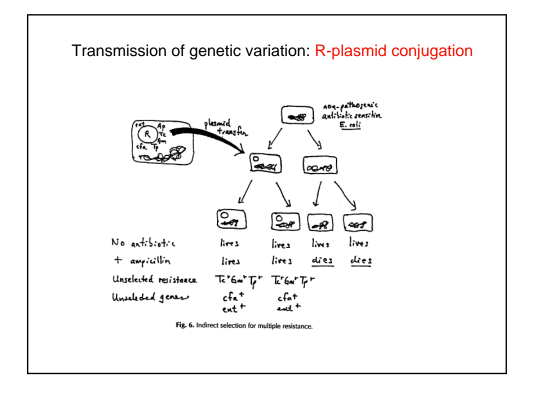
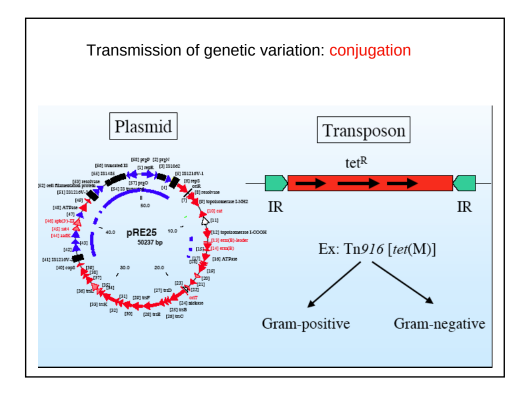
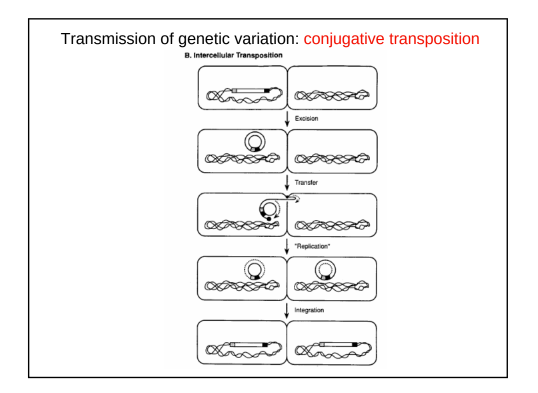
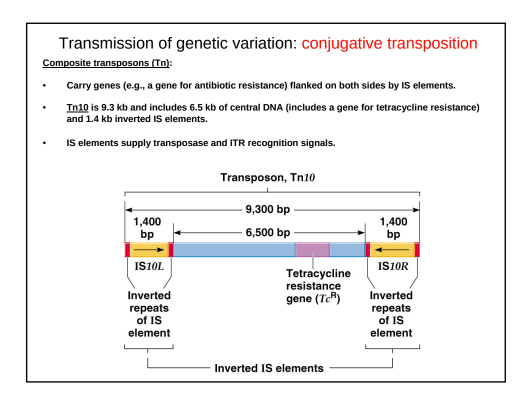


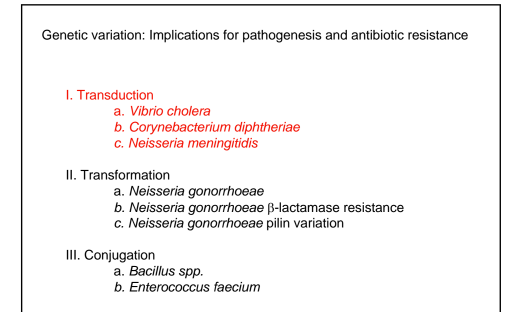
	ion of genetic	c variation: R-plasmid	oonjugat
Propertie	es of some R	l plasmids	
Plasmid	Origin	Resistances	Size (kb)
RP1	England	CbKmTc	36
R527	Spain	CbCmGmKmSmSuTcHg	49
pMG5	Japan	AkKmSuTmBorHgPmrTer	280
pMG90	France	CbCmGmKmSmSuTcTmBorHg	150
Rms149	Germany	CbGmSmSuTra <sup>-</sup>	36
pMG38	USA	CbGmKmSuTcTmHg	53
FP110	Australia	CmaPaeFp110	60
pMG25	South Africa	CbCmGmKmSmSuTmBor	66
pMG69	Ireland	CbGmKmSmSuTcTmTra-	47













I. Transduction

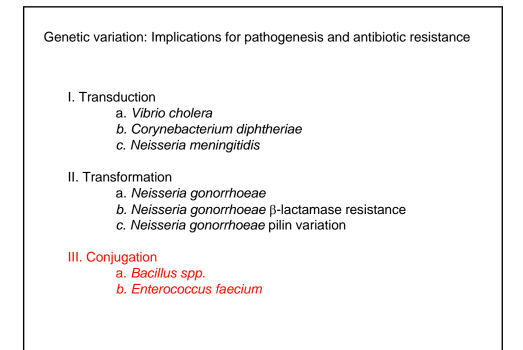
- a. Vibrio cholera
- b. Corynebacterium diphtheriae
- c. Neisseria meningitidis

## **II.** Transformation

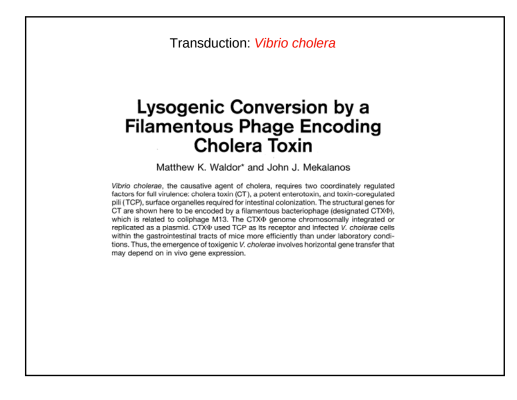
- a. Neisseria gonorrhoeae
- *b. Neisseria gonorrhoeae* β-lactamase resistance
- c. Neisseria gonorrhoeae pilin variation

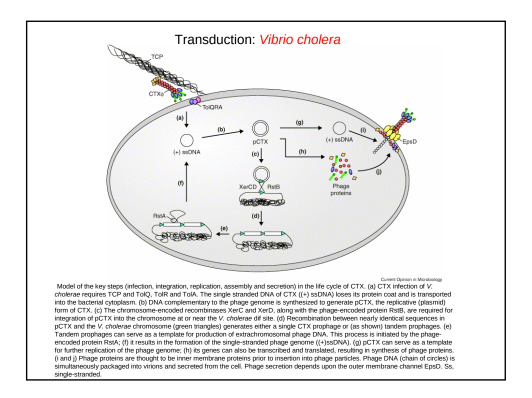
## III. Conjugation

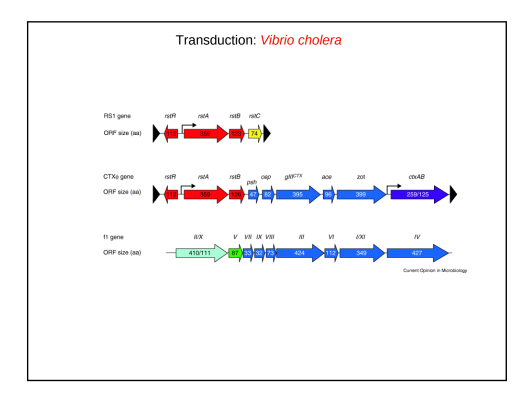
- a. Bacillus spp.
- b. Enterococcus faecium

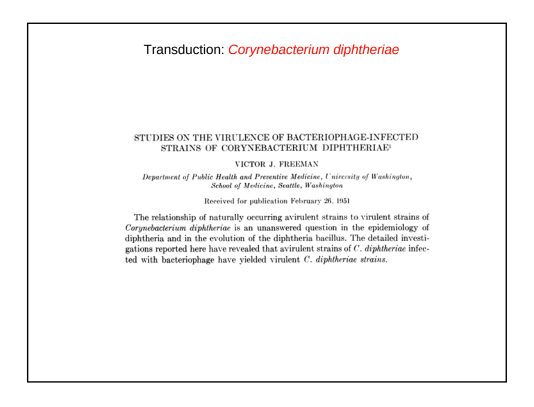


			s Carried by F	
Bacterium	Phage	Gene Product	Phenotype	
Vibrio cholerae	CTX phage	cholerae toxin	cholera	
Escherichia coli	lambda phage	shigalike toxin	hemorrhagic diarrhea	
Clostridium botulinum	clostridial phages	botulinum toxin	botulism (food poisoning)	
Corynebacterium diphtheriae	corynephage beta	diphtheria toxin	diphtheria	
Streptococcus pyogenes	T12	erythrogenic toxins	scarlet fever	









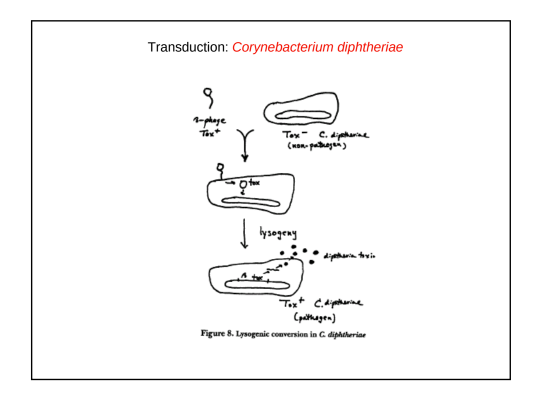
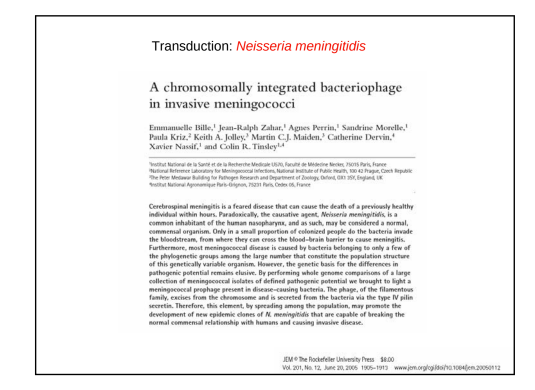


		TABLE 3		
Subcut	taneous tests of l	acteriophage lysa	tes* in guinea p	igs
STRAIN NO.	CULTURE PLUS SALINE	CULTURE PLUS PHAGE A	CULTURE PLUS PHAGE B	CULTURE PLUS PHAG B AND ANTITOXIN
444	0/3†	0/1	4/4	0/2
1174	0/1	0/1	2/2	0/1
1180	0/1	0/1	2/2	0/1
770	0/1	0/1	2/2	0/1
411	0/1	0/1	0/1	0/1
Total	0/7	0/5	10/11	0/6



	Transduction: Neisseria meningitidis						
Table I. Proper	ties of the						
			ningococcal strains				
Gene	Protein length	In 29 "invasive"	In 20 "noninvasive	Protein homologies (BlastP)*	Conserved domains (BlastP)		
ORF1, NMA1792	429	27	0	ORF C7 [Ralstonia solanacearum plasmid] (NP_052309.1), 9 × 10 <sup>-24</sup> ;	replication initiation factor (pfam02486), $6 \times 10^{-42}$ ;		
				RstA1 protein [Vibrio cholerae prophage] (NP_231106.1), 1 × 10-11	phage replication protein RstA (COG2946), 4 × 10-143		
ORF2, NMA1793	104	29	1	NSr	NS		
ORF3, NMA1794	67	29	1	NS	NS		
ORF4, NMA1795	77	29	1	NS	NS		
ORF5, NMA1796	102	29	1	NS	NS		
ORF6, NMA1797	547	29	2	NS	Neisserial TspB proteins		
ORF7, NMA1798	95	29	2	NS	NS		
ORF8, NMA1799	401	29	2	ORF C6 [Ralstonia solanacearum plasmid] (NP_052316.1), 2 × 10 <sup>-31</sup> ; phage-related protein [Xylella fastidiosa] (NP_779131.1), 3 × 10 <sup>-18</sup>	zonular occludens toxin (Zot) (pfam05707), 7 × 10 <sup>-47</sup> ; NS		
ORF9, NMA1800	323	26	0	transposase [Escherichia coli IS621] (BAC76887.1), 3 × 10-37	transposase (COG3547), 1 × 10-17		

