacteria onto surface and stabbing it into media, both aerobic and anaerobic conditions are generated



API

- Minutuarized biochemical reactions in >20 wells
- Takes 2-24 hrs
- Reaction profile (“biocode”) compared to an on-line database of >20000 isolates
- Commerical test



Tests	Active ingredients	Reaction enzymes
1. GMP	Carbonyl iron/2,4-diaminophenylhydrazine	Isolactonidase
2. ADH	Acetone	Acetone Dehydrogenase
3. LDC	L-lysine	L-lysine Decarboxylase
4. ODC	Oxidation state	Oxidation (Oxidation)
5. CIT	Citrate utilization	Citrate utilization
6. H ₂ S	Sulfide production	H ₂ S production
7. URE	Urea	Urease
8. TDA	Tryptophan	Tryptophan deaminase
9. IND	Indole	Indole production
10. VP	Voges-Proskauer	Acetoin production (Voges-Proskauer)
11. GEL	Gelatin	Gelatinase
12. GLU	Glucose	Fermentation (Glucose)
13. MAN	Mannitol	Fermentation (Mannitol)
14. INO	Inositol	Fermentation (Inositol)
15. MIB	Methyl	Fermentation (Methyl)
16. RBA	Ribitol	Fermentation (Ribitol)
17. SAC	Saccharose	Fermentation (Saccharose)
18. MEL	Mellicose	Fermentation (Mellicose)
19. AMY	Amylase	Fermentation (Amylase)
20. ARA	Arabinose	Fermentation (Arabinose)

Automated Biochemical ID systems

- Examples:

- Vitek
- Biolog
- Pheonix
- Autoscan Walkaway



- Varying capacity for:

- Number of specimens they can handle
- Size/extent of comparative database
- Interfacing with lab data program
- Turn around time
- Capacity for ID to species level



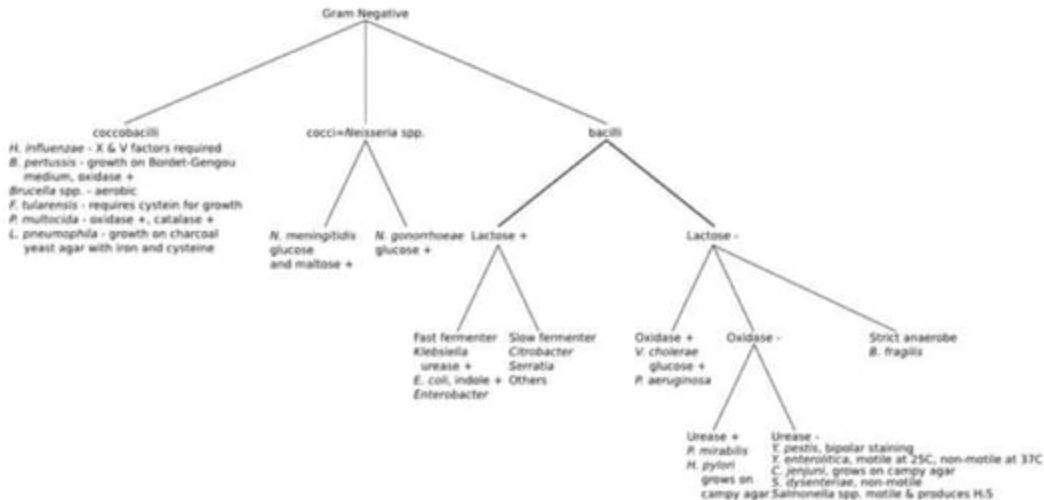
Diagnostic algorithms for bacterial ID

- Primary tests allow genus level ID (enterobacteriaceae, “non-glucose fermenters”, HACEK, etc)
 - Gram stain
 - Culture morphology
 - Basic biochemical tests
 - Eg Oxidase, indole, urease tests, etc
- Species level identification requires more complex, second line tests

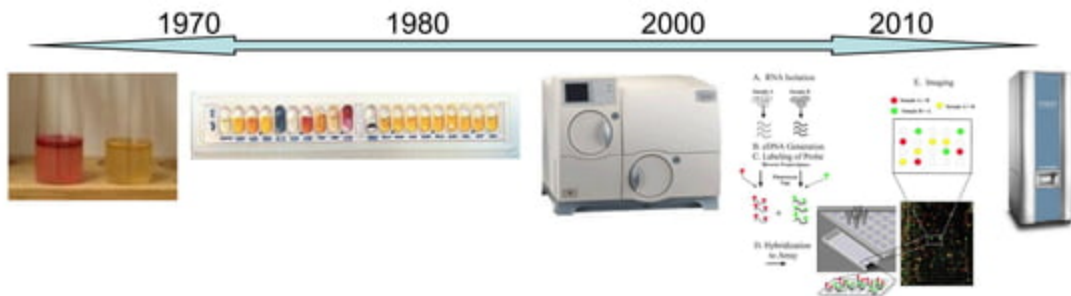
Example 1 of diagnostic algorithm

	Indole	Methyl red	Voges Proskauer	Citrate	Urease
E.coli	+	+	-	-	-
Enterobacter	-	-	+	+	-
Klebsiella pneumoniae	-	-	+	+	+
Salmonella	-	+	-	+	-
Shigella	-	+	-	-	-
Proteus mirabilis	-	+	-	+/-	+

Example 2 of diagnostic algorithm



Changes in biochemical tests for ID: past and future



- Increased automated and minituarisation
- Increasingly replaced by genotypic tests
- Is identification necessary: could we manage with susceptibility testing alone?

Conclusions

- Biochemical tests remain critical to bacterial identification
- Need to understand the principles of the common/primary tests
- Biochemical tests have limitations
- In the future they will increasingly be replaced by genotypic tests