



Plant Pathogenic Bacteria

A Basic Guide to Identification

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Module 1



Methods, processes and end-points [2]

- Phytosanitary testing
 - Pre-described test procedure for known crop / pest combinations only
 - Limited need for wider inclusion of other pest knowledge; can be achieved by non-specialists
 - Require specific, dedicated infrastructure
 - Known outcome, with statistical confidence
 - Primarily driven by EU directives

Methods, processes and end-points [3]

- Identification of unknowns
 - Receiving of plant / pest combinations of any type
 - Need to be inclusive of all pest types
[entomological, mycological, bacterial, viral]
 - Requires expert knowledge, infrastructure and access to reliable information resources
 - Driven by national demand for providing broad services in plant health as supports commercial interests

Methods, processes and end-points [4]

- Research
 - Responsive to demand
 - Need to be inclusive of all pest types [entomological, mycological, bacterial, viral]
 - Requires expert knowledge, infrastructure and access to reliable information resources
 - Driven by national demand for providing broad services in plant health as supports commercial interests

Methods for bacterial identification

- Biochemical tests – traditional methods
- Formatted biochemical tests
 - API strips
 - Biolog
- GC Fatty acid profiles – MIDI system
- Serological – immunological methods
- DNA methods
 - DNA homology
 - 16S rDNA
 - Fingerprinting

Biochemical tests

Biochemical tests – example 1

- *Pantoea stewartii*
 - Non-motile
 - Colonies do not show symplasmata or inclusion
 - Negative for production of H₂S from cysteine, acetone, phenylalanine deaminase, nitrate reductase and gelatinase
 - Acid is produced from melibiose; non-acid from dulcitol, maltose, rhamnose or starch

Biochemical tests – example 2

- *Ralstonia solanacearum*
 - Non-fluorescent pseudomonas with polar tuft flagella
 - Cells non-pigmented, but brown diffusible pigment often produced
 - PHB is accumulated
 - Levan not formed from sucrose
 - Gelatin hydrolysis weak
 - Starch and aesculin not hydrolysed
 - Nitrate reduced by nearly all strains; many produce gas [denitrifying]
 - No growth at 40C
 - Oxidase positive
 - Arginine dihydrolase negative
 - Most strains produce tyrosinase
 - Light or no growth in broth containing 2% NaCl; no growth at 40C
 - Carbon sources used for growth: acetate, aconitate, L. alanine, D-alanine, γ -aminobutyrate, asparagine, L-aspartate, benzoate, butyrate, citrate, fumarate, gluconate, D-glucose, L-glutamate, glycerol, L-histidine, β -hydroxybutyrate, α -ketoglutarate, L-malate, mucate. L-proline, proionate, pyruvate, saccharate, succinate, sucrose and trehalose

Dichotomous key – some key biochemical tests

- Gram test [Gram –ve and +ve bacterium]

- Gram –ve

- Anaerobic growth
- Yellow colonies on YDC
- Fluorescent pigment
- Urease
- Growth at 33C and 40C
- Growth on D1M agar
- Utilization of arginine and betaine

- Gram +ve

- Endospores formed
- Anaerobic growth
- Ariel mycelium

The LOPAT tests for fluorescent *Pseudomonads*

L	O	P	A	T	Group	Example
+	-	-	-	+	Ia	<i>P. syringae</i>
-	-	-	-	+	Ib	<i>P. savastanoi</i>
-	-	+	-	+	II	<i>P. viridiflava</i>
-	+	-	-	+	III	<i>P. cichorii</i>
+	+	+	+	-	IVa	<i>P. marginalis</i>
-	+	+	+	-	IVb	<i>P. fluorescens</i> complex
-	+	-	+	-	Va	<i>P. tolaasii</i>
+	+	-	+	-	Vb	<i>P. fluorescens</i> complex

Levan production / oxidase reaction / Potato Rot / Arginine dihydrolase production / tobacco hypersensitivity – LOPAT – p26 Phytobacteriology book

Biochemical tests

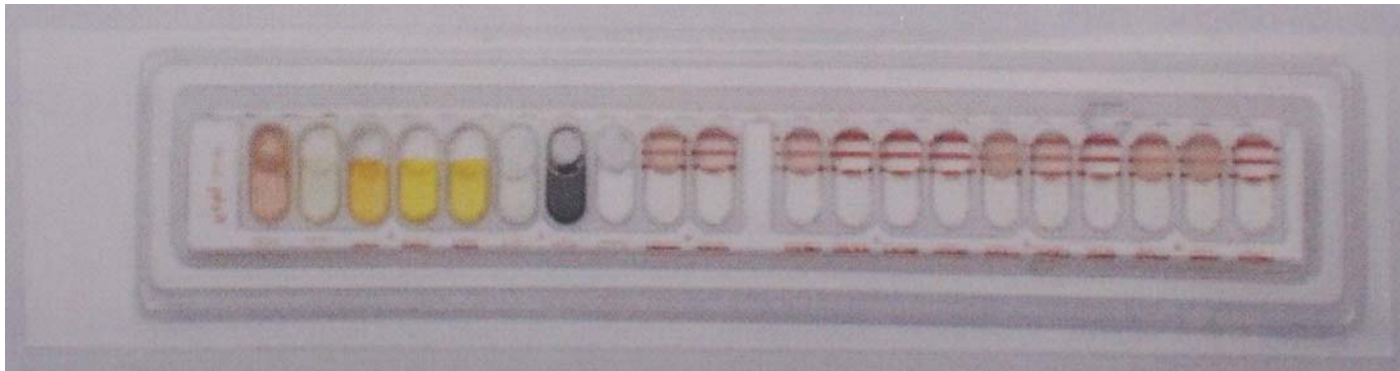
- Advantages
 - Is not requiring of expensive equipment and searchable data bases
- Limitations
 - Methods are many, some sequential and time consuming
 - Reagent list for tests is extensive and prepared media is to be aliquoted into many different formats
 - Many tests give variable strain specific results and some tests are unreliable
 - Technicians need to be very familiar with methods and competent in their use

Biochemical formatted platforms

- Takes the biochemical tests and places them on a more convenient format
- Two main commercial products
 - API strips — [<http://industry.biomerieux-usa.com/industry/food/api/index.htm>]
 - Biolog — [<http://www.biolog.com/main.html>]
- Results achieved within 48hrs
- Results [+ & -ve data] fed into library of described strains
- Similarity values on most likely identification
- Requires judgement over identifications presented

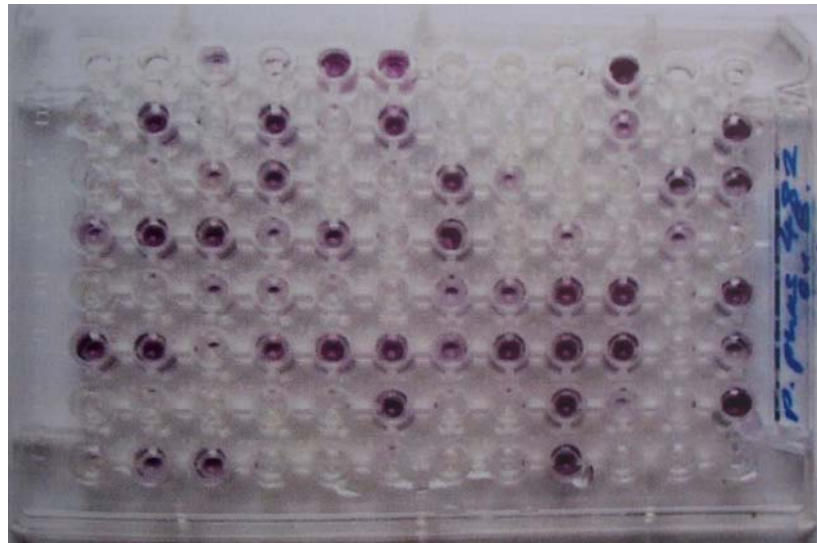


API



- Each well contains a different substrate
- Results are recorded as either a substrate colour change or as growth

The Biolog system



- The Biolog system presents an extended array of biochemical tests
- A positive result is seen as a purple colour change
- The plate can be read by eye or by a plate reader

A1 Water	A2 α -cyclodextrin	A3 dextrin	A4 glycogen	A5 tween 40	A6 tween 80	A7 N-acetyl-D-galactosamine	A8 N-acetyl-D-glucosamine	A9 adonitol	A10 L-arabinose	A11 D-arabitol	A12 cellobiose
B1 D-erythritol	B2 D-fructose	B3 L-fucose	B4 D-galactose	B5 gentiobiose	B6 α -D-glucose	B7 D-mannitol	B8 α -D-lactose	B9 lactulose	B10 maltose	B11 D-mannitol	B12 D-mannose
C1 D-melibiose	C2 β -methyl D-glucoside	C3 D-psicose	C4 D-raffinose	C5 L-rhamnose	C6 D-sorbitol	C7 sucrose	C8 D-trehalose	C9 turannose	C10 xylitol	C11 methyl pyruvate	C12 mono-methyl succinate
D1 acetic acid	D2 cis-aconitic acid	D3 citric acid	D4 formic acid	D5 D-galactonic acid lactone	D6 D-galacturonic acid	D7 D-gluconic acid	D8 D-glusaminic acid	D9 D-glucuronic acid	D10 α -hydroxy butyric acid	D11 β -hydroxy butyric acid	D12 γ -hydroxy butyric acid
E1 p-hydroxy phenylacetic acid	E2 itaconic acid	E3 α -keto butyric acid	E4 α -keto glutaric acid	E5 α -keto valeric acid	E6 D, L-lactic acid	E7 malonic acid	E8 propionic acid	E9 quinic acid	E10 D-saccharic acid	E11 sebacic acid	E12 succinic acid
F1 bromo succinic acid	F2 succinamic acid	F3 glucunoramide	F4 alaninamide	F5 D-alanine	F6 L-alanine	F7 L-alanyl-glycine	F8 L-asparagine	F9 L-aspartic acid	F10 L-glutamic acid	F11 glycyl L-aspartic acid	F12 glycyl L-glutamic acid
G1 L-histidine	G2 hydroxy L-proline	G3 L-leucine	G4 L-ornithine	G5 L-phenylalanine	G6 L-proline	G7 L-pyrogutamic acid	G8 D-serine	G9 L-serine	G10 L-threonine	G11 D, L-carnitine	G12 γ -amino butyric acid
H1 urocanic acid	H2 inosine	H3 uridine	H4 thymidine	H5 phenyl ethylamine	H6 putrescine	H7 2-amino ethanol	H8 2,3-butanediol	H9 glycerol	H10 D, L- α -glycerol phosphate	H11 glucose-1-phosphate	H12 glucose-6-phosphate

Biochemical formatted platforms [Biolog]

- Advantages

- Is not requiring of expensive equipment
- System is quick, reproducible and easy to perform
- Data can be shared between laboratories
- Can provide a reasonable identification to the genus and species level

- Limitations

- Requires investment [access] to the library
- Has limitation in resolving below species level [pathovar separation]
- Library stronger on human microbials than plant pathogenic bacteria

Fatty acid analysis – the Midi system

Fatty acids

- Gram-negatives
 - Unique hydroxy patterns
 - Some cyclopropanes
 - Few branched acids
- Gram positives
 - Many branched acids
 - Very few hydroxy and cyclopropane acids.

The Midi system

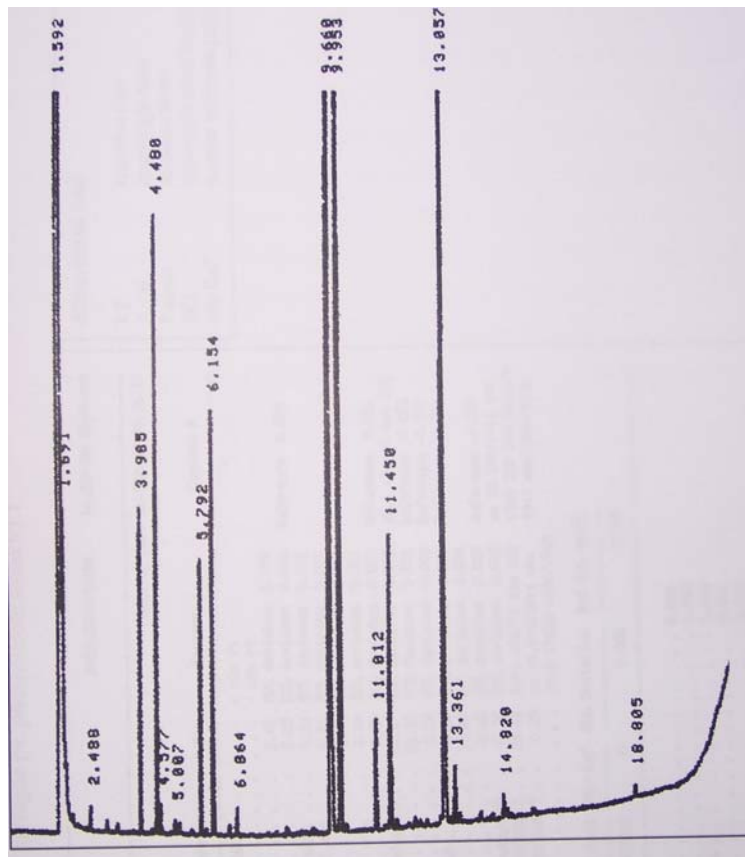
- Commercial and / or lab produced — [<http://www.midi-inc.com/>]
- Based on comparisons of types and amounts of acids
- Interrogates library for identification
- Typical “return”:

NCPPB Rev 3.0	Agrobacterium biovar 1	0.814
	Agrobacterium biovar 2	0.567
	Agrobacterium biovar 3 [vitis] ...	0.316

Fatty acid extraction process

- Culture Cells [i.e. 24hr on TSBA]
- Harvest Cells
- Saponify Lipids
- Methylate Fatty Acids [FAMEs]
- Extract and Purify
- GC analysis
- Comparison to library

MIDI system print outs



Sherlock Version: 3.10 DATA: E00915545A 16-SEP-00 09:06:00

ID: 2032 NM-P. PHAS. 95.1 Date of run: 16-SEP-00 08:35:27
 Bottle: 41 SAMPLE [TSBA40]

RT	Area	Ar/Ht	Respon	ECL	Name	%	Comment 1	Comment 2
1.604	309664512	0.031	...	7.021	SOLVENT PEAK	...	< min rt	
2.497	656	0.026	...	8.942	< min rt	
3.995	10872	0.032	1.053	11.424	10:0 3OH	2.41	ECL deviates 0.002	
4.490	23184	0.035	1.031	12.000	12:0	5.03	ECL deviates 0.000	Reference 0.001
4.587	1112	0.034	1.028	12.090	11:0 ISO 3OH	0.24	ECL deviates 0.001	
5.015	600	0.037	1.015	12.485	unknown 12.484	0.13	ECL deviates 0.001	
5.803	11480	0.038	0.996	13.177	12:0 2OH	2.40	ECL deviates 0.000	
6.164	18088	0.039	0.989	13.455	12:0 3OH	3.76	ECL deviates 0.001	
6.872	1216	0.038	0.978	14.000	14:0	0.25	ECL deviates -0.000	Reference 0.001
9.670	165296	0.046	0.954	15.819	Sum In Feature 3	33.17	ECL deviates -0.003	16:1 w7c/15 iso 2OH
9.963	141496	0.046	0.953	16.001	16:0	28.35	ECL deviates 0.001	Reference 0.002
11.021	6136	0.049	0.949	16.628	17:0 ISO	1.22	ECL deviates -0.002	Reference -0.001
11.460	16376	0.049	0.948	16.889	17:0 CYCLO	3.26	ECL deviates 0.001	Reference 0.001
13.068	94856	0.050	0.946	17.824	18:1 w7c	18.87	ECL deviates 0.001	
13.372	3184	0.050	0.946	18.000	18:0	0.63	ECL deviates -0.000	Reference -0.001
14.830	1384	0.051	0.945	18.848	Sum In Feature 7	0.28	ECL deviates 0.002	un 18.846/19:1 w6c
*****	165296	SUMMED FEATURE 3	33.17	16:1 w7c/15 iso 2OH	15:0 ISO 2OH/16:1 w7c
*****	1384	SUMMED FEATURE 7	0.28	un 18.846/19:1 w6c	19:1 w6c/.846/19cy
*****	19:0 CYCLO w10c/19w6	

Solvent Ar	Total Area	Named Area	% Named	Total Annt	Nbr Ref	ECL Deviation	Ref ECL Shift
309664512	495280	495280	100.00	425532	6	0.001	0.001

IMIBAC [Rev 1.0] * NO MATCH *

TSBA40 [Rev 4.10]	Pseudomonas	ECL
P. syringae	...	0.899
P. s. phaseolicola	...	0.899
P. s. syringae	...	0.841
P. s. glycinea	...	0.820

- GC trace [left]
- Library analysis [above]

Key acids from 4 genera

Acid	<i>Acidovorax</i>	<i>Ralstonia</i>	<i>Pseudomonas</i>	<i>Burkholderia</i>
10:0 3OH	+		+	+
12:0 2OH			+	
12:0 3OH			+	
14:0 3OH		+	+	+
16:0 2OH		+		+
16:0 3OH			+	+
16:1 2OH		+		
18:1 2OH		+		+

Fatty acid analysis

- Advantages

- System is quick, reproducible and easy to perform
- Data can be shared between laboratories
- Can provide a reasonable identification to the genus and species level

- Limitations

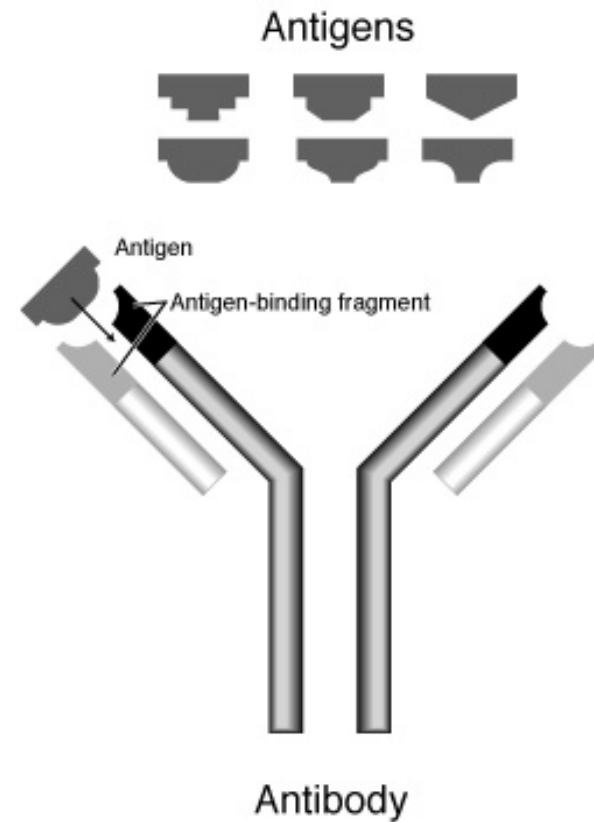
- Requires investment in GC equipment and MIDI library
- Has limitation in resolving below species level [pathovar separation]
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Serological approaches

Serological approaches

- Rapid
- Sensitive
- Specific
- Diagnose diseases



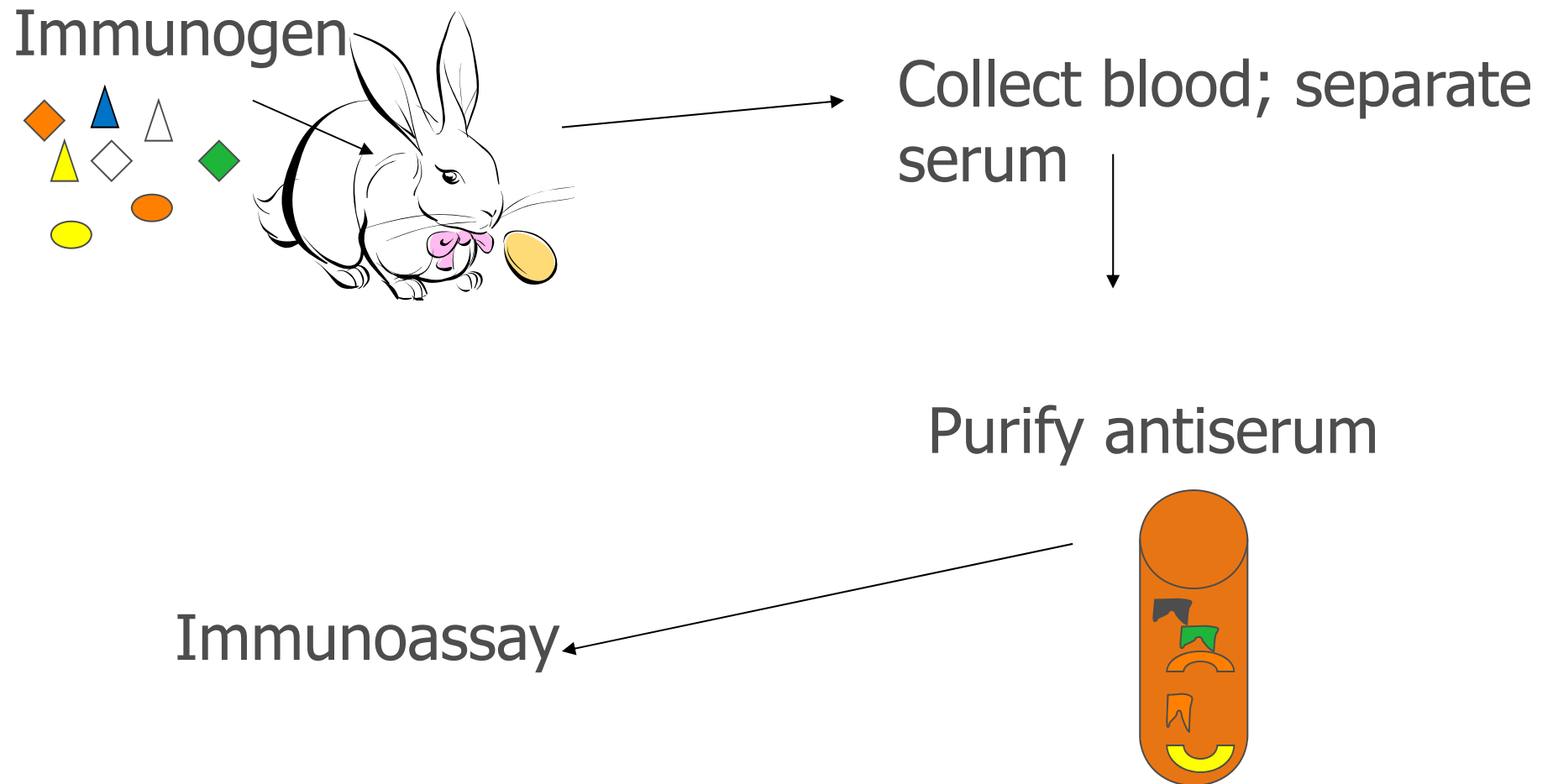
Immunoassays are based on antibodies....

- Mammals produce antibodies that specifically recognize binding sites (epitopes) on proteins, glycoproteins, lipopolysaccharides, carbohydrates (antigens)
 - Polyclonal antibodies
 - Monoclonal antibodies
- Antibodies specifically bind antigens
- Bound antibodies are detected with various markers

Polyclonal Antibodies

- Immunogens (preparations containing antigens that are used to immunize an animal)
 - Various degrees of purification of immunogens
 - Whole cells
 - Cell (surface) washings
 - Virus particles
 - Broken cells
 - Purified cell components
- Immunogens injected into animals for antibody production

Polyclonal antibody production



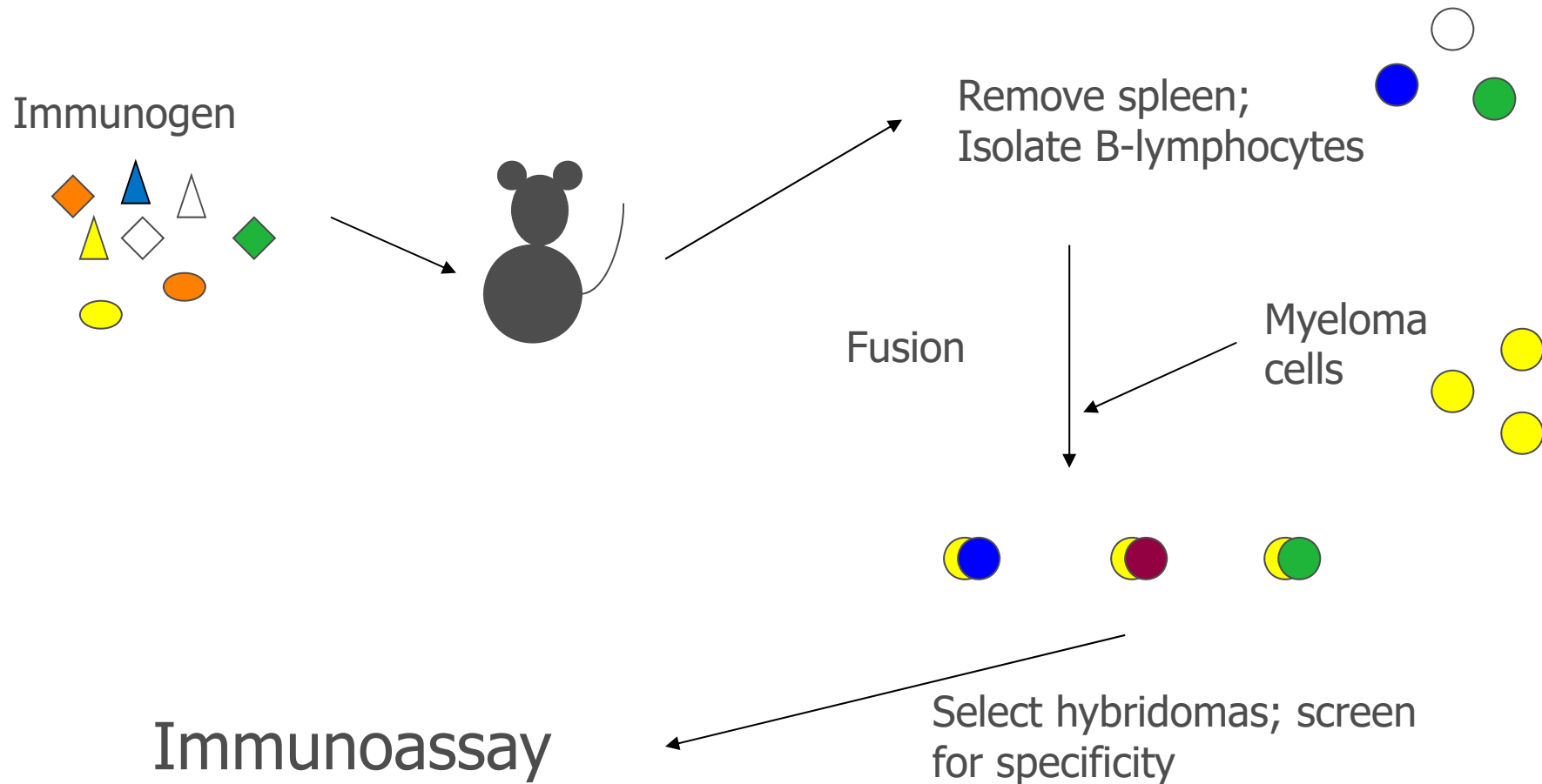
Characteristics of polyclonal antisera

- High sensitivity
- Varying specificity - depending on purity of immunogen/number of epitopes
- May vary from batch to batch

Monoclonal antibodies

- Single type of antibody
- Highly specific
 - Recognize single epitope
- Sensitivity varies
- Produced by hybridoma cell lines that are theoretically immortal

Monoclonal antibody production



Immunoassay formats

- Enzyme-linked immunosorbent assay (ELISA)
 - Enzyme conjugated to antibody = marker
 - Alkaline phosphatase
 - Peroxidase
- Lateral flow immunoassay
 - Ab-Ag binding occurs as mixture flows through solid phase in liquid
- Immunofluorescence
 - Fluorescent molecule marks Ab-Ag reaction

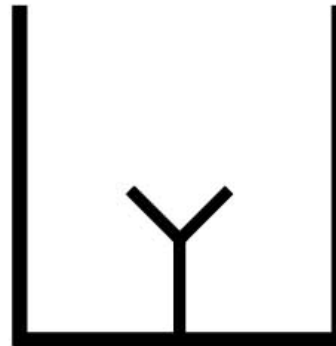
ELISA

- Positive reaction indicated by enzymatic reaction with chromogenic substrate = color change
 - Antigen capture/plate-trapped antigen
 - Antigen bound to solid phase
 - Indirect vs. direct
 - Direct = detecting antibody conjugated with enzyme
 - Indirect = enzyme conjugated to secondary antibody
 - Sandwich ELISA (double antibody, triple antibody)

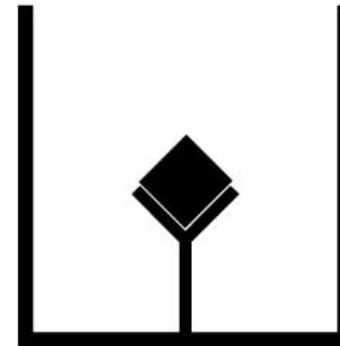
Direct, Double Ab Sandwich ELISA

DAS-ELISA

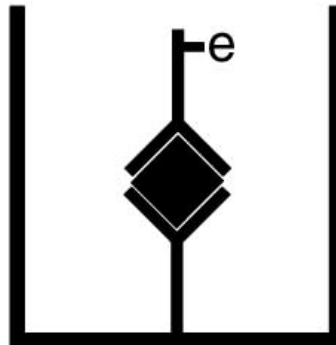
Step 1
Antigen-specific antibody is attached to a solid-phase surface



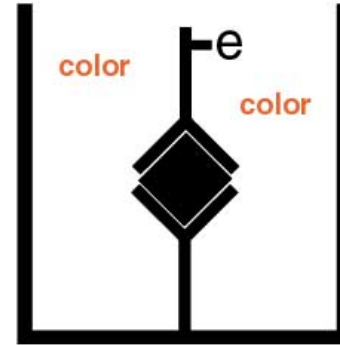
Step 2
Test specimen is added, which may or may not contain the antigen



Step 3
An enzyme-labeled antibody specific to the antigen is added (conjugate)



Step 4
Chromogenic substrate is added, which in the presence of the enzyme, changes color.

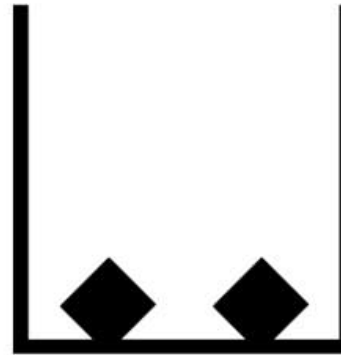


Indirect, Plate-trapped Antigen ELISA

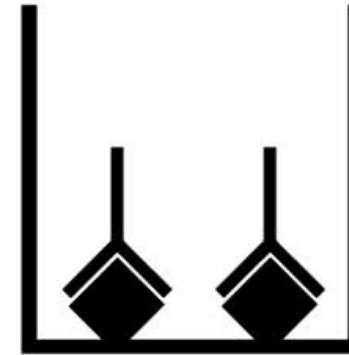
Primary Ab specific
to Antigen

Secondary Ab produced
in a different species,
e.g. goat - specific to
primary Ab

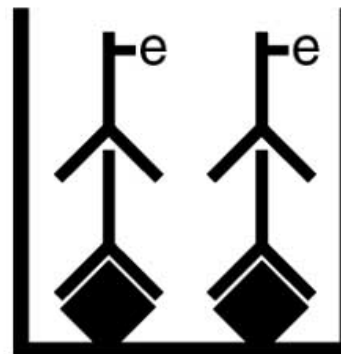
Step 1
Specific antigen is attached to
a solid-phase surface



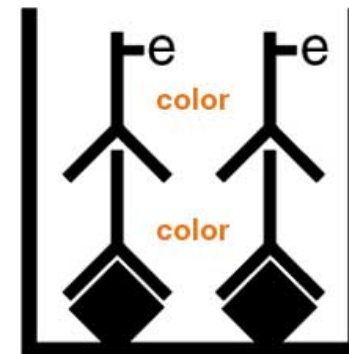
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An enzyme-labeled antibody
specific to the test antibody is
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Chromogenic substrate is added,
which in the presence of the
enzyme, changes color.



Multiwell Immunoassay



- Many commercially available
- Most detect various viruses
- Also for bacteria, fungi

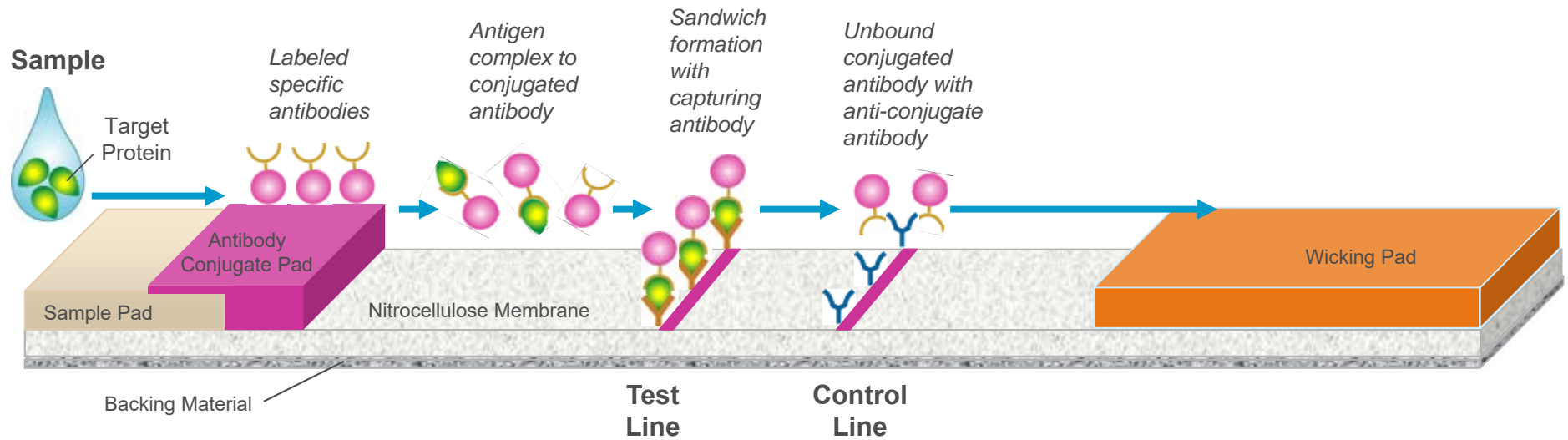
Immunostrip (Lateral Flow) Assays



- Very fast - 3-5 minutes
- Sensitive
- Some are available commercially
- Extracts diffuse through paper strips
- Marker may be gold microparticles

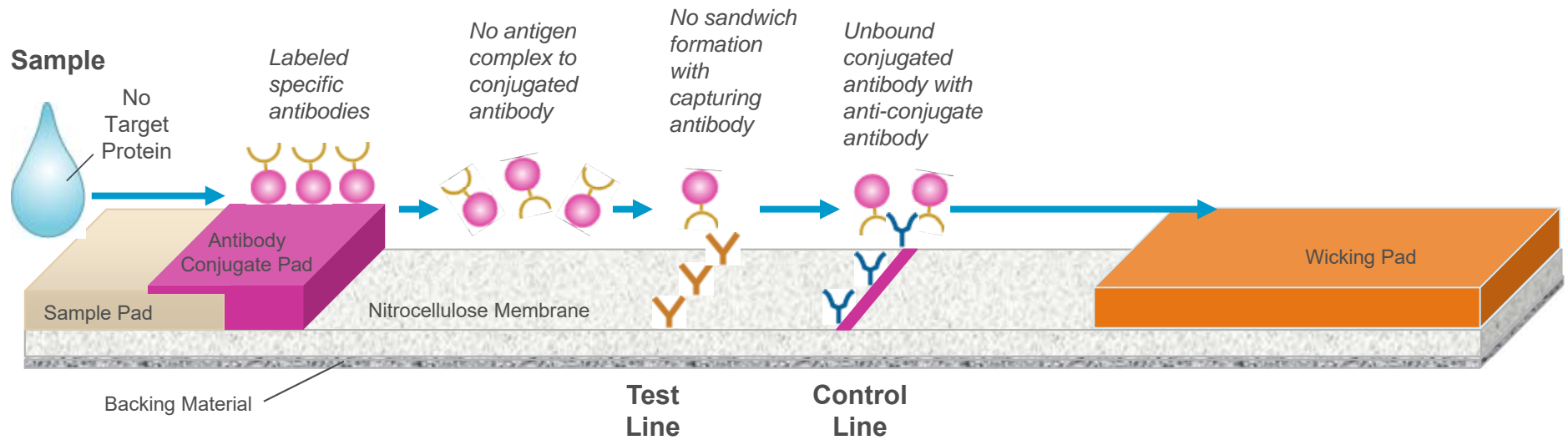
LFD Cross-Sectional View

Positive Result



LFD Cross-Sectional View

Negative Result



Environlogix, Inc.

DNA approaches

DNA sequencing

- 16s rDNA sequencing
 - One example: 27F and 1492R primers amplification followed by 518F and 800R primers
 - Stringent annealing conditions
 - BLAST search for nearest relatives
 - Assemblage of closest relatives

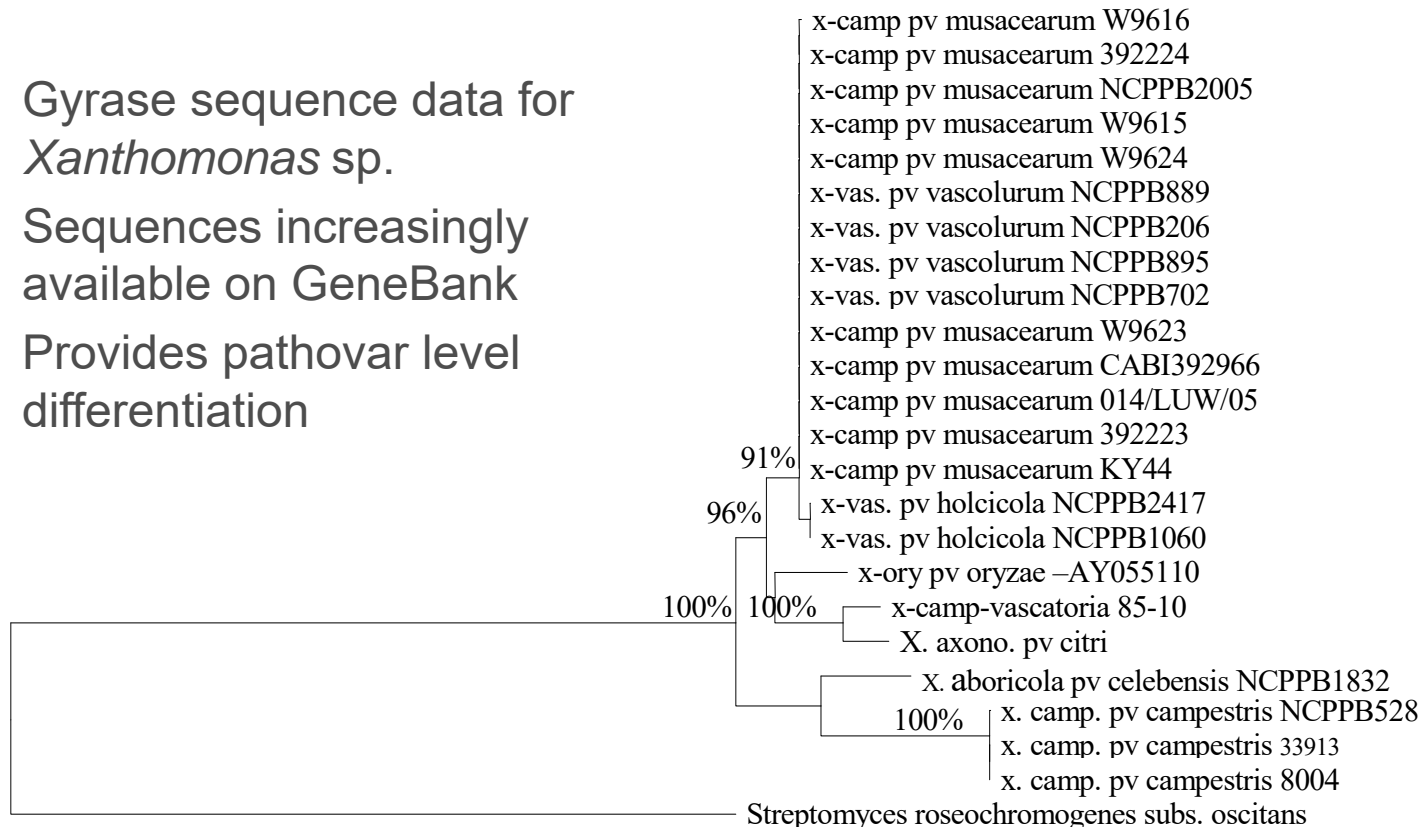
Sequencing of other genes

- Whilst 16S rDNA is the normal target for sequencing, for some bacteria insufficient variation may be present to allow differentiation below the species level [pathovar level]
- For these bacteria different target sequences can be used which present more variation
 - Examples include:
 - Hrp genes
 - Gyrase gene
 - 16-23S rDNA interspacer region

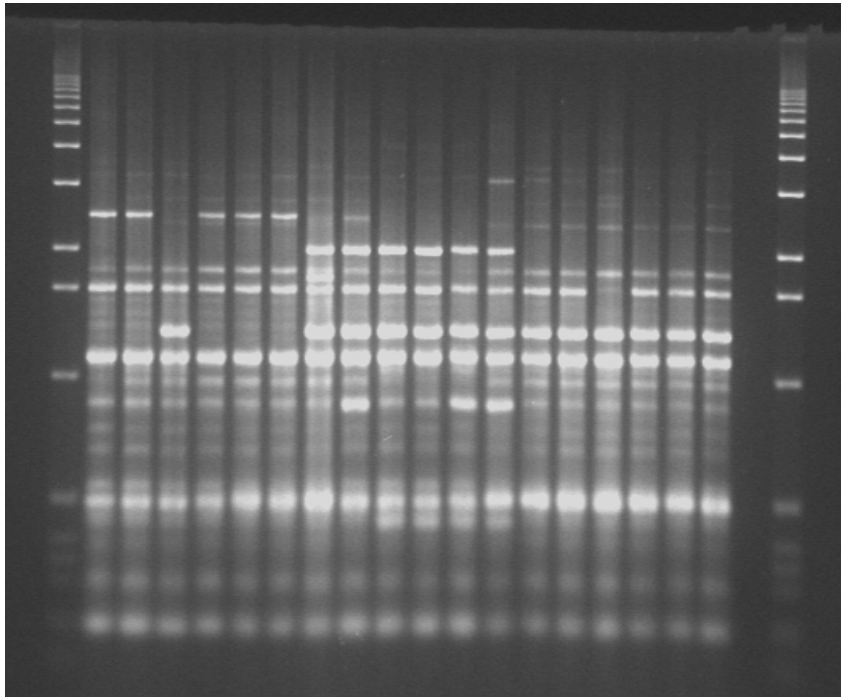
Sequence alignment

0.1

- Gyrase sequence data for *Xanthomonas* sp.
- Sequences increasingly available on GeneBank
- Provides pathovar level differentiation



DNA fingerprinting



- By comparing DNA fingerprint of unknown to known strains an identification can be achieved
- Is particularly appropriate for pathovar level identifications
- Require access to known strains [genetic resource collection]

DNA approaches to identification

- Advantages
 - Commercial services available for sequencing
 - Data can be shared between laboratories
 - By a combination of approaches identification to the genus, species and pathovar level can be achieved
- Limitations
 - Requires investment in PCR and gel equipment
 - Cost of molecular consumables is high
 - Technically demanding; PCR is notorious for 'random' problems