PRINCIPLES OF INHERITANCE AND VARIATION

BIOLOGY

	Single Correct	Answer Type				
1.	A haemophilic woman marries a normal man, then					
	a) All the children will be normal	b) All the sons will be had	emophilic			
	c) All the girls will be haemophilic	d) Half girls will be haem	ophilic			
2.	Disorder inherited as Mendel's law of inheritance called					
	a) Mendelian disorder	b) Chromosomal disorde	r			
	c) Maternal inheritance	d) Polygenic inheritance				
3.	The term 'gene' was coined by					
	a) Avery b) Bateson	c) Johanssen	d) Mendel			
4.	The phenotypic ratio in the F_2 generation of dihybrid		X			
	a) 9 : 3 : 3 : 1	b) 1 : 2 : 2 : 4 : 1 : 2 : 1 : 2	: 1			
	c) 7:1:1:7	d) 12 : 8 : 4				
5.	Chromosome is made up of					
	a) DNA +pectin b) RNA +DNA	c) DNA +histone	d) Only histone			
6.	Select the incorrect statemant from the following.					
	a) Linkage is an exception to the principle of independent assortment in heredity	b) Galactosemia is an inb	orn error of metabolism			
	c) Small population size result in random genetic drift in a population	d) Baldness is a sex-limit	ed trait			
7.	A pure tall and a pure dwarf plant were crossed to p	roduced offsprings. Offspri	ings were self crossed, then			
<i>.</i> .	find out the ratio between true breeding tall to true		ings were sen erossea, men			
	a) 1 : 1 b) 3 : 1	c) 2 : 1	d) 1 : 2 : 1			
8.	Exposure of X-rays enhances the frequency of	-) = -				
	a) Linkage	b) Crossing over				
	c) Pairing of chromosome	d) Segregation				
9.	A self-fertilizing trihybrid plant forms	, , , ,				
	a) 8 different gametes and 64 different zygotes	b) 4 different gametes an	d 16 different zygotes			
	c) 8 different gametes and 16 different zygotes	d) 8 different gametes an	d 32 different zygotes			
10.	Genotype is the					
	a) Genetic constitution	b) Genetic constitution of	f the phenotype			
	c) Trait expressed	d) Expressed genes				
11.	Failure of cytokinesis after A stage of cell division	n results in an increase in a	whole set of chromosomes			
	in an organism calledB					
	a) A-prophase, B-polyploidy	b) A-metaphase, B-polyp	loidy			
5	c) A-anaphase, B-polyploidy	d) A-telophase, B-polyplo	pidy			
12.	In previous question find out total seeds (plants) ha	ving round seed texture				
	a) 12 b) 10	c) 9	d) 11			
13.	The ratio 1 : 1 : 1 : 1 is obtained from a cross betwee	en the parents				
	a) RRYY × rryy b) RRYY × rryy	c) RRYY × Rryy	d) RrYy × rryy			
14.	Which of the following terms represent a pair of con					
	a) Homozygous b) Heterozygous	c) Allelomorphs	d) Codominant genes			
15.	Harmful mutation does not get eliminated from the		-			
	a) Dominant, which have beneficial effect on popula	tion and carried by heteroz	zygous individuals			

	b) Dominant, which have beneficial effc) Carried from one generation to anot		-	
	d) They show genetic drift	0	0	
16.	Incomplete linkage isA Complete	linkage isB	Choose correct optio	n for A and B
	a) A-common, B-rare	0	b) A-rare, B-common	
	c) A-impracticle, B-practicle		d) A-practicle, B-impra	acticle
17.	Mendelism was rediscovered by			
	I. Morgan			
	II. De Vries			
	III. Correns			
	IV. Tschermark			
	Choose the correct option a) I, III and IV b) I, II, III and	d W	c) II, III and IV	d) I, II and III
10	In gynandromorphs,	lu Iv	cj II, III allu IV	uj i, ii aliu ili
10.	a) Some cells of body contain XX and se	ome cells with	genotyne XV	
	b) All cells have XX genotype	Sine cens with	genotype XI	
	c) All cells have XY genotype			
	d) All cells with genotype XXY			
19.	Example of interagenic gene interactio	n is/are		
	a) Incomplete dominance		b) Codominant	
	c) Multiple alleles		d) All of the above	
20.	If a cross between two individuals proc	duces offspring		character (A) and 50%
	recessive character (a),then the genoty	vpes of parents	are	
	a) Sex linked genes		b) Pseudoallelic genes	
	c) Intermediate inheritance		d) Dominant and reces	ssive genes
21.	Which is correct about traits choosen b	y Mendel for h		
	a) Terminal pod was dominant		b) Constricted pod wa	
	c) Green coloured pod was dominant		d) Tall plants were rec	cessive
22.	Codominance is found in			
22	a) Plants b) Animal		c) Both (a) and (b)	d) Prokaryote
23.	During Mendel's investigation, it was fi	irst time that	.A andB were app	blied in biology. Here A and B
	refers to a) A-statistical analysis; B-mathematic	allogic		
	b) A-statistical analysis; B-physical log	-		
	c) A-statistical analysis; B-chemistry lo			
	d) A-statistical analysis; B-simple logic	•		
24.	The chromosomal denotation for heter		ale and homogametic m	ale are
	a) ZW and ZZ b) ZO-ZZ	-8	c) XX-XO	d) Both (a) and (b)
25.	Pure tall plants are crossed with pure of	dwarf plants. Ii	1 the F ₁ -generation, all	
	plants of F ₁ -generation were selfed and	d the ratio of ta	all to dwarf plants obta	ined was 3: 1. This is called
	a) Dominance b) Inheritan	ce	c) Codominance	d) heredity
26.	The best method to determine the hom	nozygosity and	heterozygosity of an ir	ndividual is
	a) Self-fertilisation b) Back cross		c) Test cross	d) Inbreeding
27.	A medical technician, while observing			
	Barr body close to the nuclear membra	ine in the WBC	. This indicates that the	e person under investigation is
	a b) Usernari	, ilia	a) Norme -1 fam. 1	d) Norma - I I -
າດ	a) Colourblind b) Haemoph		c) Normal female	d) Normal male
28.	Find out <i>A</i> , <i>B</i> and <i>C</i> in the diagram give	in below In		

	0 °						
	$XY \times XX$ Par	ents					
	(x)(y)(x) (z)	2					
	(XX) (XY)						
	A B						
	a) A-Male, B-Female,	C-Gametes]	b)	A-Male, B-Female,	C-Sperm	
	c) A-Female, B-Male,	C-Gametes		d)	A-Gametes, B-Male	e, C-Female	
29.	In Turner's syndrom	e					
	a) Female is fertile	b) Male is fe		-	Female is sterile	d) Male is sterile	
30.	=					s in insect damaging a crop is	
	a) Random mutation			-	Genetic recombina		
	c) Directed mutation			d).	Acquired heritable	changes	
31.	Lampbrush chromos	omes sre seen in					
	a) Interphase	b) Zygotene		-	Diplotene	d) metaphase	
32.	In case of incomplete						
	a) Histones	b) Hydrocar		-	Polynucleotides	d) Lipoproteins	
33.	Examples of dissimil	ar sex chromosom	es are given be	elov	V		
	$I. XX - XY \Rightarrow I$						
	II. XX – XO \Rightarrow II						
	I and II in the above s						
	Ι	II					
	a) Man and most	Cockroach and		b)	Cockroach and	Man and most	
	insect	roundworms		۲L	roundworms	insect	
24	c) Butterfly	Fishes			Bird	Reptiles	
54.	Mutations, which alter a) Frameshift mutati	-		-		tions	
	c) Both (a) and (b)	011		-	Base pair substitut None of these	lions	
25	The F_1 generation has	s all tall and E rati		-			
55.	a) Law of dominance				s Independent assor	tmont	
	c) Law of segregation			-	linkage	tillent	
36	Mendel's law were tr			uj	llikage		
50.	a) Alleles are affected			h)	Alleles shows com	nlete dominance	
	2	-		-		is determined by more than on	۱e
	ej mieres or a gene a		anerene gene	-	gene	is determined by more than on	i.C
37.	Blood group-O has				Beile		
07.	a) No antibodies	b) No antige	ns	c) :	a or b antibodies	d) A and B antigens	
38.	Wilson detected the			ey i		a) II and D antigens	
001	a) 1921	b) 1911		C)	1912	d) 1910	
39.	,	,				notypes in F_2 generation will b	е
0 71	a) 4	b) 3		c)		d) 1	
40.		,				t character (A) and 50%	
101	recessive character (-					
	a) Genic interactions					trolling a character	
	c) Expression of mar	-		-		of a gene at a given locus	
41.	A women with albini		0 0	-		0	
	a) 2 normal : 1 albini				All normal		
	c) All albinic	-		-	1 normal : 1 albini	с	
	,			,			

42.	_	nale and males are homogametic, in that condition, the s	sex
	chromosomal representation is a) ZO-ZZ b) XY-XX	c) XX-XO d) ZW-ZZ	
43	-	inherit the genes for blonde hair, while other individual	ls with
15.		ir. This can be best explained by the principle of	
	a) Dominance	b) Multiple alleles	
	c) Independent assortment	d) Incomplete dominance	
44.	In bugs and cockroaches, the sex determina		
	a) XX and XO chromosomes	b) XX and XY chromosomes	
	c) ZZ-ZW chromosomes	d) ZO-ZZ chromosomes	
45.	The twoA separate and pass into two d	aughter nuclei and cells during mitosis. Similarly, each .	B
	replicates, with one pair passing into each o	aughter cell during mitosis. This maintains the similar	C of
	all the cells.		
	Find out correct option for A, B and C		
	a) A-chromatid, B-allele pair, C-morpholog	b) A-chromatid, B-allele pair, C-genetic comp	osition
	c) A-organ, B-organ pair, C-individuality	d) A-unlinked gene, B-linked gene, C-morpho	logy
46.	The shape of chromosome is determined by		
	a) Centrosome b) Centromere	c) Chromomere d) telomere	
47.	Mendel was a		
	a) Austrian biology teacher	b) Austrian monk	
40	c) Austrian scientist	d) Austrian mathematician	
48.	Who clearly proved and define linkage?		
40	a) Morgan b) Castle Improvement of human race through hered	c) Bateson d) Punnett	
49.	a) Euthenics b) Human hered		
50	Test cross involves	ty c) futurial demography d) Eugenics	
501	a) Crossing between two genotypes with re	cessive trait	
	b) Crossing between two F ₁ -hybrids		
	c) Crossing the F_1 -hybrid with a double re-	essive genotype	
	d) Crossing between two genotypes with de	ominant trait	
51.	When a diploid female plant is crossed with	a tetraploid male, the ploidy of endosperm cells in the	
	resulting seed is		
	a) Tetraploidy b) Pentaploidy	c) Diploidy d) Triploidy	
52.	Colour blindness is		
	a) Sex-linked recessive disease		
	b) Sex-linked dominant disease		
	c) Autosomal dominant disease		
۳D	d) Autosomal recessive disease	in only a single convince diploid call is called	
53.	Four different types of gametes produed	in only a single copy in a diploid cell, is called by the F_{-}	
	a) dihybrid	by the F_1 -b) Homozygous condition of the F_1 -dihybrid	
		Four different types of gametes produed by	v the
	c) Four different types of F_1 -dihybrids	d) P_1 -parent	,
54.	If the blood group of a child is A and of mot	ner is B, then the genotype of mother and father may be	ļ
	a) $BB \times AA$ b) $AB \times AB$	c) $B0 \times 00$ d) $B0 \times A0$	
55.			
	• symbol in pedigree analysis represents		
F <	a) Still birth b) Still death	c) Still carrier d) Still mating	
56.	Which amino acids are present in histories?		
	a) Lysine and histidine	b) Valine and histidine	
	c) Arginine and lysine	d) Arginine and histidine	

57	Monosomic trisomy are represented as		
57.	a) $2n - 1 + 1$ b) $2n - 1 - 1$	c) 2 <i>n</i> −1	d) 2 <i>n</i> + 1 + 1
58	Which is a sex-influenced disease?		
50.	a) Baldness in male	b) Haemophilia	
	c) Xeroderma pigmentosa	d) Down's syndrome	
59	Thalassaemia is	aj Down 5 Synaronie	
07.	a) Autosomal recessive disease	b) Autosomal dominant d	lisease
	c) Sex-linked dominant disease	d) Sex-linked recessive di	
60.	Mutation is phenomena which results in alternation		
	a) Sequence b) Carbohydrates	c) Proteins	d) Fat
61.	A man with normal vision whose father was colourb	,	,
	colourblind. Suppose their first child is daughter the	n what are the chances of t	his child to be colourblind?
	a) 100% b) 25%	c) 50%	d) 0%
62.	Gamete mother cells of the chromosome 44 + XY suf	fers from non-disjunction	at first meiotic division.
	Which of the following set of gametes would result?		
	a) 22 + XX, 22 + XY, and 22, 22	b) 22 + XY, 22 + XY, and	22, 22
	c) 22 + X, 22 + Y, and 22 + Y, 22	d) 22 + X, 22 + XY, and 2	2 + Y, 22 + Y
63.	Law of Mendel, which is not completely applicable is	?	
	a) Codominance	b) Law of segregation	
	c) Law of independent assortment	d) Law of dominance	
64.	Low pitched voice, beared and moustaches, belong to		
	a) Sex limited traits b) Sex linked trait	c) Nullisomic traits	d) Sex influenced traits
65.	Multiple allele can be manifested only when there is		
	a) Individual organism b) Genus	c) Population	d) Phylum
66.	Nicotiana sylvestris flowers only during long days		
	raised in the laboratory under different photoperiod	=	
	can be cross fertilized to produce self-fertile offsprin	=	
	can be cross fertilized to produce self-fertile offsprin and <i>N. tobaccum</i> to be separate species?	g. What is the best reason	for considering <i>N. sylvestris</i>
	can be cross fertilized to produce self-fertile offsprin and <i>N. tobaccum</i> to be separate species?a) They are physiologically distinct	g. What is the best reason b) They are morphologica	for considering <i>N. sylvestris</i> ally distinct
67	can be cross fertilized to produce self-fertile offsprin and <i>N. tobaccum</i> to be separate species?a) They are physiologically distinctc) They cannot interbreed in nature	g. What is the best reasonb) They are morphologicad) They are reproductive	for considering <i>N. sylvestris</i> ally distinct
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68. 69.	can be cross fertilized to produce self-fertile offsprin and <i>N. tobaccum</i> to be separate species? a) They are physiologically distinct c) They cannot interbreed in nature The following diagram shows two types of chromoso Part of a EFGHIJKLMNO chromosome A EFGHIJKLMJKLMNO EFGHINO Give the name or type of mutation in respect to A and a) A-Duplication, B-Substitution c) A-Inversion, B-Deletion How many different kinds of gametes will be produc a) Three b) Four Down's syndrome and Turner's syndrome occur in h a) Monosomic and nullisomic conditions respectively What are all the chances of colourblind daughters of was colourblind?	g. What is the best reason b) They are morphologica d) They are reproductive omal mutations d B b) A-Duplication, B-Delet d) A-Inversion, B-Substitue ed by a plant having the ge c) Nine uman beings due to yb) Monosomic and trisom d) Trisomic and tetrasom a normal man marrying no	for considering <i>N. sylvestris</i> ally distinct ly distinct ion ation enotype AABbCC? d) Two hic conditions respectively ic conditions respectively ormal women whose father
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	normal	phenotypically norma	
71.	In males, pattern baldness is related to both autoso	mal genes as well as excess	sive secretion of
	a) Oestrogen b) Growth hormone	c) Testosterone	d) Inhibits
72.	Which of these is not a Mendelian disorder?		
	a) Cystic fibrosis b) Sickle-cell anaemia	c) Colourblindness	d) Turner's syndrome
73.	Which of the following is not true of haemophilia?		
	a) Royal disease	b) Bleeder's disease	
	c) X-linked disorder	d) Y-linked disorder	
74.	If heterozygous dominant (tT) crossed with homozy	ygous dwarf plant, then the	e percentage of progeny
	having dwarf character is		
	a) 60% b) 40%	c) 50%	d) 70%
75.	Mutations are generally induced by means of	-	-
	a) $\alpha - rays$ b) $\beta - rays$	c) $\gamma - rays$	d) UV radiations
76.	Two crosses between the same pair of genotypes or		sources of the gametes are
	reversed in one cross, is known as		<u> </u>
	a) Dihybrid cross b) Reverse cross	c) Test cross	d) Reciprocal cross
77.	A hereditary, disease, which is never passed on from	,	, I
	a) X-chromosomal linked disease	b) Autosomal linked dise	ease
	c) Y-chromosomal linked disease	d) None of the above	
78.	Bateson used the term coupling and repulsion for li	•	choice the correct coupling
	and repulsion combination	5 5	1 0
	Coupling Repulsion		
	a) AABB, aabb AAbb, aaBB	b) AABB, aabb AABB	, AAbb
	c) AAbb, aaBB AaBb, aabb	d) aaBB, aabb AABB	
79.	In blood group typing in human, if an allele contribu	-	
	other parent is <i>i</i> , the resulting blood group of the of		-
	a) A b) B	c) AB	d) 0
80.	A person having 45 chromosomes and Y-chromosom	me absent. Is suffering fror	n
	a) Down's syndrome	b) Klinefelter's syndrom	le
	c) Turner's syndrome	d) gynandromorph	
81.	Linkage and crossing over are		
	a) Same phenomena	b) Different phenomena	
	c) Opposite phenomena	d) Identical phenomena	
82.	The modern concept of gene is		
	a) A segment of DNA, capable of crossing over	b) Functional unit of DN	A
	c) A segment of RNA	d) A segment of chromo	some
83.	Females in haplodiploidy sex determination are		
	a) <i>N</i> b) 2 <i>n</i>	c) $\frac{1}{2}n$	d) 3 <i>n</i>
04	Using imprints from a plate with complete medium	2	ning you can calact
04.	Using imprints from a plate with complete medium		-
	streptomycin resistant mutants and prove that such	i mutations do not origina	te as adaptation. These
	imprints need to be used	h) On platas with minim	almadium
	a) Only on plates with streptomycin	b) On plates with minim	
05	c) Only on plates without streptomycin	d) On plates with and wi	
85.	Phenylketonuria, Huntington's disease and sickle ce	en anaemia are caused resp	bectively due to disorders
	associated with	10	
	a) Chromosome-7, chromosome-11 and chromosom		
	b) Chromosome-11, Chromosome-4 chromosome-1		
	c) Chromosome-7, chromosome-12 and chromosom		
01	d) Chromosome-12, chromosome-4 and chromosom	ne-11	
86.	The arrangement of genes on chromosome is		

86. The arrangement of genes on chromosome is

	a) Linear b) Oviod	c) Diffused	d) Spiral
07	When two genetic loci produce identical phenotypes		
07.	a) Pseudoalleles	-	they are considered to be
	-	b) Multiple alleles	
00	c) The part of same gene	d) Different genes	
88.	Which of the following matches correctly?		
	a) Factor –II - Thromboplastin	b) Factor –III – Prothro	
00	c) Factor – VIII - Antihaemophilic globulin	d) Factor –XII - Haemo	opnilic
89.	The longest chromosomes is seen in		
0.0	a) <i>Allium</i> b) <i>Lilium</i>	c) <i>Trillium</i>	d) <i>Zea mays</i>
90.	Mendel observed that generation shows always p		
~ 1	a) F_4 b) F_2	c) F ₁	d) F_0
91.	A genes are those which occurs on the same chron	mosome andB genes a	re those, which are present
	on different chromosome.		
	Choose correct choice for A and B		
	a) A-linked; B-unlinked gene	b) A-unlinked; B-linked	
	c) A-identical; B-non-identical	d) A-non-identical; B-ide	
92.	Allelic sequence variations where more than one var	iant (allele) at a locus in a	human population with a
	frequency greater than 0.01 is refered to as		
	a) Incomplete dominance	b) Multiple allelism	
	c) SNP	d) DNA polymorphism	
93.	The possibility of a female becoming a haemophilia i	sA rare because mothe	er of such a female has to be
	at leastB and the father should beC		
	Choose the correct option for A, B and C		
	a) A-extremely, B-carrier, C-haemophilic		
	b) A-extremely, B-carrier, C-carrier		
	c) A-extremely, B-haemophilic, C-carrier		
	d) A-extremely, B-haemophilic, C-haemophilic		
94.	If the foetus is Rh ⁺ and mother is Rh ⁻ , then		
	a) Foetus will transmit antigen to mother blood		
	b) Foetus will transmit antibody to mother blood		
	c) Foetus is attacked by antibodies to mother blood		
05	d) Foetus is attacked by antigen to mother blood		
95.	The most popularly known blood grouping is the AB	O grouping. It is named AE	30 and not ABC, because '0'
	in it refers to having		- I
	a) Other antigens besides A and B on RBCs	b) Over dominance of thi	s type on the genes for A
	One entitle du entre sitter entite en entit her the	and B types	
	c) One antibody only–either anti-a or anti-b on the RBCs	u) No anugens A and B o	II KBUS
07			
96.	Alleles are	h) Homelegeus chromes	
	a) Alternate forms of a gene	b) Homologous chromos	ome
07	c) Pair of sex chromosome	d) None of the above	comos hospinas they
97.	Telomere repetitive DNA sequences control the func	-	
	a) Act as replicons	b) Are RNA transcription	
00	c) Help chromosome pairing	d) Prevent chromosome	1055
98.	Genotypic and phenotypic ratios remains the same in		
	a) Sex-linked genes	b) Pseudoallelic genes	• • • • • • • • •
00	c) Intermediate inheritance	d) Dominance and recess	sive genes
99.	Mendelian disorder may be of	a) Dath $(-) (1)$	d) Conthe Jeter
100	a) Recessive b) Dominant	c) Both (a) and (b)	d) Can't be determined
100	. Sickle –cell anaemia has not been eliminated from th		use it
	a) Is controlled by recessive genes	b) Is not a fatal disease	

- d) Is controlled by dominant genes
- 101. A condition characterized by not having an exact number of chromosomes in a multiple of haploid set is called

a) Polyploidy b) Synploidy c) aneuploidy d) None of these 102. Choose correct option for A, B, C and D Τt TΤ Х a) A-tt, B-TT, C-TT, D-TT b) A-Tt, B-Tt, C-Tt, D-Tt c) A-TT, B-TT, C-Tt, D-TT d) A-Tt, B-Tt, C-Tt, D-TT 103. When a cross is conducted between black feathered hen and a white feathered cock, blue feathered fowls are formed. When these fowls are allowed for interbreeding, in F₂- generation, there are 20 blue fowls. What would be the number of black and white fowls? a) Black 20, white 10 b) Black 20, white 20 c) Black 10, white 10 d) Black 10, white 20 104. Chromosomes are made up of a) DNA are protein b) RNA and DNA c) DNA and histone d) Only histones 105. In pedigree analysis, the square, blackened and horizontal lines represents a) Female, healthy individual, parents b) Female, affected individual, parents c) Male, affected individual, parents d) Male, affected individual, progeny 106. Following pedigree chart shows a) Character is carried by Y-chromosome b) Character is sex-linked recessive c) Character is sex-linked dominant d) Character is recessive autosomal 107. Mr. Sidd is suffering from hypertrichosis and phenylketonuria. His father is heterozygous for phenylketonuria. The probability of Sidd's sperm having one recessive autosomal allele and holandric gene is a) $\frac{1}{2}$ c) $\frac{1}{10}$ d) $\frac{1}{4}$ b) $\frac{1}{8}$ 108. F₃-generation is obtained by b) Selfing of F₂ c) Crossing of F_1 and F_2 d) None of these a) Selfing of F_1 109. In which one of the following, complementary gene interaction rato of 9 : 7 is observed? b) Coat colour in mouse a) Fruit shape in Shepherd's purse c) Feather colour in fowl d) Flower colour in pea 110. Starch synthesis gene in pea plant is the example of a) Single gene produce more than one effects b) Multiple genes produce more than one effects c) Two genes produce more than one effects d) Multiple genes produce less than one effects 111. In Drosophila, the sex is determined by a) The ratio of pairs of X-chromosomes to the pairs of autosomes b) Whether the egg is fertilized or develops parthenogenetically c) The ratio of number of X-chromosomes to the set of autosomes d) X and Y-chromosomes 112. The 1:2:1 ratio with the pink flower in the F_2 -generation indicate the phenomenon of Page 8

a) Dominance	b) Codominance	
c) Incomplete dominance	d) Segregation	
113. Sexual reproducation leads to		
a) Genetic recombination	b) Polyploidy	
c) Aneuploidy	d) Euploidy	
114. Husband has blood group-A and wife has blood gro		
a) A b) B	c) AB	d) A, B, AB and O
115. Study the following figure and find out the most pr	obable position at which t	he crossing over takes place
W X Y Z		
W X Y Z		
a) w and W b) X and y	c) y and Z	d) w and z
116. Given diagram shows certain type of traits in huma	n. Which one of the follov	ving option could be an
example of this pattern?		
Female Male		
mother father		
Daughter Son		
a) Haemophilia b) Anaemia	c) Phenylketonuria	d) Thalassaemia
117. In case of incomplete dominance, what will be the	· ·	•
a) 3 : 1 b) 1 : 2 : 1	c) 1:1:1:1	d) 2 : 2
118. Haemophilia, a X-linked recessive disease is caused	-	
a) Blood plasma and vitamin–K	b) Blood platelets and	haemoglobin
c) Lack of clotting material and vitamin-K	d) All of the above	
119. All of this obeys Mendel's laws except		
a) Codominance	b) Independent assorti	nent
c) Dominance	d) Purity of gametes	
120. in β -thalassaemia, the affected chromosome is		
a) 16th b) 14th	c) 13th	d) 19th
121. In pea plants, yellow seeds are dominant to green.		-
groop good and a what ratio of wellow and groop		
green seeded plant, what ratio of yellow and green		—
a) 50 : 50 b) 9 : 1	<pre>seeded plants would you c) 1:3</pre>	expect in <i>F</i> ₁ generation? d) 3 : 1
a) 50 : 50 b) 9 : 1 122. Who was fly men of genetics?	c) 1:3	d) 3 : 1
a) 50 : 50 b) 9 : 1 122. Who was fly men of genetics? a) Sutton b) Pasteur		—
 a) 50: 50 b) 9: 1 122. Who was fly men of genetics? a) Sutton b) Pasteur 123. Mendel's contribution for genetic inheritance was 	c) 1:3	d) 3 : 1
 a) 50: 50 b) 9: 1 122. Who was fly men of genetics? a) Sutton b) Pasteur 123. Mendel's contribution for genetic inheritance was a) The idea that genes are found on chromosomes 	c) 1:3 c) Robert Hooke	d) 3 : 1
 a) 50: 50 b) 9: 1 122. Who was fly men of genetics? a) Sutton b) Pasteur 123. Mendel's contribution for genetic inheritance was a) The idea that genes are found on chromosomes b) Providing a mechanism that explains patterns or 	c) 1 : 3 c) Robert Hooke f inheritance	d) 3 : 1
 a) 50: 50 b) 9: 1 122. Who was fly men of genetics? a) Sutton b) Pasteur 123. Mendel's contribution for genetic inheritance was a) The idea that genes are found on chromosomes b) Providing a mechanism that explains patterns o c) Describing how genes are influenced by the environment 	c) 1 : 3 c) Robert Hooke f inheritance ironment	d) 3 : 1
 a) 50: 50 b) 9: 1 122. Who was fly men of genetics? a) Sutton b) Pasteur 123. Mendel's contribution for genetic inheritance was a) The idea that genes are found on chromosomes b) Providing a mechanism that explains patterns o c) Describing how genes are influenced by the envior d) Determining that the information contained in Environment of the second s	c) 1 : 3 c) Robert Hooke f inheritance ironment DNA codes for proteins	d) 3 : 1
 a) 50: 50 b) 9: 1 122. Who was fly men of genetics? a) Sutton b) Pasteur 123. Mendel's contribution for genetic inheritance was a) The idea that genes are found on chromosomes b) Providing a mechanism that explains patterns o c) Describing how genes are influenced by the env d) Determining that the information contained in II 124. The genotypic ratio of a monohybrid cross in F2-genetic for the second second	 c) 1:3 c) Robert Hooke f inheritance ironment DNA codes for proteins eneration is 	d) 3 : 1 d) TH Morgan
 a) 50: 50 b) 9: 1 122. Who was fly men of genetics? a) Sutton b) Pasteur 123. Mendel's contribution for genetic inheritance was a) The idea that genes are found on chromosomes b) Providing a mechanism that explains patterns o c) Describing how genes are influenced by the env d) Determining that the information contained in II 124. The genotypic ratio of a monohybrid cross in F2-general 3: 1 b) 1: 2: 1 	 c) 1:3 c) Robert Hooke f inheritance ironment DNA codes for proteins eneration is c) 2:1:1 	d) 3 : 1 d) TH Morgan d) 9 : 3 : 3 : 1
 a) 50: 50 b) 9: 1 122. Who was fly men of genetics? a) Sutton b) Pasteur 123. Mendel's contribution for genetic inheritance was a) The idea that genes are found on chromosomes b) Providing a mechanism that explains patterns o c) Describing how genes are influenced by the env d) Determining that the information contained in II 124. The genotypic ratio of a monohybrid cross in F2-general 3: 1 b) 1: 2: 1 125. Baldness is more common in men than in woman. 	 c) 1:3 c) Robert Hooke f inheritance ironment DNA codes for proteins eneration is c) 2:1:1 It could be explained on the 	d) 3 : 1 d) TH Morgan d) 9 : 3 : 3 : 1
 a) 50: 50 b) 9: 1 122. Who was fly men of genetics? a) Sutton b) Pasteur 123. Mendel's contribution for genetic inheritance was a) The idea that genes are found on chromosomes b) Providing a mechanism that explains patterns o c) Describing how genes are influenced by the env d) Determining that the information contained in II 124. The genotypic ratio of a monohybrid cross in F2-general 3: 1 b) 1: 2: 1 	 c) 1:3 c) Robert Hooke f inheritance ironment DNA codes for proteins eneration is c) 2:1:1 It could be explained on the 	d) 3 : 1 d) TH Morgan d) 9 : 3 : 3 : 1

c) Genes of baldness are autosomal bu d) None of the above	it influenced by androgens	
-	tore in non-nod word chosen by Mondal	n
126. How many pairs of contrasting charac		
a) 3 b) 5	c) 7	d) 9
127. A mutagen pollutant is		
a) Organophosphates	b) Resins	
c) Chlorinated hydrocarbons	d) Nitrogen oxides	
128. Both chromosome and gene (Mendelia		ive are transmitted from
generation to generation in which form a) Changed b) Unaltere		d) Disintegrated
, , , , , , , , , , , , , , , , , , , ,	-	uj Disintegrateu
129. Pedigree analysis is very important in		
a) It helps genetic counselers to avoid	uisoi dei s	
b) It shows origin of traits		
c) It shows the flow of traits in family		
d) All of the above	andition non-stable program	the feater reconcided for
130. Genes when present in homozygous co	Shaluon results in non – viable progeny	, the factor responsible for
such conditions are		
a) Polygenes b) Linked g	,	d) Epistatic genes
131. Turner's syndrome caused due to the		
a) One X-chromosome (44 with XO)	b) One Y-chromosome	
c) One X-and Y-chromosome	d) Two X-chromosome	
132. The recessive genes located on X-chro	-	
a) Lethal b) Sub-leth		d) Expressed in females
133. Strength of the linkage between the tw		
a) Proportionate to the distance betwee		
b) Inversely proportionate to the dista	ance between them	
c) Depend on the chromosomes		
d) Depend upon the size of chromosor		
134. Fruitfly is excellent model for genetics	because of	
I. Small life cycle (two week)	1.	
II. Can be feed on simple synthesis me III. Single mating produce large numbe		
IV. Clear differentiation of sexes	er of progeny	
V. Many heredity variation can be seen	n with low power microscopes	
Choose the correct option	1 1	
a) I, II and III b) III, IV an	d V c) I, IV and V	d) All of these
135. In Guinea pigs, black short hair (BBSS)) is dominant over white long hair (bbs	s). During a dihybrid cross,
the F_2 -generation individuals with ger	notypes BBSS, BbSS, BBSs and BbSs are	in the ratio of
a) 9:3:3:1 b) 4:2:1:	2 c) 1:2:1:2	d) 1 : 2 : 2 : 4
136. When both parents are of blood type A	AB, they can have children with	
a) A, B, AB and O blood types	b) A, B, and AB blood ty	pes
c) A and B blood types	d) A, B and O blood type	es
137. Test cross is		
a) Recessive F ₁ -plant crosses with dor	ninant F ₂ -plant	
b) Recessive F ₂ -plant crosses with do	ninant F ₃ -plant	
c) Dominant F_2 -plant crosses with rec	essive parent plants	
d) Dominant F ₂ -plant crosses with het	terozygous parent plants	
138. The phenomenon of a single gene regu	llating several phenotypes is called	
a) Multiple allelism	b) epistasis	
c) Incomplete dominance	d) Pleiotropism	
139. If two pea plants having red (dominan	t) coloured flowers with unknown gene	otypes are crossed, 75% of
the flowers with unknown genotypes	are crossed, 75% of the flowers are red	and 25% are white. The

construic constitution of	f the parents having red co	lourad flowers will be	
a) Both homozygous	i the parents having red to	b) One homozygous and	other beterozygous
c) Both heterozygous		d) Both hemizygous	other neterozygous
140. A woman has a haemopl	ailic son and three normal (,	that of her husband with
respect to this gene wou		children. Her genotype and	that of her husband with
	b) $X^h X^h$ and $X^h Y$	c) X ^h X ^h and XY	d) ^{Xh} X and XY
a) XX and X ^h Y	b) A A and A I	c) $X X$ and $X I$	d) A A and Al
141. The proportion of plants	s that were dwarf and tall in	n F ₂ - generation ofo Mende	el experiment
a) $\frac{1}{4}$ th and $\frac{3}{4}$ th	b) $\frac{3}{4}$ th and $\frac{1}{4}$ th	c) $\frac{2}{3}$ rd and $\frac{1}{3}$ rd	d) $\frac{1}{3}$ rd and $\frac{4}{3}$ rd
142. Night blindness is	4 4	- 3 3	- 3 - 3
a) Genetic disease		b) Nutritional deficiency	<i>v</i> disease
c) Generally found in ma	ale	d) Generally found in fer	
143. Two genes R and Y are lo		· ·	
_	hybridized, then F_2 -segregation		1
a) 1:2:1	b) 3 : 1	c) 9:3:3:1	d) 1 : 1 : 1
144. Who argued that pairing	and separation of chromo	somes would lead to the se	gregation of a pair of factor
they carried?	_		
a) Sutton	b) Boveri	c) Both (a) and (b)	d) Morgan
145. Sex chromosomes of ma	le are		
a) Homozygous	b) Heterozygous	c) Hemizygous	d) autosomes
146. Trisomy of which chrom	osome is involved in Dowr	n's syndrome?	
a) 15 th	b) 21 st	c) 20 th	d) 19 th
147. Which of the following s	ymbols are used for repres	enting chromosome of bird	ds?
a) ZZ-ZW	b) XX-XY	c) XO-XX	d) ZZ-WW
148. Sudden and heritable ch	ange in a character of an oi	rganism is called	
a) Mutation	b) Heterosis	c) Inbreeding	d) selection
149. Heterozygous purple flo	wer is crossed with recessi		eny has the ratio
a) All purple		b) All white	
c) 50% purple, 50% wh		d) 75% purple, 25% wh	
150. The Mendel crossed true	0 1	-	
	character was dwarf. The r		
a) F_1	b) F ₂	c) F_3	d) F_2 and F_3
151. Night blindness can be c	orrected by giving vitamin-	A but colour blindness	s can't be cured because it is
B disease.	n fan A an J D		
Choose the correct optio			
a) A-A; B-genetic 152. Heredity is	b) A-B; B-autosomal	c) A-C; B-non-genetic	d) A-D; B-genetic
a) Transmission of chara	actors	b) Mixing of characters	
c) Blending of inheritan		d) Deleting of characters	c.
153. Which of these statemen		, 0	3
	-		n's disease do not exist at this
time	t the presence of the unere	responsible for francingtor	
	on's disease is typically be	tween birth and three vear	s of age
	effective treatment of Hun		
	is caused by the expressior		
154. Centromere is required			
a) Transcription		b) Crossing over	
c) Cytoplasmic cleavage		d) Movement of chromo	somes towards poles
155. Which of the following c			
abnormality/linkage?			

a) Klinefelter's syndrome			
aj minerener 5 Synaronne	-44 autosomes + XXY	b) Colour blindness	–Y- linked
c) Erythroblastosis foetalis		d) Down' s syndrome	
156. Rrrr progeny : Red (dominar	nt) flowered heterozygo	us crossed with white flowe	er
a) $350 \rightarrow \text{red} : 350 \rightarrow \text{white}$		b) $450 \rightarrow \text{red} : 250 \rightarrow \text{whi}$	te
c) $380 \rightarrow \text{red} : 250 \rightarrow \text{white}$		d) None of these	
157. A hereditary disease which is	s never passed on form	Father to son is	
a) X- chromosomal linked di	-	b) Autosomal linked disea	50
c) Y- chromosomal linked di		d) None of the above	
158. A man with blood group-B m		,	having blood group-B
What is the genotype of child			nuting block group Di
) ^{IaIo}	c) _{IpIo}	d) ^{IbIb}
)	C)	a)
159. Linked gene are present on			
a) Same chromosome		b) Different chromosome	
c) Heterologous chromosom		d) Paired chromosome	
160. The structure that become d		•	
5) Centriole	c) DNA	d) None of these
161. Genetics is the branch of biol			
) Inheritance	c) Both (a) and (b)	d) Study of characters
162. Giant chromosomes are foun	id inside		
a) nucleus of man	.1.	b) oocytes of frog	1. *1 .
c) salivary glands of silk mot		d) salivary glands of <i>Drose</i>	
163. Who is known as father of ph		-	
) Charles Elton	c) Taylor	d) Archibald Garrod
164. The graphical representation cross, is called	ii to calculate the probab	inty of all possible genotyp	les of offspring in a genetic
) Karvotvne	c) Punnett square	d) Chromosome man
a) Pedigree analysis b)		c) Punnett square	d) Chromosome map
a) Pedigree analysis b) 165. Rh factor can produce diseas			d) Chromosome map
a) Pedigree analysis b) 165. Rh factor can produce diseas a) AIDS		b) Turner's syndrome	d) Chromosome map
a) Pedigree analysis b) 165. Rh factor can produce diseas a) AIDS c) Erythroblastosis foetalis	56	b) Turner's syndrome d) Sickle-cell anaemia	d) Chromosome map
 a) Pedigree analysis b) 165. Rh factor can produce diseas a) AIDS c) Erythroblastosis foetalis 166. To determine heterozygousit 	se ty of a cross, one has to j	b) Turner's syndrome d) Sickle-cell anaemia perform	
 a) Pedigree analysis b) 165. Rh factor can produce diseas a) AIDS c) Erythroblastosis foetalis 166. To determine heterozygousit a) Back cross b) 	se ty of a cross, one has to j) Reciproacal cross	b) Turner's syndrome d) Sickle-cell anaemia perform c) Test cross	d) Any of these
 a) Pedigree analysis b) 165. Rh factor can produce diseas a) AIDS c) Erythroblastosis foetalis 166. To determine heterozygousit 	se ty of a cross, one has to j) Reciproacal cross	b) Turner's syndrome d) Sickle-cell anaemia perform c) Test cross	d) Any of these
 a) Pedigree analysis b) 165. Rh factor can produce diseas a) AIDS c) Erythroblastosis foetalis 166. To determine heterozygousit a) Back cross b) 167. Which of the following type of 	se ty of a cross, one has to j) Reciproacal cross	b) Turner's syndrome d) Sickle-cell anaemia perform c) Test cross reverse order of genes in a	d) Any of these chromosome?
 a) Pedigree analysis b) 165. Rh factor can produce diseas a) AIDS c) Erythroblastosis foetalis 166. To determine heterozygousit a) Back cross b) 167. Which of the following type of a) Deletion 	se ty of a cross, one has to p) Reciproacal cross of mutation involves the	 b) Turner's syndrome d) Sickle-cell anaemia perform c) Test cross reverse order of genes in a b) Duplication d) Reciprocal translocation 	d) Any of these chromosome?
 a) Pedigree analysis b) 165. Rh factor can produce diseases a) AIDS c) Erythroblastosis foetalis 166. To determine heterozygousite a) Back cross b) 167. Which of the following type of a) Deletion c) Inversion 168. The chromosomal number in 	se ty of a cross, one has to p) Reciproacal cross of mutation involves the	 b) Turner's syndrome d) Sickle-cell anaemia perform c) Test cross reverse order of genes in a b) Duplication d) Reciprocal translocation 	d) Any of these chromosome?
 a) Pedigree analysis b) 165. Rh factor can produce diseases a) AIDS c) Erythroblastosis foetalis 166. To determine heterozygousite a) Back cross b) 167. Which of the following type of a) Deletion c) Inversion 168. The chromosomal number in 	se ty of a cross, one has to p) Reciproacal cross of mutation involves the n the meiocytes of house) 12	 b) Turner's syndrome d) Sickle-cell anaemia perform c) Test cross reverse order of genes in a b) Duplication d) Reciprocal translocation 	d) Any of these chromosome? n
 a) Pedigree analysis b) 165. Rh factor can produce diseases a) AIDS c) Erythroblastosis foetalis 166. To determine heterozygousite a) Back cross b) 167. Which of the following type of a) Deletion c) Inversion 168. The chromosomal number in a) 8 	se ty of a cross, one has to p) Reciproacal cross of mutation involves the n the meiocytes of house) 12	 b) Turner's syndrome d) Sickle-cell anaemia perform c) Test cross reverse order of genes in a b) Duplication d) Reciprocal translocation 	d) Any of these chromosome? n
 a) Pedigree analysis b) 165. Rh factor can produce diseases a) AIDS c) Erythroblastosis foetalis 166. To determine heterozygousite a) Back cross b) 167. Which of the following type of a) Deletion c) Inversion 168. The chromosomal number in a) 8 b) 169. The alternate forms of a general 	se ty of a cross, one has to p) Reciproacal cross of mutation involves the n the meiocytes of house) 12	 b) Turner's syndrome d) Sickle-cell anaemia perform c) Test cross reverse order of genes in a b) Duplication d) Reciprocal translocation fly is c) 21 	d) Any of these chromosome? n
 a) Pedigree analysis b) 165. Rh factor can produce diseases a) AIDS c) Erythroblastosis foetalis 166. To determine heterozygousitive a) Back cross b) 167. Which of the following type of a) Deletion c) Inversion 168. The chromosomal number in a) 8 b) 169. The alternate forms of a general Recessive character c) Alleles 170. Haemophilia is related to 	ty of a cross, one has to p) Reciproacal cross of mutation involves the n the meiocytes of house) 12 e is called	 b) Turner's syndrome d) Sickle-cell anaemia perform c) Test cross reverse order of genes in a b) Duplication d) Reciprocal translocation fly is c) 21 b) Dominant character d) Alternative gene 	d) Any of these chromosome? n d) 23
 a) Pedigree analysis b) 165. Rh factor can produce diseases a) AIDS c) Erythroblastosis foetalis 166. To determine heterozygousite a) Back cross b) 167. Which of the following type of a) Deletion c) Inversion 168. The chromosomal number in a) 8 b) 169. The alternate forms of a general Recessive character c) Alleles 170. Haemophilia is related to a) Albinism b) 	ty of a cross, one has to p) Reciproacal cross of mutation involves the n the meiocytes of house) 12 e is called) Sickle-cell anaemia	 b) Turner's syndrome d) Sickle-cell anaemia perform c) Test cross reverse order of genes in a b) Duplication d) Reciprocal translocation fly is c) 21 b) Dominant character 	d) Any of these chromosome? n
 a) Pedigree analysis b) 165. Rh factor can produce diseases a) AIDS c) Erythroblastosis foetalis 166. To determine heterozygousitive a) Back cross b) 167. Which of the following type of a) Deletion c) Inversion 168. The chromosomal number in a) 8 b) 169. The alternate forms of a general a) Recessive character c) Alleles 170. Haemophilia is related to a) Albinism b) 171. Identify a Mendelian disorder 	ty of a cross, one has to p) Reciproacal cross of mutation involves the n the meiocytes of house) 12 e is called) Sickle-cell anaemia	 b) Turner's syndrome d) Sickle-cell anaemia perform c) Test cross reverse order of genes in a b) Duplication d) Reciprocal translocation fly is c) 21 b) Dominant character d) Alternative gene c) Colour blindness 	d) Any of these chromosome? n d) 23
 a) Pedigree analysis b) 165. Rh factor can produce diseases a) AIDS c) Erythroblastosis foetalis 166. To determine heterozygousinal Back cross a) Back cross b) 167. Which of the following type of a) Deletion c) Inversion 168. The chromosomal number in a) 8 b) 169. The alternate forms of a general Back character c) Alleles 170. Haemophilia is related to a) Albinism b) 171. Identify a Mendelian disorder a) Down's syndrome 	ty of a cross, one has to p) Reciproacal cross of mutation involves the n the meiocytes of house) 12 e is called) Sickle-cell anaemia	 b) Turner's syndrome d) Sickle-cell anaemia perform c) Test cross reverse order of genes in a b) Duplication d) Reciprocal translocation fly is c) 21 b) Dominant character d) Alternative gene c) Colour blindness b) Turner's syndrome 	d) Any of these chromosome? n d) 23
 a) Pedigree analysis b) 165. Rh factor can produce diseases a) AIDS c) Erythroblastosis foetalis 166. To determine heterozygousitive a) Back cross b) 167. Which of the following type of a) Deletion c) Inversion 168. The chromosomal number in a) 8 b) 169. The alternate forms of a general 8 b) 169. The alternate forms of a general 8 c) Alleles 170. Haemophilia is related to a) Albinism b) 171. Identify a Mendelian disorderal 3 c) Phenylketonuria 	ty of a cross, one has to p) Reciproacal cross of mutation involves the n the meiocytes of house) 12 e is called) Sickle-cell anaemia er from the following.	 b) Turner's syndrome d) Sickle-cell anaemia perform c) Test cross reverse order of genes in a b) Duplication d) Reciprocal translocation fly is c) 21 b) Dominant character d) Alternative gene c) Colour blindness b) Turner's syndrome d) Klinefelter's syndrome 	 d) Any of these chromosome? n d) 23 d) thalassemia
 a) Pedigree analysis b) 165. Rh factor can produce diseases a) AIDS c) Erythroblastosis foetalis 166. To determine heterozygousitive a) Back cross b) 167. Which of the following type of a) Deletion c) Inversion 168. The chromosomal number in a) 8 b) 169. The alternate forms of a general a) Recessive character c) Alleles 170. Haemophilia is related to a) Albinism b) 171. Identify a Mendelian disorderal bown's syndrome c) Phenylketonuria 172. When a tall plant with round 	ty of a cross, one has to p) Reciproacal cross of mutation involves the n the meiocytes of house) 12 e is called) Sickle-cell anaemia er from the following.	 b) Turner's syndrome d) Sickle-cell anaemia perform c) Test cross reverse order of genes in a b) Duplication d) Reciprocal translocation fly is c) 21 b) Dominant character d) Alternative gene c) Colour blindness b) Turner's syndrome d) Klinefelter's syndrome 	 d) Any of these chromosome? n d) 23 d) thalassemia hkled seeds (ttrr), the F₁-
 a) Pedigree analysis b) 165. Rh factor can produce diseases a) AIDS c) Erythroblastosis foetalis 166. To determine heterozygousite a) Back cross b) 167. Which of the following type of a) Deletion c) Inversion 168. The chromosomal number in a) 8 b) 169. The alternate forms of a generation disorder a) Albinism b) 171. Identify a Mendelian disorder a) Down's syndrome c) Phenylketonuria 172. When a tall plant with round generation consists of tall plat 	ty of a cross, one has to p) Reciproacal cross of mutation involves the n the meiocytes of house) 12 e is called) Sickle-cell anaemia er from the following. I seeds (TTRR) crossed w ants with round seeds. V	 b) Turner's syndrome d) Sickle-cell anaemia perform c) Test cross reverse order of genes in a b) Duplication d) Reciprocal translocation fly is c) 21 b) Dominant character d) Alternative gene c) Colour blindness b) Turner's syndrome d) Klinefelter's syndrome 	 d) Any of these chromosome? n d) 23 d) thalassemia hkled seeds (ttrr), the F₁-
 a) Pedigree analysis b) 165. Rh factor can produce diseases a) AIDS c) Erythroblastosis foetalis 166. To determine heterozygousite a) Back cross b) 167. Which of the following type of a) Deletion c) Inversion 168. The chromosomal number int a) 8 b) 169. The alternate forms of a general a) Recessive character c) Alleles 170. Haemophilia is related to a) Albinism b) 171. Identify a Mendelian disorder a) Down's syndrome c) Phenylketonuria 172. When a tall plant with round generation consists of tall plat wrinkled seeds in F₁-generate 	ty of a cross, one has to p) Reciproacal cross of mutation involves the n the meiocytes of house) 12 e is called) Sickle-cell anaemia er from the following. I seeds (TTRR) crossed v ants with round seeds. V tion?	 b) Turner's syndrome d) Sickle-cell anaemia perform c) Test cross reverse order of genes in a b) Duplication d) Reciprocal translocation fly is c) 21 b) Dominant character d) Alternative gene c) Colour blindness b) Turner's syndrome d) Klinefelter's syndrome with a dwarf plant with write 	 d) Any of these chromosome? n d) 23 d) thalassemia hkled seeds (ttrr), the <i>F</i>₁-on of dwarf plant with
a) Pedigree analysis b) 165. Rh factor can produce diseases a) AIDS c) Erythroblastosis foetalis 166. To determine heterozygousit a) Back cross b) 167. Which of the following type of a) Deletion c) Inversion 168. The chromosomal number in a) 8 b) 169. The alternate forms of a generic a) Recessive character c) Alleles 170. Haemophilia is related to a) Albinism b) 171. Identify a Mendelian disorder a) Down's syndrome c) Phenylketonuria 172. When a tall plant with round generation consists of tall plat wrinkled seeds in F_1 -generat	ty of a cross, one has to p) Reciproacal cross of mutation involves the n the meiocytes of house) 12 e is called) Sickle-cell anaemia er from the following. I seeds (TTRR) crossed w ants with round seeds. V	 b) Turner's syndrome d) Sickle-cell anaemia perform c) Test cross reverse order of genes in a b) Duplication d) Reciprocal translocation fly is c) 21 b) Dominant character d) Alternative gene c) Colour blindness b) Turner's syndrome d) Klinefelter's syndrome 	 d) Any of these chromosome? n d) 23 d) thalassemia hkled seeds (ttrr), the F₁-

173	The term	'Genetics'	was proposed by	
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a) Mendel b) Bateson

174. Sex chromosomes are also known as

a) Autosomes b) Allosomes

d) karyotype 175. Mendel obtained recessive character in F₂ by ...A... the ...B... plants. Here A and B refers to

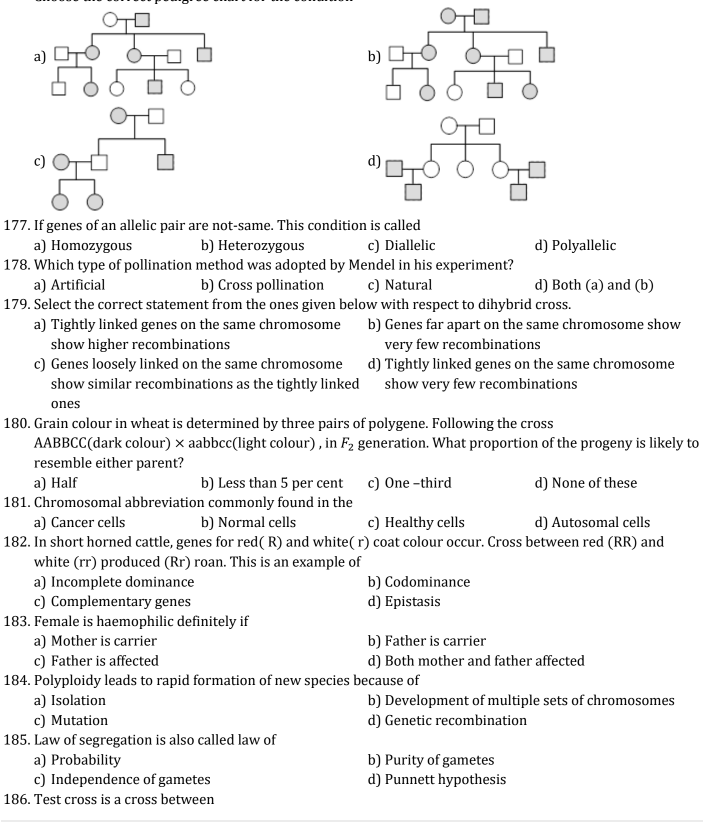
- a) A-self-pollinating; $B-F_1$
- c) A-cross-pollinating; $B-F_1$
- b) A-self-pollinating; $B-F_2$ d) A-cross-pollinating; $B-F_2$

d) Johanssen

176. In a family father had a trait but mother did not. All their sons and daughter had this trait. The same trait was found in some grand daughters, through daughter were married to the normal persons. Choose the correct pedigree chart for the condition

c) Motgan

c) Genome



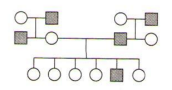
a) Urbrid V Daminan	h w a waw t	h) Urihnid v Dogogirov	
a) Hybrid × Dominant parent c) Hybrid × Hybrid parent		b) Hybrid × Recessive parentd) Two distantly related species	
,	nal sex determination, female		aspecies
a) Homogametic	iai sex ueter initiation, remate	b) Heterogametic	
c) Can not determine		d) All of the above	
188. Heterogametic male c	andition door not occur in	uj Ali ol tile above	
a) Birds		c) <i>Drosophila</i>	d) Honoy boo
	b) Humans cross which is a dihybrid cro	, 1	d) Honey bee
	homozygous for both recessi		
-	ombinations appear. The phe		-
recombinations, is	onionations appear. The phe	anotypic ratio of parental c	
a) 10:6	b) 12:4	c) 9:7	d) 15:1
	it showing the dominant phe	,	uj 13.1
a) Test cross			d) Back cross
,	b) Dihybrid cross blind marries a women, who	· ·	-
sons have colour blind		is pure normarior colour	vision, the chances of them
a) 100%	b) 50:50	c) 0%	d) 75 : 25
,	(TT) is crossed with dwarf pl	,	,
a) All tall plants	(11) is crossed with dwarf pi	b) All dwarf plants	2-generation:
c) Both tall and dwarf	nlants in 1 · 1 ratio	d) Both tall and dwarf p	lants in 3 · 1 ratio
	sorders may be classified in .		
	e mainly determined byC.		
Choose the correct op		in single gene.	
a) A-two, B-chromoso		b) A-two, B-chromosom	al C-inversion
c) A-two, B-chromoso		d) A-three, B-chromoso	
	B phenotype but they are		-
transmission of muta			
Choose the correct op			
a) A-homozygous, B-a		b) A-homozygous, B-un	affected. C-carrier
c) A-heterozygous, B-		d) A-heterozygous, B-af	
	le is tt than they contribute p		
a) T and T gametes	b) tt and TT gametes	c) TT and tt gametes	d) T and t gametes
196. Number of linkage gro		, 0	, ,
a) 2	b) 5	c) 7	d) 9
197. In Mendel's experime	nts with garden pea, round se	eed shape (RR) was domin	ant over wrinkled seeds (rr),
yellow cotyledon (YY)	was dominant over green co	otyledon (yy). What are the	e expected phenotypes in the
F_2 - generation of the o	cross RRYY × rryy?		
a) Only round seeds v	vith green cotyledons	b) Only wrinkled seeds	with yellow cotyledons
c) Only wrinkled see	ls with green cotyledons	d) Round seeds with ye	llow cotyledons and wrinkled
		seeds with yellow co	tyledons
198. BB = for black colour	alleles		
bb = for brown colour	alleles		
Offspring of a cross be	etween a black mouse and bro	own mouse allowed to inte	erbreed than find out the
percentage of black co	at in them		
a) 75%			
b) 50%			
c) Cross is not possibl	e because black and brown n	nouse are different species	
d) 100%			
199. Given pedigree chart i	ndicates		

<u></u>	-		
	-0		
]		
a) Autosomal recessiv	ve trait	b) Y-linkage trait	
c) Autosomal domina		d) Sex linkage recessive	e trait
	bin molecule undergoes poly		
	f RBC from biconcave to elon		
a) Haemophilia	b) Colour blindness	c) Phenylketonuria	d) B-thalassaemia
201. XO type of sex determ	ination is seen in		
a) Man	b) Grasshopper	c) Drosophila	d) Birds
202. TtRr represents (hete	rozygous tall, heterozygous j	pink). If this plant is self cr	ossed then
(T-dominant, t-recess	ive, R-dominant, r-recessive))	
I. 25% plant have red			
II. 25% plant have wh			
III. 50% plant have pi	nk flower		
IV. 50% plant are tall			
Choose the correct op			
a) I and II	b) I, II and III	c) II, III and IV	d) I, II, III and IV
203. Chimera is produced of	lue to	h) Dovorao mutationa	
a) Somatic mutationsc) Lethal mutations		b) Reverse mutationsd) Pleiotropic mutation	
-	ie breeding varieties were se		
a) 12	b) 13	c) 7	d) 15
205. Syndrome stands for	0) 13		uj 15
a) A group of sympton	ns	b) Viral disease	
c) Diseased condition		d) Dwarf organism	
	oup-A and AB will not produ	, 0	oup
a) A	b) AB	c) B	d) 0
207. The genetic deficiency			-
a) Diabetes mellitus		b) Glycosuria	
c) Diabetes insipidus		d) Nephrogenic diabete	es
208. Which of the following	g observation made Mendel i	n refutation of the blendin	g theory of inheritance?
a) Red plant crossed v	with white-the resulting prog	geny was pink	
	ng are not intermediate		
	lifferent type of alleles could	-	
	copies of given gene end up i	n the same gamete	
209. Mutations are general	-		
a) Recessive	b) Polymorphic	c) Lethal	d) dominant
a) Deletion	rome is caused by the chang b) Duplication	c) Inversion	d) translocation
	cated that Mendel's principa		
	: likeB inheritance, sex lii	= =	
Choose the correct op			5.
a) A-animal; B-quanti		b) A-human; B-qualitat	ive
c) A-human; B-quanti		d) A-animal; B-qualitat	
,	wing traits of garden pea stu	<i>,</i> .	
a) Green pod colour	b) Round seed colour	c) Axial flower position	
, ,	male sterility in plants are g	, .	

 214. A distinct mechanism that usually involves a short segment of DNA with remarkable capacity to move from one location in a chromosome to another is called a) DNA replication b) DNA hybridization c) DNA recombination d) DNA transposition 215. When F₁-generation progeny resembles both the parents this is called a) Condominance b) Incomplete dominance c) Both (a) and (b) d) Complete dominance c) Both (a) and (b) d) Complete dominance c) Both (a) and (b) d) Complete dominance c) Both (a) and (b) d) Propositus c) Both (a) and (b) d) Origin 217. Plant which used by Hugo de Vries for mutation experiment was a) Oenothera lamarckiana b) Solanum tuberosum c) Ficus elastica d) None of the above 218. A person is suffering from disease phenylketonuria, which is an autosomal recessive disease. Which of these is lacking in the person? a) Homogentisic acid b) Phenylalanine hydroxylase c) Caeruloplasmin d) Cystine 219. Haemophilia in man is due to a) Sex-linked inheritance b) Sex-linited inheritance c) Sex-inked inheritance d) Primary non-disjunction 220. When a dihybrid cross is fit into a Punnett square with 16 boxes, the maximum number of different phenotypes available, are a) Monosomic b) Trisomic c) Nullisomy d) Polyploidy 222. A man and a woman, who do not show any apparent sign of a certain inherited disease, have seven children (two daughter and five sons). Three of the sons suffer from the given disease but none of the daughters are affected. Which of the following mode of inheritance do you suggest for this disease? a) Autosomal dominant b) Sex -linked dominant <lic) -linited="" li="" recessive<="" sex=""> <li< th=""></li<></lic)>
a) DNA replication b) DNA hybridization c) DNA recombination d) DNA transposition 215. When F1-generation progeny resembles both the parents this is called a) Condominance b) Incomplete dominance c) Both (a) and (b) d) Complete dominance 216. The individual from which a pedigree analysis initiated is called a) Probend b) Propositus c) Both (a) and (b) d) Origin 217. Plant which used by Hugo de Vries for mutation experiment was a) <i>Oenothera lamarckiana</i> b) <i>Solanum tuberosum</i> c) <i>Ficus elastica</i> d) None of the above 218. A person is suffering from disease phenylketonuria, which is an autosomal recessive disease. Which of these is lacking in the person? a) Homogentisic acid b) Phenylalanine hydroxylase c) Caeruloplasmin d) Cystine 219. Haemophilia in man is due to a) Sex-linked inheritance b) Sex-limited inheritance c) Sex-influenced inheritance d) Primary non-disjunction 220. When a dihybrid cross is fit into a Punnett square with 16 boxes, the maximum number of different phenotypes available, are a) 8 b) 4 c) 2 d) 16 221. 2 <i>n</i> -2 is known as a) Monosomic b) Trisomic c) Nullisomy d) Polyploidy 222. A man and a woman, who do not show any apparent sign of a certain inherited disease, have seven children (two daughter and five sons). Three of the sons suffer from the given disease but none of the daughters are affected. Which of the following mode of inheritance do you suggest for this disease? a) Autosomal dominant b) Sex – linked dominant c) Sex – linked dominant c) Sex – linked dominant c) Sex – linked recessive d) Sex – linked dominant c) Sex – linked recessive d) Sex – linked dominant c) Sex – linked se suggest for this disease? 223. Colourblindness is caused due to
a) Condominance b) Incomplete dominance c) Both (a) and (b) d) Complete dominance dominance c) Both (a) and (b) d) Complete dominance dominance dominance dominance c) Both (a) and (b) d) Origin d) d) Propositus c) Both (a) and (b) d) Origin d) d) Propositus c) Both (a) and (b) d) Origin d) d) One of the above d) d) One of the above d) None of the above d) d) Origin d)
c) Both (a) and (b) d) Complete dominance 216. The individual from which a pedigree analysis initiated is called a) Probend b) Propositus c) Both (a) and (b) d) Origin 217. Plant which used by Hugo de Vries for mutation experiment was a) <i>Oenothera lamarckiana</i> b) <i>Solanum tuberosum</i> c) <i>Ficus elastica</i> d) None of the above 218. A person is suffering from disease phenylketonuria, which is an autosomal recessive disease. Which of these is lacking in the person? a) Homogentisic acid b) Phenylalanine hydroxylase c) Caeruloplasmin d) Cystine 219. Haemophilia in man is due to a) Sex-linked inheritance d) Cystine 210. When a dihybrid cross is fit into a Punnett square with 16 boxes, the maximum number of different phenotypes available, are a) 8 b) 4 c) 2 d) 16 221. 2 <i>n</i> -2 is known as a) Monosomic b) Trisomic c) Nullisomy d) Polyploidy 222. A man and a woman, who do not show any apparent sign of a certain inherited disease, have seven children (two daughter and five sons). Three of the sons suffer from the given disease but none of the daughters are affected. Which of the following mode of inheritance do you suggest for this disease? a) Autosomal dominant c) Sex -linked dominant c) Sex -linked dominant c) Sex -linked recessive disease due to
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217. Plant which used by Hugo de Vries for mutation experiment was a) 0 a) 0enothera lamarckiana b) Solanum tuberosum c) Ficus elastica d) None of the above 218. A person is suffering from disease phenylketonuria, which is an autosomal recessive disease. Which of these is lacking in the person? a) Homogentisic acid b) Phenylalanine hydroxylase c) Caeruloplasmin d) Cystine 219. Haemophilia in man is due to a) Sex-linked inheritance b) Sex-limited inheritance c) Sex-linked inheritance d) Primary non-disjunction 220. When a dihybrid cross is fit into a Punnett square with 16 boxes, the maximum number of different phenotypes available, are a) 8 b) 4 c) 2 d) 16 221. 2n-2 is known as a) Monosomic b) Trisomic c) Nullisomy d) Polyploidy 222. A man and a woman, who do not show any apparent sign of a certain inherited disease, have seven children (two daughter and five sons). Three of the sons suffer from the given disease but none of the daughters are affected. Which of the following mode of inheritance do you suggest for this disease? a) Autosomal dominant b) Sex -linked dominant c) Sex -linked dominant c) Sex -linked dominant c) Sex -linked d
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c) Sex –limited recessive d) Sex –linked recessive 223. Colourblindness is caused due to
223. Colourblindness is caused due to
a) Recessive female chromosome h) Dominant female chromosome
c) Dominant male chromosome d) linkage
224. Which principle/law has been called the 2nd law of inheritance?
a) Law of independent assortment b) Law of segregation
c) Law of dominance d) Law of paired factor
225. Mendel's experiment were based on hybridization between two plants differing in
a) A pair of contrasting character
b) Three pairs of contrasting character
c) Many pairs of contrasting character d) None of the above
226. Alleles can be similar as in the case ofA likeB or can be dissimilar as in the case ofC likeD
Choose the correct option for A,B,C and D
a) A-heterozygous, B-T T or T t, C-homozygous, D-T T
b) A-homozygous, B-T T or t t, C-heterozygous, D-T t
c) A- homozygous, B-T t , C- heterozygous , D-T T
d) A- homozygous, B-T t , C- heterozygous , D-t t
227. The Barr body is observed in
a) Basophils of male b) Neutrophils of female
c) Basophils of female d) Eosinophils
228. The phenotypic ratio of a monohybrid cross in F_2 -generation is

- a) 3:1
 b) 1:2:1
 c) 2:1:1
 d) 9:3:3:1

 229. Total number of wrinkled seed in previous question
 c) 2
 d) 1
- 230. This pedigree is of a rare trait, in which children have extra fingers and toes. Which one of the following patterns of inheritance is consistent with this pedigree?



a) Autosomal recessive

c) Y-linkage

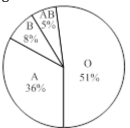
b) Autosomal dominant

d) Sex -linked recessive

231. If a colourblind woman marries a normal visioned man, their sons will be

- a) All normal visioned
- b) One half colourblind and one half normal
- c) Three-fourth colourblind and one-fourth normal
- d) All colourblind
- 232. Barr body is produced due to partial inactivation of one X-chromosome in female. This is called
 - a) Dosage compensation

- b) Facultative heterochromatisation
- c) Both (a) and (b) d) None of the above
- 233. Percentage of blood groups in India is given in the diagram below. Choose the correct option from the given statements



- a) Only 10% of individuals are heterozygous for blood group alleles
- b) Group A is the most common as it is the homozygous recessive group
- c) The alleles for blood group A and O are dominant to the allele for blood group O
- d) Any individual, selected at random from the sample population, has a 1 in 20 chance of being blood group AB

234. find out the genotype of father and mother is the given pedigree chart

	1 0
Mother Father	
a) AA AA	b) Aa Aa
c) AA aa	d) aa Aa
235. Analysis of traits of several generation of a family i	n the form of diagram is called
a) Gene analysis	b) Chromosome analysis
c) Allele analysis	d) Pedigree analysis
236. Among the following which one is the mutagenic ag	gent?
a) Visible light b) Penicillin	c) Formalin d) Water vapour
237. Frameshift mutation and base pair substitution cha	anges the
a) Nucleotide structure	b) Nucleotide sequence
c) Nucleoside sequence	d) Sugar phosphate sequence
238. A women with blood-O has a child with blood grou	p-0. She claims that a man with blood group-A is the

a) 100 b) 1 ^A 1 ^B c) 1 ^A 1 ^B c) 1 ^A 1 ^D d) 1 ^B 1 ^D 239. The terminal end of chromosomes is called a) Centromere b) Telomere c) Chromomere d) metamere 240. Mendel conducted experiments for a) 1 years b) 6 years c) 5 years d) 4 years 241. Cross between unrelated group of organisms, is called a) Hybridization b) Test cross c) Back cross d) heterosis 242. If <i>AAbb x aabB</i> , then phenotypic ratio of its progeny will be a) 9:3:3:1 b) 1:2:1 c) 1:1:1:1 d) 4:1 243. I. Short statured with small round head II. Furrowed tongue and partially opened mouth III. Partowed tongue and partially opened mouth III. Furrowed tongue and partially opened mouth III. Partowed tongue and partially opened mouth III. Partowed tongue and partially opened mouth III. Partowed to haracteristic pain crease IV. Slow physical, psycomotor and mental development These are the characters of a) Down's syndrome b) Turner's syndrome c) Klinefelter's syndrome d) Edward syndrome 244. Which of the following statements are false? II. A recessive allele do to shows its effects when paired with a dominant allele III. A recessive allele do not shows its effects when paired with a dominant allele III. A recessive allele on thows its effects when paired with a dominant allele III. A recessive allel do not shows its effects when paired with a dominant allele III. A recessive allel as ways better for an organism a) II. I and IV b) II, III and IV c) I, II and III d) I, III and IV 245. Following pedigree chart shows 247. Which of the following is not a hereditary disease? a) Certhism Allows and Inkaegs m) dexcual dimorphism d) Genotype and environment interactions 247. Which of the following is not a hereditary disease? a) Certhism b) Dy Cystic Ibroosis c) Thalassenia d) Ilaemophilia 248. F.progeny of a cross between pure tall and dward plant is slaways a) Tall b) Short c) Cystic Ibroosis c) of flasteris syndrome c) EWitonomental tends gene is one which a) Allows the organism to survive but not reproduce b) Determines ex of of	father of her child. What we	ould be the genotype of th		ıt?
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III. Palm is broad with characteristic palm crease IV. Slow physical, psycomotor and mental development These are the characters of a) Down's syndrome b) Turner's syndrome c) Klinefelter's syndrome d) Edward syndrome 2344. Which of the following statements are false? I. A Dominant allele determines the phenotype when paired with a recessive allele II. A recessive allele is weaker than a dominant allele II. A recessive allele do not shows its effects when paired with a dominant allele II. A recessive allele is always better for an organism a) I. I and IV b) II. III and IV c) J. II and III d) I, III and IV 245. Following pedigree chart shows II. Processive and autosomal b) Recessive and autosomal b) Recessive and autosomal c) Dominant and asex-linked d) Dominant and asex-linked d) Dominant and asex-linked d) Dominant and autosomal 246. Phenotype of an organism is the result of a) Mutations and linkages b) Cytoplasmic effects and nutrition c) Environmental changes and sexual dimorphism d) Genotype and environment interactions 247. Which of the following is not a hereditary disease? a) Tall b) Short	II. Furrowed tongue and pa	artially opened mouth		
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(out of total 16 resulted)		=		
		d in a cross between pure	yellow round and pure gre	een wrinkled seeds in F_2 is
a) 9 b) 12 c) 11 d) 10		1) 10) 11	D 10
	aj 9	DJ 12	CJ 11	aj 10

	ed toA and unlinked gen on for A and B	e is related toB	
a) A-linkage; B-cross		b) A-crossing over; B-l	inkage
c) A-crossing over; E	-	d) A-recombination; B	_
253. The linkage map of X	K-chromosomes of fruit fly h	nas 66 units, with yellow bod mbination frequency betwee	
should be			
a) ≤ 50%	b) 100%	c) 66%	d) >50%
-	,	henotypes may be the correc	,
a) 22 pairs+XXY ma	les	b) 22 pairs+XX female	S
c) 22 pairs+XXXY fe	males	d) 22 pairs+X females	
255. Experimental evider	ice of chromosomal theory	of inheritance was given by	
a) HT Morgan	b) TH Morgan	c) H de Vries	d) DH Vries
, ,	, 0	,	another allele may function as
a) Normal allele		b) Non-functional allel	
c) Normal but less ef	fficient allele	d) All of the above	
			group of their children will be
a) Only A	b) A or B or AB or O	c) Only O	d) Only B
, ,	inheritance were enunciate		
a) Mendel	b) Morgan	c) Bateson	d) Punnett
259. Mendel's law was ex	, ,	,	5
a) Meiosis	b) Mitosis	c) Both (a) and (b)	d) None of these
260. Which statement abo	,		
a) His discoveries co time	oncerning genetic inheritand	ce were generally accepted b	y scientific community at his
b) He discovered lin	lago		
	Kage		
c) He believed that g	0	usually blend in their childr	en
	0	-	en
	genetic traits of parents will out genetics apply usually to	-	en
d) His principles abo	genetic traits of parents will out genetics apply usually to	-	en d) Inversions
d) His principles abo 261. The loss of chromoso	genetic traits of parents will out genetics apply usually to omal segment is due to b) Deletions	plants and animals	
d) His principles abo 261. The loss of chromoso a) Polyploidy	genetic traits of parents will out genetics apply usually to omal segment is due to b) Deletions	plants and animals	
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d) His principles above 261. The loss of chromoso a) Polyploidy 262. Symbol <i>A</i> , <i>B</i> and <i>C</i> in $\bigcirc_A \bigotimes_B ($ a) Carrier female 263. The chromosomal co	genetic traits of parents will out genetics apply usually to omal segment is due to b) Deletions dicates b) Effected female ondition in Turner's syndro	c) Duplications c) Duplications c) Death of female me is	d) Inversions d) Normal female
d) His principles above 261. The loss of chromoso a) Polyploidy 262. Symbol <i>A</i> , <i>B</i> and <i>C</i> in A B C in <i>A</i> B C a) Carrier female 263. The chromosomal co a) 21 trisomy with X	genetic traits of parents will out genetics apply usually to omal segment is due to b) Deletions idicates b) Effected female ondition in Turner's syndro	c) Duplications c) Duplications c) Death of female me is b) 44 autosomes + XX	d) Inversions d) Normal female
d) His principles above 261. The loss of chromoso a) Polyploidy 262. Symbol <i>A</i> , <i>B</i> and <i>C</i> in A B C in <i>A B B C</i> a) Carrier female 263. The chromosomal co a) 21 trisomy with X c) 44 autosomes + X	genetic traits of parents will out genetics apply usually to omal segment is due to b) Deletions dicates b) Effected female ondition in Turner's syndro cy	c) Duplications c) Death of female me is b) 44 autosomes + XX d) 18 trisomy with XY	d) Inversions d) Normal female
d) His principles above 261. The loss of chromoso a) Polyploidy 262. Symbol <i>A</i> , <i>B</i> and <i>C</i> in A B C in <i>A B B C</i> a) Carrier female 263. The chromosomal co a) 21 trisomy with X c) 44 autosomes + X	genetic traits of parents will out genetics apply usually to omal segment is due to b) Deletions dicates b) Effected female ondition in Turner's syndro (Y (YY) d group and his father is of	c) Duplications c) Death of female me is b) 44 autosomes + XX d) 18 trisomy with XY B blood group, the genotype	d) Inversions d) Normal female Y of father is
d) His principles above 261. The loss of chromoso a) Polyploidy 262. Symbol <i>A</i> , <i>B</i> and <i>C</i> in A B C in <i>A B B C</i> a) Carrier female 263. The chromosomal co a) 21 trisomy with X c) 44 autosomes + X	genetic traits of parents will out genetics apply usually to omal segment is due to b) Deletions dicates b) Effected female ondition in Turner's syndro cy	c) Duplications c) Death of female me is b) 44 autosomes + XX d) 18 trisomy with XY B blood group, the genotype	d) Inversions d) Normal female
d) His principles above 261. The loss of chromoson a) Polyploidy 262. Symbol <i>A</i> , <i>B</i> and <i>C</i> in A B $Ca) Carrier female263. The chromosomal cona) 21 trisomy with Xc) 44 autosomes + X264. If a child is of 0 blooda) I0I0$	 b) Effected female c) ondition in Turner's syndro c) Y c) Y c) Y c) Y c) A c) A	c) Duplications c) Duplications c) Death of female me is b) 44 autosomes + XX d) 18 trisomy with XY B blood group, the genotype c) ^{IO} I ^B	d) Inversions d) Normal female Y of father is
d) His principles above 261. The loss of chromoson a) Polyploidy 262. Symbol <i>A</i> , <i>B</i> and <i>C</i> in \bigcirc_A \bigotimes_B \bigcirc_Q a) Carrier female 263. The chromosomal con a) 21 trisomy with X c) 44 autosomes + X 264. If a child is of 0 blood a) I ⁰ I ⁰ 265. Work of Beadle and C	b) Effected female bout group and his father is of b) I and his father is of b)	c) Duplications c) Duplications c) Death of female me is b) 44 autosomes + XX d) 18 trisomy with XY B blood group, the genotype c) ^{IO} I ^B ssa proved that	d) Inversions d) Normal female Y of father is d) ^{I⁰I^A}
d) His principles above 261. The loss of chromoson a) Polyploidy 262. Symbol <i>A</i> , <i>B</i> and <i>C</i> in \bigcirc <i>A</i> \bigotimes <i>B</i> \bigcirc <i>G</i> a) Carrier female 263. The chromosomal condition a) 21 trisomy with X c) 44 autosomes + X 264. If a child is of O blood a) I ^O I ^O 265. Work of Beadle and C	b) Effected female bout group and his father is of b) I and his father is of b)	c) Duplications c) Duplications c) Death of female me is b) 44 autosomes + XX d) 18 trisomy with XY B blood group, the genotype c) ^{IO} I ^B ssa proved that b) Blending inheritanc	d) Inversions d) Normal female Y of father is d) ^{I⁰I^A}
d) His principles above 261. The loss of chromoson a) Polyploidy 262. Symbol <i>A</i> , <i>B</i> and <i>C</i> in $\bigcirc A$ $\bigotimes B$ $\bigcirc A$ a) Carrier female 263. The chromosomal condition a) 21 trisomy with X c) 44 autosomes + X 264. If a child is of O blood a) I ^O I ^O 265. Work of Beadle and C a) Complementary g c) Multiple allels	b) Effected female bout group and his father is of b) I and his father is of	c) Duplications c) Duplications c) Death of female me is b) 44 autosomes + XX d) 18 trisomy with XY B blood group, the genotype c) ^{IO} I ^B ssa proved that b) Blending inheritanc d) psedoalleles	d) Inversions d) Normal female Y of father is d) ^{I⁰I^A}
d) His principles above 261. The loss of chromoson a) Polyploidy 262. Symbol <i>A</i> , <i>B</i> and <i>C</i> in $\bigcirc A$ $\bigotimes B$ $\bigcirc A$ a) Carrier female 263. The chromosomal condition a) 21 trisomy with X c) 44 autosomes + X 264. If a child is of 0 blood a) I ^O I ^O 265. Work of Beadle and C a) Complementary g c) Multiple allels 266. The F ₂ -generation of	genetic traits of parents will out genetics apply usually to omal segment is due to b) Deletions idicates b) Effected female ondition in Turner's syndro cy CY d group and his father is of b) ^{IAIB} Tatum on <i>Neurospora cras</i> genes	c) Duplications c) Duplications c) Death of female me is b) 44 autosomes + XX d) 18 trisomy with XY B blood group, the genotype c) ^{IO} I ^B ssa proved that b) Blending inheritanc d) psedoalleles incomplete dominance, exhil	d) Inversions d) Normal female Y of father is d) ^{IO} I ^A e
d) His principles above 261. The loss of chromoson a) Polyploidy 262. Symbol <i>A</i> , <i>B</i> and <i>C</i> in A B $Ca) Carrier female263. The chromosomal conditiona) 21 trisomy with Xc) 44 autosomes + X264. If a child is of 0 blooda) IOIO265. Work of Beadle and Ca) Complementary gc) Multiple allels266. The F2-generation ofa) Variable genotypi$	 genetic traits of parents will out genetics apply usually to omal segment is due to b) Deletions dicates b) Effected female ondition in Turner's syndro (Y) (YY) d group and his father is of b) I^AI^B Tatum on <i>Neurospora cras</i> genes ffspring in a plant showing c and phenotypic ratio 	c) Duplications c) Duplications c) Death of female me is b) 44 autosomes + XX d) 18 trisomy with XY B blood group, the genotype c) ^{IO} I ^B ssa proved that b) Blending inheritanc d) psedoalleles incomplete dominance, exhil b) a genotypic ratio of	d) Inversions d) Normal female Y of father is d) ^{IO} I ^A e
d) His principles above 261. The loss of chromoson a) Polyploidy 262. Symbol <i>A</i> , <i>B</i> and <i>C</i> in $\bigcirc A$ $\bigotimes B$ $\bigcirc A$ a) Carrier female 263. The chromosomal condition a) 21 trisomy with X c) 44 autosomes + X 264. If a child is of 0 blood a) I ^O I ^O 265. Work of Beadle and C a) Complementary g c) Multiple allels 266. The F ₂ -generation of	b) Effected female b) Effected female b) Deletions b) Effected female b) Effected female b) Effected female b) Effected female b) Effected female b) I ^A I ^B Tatum on <i>Neurospora cras</i> genes ffspring in a plant showing c and phenotypic ratio b) of 3 : 1	c) Duplications c) Duplications c) Death of female me is b) 44 autosomes + XX d) 18 trisomy with XY B blood group, the genotype c) ^{IO} I ^B ssa proved that b) Blending inheritanc d) psedoalleles incomplete dominance, exhil b) a genotypic ratio of	d) Inversions d) Normal female Y of father is d) ^{IO} I ^A e

a) Paternal characters		b) Maternal characters		
c) Parental characters	c) Parental characters		d) Little paternal and more maternal characters	
268. Type of gamete participat	ing in selfing of members i	in monohybrid cross are of		
a) One type	b) Two type	c) Four type	d) Many type	
269. In sickle-cell anaemia, GA	G is replaced by			
a) GGA	b) GUG	c) AAG	d) GGG	
270. Genes, when close togethe	er on a chromosome, are k	nown as	-	
a) Linkage	b) Mutation	c) Translation	d) transscription	
271. If a character is always tra	ansmitted directly from a f	ather to all his sons and fro	m their sons to all their	
sons, then which chromos	some carries the gene for t	he character?		
a) Autosome	b) X-chromosome	c) Y-chromosome	d) None of these	
272. Hugo de Vries is famous fo	or			
a) Natural selection theor	У	b) Mutation theory		
c) Organic theory		d) Chemical theory		
273. Jumping genes in maize w	vere discovered by			
a) Hugo de Vries	b) Barbara McClintock	c) T H Morgan	d) Mendel	
274. A plant of F ₁ - generation l	has genotype 'AABbCC'. Or	n selfing of this plant, the ph	enotypic ratio in F ₂ -	
generation will be				
a) 3:1		b) 1 : 1		
c) 9:3:3:1		d) 27 : 9 : 9 : 9 : 3 : 3 : 3 :1		
275. The character that is expr	essed in the F ₁ -generation	is called the		
a) Recessive character		b) Dominant character		
c) Codominant character		d) None of these		
276. Chromosomal theory of ir	heritance was proposed b	у		
a) Gregor Mendel	b) Hugo de Vries	c) Bridges	d) Sutton and Boveri	
277. In sex linkage, the special	ity is			
a) Atavism		b) Criss-cross inheritance	<u>)</u>	
c) Reversion		d) Gene flow		
278. Mother = A blood group				
Father = AB blood group				
The child will not have				
a) A blood group	b) O blood group	c) B blood group	d) A blood group	
279. The chromosome constitu	-	-		
a) Monosomic	b) Nullisomic	c) Haploid	d) trisomic	
280. Phenylalanine does not ch	nanged to tyrosine. This co			
a) Sickle-cell anaemia		b) Phenylketonuria		
c) Thalassaemia		d) Haemophilia		
281. When the chromosome nu		has one additional chromo	osome in one of the	
homologous pairs, the add				
a) Trisomy	b) Monosomy	c) Polyploidy	d) nullisomy	
282. What type of gametes will				
a) ry, rY	b) RY, Ry	c) Ry, Yy	d) RR, Yy	
283. The term 'genetics' was in		24004	D 4000	
a) 1906	b) 1905	c) 1904	d) 1903	
284. Mutant genes that give sli				
a) Heteroalleles	b) Recessive alleles	c) Isoalleles	d) Dominant alleles	
285. Which of the following is a	-		d) Conorribation	
a) AIDS	b) Colour blindness	c) Syphilis	d) Gonorrhoea	
286. Mutations can be induced		c) Camma radiationa	d) Infra rad radiations	
a) IAA 287. Which of the following is a	b) Ethylene	c) Gamma radiations	d) Infra red radiations	
207. WINCH OF THE IOHOWING IS a				

a) Leprosy	b) Goitre	c) AIDS	d) Albinism
288. The title of Mendel's pap	per, while presenting at Bru	ınn Natural History Society	r in 1865 was
a) Laws of inheritance		b) Laws of heredity	
c) Experiments on pea p	lants	d) Experiments on plant	hybridisation
289. XO chromosomal abnor	nality in humans causes		
a) Turner's syndrome		b) Down's syndrome	
c) Darwin's syndrome		d) Klinefelter's syndrom	e
290. Milk secretion and baldr	ess, both the traits belong		
a) Sex limited	b) Sex linked	c) Sex influenced	d) Autosomal traits
291. The daughter born to ha	emophilic father and norm		,
a) normal	b) Carrier	c) Haemophilic	d) None of these
292. Blood grouping is the ex		, I	,
a) Multiple allele	1	b) Condominance	
c) Both (a) and (b)		d) Independent assortm	ent
293. A true breeding plant pr	oducing red flowers is cros	, ,	
01 1		he plants of first filial gener	
	flowers in the progeny wou		
			1
a) $\frac{3}{4}$	b) $\frac{1}{4}$	c) $\frac{1}{3}$	d) $\frac{1}{2}$
294A is sex linked recess	sive disease. Which shows i	5	female toC progeny.
Choose the correct optic			
a) A-haemophilia, B-cari		b) A-cystic fibrosis, B-ca	rrier, C-male
c) A-sickle-cell anaemia		d) A-phenylketonuria, B	
295. Crossing over is advanta			
a) Variation	b) Linkage	c) Inbreeding	d) Stability
296. Father of 'genetics' is	o) zimege	oj morecumg	a) outonity
a) De Vries	b) Mendel	c) Bateson	d) Robert Hooke
297. The recessive gene that	•	-	aj 11000101100110
a) Pleiotropic gene		b) Complementary gene	
c) Holandric gene		d) Supplementary gene	
298. When different alleles of	f the same gene are presen	, 11 , 0	ridual is a
a) Heterozygous	b) Diploid	c) Homozygous	d) mosaic
299. Sex linked traits are the		0) 1101110298040	aj modulo
a) Sex chromosome	b) Autosomes	c) Allosomes	d) All of these
300. Number of Barr body in	,		
a) 1	b) 2	c) 3	d) 4
301. 21 trisomy in humans ca			uj i
a) Klinefelter's syndrom		b) Down's syndrome	
c) Tumer's syndrome		d) Patau's syndrome	
302. Paternal baldness, mous	taches and heard in human	· ·	
a) Sex differentiating tra		b) Sex determining trait	c
c) Sex-linked traits	1105	d) Sex-influenced traits	5
303. Polytene chromosomes	in calivary glands of Droso	-	tof
a) Endoduplication	in Sanvary gianus of DT050	b) Duplication without s	
c) Replication of DNA w	ithout call division	d) All of the above	
304. A normal women whose			The cone would be
a) 75% colourblind	b) 50% colourblind	c) All normal	d) All colourblind
305. Match the symbols with	Statement		

1. □ 5. ☉

2.0 <u>6.</u>				
2.0 6. 3. 				
4.				
A. Diseased (death)				
B. Carrier (female) of X-link	ked recessive gene			
C. Marriage in blood relative	es			
D. Unknown sex				
Codes				
A B C D				
a) 1 2 3 4		b) 6 5 7 4		
c) 2 1 3 4		d) 6 2 3 4		
306. The female children of haen	nophilic father and carrie			
a) All haemophilic		b) Half haemophilic, half	carrier	
c) All normal	ntify hotopogygous indivi	d) All carrier		
307. Genetic counsellors can ideaa) Height of individuals	nuly neterozygous mulvi	b) Colour of individuals		
c) Screening procedures		d) All of these		
308. How many conditions exhibit	nit in dissimilar sex chron	,		
-	b) 3	c) 4	d) 5	
309. Number of chromosomes in	,		uj b	
	b) 47	c) 48	d) 49	
310. Which phenomena leads to	,	,	,	
-	b) Linkage	c) Both (a) and (b)	d) Mitosis	
311. External morphology or app	pearance or descriptive te	erm of an genotype is calle	d	
a) Genotype b	b) Phenotype	c) Both (a) and (b)	d) None of these	
312. In which of the following, th	here is no defect in the sex	x chromosome?		
a) Turner's syndrome		b) Down's syndrome		
c) Colour blindness		d) Klinefelter's syndrome		
313. The traits which are not exp are	pressed due to a particula	ar gene but are expressed b	y products of sex hormones	
a) Sex influenced traits	b) Autosomal traits	c) Allosomic traits	d) Sex linked traits	
314. Choose the correct option for	or the chromosomal disor	rders		
I. Colour blindness				
II. Down's syndrome				
III. Phenylketoria				
IV. Turner's syndrome				
V. Thalassaemia				
-	b) II, IV and V	c) III, IV and V	d) II and IV	
315. First time who used the term a) Alfred Sturtevant	b) Alfred Nobel		d) Mondol	
316. Who postulated the mutation	,	c) Pasteur	d) Mendel	
_	b) Darwin	c) Lamarck	d) Hugo de Vries	
317. Choose the chemical used in			aj mago ac vinco	
a) Polyethylene glycol	in an entiteral polypiolay	b) Sodium alginate		
c) Acenaphthene		d) Sodium hypochlorite		
318. Linkage groups are always	present on the	,		
a) Homologous chromosom				
b) Analogous chromosomes				

c) Sex chromosomes		
d) Heterologous chromosomes		
^{319.} Sex determination in an organism is given by $\frac{x}{A} = 1$.5, then organism will be	
a) Male b) Female	c) Super female	d) Intersex
320. Emasculation is the removal of		
a) Flower buds	b) Anthers before dehi	scence
c) Carpels before dehiscence	d) Mature flowers	
321. The genes, which remain confined to differential re	gion of Y-chromosome, a	re
a) Autosomal genes b) Holandric genes	c) Sex-linked genes	d) Mutant genes
322. Study the pedigree chart given below.		
What does it show?		
a) Inheritance of a sex- linked inborn error of	b) Inheritance of a con	dition like phenylketonuria as
metabolism like phenylketonuria	an autosomal recess	
c) The pedigree chart is wrong as this is not possib		
323. Mutation cannot change	1	
a) RNA b) Environment	c) Enzyme	d) DNA
324. One of the parents of a cross has mutation in its mit		•
During segregation of F ₂ -progenies that mutation is		-
a) One –third of the progenies	b) None of the progeni	es
c) All of the progenies	d) Fifty per cent of the	progenies
325. Mendel does not get linkage due to		
a) Dominance	b) Independent assort	ment
c) Segregation	d) Genes on same chro	omosome
326. Frameshift mutation arises due to		
a) Deletion of base pair of DNA	b) Insertion of base pa	ir of DNA
c) Both (a) and (b)	d) Change in single bas	se pair of DNA
327. Genes A, B and C are linked. Genes A and B are mor	e close than A and C. Find	l out the correct option for the
given statements		
I. A might be before B and C		
II. B might be between A and C		
III. C might be between A and B		
IV. More crosses has occurred between A and C tha		
a) I and II b) II and III	c) III and IV	d) I, II and IV
328. In previous question find out the ratio between rou		
a) 3 : 1 b) 2 : 2	c) 1:1	d) 9 : 6 : 1
329. Which of the following blood groups' person can no		
a) AB blood group b) O blood group	c) A blood group	d) B blood group
330. Which of the following is not related to sex chromo		
a) Turner's syndrome	b) Klinefelter's syndro	
c) Down's syndrome	d) Haemophilia and co	
331. Inheritance of characters not located in the gene bu to	it the young one resembli	ing only the female part is due
a) Cytoplasmic inheritance	b) Chromosomal inher	ritance
c) Plastid inheritance	d) epigenesis	
332. Mendel found the phenotype of the F ₁ heterozygote	e Tt was to be exactly like	theA parent in

appearance, he proposed that in a pair of dissimilar	factors, one dominates the other (as in the F_1) and
hence is called theB factor, while the other factor	,
Choose the correct option for A, B and C	
a) A-T T, B-dominant, C-recessive	b) A-T t, B-dominant, C-recessive
c) A-t t, B-dominant, C-recessive	d) A-T t, B-Recessive , C-dominant
333. Which of the following pairs of features is a good ex	ample of polygenic inheritance?
a) Human height and skin colour	ABO blood groups in humans and flower colour of <i>Mirabilis jalapa</i>
c) Hair pigment of mouse and tongue rolling in	d) Humans eye colour and sickle cell anaemia
humans	
334. Find the phenotype of A, B, C, D from given cross (R	-Red and $r =$ white)
$R r \times R r$	
(A) (B) (C) (D)	
a) A-Red, B-Red, C-Red, D-White	b) A-Red, B-Red, C-White , D-White
c) A-Pink, B-Red, C-White, D-White	d) A-Pink, B-Red, C-Red, D-White
335. Incomplete dominance is shown by	
a) Primrose b) <i>Mirabilis</i>	c) <i>Helianthus</i> d) China rose
336. Which of the following genes show the hetertozygou	us condition?
a) Rr b) RR	c) Rr d) None of these
337. Rrrr (progeny): Red (dominant) flowers (heterozyg	gous) were crossed with white flower. The result will
be	
a) $350 \rightarrow \text{Red} : 350 \rightarrow \text{white}$	b) $450 \rightarrow \text{Red} : 250 \rightarrow \text{white}$
c) $380 \rightarrow \text{Red} : 320 \rightarrow \text{white}$	d) None of the above
338. A common test to find the genotype of a hybrid is by	
a) Crossing of one F_2 -progeny with male parent	b) Crossing of one F_2 -progeny with female parent
c) Studying the sexual behaviour of F_1 -progenies	d) Crossing of one F_1 -progeny with male parent
339. Which of the following has the least number of chro	
a) <i>Amoeba</i> c) <i>Pheretima</i>	b) <i>Drosophila</i> d) <i>Ascaris megalocephala</i>
340. In given genetic basis of human blood group table fi	, , , , , , , , , , , , , , , , , , , ,
S.no. Allele Allele Genotype	nd out which belongs to blood group A, B, Ab and O
from from of	
Parent Parent Offspring	
1 2 s	
I. IA IA IAIA	
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	
VII. i i ii	
A B AB O	
a) I,III V,VI II,IV VII	b) I,IV VI,II II,III V
c) VII II,IV V,VI I,II	d) I,III II,IV V,VI VII

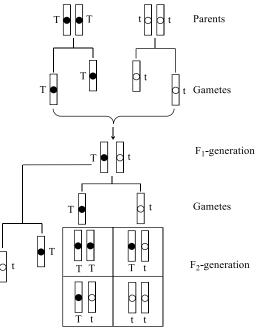
341. The chemical nature of chromatin is as follows

Ι

a) Nucleic acids

b) Nucleic acid and histone proteins

- c) Nucleic acids, histone and non-histone proteins
- d) Nucleic acids and non-histone proteins
- 342. What does this diagram indicate?



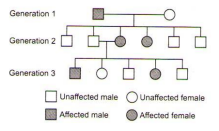
a) Law of dominance interpretated on basis of genes

b) Law of segregation interpretated on basis of genes

c) Law of independent assortment interpretated on basis of genes

d) Simply gamete genes

343. Given below is a pedigree chart showing the inheritance of a certain sex- linked trait in humans.



The trait traced in the above pedigree chart is

a) Dominant X-linked
 b) Recessive X-linked
 c) Dominant Y-linked
 d) Recessive Y-linked
 344. Mendel observed that certain character did not assort independently. Later, scientist found that this is due to

a) Linkage in traits

b) Crossing over

d) Dominance of one trait over the other

345. Identify the symbols given below and the correct option with respect to *A*, *B* and *C*

c) Both (a) and (b)

 a) A-Male, B-Female, C-Sex unspecified 		b) A-Male, B-Female, C-Sterile		
c) A-Male, B-Female, C-Fertile		ertile	d) A-Female, B-Male, C-Sex unspecified	
346. Mendel investigated characters in garden pea plant that were manifested in two trait			o trait	
a) Simi	lar	b) Non-zygote	c) Identical	d) Opposite
347. Phenyl	ketonuria disease	is a		
a) Autosomal dominant			b) Autosomal recessive	
c) Sex linked recessive			d) Sex linked dominant	
348. The lite	eral meaning of ch	romosome is		
a) Pain	ted body	b) Coloured body	c) Doubling body	d) Thread like body
349. The <i>F</i> ₂ genotypic ratio of monohybride cross is				

a) I ^A I ^A and I ^A I ^O b) I ^O I ^O and I ^O I ^O	O or A blood groups. Whi	1 C I C II I			
		350. The offspring produced from a marriage have only O or A blood groups. Which of the following genotypes			
ajii anali Dj	c) $I^{A}I^{A}$ and $I^{A}I^{O}$	d) $I^{A}I^{O}$ and $I^{A}I^{O}$			
351. In order to find out the different types of gametes	produced by a pea plant h	aving the genotype AaBb, it			
should be crossed to a plant with the genotype					
a) aaBB b) AaBb	c) AABB	d) aabb			
352. The lowest number of chromosomes is found, in w	-				
a) <i>Haplopappus gracilis</i> b) <i>Poa litorosa</i>	c) <i>Salix tetrasperma</i>	d) <i>Ageratum coigzoides</i>			
353. The genes for seven characters of pea plant that we	ere considered in Mendel	nybridisation experiment are			
present on					
a) 4 chromosome b) 5 chromosome	c) 7 chromosome	d) 8 chromosome			
354. Chromosome diagram of the given fruitfly tick the	correct choice for autoson	ie labelled			
00 00					
a) A b) C	c) D	d) B			
355. Identify the wrong statement.	a an abromacama				
a) In male grasshoppers, 50% of the sperms have 1 b) Usually , female birds produce two types of gam		some			
c) The human males have one of their sex chromos					
d) In domesticated fowls, the sex of the progeny de					
356. The chromosome shown in the diagram below is b					
and the genes between these points became invert	-				
	cu				
Chromosome PQRSTUVW consisting					
$\begin{array}{c} P \ Q \ R \ S \ T \ U \ V \ W \end{array} \xrightarrow{\begin{subarray}{c} Chromosome \\ consisting \\ of eight genes \end{array}} \\ \hline \end{array}$					
$\begin{array}{c c} P & Q & R & S & T & U & V & W \\ \hline & \uparrow & \uparrow & \\ Break & Break & \\ \end{array}$					
$\begin{array}{c c} P & Q & R & S & T & U & V & W \\ \hline P & Q & R & S & T & U & V & W \\ \hline \uparrow & \uparrow & \uparrow & \\ Break & Break & \\ \end{array}$ The resulting order of the genes will be		d) WWWTSDDO			
PQRSTUVW Chromosome consisting of eight genes Break Break Break Break Description Description Description Description	c) PQTURSVW	d) VWUTSRPQ			
Chromosome consisting of eight genes Break The resulting order of the genes will be a) PQUTSRVW b) WVUTSRQP 357. Which of these is a dominant factor?	c) PQTURSVW				
Chromosome consisting of eight genes Break The resulting order of the genes will be a) PQUTSRVW b) WVUTSRQP 357. Which of these is a dominant factor? a) Rh factor b) Haemophilia	c) PQTURSVW c) Albinism	d) Colour blindness			
Chromosome consisting of eight genes Break The resulting order of the genes will be a) PQUTSRVW b) WVUTSRQP 357. Which of these is a dominant factor? a) Rh factor b) Haemophilia 358. A person with unknown blood group under ABO sy	c) PQTURSVW c) Albinism vstem, has suffered much l	d) Colour blindness blood loss in an accident and			
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PQRSTUVW Chromosome consisting of eight genes Break Break The resulting order of the genes will be a) PQUTSRVW b) WVUTSRQP 357. Which of these is a dominant factor? a) Rh factor b) Haemophilia 358. A person with unknown blood group under ABO sy needs immediate blood transfusion. His one friend for blood donationwithout delay. What would have a) Type AB b) Type 0	 c) PQTURSVW c) Albinism vstem, has suffered much l who has a valid ceruficate been the type of blood gr c) Type A 	d) Colour blindness blood loss in an accident and e of his own blood type, offers oup of the donor friend? d) Type B			
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- a) Autosomal linked diseasec) Y-chromosomal linked disease
- 363. Two genes R and Y are located very close on the chromosomal linkage map of maize plant. When RRYY and rryy genotypes are hybridized, then F_2 -segregation wii show
 - a) Higher number of the recombinant types
 - c) Segregation in 3 : 1 ratio
- 364. During ...A... both members of chromosome pair as well as ...B... separate and pass to different gametes. Choose the correct option for A and B
 - a) A-mitosis; B-allele pair
 - c) A-allele pair; B-meiosis
- 365. Genetic map is one that
 - a) Shows the stages during the cell division
 - b) Shows the distribution of various species in a region
 - c) Establishes sites of the genes on a chromosome
 - d) Establishes the various stages in gene evolution
- 366. Given below is a highly simplified representation of the human sex chromosomes from a karyotype. The gene 'a' and 'b' could be of
 - N K

Gene 'a'

a) Colour blindness and body height

- c) Haemophilia and red-green colourblindness
- 367. Human females have

from the given cross

- a) 22 pairs of autosomes and one pair of sex chromosome
- b) 21 pairs of autosomes and two pairs of sex chromosome
- c) 23 pairs of autosomes and one pair of sex chromosome
- d) 20 pairs of autosomes and one pair of sex chromosome
- 368. The progenies are found to be male sterile after crossing two plants. This is due to some genes, which are present in

	a) Mitochondria	b) Cytoplasm	c) Nucleus	d) chloroplast
369	. Mutation may results in t	the		
	a) Change in genotype b) Change in phenotype			
	c) Change in metabolism		d) All of these	
370	. In cross between yellow	round (YYRR) and pure bre	eeding pea plants having gro	een wrinkled (yyrr) find
	out the total seeds (plant	s) having yellow colour in	F_2 -generation	
	a) 12	b) 10	c) 14	d) 11
371. A cross in which parents differ in a single pair of contrasting character is callled				
	a) Monohybrid cross	b) Dihybrid cross	c) Trihybrid cross	d) Tetrahybrid cross
372. Calvin bridges demonstrated sex determining factor is the ratio of number of				
a) X-chromosome to autosome b) Autosome to X-chromosome				
c) Y-chromosome to X-chromosome d) Y-chromosome to autosome				
373. Find out the genotype and phenotype of F_1 -generation (R = dominant and red, r = recessive and white)				

- b) A-meiosis; B-allele pair
- d) A-allele pair; B-mitosis
- b) Segregation in the expected 9:3:3:1 ratiod) Higher number of the parental types

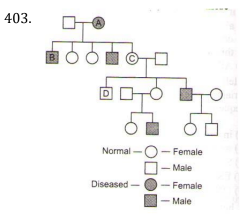
b) Attached ear lobe and rhesus blood group

d) Phenylketonuria and haemophilia

- b) X-chromosomal linked disease
- d) None of the above

с" Р				
$ \begin{array}{ccc} $	ation			
(R) (r) Gameter	5			
F_1 -generation				
a) Rr and white	b) Rr and red	c) Rr and pink	d) Can not predict	
374. Which one of the follo	wing conditions correctly d	lescribes the manner of	determining the sex in the given	
example?				
	omosomes determine male s			
	mans as found in Turner's s		emale sex	
	hromosomes (XX) produce i	=		
	hromosomes, (ZZ) determin ybrid cross (phenotypically			
a) 3 : 1	b) $1:2:1$	c) 9:7	d) 9 : 3 : 3 :1	
376. Trisomy stands for	0,1.2.1	cj 9.7		
a) 2 <i>n</i> – 1	b) 2 <i>n</i> + 2	c) 2 <i>n</i> + 3	d) 2 <i>n</i> + 1	
377. Klinefelter's syndrom	-	-	-	
a) XX egg of Y sperm		b) XX egg and XY s	perm	
c) X egg and YY spern		d) XY egg and X spe		
-	are colourblind with AB blo		_	
=	d with A blood group, and fa			
	th blood group-A, and father			
-	d with blood group-B, and fa			
	ch blood group-A, and fathe g chromosomal mutation ar			
chromosomes are und	-	e most inkely to take pla	ace when homologous	
a) Inversion and tran		b) Deletion and du	plication	
c) Inversion and dele		d) Translocation ar	-	
380. What percentage of h	omozygous Rh ⁻ will be born	n amongst four children	of a couple where the husband is	
heterozygous for Rh ⁺	and wife is homozygous for	r Rh ⁻ gene?		
a) 25%	b) 50%	c) 75%	d) 100%	
381. Mendel could not find	•			
-	ed but they are too far apar	t for crossing over to be	distinguished from independent	
assortment	never tested for the same ti	in an ana		
0	ere present on the same chr			
•	ere present on 4 chromosom		t far apart	
Find out the correct o	=			
a) I and II	b) II and III	c) III and IV	d) IV only	
382. Haemophilia is also ca	alled	-		
a) Bleeders disease	b) Blood disease	c) RBC disease	d) All of these	
_	the same chromosome do no	ot separate and are inhe	erited together over its	
generations due to the phenomenon of				
a) Complete linkage	·	b) Incomplete linka	-	
c) Incomplete recombination d) Complete recombination				
384. Universal donor is	$\dots \Omega Rh^{-}$	AR Rh ⁺	$a AB Bh^{-}$	
a) O Rh ⁺	b) ^{O Rh-}	c) ^{AB Rh+}	d) $^{AB} Rh^{-}$	
	ter's syndrome have chrome	osomes		
a) XX	b) XY	c) XXY	d) XYY	

386. Mendel crossed tall and	dwarf plant In Fa-generati	on both the tall and dwarf	nlants were produced. This
shows	awari plant. In 12 generati	on both the tail and awarr	plants were produced. This
a) Blending of character	S	b) Atavism	
c) Non-blending of char		d) Intermediate charact	ers
387. Sex- limited and sex- lin		,	
a) Autosomes	b) X-chromosome	c) Y-chromosome	d) Both (b) and (c)
388. How many different typ		-	
AA BB CC \times aa bb cc			
a) 3	b) 8	c) 27	d) 64
389. Point mutation involves	•	C) 27	
a) Insertion		b) Change in single base	nair
c) Duplication		d) deletion	pan
390. A person with type A blo	od group may safely receiv	•	
a) Type-AB	Sou group may salely received	b) Type-A and type –0	
c) Type-A and type –AB		d) Type-AB and type –0	
391. In which cross will you	tot most piple flowors?	uj Type-AD allu type =0	
a) Red × red	b) Red \times pink	c) Diple v piple	d) Pod x white
	, ,	c) Pink × pink	d) Red \times white
392. <i>Triticale</i> has been prod			d) Dias and mains
a) Wheat and rice	b) Wheat and rye	c) Wheat and aegilops	
393. Which one of the follow	ing characters studied by M		
a) Green seed colour		b) Terminal flower posit	tion
c) Green pod colour		d) Wrinkled seed	
394. Mendel's experimental r			
a) <i>Pisum sativum</i>	b) <i>Lathyrus odoratus</i>	c) <i>Oryza sativa</i>	d) <i>Mirabilis jalappa</i>
395. Which of the following is	s not considered as mutage		
a) Lower temperature		b) X-rays	
c) Higher temperature		d) UV rays	
396. The physical expression	••		
a) Morphology	b) Genotype	c) Phenotype	d) Ecotype
397. Carrier organism refers			
a) Dominant gene, that i		b) Recessive gene, that i	-
c) Recessive gene, that i		d) Dominant gene, that is expressed	
398. In previous question, fin	d out which alphabete (A-I	D) labelled for X and Y-chro	omosome
X Y			
a) A D		b) A,C D	
c) C D		d) B D	
399. In amniocentesis of a pr	egnant woman, it is found t	hat the embryo contains b	oth, Barr body and F-body.
The syndrome likely to l	be associated with the emb	ryo is	
a) Edward' syndrome		b) Down's syndrome	
c) Klinefelter's syndrom	ie	d) Patau's syndrome	
400. In the previous question	, find out the chances of fif	th child to be albino	
a) 1 in 2	b) 1 in 4	c) 1 in 3	d) 1 in 5
401. Three children in a fami	ly have blood types O, AB a	nd B respectively. What are	e the genotypes of their
parents?			
a) I ^A i and I ^B i	b) ^{IAIB} and i i	c) ^{IB} I ^B and I ^A I ^A	d) ^{IA} I ^A and I ^B i
-	,	C)	uj
402. The chromosomal arran	•		
a) Euploidy	b) Aneuploidy	c) Duplication	d) polyploidy



In the above pedigree, assume that no outsider marrying in, carry a disease. Write the genotypes of C and D.

D.	C .			
a) X ^C Y and X ^C X ^C	b) ^{XXC} and XY	c) ^{XY} and X ^C X ^C	d) ^{X^CX^C and X^CX}	
404. The specific pair of chro	mosomes which determine	the sex of the individual ca	lled	
a) Sex chromosomes	b) Allosomes	c) Heterosomes	d) All of these	
405. The 'Cri-du-chat' syndro	ome is caused by change in c	hromosome structure invo	lving	
a) Deletion	b) Duplication	c) Inversion	d) translocation	
406. During his experiments,	Mendel used the term facto	or for		
a) Genes	b) Traits	c) Characters	d) Qualities	
407. In a monohybrid cross in	nvolving incomplete domina	ance, the phenotypic ratio e	equals the genotypic ratio in	
F ₂ - generation. The ratio	o is			
a) 3 : 1	b) 1 : 2 : 1	c) 1:1:1:1	d) 9 : 7	
408. The genome of Caenorh	abditis elegans consists of			
a) 3 million base pairs a	nd 30,000 genes	b) 180 million base pairs	and 13,000 genes	
c) 4.7 million base pairs and 4,000 genes d) 97 million base pairs and 18,000 genes				
409. Albinism is caused by th	e deficiency of			
a) Amylase	b) Tyrosinase	c) Phenylalanine	d) Xanthene oxidase	
410. The ABO blood grouping in human beings is an example for				
I.Dominance				
II.Incomplete dominance				
III.Codominance				
IV.Multiple alleles				
a) I and II	b) II , III and IV	c) I , III and IV	d) III and II	
411. Sickle-cell anaemia is an autosomal linked recessive trait can be transmitted from parents to the offspring				
when both the partners are carrier for all the genes or heterozygous. The disease is controlled by a single				
pair of allele, Hb ^A and H	b ^S . Identify X, Y and Z			

Normal Hb (A) Gene	
····GAC····	
mRNA ···· GÅG ····	
\downarrow	
(Val)-(His)-(Leu)-(Thr)-(Pro)-(Glu)-(Glu)	
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	
Hb ^A Peptide	
·····X····	
Sickle-cell Hb (B)	
GeneY	
mRNA ···· GUG ····	
\sim	
(Val)-(His)-(Leu)-(Thr)-(Pro)-(Z)-(Glu)	
1 2 3 4 5 6 7	
Hb ^S Peptide	
a) GTG GAC Val (GUG)	b) CAC CTC val (GUG)
c) GTA GAG val (GUG)	d) GTC GAC val (GUG)
412. Diploid cells have	
a) Two chromosomes	b) One set of chromosomes
c) Two pairs of homologous chromosomes	d) Two sets of chromosomes
413. Single gene can produce more than one effect. Like	starch synthesis in pea plant. It has two alleles (B and
b) for starch synthesis the phenotypes of which are	e also given below
B B, b b, B b	-
I. BB – round seed, large starch synthesis	
II. bb – wrinkle seed, less starch synthesis	
III. Bb – intermediate size seed, intermediate less s	tarch synthesis
Choose the correct option	
a) I and II b) II and III	c) III and I d) I, II and III
414. After examining the blood groups of husband and	wife, the doctor advised them not to have more than one
child, the blood group of the couple are likely to be	
a) male Rh ⁻ and female Rh ⁺	b) Female Rh ⁻ and male Rh ⁺
c) Male and female Rh ⁺	d) Male and female Rh^-
415. A person with blood group-A has	
a) Antigen-A and antibody-b	b) Antigen-B and antibody-a
c) Both antibodies	d) No antibody and no antigen
416. Which of the following is not a correct match?	
a) Sex determination	– A chromosomal phenomenon
b) Y-chromosome –	Autosomal
c) Red-green colour blindness in human –	A sex-linked character
d) An abnormal chromosome number in each cell -	 A case of polyploidy
417. In law of independent assortment. How many factor	ors are involved? (for a dihybrid cross)
a) 2 b) 3	c) 4 d) 1
418. Mother B homozygous, father A unknown, therefor	e, possible blood group in progeny is
a) AB and B possible b) AB and A possible	c) A and B possible d) O possible
419. Consider the following four statements I, II, III and	IV and select the correct statements
I. Mendelian experiments has a large sampling size	, which gave greater credibility to the data that he
collected	
II. Recessive allele influences the appearance of the	e phenotype even in the presence of a dominant allele

III. Multiple alleles can be found only when population studies are made IV. In F₂-generation of a Mendelian monohybrid cross, the tall and dwarf traits were identical to their parental types and shows blending inheritance The correct statements are a) I and III b) III and IV c) II and IV d) II and III 420. When released from ovary, human egg contain b) Two X-chromosome a) One Y-chromosome c) One X-chromosome d) XY-chromosome 421. The tendency of offsprings to differ from their parents is called a) Variation b) Heredity c) Inheritance d) Resemblance 422. The gene, which controls many characters, is called b) Polygene a) Codominant gene d) Multiple gene c) Pleiotropic gene 423. The given diagram A and B indicates a) A-Zygotic twins; B-Dizygotic twins b) A-Dizygotic twins; B-Identical twins c) A-Zygotic twins; B-Identical twins d) A-Identical twins; B-Dizygotic twins 424. Which of the following statement is/are correct regarding law of segregation? a) Alleles separate with each other during gametogenesis b) The segregation of factors is due to the segregation of chromosomes during meiosis c) Law of segregation is called as law of purity of gametes d) All of the above 425. Which of the following discoveries resulted in a Nobel Prize? a) Recombination of linked genes b) Genetic engineering c) X-rays induce sex-linked recessive lethal d) Cytoplasmic inheritance mutations 426. When alleles of two contrasting characters are present together, one of the character expresses itself during the cross while the other remains hidden. This is the a) Law of purity of gametes b) Law of segregation c) Law of dominance d) Law of independent assortment 427. In which phase of meiosis-I the two chromosome can align at the metaphase plate independently of each other a) Metaphase-II b) Metaphase-I c) Anaphase-I d) Telophase-I 428. When a mutation is limited to the substitution of one nucleotide for another, it is called a) Translocation b) Point mutation c) Base inversion d) Sugar phosphate deletion 429. Types of genotype observed in a dihybrid cross are a) 9 b) 12 c) 4 d) 6 430. In Morgan's experiments on linkage, the percentage of white eyed, miniature-winged recombinants in F₂generation is a) 1.3 b) 37.2 c) 62.8 d) 73.2 431. Which cross was used to study the independent assortment? b) Dihybrid cross a) Monohybrid cross c) Trihybrid cross d) Tetrahybrid cross 432. Hyperdactyly (the possession of more than 12 finger) is determined by the dominant allele (H) and normal condition by recessive allele (h). The diagram shows a family tree in which some members of the family are hyperdactylus

	erdactylus male		
	mal male		
	mal female		
	erdactylus female		
Find out the genotype of A	A, B and C		
a) A-Hh, B-Hh, C-hh	b) A-HH, B-Hh, C-hh	c) A-Hh, B-HH, C-hh	d) A-Hh, B-HH, C-hh
433. Which of the following sta	atements about mutation a	are true?	
I. Mutations are the sourc	e of new alleles for genes		
_	create mutations to meat t	=	
	events and can happen in		
	o be harmful or have no ef	-	
a) I, II and III	b) I, II, III and IV	c) I, III and IV	d) I and III
434. Centromere is also called			
a) Chromomere		b) Secondary constriction	n
c) Primary constriction		d) chromocentre	
435. Which of the following sta			
-	equired because they are i		1
		ance than dominant alleles	do
	es of each gene from each	-	
	t regard to which alleles th		d) I II and IV
a) II and III 436. Which contributed to Me	b) II and IV	c) II, III and IV	d) I, II and IV
	nuel's success?		
I. Selection of pea plant II. Knowledge of history			
III. One character at one t	ime		
IV. His statistical knowled			
Choose the correct option	1		
a) I, II, III and IV	b) II and III	c) I, III and IV	d) IV, III and II
437. In XX and XO chromosom	al sex determination there	is absence of one chromos	ome in
a) Male	b) Female	c) Both (a) and (b)	d) None of these
438. Which of the following is	-		
•		e produced in F ₂ -generatio	n
	ation are produced in F ₂ -g		
		ear in high frequency in F_2 .	generation
	which two chromosome a		
a) Only I	b) Only II	c) I and III	d) III and IV
439. The total number of prog many are recombinants?	eny obtained through dihy	brid cross of Mendel is 128	$10 \text{ in } F_2$ -generation. How
a) 240	b) 360	c) 480	d) 720
440. A child of blood group-O	cannot have parents of blo	od groups	
a) A and A	b) AB and O	c) A and B	d) B and B
441. Rh factor is present in			
a) All vertebrates		b) All mammals	
			1

443. A man of blood group–A marries woman of blood group-AB, which type of progeny would indicate that man is heterozygous?

a) 0 b) B c) A d) AB

444. The children of a haemophilic man and a normal women are

- a) All haemophilic
- b) Only daughters are haemophilic
- c) Only sons are haemophilic
- d) Neither sons nor daughter are haemophilic

445. In man, four phenotypes of blood groups are due to the presence of antigen-A and antigen-B on the RBC. The chromosome that has the gene to control these antigens is

c) ^{9th} chromosome

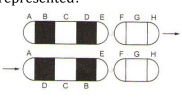
- a) X-chromosome
 - b) 21st chromosome

d) 7th chromosome

446. More men suffer from colourblindness than women because

- a) Women are more resistant to disease than men
- b) The male sex hormone testosterone causes the disease
- c) The colourblind gene is carried on the 'Y' chromosome
- d) Men are hemizygous and one defective gene is enough to make them colourblind
- 447. 'Cri-du-chat' syndrome in humans is caused by the
 - a) Fertilization of an XX egg by a normal Y-bearing sperm
 - b) Loss of half of the short arm of chromosome 5
 - c) Loss of half of the long arm of chromosome 5
 - d) Trisomy of 21st chromosome

448. Given below is representation of a kind of chromosomal mutation. What is the kind of mutation represented?



a) Deletion c) Inversion b) Duplication

d) Reciprocal translocation

449. Which of the following symbols and its representation, used in human pedigree analysis is correct?

- = Mating between relatives
- \bigcirc = Unaffected male b)

= Affected male

d) Diabetes

d) antibody-a

- = Unaffected female c)
- 450. Ischihara chart is used to detect
- a) Tuberculosis
- b) Eve sight 451. Genes exibiting multiple effects are known as
 - a) Complementary genes
 - c) Cistrons

a) AB antigen

- 452. A person with blood group –AB has
- c) no antigen

d) Pseudogenes

c) Colour blindness

b) Pleiotropic genes

- b) a and b antibodies 453. Excessive growth of hair on the pinna is a feature found only in males because
 - a) The female sex hormone oestrogen suppresses theb) The gene responsible for the character is present character in females on the Y-chromosomes only
 - c) The gene responsible for the character is recessived) The character is induced in males as males in females and dominant only in males produce testosterone

454. 3:1 ratio in F₂-generation is explained by

- a) Law of partial dominant b) Law of dominant c) Law of incomplete dominant d) Law of purity of gametes
- 455. Incomplete dominance is different from complete dominance in having
- a) Phenotypic ratio b) Genotypic ratio c) Both (a) or (b)
- d) None of these 456. A true breeding plant producing red flowers is crossed with a pure plant producing white flowers. Allele

for red colour of fl	ower is dominant. After selfing	the plants of first filial a	eneration the proportion of
	white flowers in the progeny wo		eneration, the proportion of
a) 9:3:3:1	b) 12 : 3 : 1	c) 9:3:4	d) 9 : 6 : 1
•	sex-linked trait shows that		
a) Male are affecte		b) Female are carrie	er mostly
c) Both (a) and (b)		d) Neither (a) or (b)	-
	, two individuals produces offsp		-
	r (a),then the genotypes of pare	-	
a) $Aa \times Aa$	b) $Aa \times aa$	c) $AA \times aa$	d) $AA \times Aa$
,	e garden pea plant for his exper		-
a) Artificial pollina	• • • •	b) Cross-pollination	
c) Self and artificia		d) None of the abov	
-	nt assortment of two genes 'A' a	-	
a) Repulsion	b) Recombination	c) Linkage	d) Crossing over
· ·	ig is not the type of blood group	, ,	
a) Lewis and Duffy		c) ABO and Rh	d) Rh and MN
,	ame genotype have different pl	,	
=	entical genotype give identical		
b) No – because of		F	
•	lifferent environment can prod	uce different phenotype	of the same genotype
	henotype decides the genotype		0
			es – I^A , I^B and i. since there are
	eles, six different genotypes are		
a) Three	b) One	c) Four	d) Two
	otype TTrr in F_2 -generation of a		
			. 6
a) $\frac{1}{16}$	b) $\frac{3}{16}$	c) $\frac{9}{16}$	d) $\frac{6}{16}$
465. In a cross between	individuals with genotypes Tt	Rr, if the resulting numb	per of offsprings is 16, then
identify the numbe	er of genotypes with TtRr and T	'tRR amongst them.	
a) 1 and 2	b) 2 and 3	c) 3 and 1	d) 4 and 2
466. Which of the follow	ving genotypes does not produ	ce any sugar polymer on	n the surface of the RBC?
a) I ^A I ^A	b) I ^B i	c) I ^A I ^B	d) i i
467. The diagrammatic	representation of the chromos	omes of an individual is	called
a) Idiogram	b) Karyotype	c) Phenotype	d) diploidy
468. In <i>Mirabilis</i> , a hyb	rid for red (RR) and white (rr)	flower produces pink (R	r) flower. A plant with pink
flower is crossed v	vith white flower, the expected	phenotypic ratio is	
a) Red : pink : whi	te(1:2:1)	b) Pink : white(1 : 1)
c) Red : pink (1 : 1)	d) Red : white (3 : 1)
469. A marriage between normal visioned man and colourblind woman will produce, which of the following			
types of offsprings	?		
a) Normal sons an	d carrier daughters	b) Colourblind sons	and carrier daughters
c) Colourblind sor	s and 50%carrier daughters	d) 50% colourblind	sons and 50%carrier daughters
470. Given below is a p	edigree chart of a family with fi	ve children. It shows the	e inheritance of attached ear
lobes as opposed to the free ones. The squares represent the male individuals and circles the female			
individuals. Which	of the following conclusions di	rawn is correct?	
()	1		

a) The parents are homozygous recessive

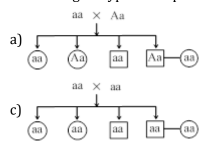
Attached

Ear lobes Ear lobes

Free

b) The trait is Y-linked

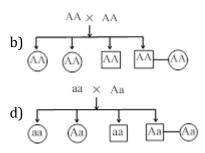
a) The percents are home	arragua dominant	d) The percents are beter		
c) The parents are homozygous dominant d) The parents are heterozygous 471. I. Myotonic dystrophy is an autosomal dominant trait				
	an autosomal recessive trai			
	n of alleles results in chrom			
	n of allele result in chromos	iomal gain		
V. Cystic fibrosis is a Mer	ndelian disorder			
Correct statements are		· · · · · · · · · · · · · · · · · · ·		
a) I, II, III and IV	b) I, III, IV and V			
472. Haemophilia is more cor				
a) This disease is due to mutation	a X-linked dominant	b) A greater proportion	of girls die in infancy	
c) This disease is due to	a X-linked recessive	d) This disease is due to	a Y- linked recessive	
mutation		mutation		
473. Which one of the followi	ng was the rediscoverer of	Mendel's work?		
a) Muller	b) Morgan	c) Correns	d) Bridge	
474. $\frac{1}{4}$: $\frac{1}{2}$: $\frac{1}{4}$ ratio of TT : Tt : tt	can be depicted mathemati	cally binomial expression	as (ideally)	
	b) $(ax + by)^3$		d) $ax + by$	
475. Pure red flowers was cro			selfing of F_1 -generation, the	
	oducing white flowers in pr			
a) ³ ⁄ ₄	b) ¼	c) 1/3	d) ½	
476. Which of the following abnormalities, results from an unnatural presence of a Barr body?				
a) Turner's syndrome b) Down's syndrome				
c) Klinefelter's syndrome d) All of these				
477. When normal and mutar		•	nologous pair, the	
heterozygotes are called	as			
a) <i>cis</i> heterozygotes		b) Homologous heterozy	/gotes	
c) <i>trans</i> heterozygotes			d) None of the above	
478. When two unrelated ind	ividuals or lines are crosse	d, the performance of F_1 hy	/brid is often superior to	
both its parents. This ph				
a) Transformation	b) Splicing	c) Metamorphosis	d) heterosis	
479. The types of gametes pro	oduced by a heterozygous a	llelic pair is/are		
a) 1	b) 2	c) 3	d) Many	
480. Prokaryotic genetic systematic	em has			
a) DNA and histone		b) DNA and no histone		
c) No DNA and histone		d) No DNA and no histor	ıe	
481. A chromosome in which the centromere is situated close to its end so that one arm is very short and the				
other very long is				
a) Acrocentric	b) Metacentric	c) Sub- metacentric	d) telocentric	
482. Write the genotype of th	e previous questions			



483. Sickle cell anaemia is

a) An autosomal linked dominant trait

c) Caused by a change in base pair of DNA



b) Caused by substitution of valine by glutamic acid in the β -globin chain of haemoglobin

d) Characterized by elongated sickle like RBCs with a

nucleus

- 484. Improvement of human race through hereditary qualities is called
 - a) Disruptive b) Directional c) Stabilizing d) Coevolution
- 485. ...A... gene produces all gametes that are similar, while aB... produces two kinds of gametes each having one allele with equal proportion
 - Choose the correct option for A and B
 - a) A-homozygous; B-heterozygous
 - c) A-homozygous; B-recessive

- b) A-homozygous; B-dominance
- d) A-heterozygous; B-homozygous
- 486. In which one of the following combinations (a-d) of the number of the chromosomes is the present day hexaploid wheat correctly represented?

Combi	Mono	Нар	Nullis	Tris
nation	somic	loid	omic	omi
				С
a) 27-28-	42-43		b) 7-82	2-40-42

c) 21-7-42-43

b) Phenylalanine reductase

d) Phenylalanine oxidoreductase

d) 41-21-40-43

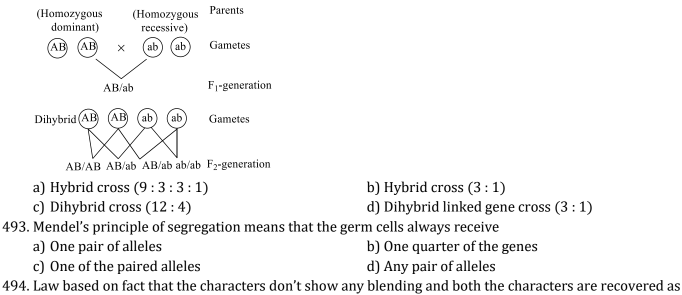
- 487. When the number of recombinant progeny is usually less than the number expected in independent assortment it is called
 - a) Complete linkage
 - b) Incomplete linkage
 - c) Complete recombination
 - d) Complete independent assortment
- 488. The enzyme missing in phenylketonuria is
 - a) Phenylalanine hydroxylase
 - c) Phenylalanine oxidase
- 489. Gene is
 - a) One pair of allele
 - b) Alternative form of a gene
 - c) Present in allelic form on homologous
 - d) Both (a) and (c) are correct

490. The telomeres of eukaryotic chromosomes consist of short sequences of

- a) Thymine rich repeats b) Cytosine rich repeats
- c) Adenine rich repeats d) Guanine rich repeats
- 491. In Mendelian dihybrid cross when heterozygous Round Yellow are self crossed. Round Green offsprings are represented by the genotype
 - a) RrYy,RrYY and RRYy b) Rryy,RRyy, and rryy c) rrYy andrrYY

d) Rryy and RRyy

492. Study the given test cross and choose the correct option for F_2 -generation



such in F ₂ -generation although one character was a	bsent in F₁-progeny, is	
a) Law of purity of gametes	b) Law of independent a	assortment
c) Law of incomplete dominance	d) Law of dominance	
495. In <i>Melandrium</i> , the sex determination type is	uj haw of dominance	
a) XX-XY b) XX-XO	c) ZZ-ZW	d) XY-XO
496. The effect of todays radioactive fall out will probabl	,	
than to children now living because	y be more narmini to chin	il ell ol luture generation
a) Infants are more susceptible to radiations		
b) Susceptibility to radiation increase with age		
c) Mutated genes are frequently recessive		
d) Contamination of milk supply is not cummulative 497. Select the statement which is not correct.	<u>,</u>	
	h) In coco of nolyconic is	haritan as thousands of
a) Polygenic character is controlled by multiple		nheritance, thousands of
alleles		pes are found between two
	extreme ones	
c) Height, weight, skin colour are polygenic		tem is an example of multiple
	allelism	
498. Linkage was first suggested by		
a) Sutton and Boveri b) Morgan	c) De Vries	d) Pasteur
499. X-linked recessive gene is		
a) Always expressed in male	b) Always expressed in t	temale
c) Lethal	d) Sub-lethal	
500. Gene for colour blindness is located on		1) 24 St 1
a) Y-chromosome b) 13 th chromosome	c) X-chromosome	d) 21 st chromosome
501. A. $\frac{x}{A} = 1$		
B. $\frac{X}{A}$ > more than 1		
$C.\frac{x}{A} = 0.5$		
Here, $X =$ number of X-chromosome		
A = set of autosomal pair		
Choose the correct option for A, B and C result		
a) A-female B-meta female C-male	b) A-female B-meta fem	
c) A-female B-female C-male	d) A-meta female B-fem	ale C-male
502. The ABO blood group are controlled by		
a) I-gene	b) c-gene	
c) B-gene	d) n-gene	
503. Which of the following is considered as a recessive of	character of Mendel?	
a) Round seed b) Wrinkled seed	c) Axial flower	d) Green pod
504. When an animal has both the characters of male and	l female, it is called	
a) Intersex b) Superfemale	c) Supermale	d) gynadromorph
505. Point mutation arises due to change in		
a) Single base DNA	b) Single base pair of DN	NA
c) Segment of DNA	d) Double base pair of D	NA
506. colour blindness is more observed inhumans		
a) Male b) Female	c) Infent	d) In old age
507. Mendel cross tall and dwarf plant. In F_2 -generation	the observed ratio was 3:2	1 (tall: short). From this
result, he deduced		
I. law of dominance		
II. law of independent assortment		
III. law of segregation		

IV. incomplete dominance				
Choose the correct option				
a) I, II, III and IV b) I and III	c) II, III and IV	d) I, II and III		
508. Genetic or chromosomal symbol used for the perso	n who is having sickle-cell			
a) Hb ^s Hb ^s b) Hb ^a Hb ^a	c) Hb ^g Hb ^g	d) Hb ^m Hb ^m		
509. Which of the following is true regarding human gen	etics?			
a) Most characters are controlled by one gene				
b) Same characters are controlled by more then two	o genes			
c) Same characters are not inherited according to M	Iendel's law			
d) All of the above				
510. Foetal sex is determined by examining cells from an	nniotic fluid looking for			
a) Chiasmata b) Barr bodies	c) Sex chromosomes	d) None of these		
511. Sex-linked allele or disease never passes from				
a) Women to her daughter				
b) Man to daughter				
c) Women to grand daughter				
d) Man to his son				
512. What is genotypic ratio in a dihybrid cross?				
a) 1:2:1:2:4:2:1:2:1	b) 2:4:2:1:2:1:1:2			
c) 1:4:2:1:1:2:1	d) 4 : 2 : 1 : 1 : 1 : 1 : 2 : 1			
513. The law of segregation of characters is also called the				
 a) Gametes have only one of the two alleles for each characters 	i b) Gametes cannot be co	mammated		
c) Gametes are very different types of cells	d) It was just another na	me adopted accidentally		
514. Four children belonging to the same parents have the following blood groups A, A, AB and O. Hence, the				
		, , , ,		
genotypes of the parents are				
a) Both parents are homozygous for 'A' group				
a) Both parents are homozygous for 'A' groupb) One parent is homozygous for 'A' and another pa	rent is homozygous for 'B'			
a) Both parents are homozygous for 'A' groupb) One parent is homozygous for 'A' and another pactor of the parent is heterozygous for 'A' and another parent is heterozygous for 'A' and another parent parent is heterozygous for 'A' and another parent parent is heterozygous for 'A' and another parent p	rent is homozygous for 'B'			
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another parent is heterozygous for 'A' and another p d) Both parents are homozygous for 'B' group 	rent is homozygous for 'B'			
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'A' and another parents are homozygous for'B' group 515. Mendel work later formulated into laws of 	rent is homozygous for 'B'			
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another parent is heterozygous for 'A' and another p d) Both parents are homozygous for'B' group 515. Mendel work later formulated into laws of I. Linkage 	rent is homozygous for 'B'			
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'A' and another parent's are homozygous for 'B' group 515. Mendel work later formulated into laws of Linkage Segregation 	rent is homozygous for 'B'			
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another parent is heterozygous for 'A' and another p d) Both parents are homozygous for'B' group 515. Mendel work later formulated into laws of I. Linkage 	rent is homozygous for 'B'			
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'A' and another parent's are homozygous for 'B' group 515. Mendel work later formulated into laws of Linkage Segregation Incomplete dominance IN Independent assortment Choose the correct option 	rent is homozygous for 'B'			
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'A' and another parents are homozygous for 'B' group 515. Mendel work later formulated into laws of Linkage Segregation III. Incomplete dominance IV. Independent assortment Choose the correct option a) I, III and IV b) II and IV 	rent is homozygous for 'B'			
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'B' group 515. Mendel work later formulated into laws of Linkage Segregation II. Segregation III. Incomplete dominance IV. Independent assortment Choose the correct option a) I, III and IV 516. Barr body is associated with 	arent is homozygous for 'B' arent is heterozygous for 'I c) II, III and IV	B' d) I, II and III		
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'B' group 515. Mendel work later formulated into laws of Linkage Segregation III. Incomplete dominance IV. Independent assortment Choose the correct option I, III and IV 516. Barr body is associated with Sex chromosome of female 	arent is homozygous for 'B' arent is heterozygous for 'J c) II, III and IV b) Sex chromosome of m	B' d) I, II and III		
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'A' and another parents are homozygous for 'B' group 515. Mendel work later formulated into laws of Linkage Segregation II. Segregation III. Incomplete dominance IV. Independent assortment Choose the correct option a) I, III and IV 516. Barr body is associated with Sex chromosome of female Autosome of female 	arent is homozygous for 'B' arent is heterozygous for 'I c) II, III and IV	B' d) I, II and III		
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'B' group 515. Mendel work later formulated into laws of Linkage Segregation II. Incomplete dominance IV. Independent assortment Choose the correct option I, III and IV 516. Barr body is associated with Sex chromosome of female Autosome of female 	arent is homozygous for 'B' arent is heterozygous for 'l c) II, III and IV b) Sex chromosome of m d) Autosome of male	B' d) I, II and III		
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'B' group 515. Mendel work later formulated into laws of Linkage Segregation II. Incomplete dominance IV. Independent assortment Choose the correct option I, III and IV 516. Barr body is associated with Sex chromosome of female Autosome of female 517. A man can inherit his X-chromosome from his Maternal grandmother or maternal grandfather 	arent is homozygous for 'B' arent is heterozygous for 'l c) II, III and IV b) Sex chromosome of m d) Autosome of male b) Father	B' d) I, II and III		
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'B' group 515. Mendel work later formulated into laws of Linkage Segregation Incomplete dominance IV. Independent assortment Choose the correct option I, III and IV 516. Barr body is associated with Sex chromosome of female Autosome of female 517. A man can inherit his X-chromosome from his Maternal grandfather 	c) II, III and IV b) Sex chromosome of m d) Autosome of male b) Father d) Paternal grandfather	B' d) I, II and III		
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'B' group 515. Mendel work later formulated into laws of Linkage Segregation Incomplete dominance IV. Independent assortment Choose the correct option I, III and IV 516. Barr body is associated with Sex chromosome of female Autosome of female 517. A man can inherit his X-chromosome from his Maternal grandmother or maternal grandfather Maternal grandfather 	c) II, III and IV b) Sex chromosome of m d) Autosome of male b) Father d) Paternal grandfather	B' d) I, II and III hale		
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'A' and another parent parents are homozygous for 'A' and another parents' group 515. Mendel work later formulated into laws of I. Linkage II. Segregation III. Incomplete dominance IV. Independent assortment Choose the correct option a) I, III and IV b) II and IV 516. Barr body is associated with a) Sex chromosome of female c) Autosome of female 517. A man can inherit his X-chromosome from his a) Maternal grandmother or maternal grandfather c) Maternal grandfather 518. The types of gametes formed by the genotype Rr Yy a) RY, Ry, rY, ry b) RY, Ry, ry, ry 	c) II, III and IV b) Sex chromosome of m c) Father d) Paternal grandfather v are c) Ry, Ry, Yy, ry	B' d) I, II and III hale d) Rr, RR, Yy, YY		
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another parent is heterozygous for 'B' group 515. Mendel work later formulated into laws of Linkage Segregation II. Incomplete dominance IV. Independent assortment Choose the correct option I, III and IV 516. Barr body is associated with Sex chromosome of female Autosome of female 517. A man can inherit his X-chromosome from his Maternal grandfather 518. The types of gametes formed by the genotype Rr Yy RY, Ry, rY, ry RY, Ry, rY, ry 	c) II, III and IV b) Sex chromosome of m d) Autosome of male b) Father d) Paternal grandfather v are c) Ry, Ry, Yy, ry er to determine whether it i	B' d) I, II and III hale d) Rr, RR, Yy, YY		
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'A' and another parent is heterozygous for 'B' group 515. Mendel work later formulated into laws of Linkage Segregation II. Incomplete dominance IV. Independent assortment Choose the correct option I, III and IV 516. Barr body is associated with Sex chromosome of female Autosome of female 517. A man can inherit his X-chromosome from his Maternal grandmother or maternal grandfather Maternal grandfather 518. The types of gametes formed by the genotype Rr Yy RY, Ry, rY, ry Rting of an organism to a double recessive in order heterozygous for a character under consideration, i 	c) II, III and IV b) Sex chromosome of m d) Autosome of male b) Father d) Paternal grandfather v are c) Ry, Ry, Yy, ry er to determine whether it is s called	B' d) I, II and III hale d) Rr, RR, Yy, YY is homozygous or		
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another pac) One parent is heterozygous for 'A' and another pac) One parent is heterozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) One parent is heterozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) One parent is heterozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) One parent is heterozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for a character under consideration, if a) Reciprocal cross 	c) II, III and IV b) Sex chromosome of m d) Autosome of male b) Father d) Paternal grandfather v are c) Ry, Ry, Yy, ry er to determine whether it i	B' d) I, II and III hale d) Rr, RR, Yy, YY		
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another pac c) One parent is heterozygous for 'A' and another pad) Both parents are homozygous for'B' group 515. Mendel work later formulated into laws of Linkage Segregation III. Incomplete dominance IV. Independent assortment Choose the correct option a) I, III and IV 516. Barr body is associated with Sex chromosome of female Autosome of female 517. A man can inherit his X-chromosome from his Maternal grandmother or maternal grandfather Maternal grandfather 518. The types of gametes formed by the genotype Rr Yy RY, Ry, rY, ry RY, Ry, rY, ry Rting of an organism to a double recessive in order heterozygous for a character under consideration, if a) Reciprocal cross Test cross 	c) II, III and IV b) Sex chromosome of m d) Autosome of male b) Father d) Paternal grandfather r are c) Ry, Ry, Yy, ry er to determine whether it is s called c) Dihybrid cross	B' d) I, II and III hale d) Rr, RR, Yy, YY is homozygous or d) Back cross		
 a) Both parents are homozygous for 'A' group b) One parent is homozygous for 'A' and another pac) One parent is heterozygous for 'A' and another pac) One parent is heterozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) One parent is heterozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) One parent is heterozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) One parent is heterozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for 'A' and another pac) Both parents are homozygous for a character under consideration, if a) Reciprocal cross 	c) II, III and IV b) Sex chromosome of m d) Autosome of male b) Father d) Paternal grandfather v are c) Ry, Ry, Yy, ry er to determine whether it is s called	B' d) I, II and III hale d) Rr, RR, Yy, YY is homozygous or d) Back cross		

have any gene for colour blindness. The pro a) 50% b) 75%	c) 25%	d) None of these
522. L –shaped chromosomes are also called	,	
a) Acrocentric b) Telocentric	c) Sub-metacentric	d) None of these
523. A homozygous sweet pea plant with blue flo	owers (RR) and long pollen (R_0	$_0 R_0$) is crossed with a
homozygous plant having red flowers (rr) a		
Which of the following genotype does not a	ppear in its progeny?	
a) <i>Rrrr</i> ₀ b) <i>RrRr</i> ₀	c) Rrr_0r_0	d) rrR_0r_0
524. A diseased man marries a normal woman an	nd they get three daughters an	d five sons. All the daughters
were diseased and sons were normal. The g	ene of this disease is	
a) Sex-linked dominant	b) Sex-linked recess	ive
c) Sex-limited character	d) Autosomal domin	nant
525. A polygenic trait is controlled by 3 genes A,	B and C. In a cross AaBbCc× A	aBbCc, the phenotypic ratio of
the offsprings was observed as $1:6:x:20:$	<i>x</i> : 6 : 1.	
What is the possible value of <i>x</i> ?		
a) 3 b) 9	c) 15	d) 25
526. Chromosomal mutations occurs due to		
I. Deletion II. Duplication		
III. Translocation IV. Inversion		
Choose the correct option		
a) I, II and III b) II, III and IV	c) I, III and IV	d) All of these
527. The allele which expresses itself in both hor		ondition is called
a) Dominant allele	b) Recessive allele	
c) Incomplete dominant allele	d) Split allele	
528. Equatorial division and reductional division		
a) Meiosis, mitosis b) Mitosis, meios	is c) Both (a) and (b)	d) Amitosis, meiosis
529. Monohybrid test cross ratio is		
a) 3:1 b) 2:1	c) 1:1	d) 9 : 3 : 3 : 1
530. Who gave the term 'genetics'?		
a) Mendel b) Robert Hooke	,	d) Purkinje
531. In which of the following disorders, blood h	•	
a) Haemophilia b) Haematuria	c) Haematoma	d) Sickle cell anaemia
532. In sickle cell anaemia, the glutamic acid is re		
a) Proline b) Alanine	c) Serine	d) Valine
533. Which of the following cannot be explained		
 a) The discrete unit controlling a particular is called a factor 	other is recessive	
, Alleles do not show any blending and bot		
c) $\frac{1}{2}$ characters recover as such in F_2 generation		Jans
534. Find out the percentage of dominant phenot		Pn P-dominant n-recessive
a) 25% b) 50%	c) 75%	d) 100%
535. Gametes produced by a homozygous individ	•	uj 10070
a) 1 b) 2	c) 3	d) Many
536. What will be the gametic chromosomes nun		
a) 10 b) 20	c) 30	d) 40
537. Human female possesses 44+XX chromosor	,	,
a) $44 + XX$ b) $22 + X$	c) 22	d) 44
538. Select the correct statement from the ones g	•	2
a) Antirrhinum b) Pisum	c) Solanum	d) Hibiscus
,	.,	.,

a) F_1 crossed with heterozygous parent	b) F ₁ crossed with homo	ozygous dominant parent
c) F_1 crossed with homozygous recessive parent	d) F ₁ crossed with home	
540. Sex chromosomes are also known as		
a) Autosomes b) Allosomes	c) Genome	d) karyotype
541. Euploidy is best explained by		
a) Exact multiple of a haploid set of chromosomes		
b) One chromosome less than the haploid set of chr	romosomes	
c) One chromosome more than the haploid set of cl		
d) One chromosome more than the diploid set of ch		
542. In which year Mendel's work rediscovered		
a) 1900 b) 1901	c) 1902	d) 1903
543. Which of the following diseases is governed by plei		
a) Sickle cell anaemia b) Haemophilia	c) Colour blindness	d) None of these
544. Inheritance of skin colour in human is an example of		a) None of these
a) Chromosomal aberration	b) Codominance	
c) Point mutation	d) Polygenic inheritance	
545. Heterochromatin remains condensed in which part	, ,,	
a) Secondary construction–I	b) Secondary construction	on-II
	d) Both (a) and (b)	011-11
c) Telomeres 546. A plant of F ₁ - generation has genotype 'AABbCC'. O		honotunic ratio in E
	n sennig of this plant, the p	menotypic ratio in F ₂ -
generation will be	h) In complete deminen	
a) Polyploidy	b) Incomplete dominant	Le la
c) Multiple allelism	d) polygeny	
547. Which have great importance in genetics?		
a) Penicillium b) Claviceps	c) Neurospora	d) None of these
548. Number of Barr bodies in XXXXY is		
a) 1 b) 2	c) 3	d) 4
549. Dihybrid ratio of the linked gene is		
a) 1:1 b) 1:1:1:1	c) 9:3:3:1	d) 3 : 1
550. Polyploidy can be induced by the application of		
a) Auxin b) Kinetin	c) Colchicine	d) ethylene
551. If a plant having yellow or round seeds was crossed	l with another plant having	green and wrinkled seeds
then F_1 -progeny are in the ratio		
a) 15 : 1	b) 1 : 15	
c) 1:13	d) All yellow and round	seeds
552. Which of the following, can be used to describe the	•	
a) Alternative form of a gene	b) Dominant form of ger	ne
c) Recessive form of a gene	d) One gene pair	
553. Which of the following animals is mostly used in ge	netics experiments?	
a) Butterfly b) Fruit fly	c) Housefly	d) Dragon fly
554. Which of the following diseases results from the ge	netic inability to synthesize	e a single enzyme?
a) Colour blindness b) Down's syndrome	c) Phenylketonuria	d) Diabetes
555. Balbiani rings are the structural features of		
a) Allosomes		
b) Polytene chromosomes		
c) Autosomes		
d) Lampbrush chromosomes		
556. Sometimes, there are more than two alleles for a gi	ven chromosome locus. In	this case, a trait is controlled
by		,
a) Codominance	b) Pseudodominance	
c) Incomplete dominance	d) Multiple alleles	
-,	.,	

557. Two pea plants were subjected for cross pollination	on. Of the 183 plants produ	uced in the next generation.94
plants were found to be tall and 89 plants were for	• •	0
plants are likely to be	0	
a) TT and tt b) Tt and Tt	c) Tt and tt	d) TT and TT
558. In haplodiploidy determination of sex, males is		2
a) Haploid b) Diploid	c) Haplodiploid	d) Diplohaploid
559. A cross between F_1 -hybrid and a heterozygous particular	, , ,	
a) 1 : 1 b) 3 : 1	c) 2:1	d) $4:1$
560. When mutation is confined to only one substitutio		,
a) Translocation b) Point mutation	c) Base inversion	d) Frame shift
561. Letter symbol refers to the dominant factors give a	aA or upper case latte	-
correspondingB or lower case letter is used fo	= =	_
a) A-capital; B-small	b) A-small; B-capital	
c) A-capital; B-capital	d) A-small; B-small	
562. In a gene pool, along with beneficial mutations the	se mutations also exists v	vhich are damaging to an
individual. It has been found that these mutations		
I. they have survival value		
II. they are acquired		
III. they are recessive and carried by heterozygous	individuals only	
IV. they show genetic drift		
Choose the incorrect option for given statement		
a) I and III b) I and II	c) II and IV	d) Only III
563. In a medicolegal case of accidental interchange be	tween two babies in a hos	pital, the baby of the blood
group-A could not be rightly given to a couple with	1	
a) Husband of O group and wife of AB group	b) Husband of A group	and wife of O group
c) Husband of B group and wife of O group d) Husband of AB group and wife of A group		
564. The plasma membrane of the red blood cells has	.A polymers that protru	de from its surface and the
kind of sugar is controlled by the gene. The gene I	has three allelesB The	e alleles I ^A and I ^B produce a
slightly different form of the sugars, while allele i	loesn't produce anyC	
Choose the correct option for A, B and C		
a) A-protein, B — I ^A I ^B I ^O , C-protein	b) A-protein, B – I ^A I ^B	³ I ⁰ , C-sugar
c) A-sugar, B – I ^A I ^B I ^O , C-protein	d) A-sugar, B – I ^A I ^B I,	C-sugar
565. The person famous for experimental genetics		
a) TH Morgan b) Sutton	c) Boveri	d) Robert Hooke
566. Morgan worked with tiny fruit fly names as		
a) Drosophila melanogaster	b) Mangifera indica	
c) Mirabilis jalapa	d) Drosophila indica	
567. <u>o</u> <u>9</u>		
1st child 2nd child 3rd child 4th child (Albino) (Normal) (Mormal) (Albino)		
A=Normal allele, a = Albino allele.		
Find out genotype of $\Box \circ$ and		
(father and mother)		
Father Mother ♂ [*] ○		
a) <u>Aa AA</u>	b) AA Aa	
c) AA AA	d) Aa Aa	
568. Linkage group is		
a) Linearly arranged group of linked gene	b) Non-linearly arrang	ged group of linked gene

c) Non-linearly arranged group of unlinked gene	d) Non-linearly arranged	group of single gene
569. Some individuals with blood group –A may inherit t	he genes for blonde hair, w	hile other individuals with
blood group – A may the gene for brown hair. This c	an be best explained by th	e principle of
a) 3 : 1 b) 9 : 3 : 3 : 1	c) 1:1	d) 1 : 1 : 1 : 1
570. I. 100% parental combinations are found in F_2 -gene	eration	-
II. F_2 phenotypic ratio is 3 : 1 in dihybrid cross		
III. Dihybrid test cross ratio is $1:1$ in F_2 -generation		
IV. Linked genes tends to separate frequently		
Choose the correct options from the above given sta	tements	
a) I, II and IV b) I, III and IV	c) II, III and IV	d) I, II and III
571. The following diagram shows two chromosomes and		•
Chromosome 1 Chromosome 2	u ille letter eu llulliber Tepr	esents the genes
PQRSTUVW EFGH		
Which of the following would result if a translocatio		
a) PQRSWVUT EFGH	b) PQRS TUVW	EFGH
c) PQRSTUVW EFH	d) PQRSTUVW E	FGHGH
572. Experimental verification of the chromosomal theor	y of inheritance was given	by
a) Gregor Johann Mendel	b) Hugo de Vries	-
c) Langdon Down	d) Thomas Hunt Morgan	
573. A gene that masks another gane's expression, is calle		
a) Dominant b) Recessive	c) Epistatic	d) Assorted
574. Transposons are	-) _ _F	
a) House- keeping genes	b) Jumping genes	
c) Transporting genes	d) Stationary genes	
575. Which of the following law was discovered first by N		
a) Law of dominance	b) Law of segregation	
c) Law of independent assortment	d) Law of sex determinat	ion
576. Unit of inheritance that required to express a partici	,	
a) Factors b) Genes	c) Phenotype	d) Genotype
577. Sex limited traits are the	cj i nenotype	uj denotype
a) Traits appeard in particular sex		
b) Traits which governed by genes present in both s	20200	
c) Traits which influenced by the sex hormones	JEXES	
d) All of the above		
-	a from	
578. Variation stands for differences in traits of progenie a) Each other		
	b) Parents	
c) Both (a) and (b)	d) From mother only	
579. In which mode of inheritance, do you expect more m	-	
a) Autosomal b) Cytoplasmic	c) Y-linked	d) X-linked
580. Mutagens are		
a) Chemical agents which cause change in DNA		
b) Physical agents which cause mutation		
c) Cancer producing agents		
d) Both (a) and (b)		
581. Which is incorrect for Mendelism?		
a) Works on garden pea		oved by monohybrid cross
c) Discovered linkage	d) All of the above	
582. In heterozygous condition, the individual expression		
a) Colourblindness b) AB blood group	c) Rh factor	d) A and B blood group
583. Polyploid derived from two different species is calle	ed	

	o) Triploid	c) Allopolyploid	d) monoploid	
584. Walter Sutton is famous for	his contribution to			
a) Genetic engineering		b) Totipotency		
c) Quantitative genetics		d) Chromosomal theory of		
585. Humans knew from as early				
		ally present in wild populat	tion, A, B and C here refer to	
a) A-8000-1000 BC, B-sexua				
b) A-8000-15000 BC, B-sexu	-			
c) A-8000-15000 BC, B-sexu	-			
d) A-20000-25000 BC, B-sez	=			
586. Punnett square was develop a) RC Punnett b) RB Punnett	c) RD Punnett	d) RE Punnett	
587. Female is heteromorphic an		•	uj KE Fulliett	
a) Fishes and bird		b) Reptiles		
c) Butterflies and moth		d) All of these		
588. Absence of one sex chromos	some causes	a) fin of these		
a) Turner's syndrome	Joine eauses	b) Klinefelter's syndrome	2	
c) Down's syndrome		d) Tay-Sach's syndrome	-	
589. Mendelian recombinations a	are due to	aj raj saon e egnarenne		
a) Linkage		b) Mutations		
c) Dominant characters		d) Independent assortment		
590. The important things to rem	nember are that chromos			
of a gene pair are located on	n homologous sites on	B chromosomes		
Choose the correct choice fo	or A and B			
a) A-single, B-analogous		b) A-pair, B-analogous		
c) A-pair, B-homozygous d) A-single, B-heterozygous				
591. The type of chromosomal aberration indicated in the diagram shows				
A B C D E F G				
•				
A B D C E F G				
a) Interstitial translocation		b) Reciprocal translocation	on	
c) Pericentric inversion		d) Paracentric inversion		
592. Who proposed chromosoma	al theory of linkage?			
a) Morgan b	o) Castle	c) Both (a) and (b)	d) Bateson	
593. Which factor expresses itsel	lf in homozygous and eve	en in heterozygous conditi	on?	
-) Weak factor	c) Recessive factor	d) Incomplete factor	
594. Number of autosomes in hu	man are			
a) 23 pairs		b) 22 pairs		
c) 46 chromosomes		d) 33 pairs of chromoson		
595. A tall plant was grown in nu	trient deficient soil and	remained dwarf. When it is	s crossed with dwarf plant	
then				
a) All hybrid plants are dwa	nrt	b) All hybrid plants are ta		
c) 50% tall and 50% dwarf		d) 75% tall and 25% dwa		
596. A man of blood group-A, ma			heterozygous for blood	
group, chances of their first			J) 1000/	
	o) 50% o aro applicable only for	c) 75%	d) 100%	
597. Mendel's laws of inheritance a) Protista b	e are applicable only for) Monera	c) Diploid organism	d) Both (a) and (b)	
598. The factors which expresses	-		uj boui (a) allu (b)	
_) Recessive	c) Hidden	d) Cryptic	
.,	,	,	J - J F	

599. Human skin colour is the example of		
I. multiple gene inheritance		
II. three separate genes controlling this trait		
III. single gene controlling this trait		
IV. two gene controlling this trait		
V. environment plays a significant role in this trait		
Choose the correct option		
a) I, II and III b) II, III and IV	c) III, IV and V	d) I, II and V
600. In haemophilia, a single protein that is a part of case	cade of protein involved in	A ofB is affected.
Single cut will result inC bleeding.		
Choose the correct option for A, B and C		
a) A-coagulation, B-RBC, C-continuous	b) A-coagulation, B-WBC	
c) A-clotting, B-blood, C-continuous	d) A-coagulation, B-bloo	
601. In <i>Drosophila</i> , the allele for a normal grey body col	our G is dominant to ebony	y body g. The following table
summarises the results of several crosses		
S.No Cross Result		
I. Strain 1 × All wild gg type		
ggtypeII.Strain 2 ×1 wild type		
gg : 1 ebony		
III. Strain $3 \times$ All ebony		
gg		
IV. Strain 4×3 wild type		
gg : 1 ebony		
Which strains both have the genotype Gg? a) I and III b) I and IV	c) II and III	d) II and IV
602. An Rh ⁻ individual receives Rh ⁺ blood. The recipient	,	d) II and IV
a) Sterile b) Dead	c) No reaction	d) isoimmunized
603. In a mutational event, when adenine is replaced by	,	d) isoinintunized
a) Frameshift mutation	b) Transcription	
c) transition	d) transversion	
604. Recessive characters are expressed	uj transversion	
a) On any autosome	b) On both the chromoso	omes of female
c) When they are present on X-chromosomes of ma	,	
ef when they are present on X enrollosomes of ma	female	it off A chromosomes of
605. The crossing of F_1 to any one of the parents is called		
a) Back cross b) Test cross	c) F ₁ cross	d) All of these
606. In cross between yellow round (YYRR) and green w		•
yellow and green seed colour	Timiea (JJTT) mia oue me	
a) 3 : 2 b) 3 : 1	c) 9:7	d) 7 : 9
607. Genes for colour blindness is carried by		
I. Abnormal development II. Father		
III. Mother IV. Autosomes		
a) I and II b) II and III	c) III and I	d) I and IV
608. Monosomy and trisomy are respectively	,	, ,
a) $n-1, n+2$ b) $2n+2, 2n+1$	c) 2 <i>n</i> −1, 2 <i>n</i> +1	d) <i>n</i> − 2, 2 <i>n</i> + 1
609. I. Haemophilia	· ·	
II. Cystic fibrosis		
III. Sickle-cell anaemia		
IV. Colour blindness		
V. Cancer		

VI. Dlagua		
VI. Plague VII. Phenylketonuria		
VIII. Thalassaemia		
Choose the correct options for Mendelian disorder	a	
a) I, II, III, IV, VI, VIII b) I, II, III, IV, VII, VIII	c) I, II, III, IV, V, VI	d) I, II, III, IV, V, VIII
	CJ 1, 11, 111, 1V, V, VI	uj 1, 11, 111, 1V, V, V111
610. in α-thalassaemia, the affected chromosomes is	a) 10th	d) 10+b
a) 16th b) 17th	c) 18th	d) 19th
611. The first hybrid progenies obtained by Mendel wer		
a) F_1 - progeny b) F_0 - progeny	c) F ₂ - progeny	d) F ₃ - progeny
612. What type of gametes will form by genotype RrYy?		
a) RY, Ry, rY, ry b) RY, Ry, ry, ry	, , , , , , ,	d) Rr , RR , Yy , YY
613. A condition, where a certain gene is present in only		
a) Heterozygous b) Monogamous	c) Homozygous	d) hemizygous
614. Frequency of crossing over isA in linked gene .	B in unlinked gene.	
Choose correct combination for A and B		
a) A-more; B-less		
b) A-less; B-more		
c) A-same; B-same		
d) A-same; B-happened		
615. Find out the phenotypic and genotypic ratios in pre	=	
a) 1:2:1,1:3 b) 1:2:1,3:1	c) 1:2:1,1:2:1	d) 1 : 3 : 1, 1 : 2 : 1
616. Which one of the following is necessary to start clo	-	
a) Heparin	b) Serotonin	
c) Thromboplastin and Ca^{2+}	d) Fibrinogen and proth	rombin
617. The organism chosen by Mendel to explain the law	of inheritance is	
a) Drosophila melanogaster	b) Antirrhinum majus	
c) Pisum sativum	d) Homo sapiens	
618. A woman is married for the second time. Her first h	usband was ABO blood ty	pe A, and her child by that
marriage was type O. Her new husband is type B ar	nd their child is type AB.	
What is the women's ABO genotype and blood type		
a) I ^A I ^O ; Blood type A b) I ^A I ^B ; Blood type AB	c) I ^B I ^O ; Blood type B	d) I ^O I ^O ; Blood type O
619. A couple has 6 children-5 are girls and 1 is boy. The	e percentage of having a gin	rl on next time is
a) 10% b) 20%	c) 50%	d) 100%
620. On selfing RrTt, we produce 400 plants, find out nu	mber of plants with genoty	/pe RrTt.
a) 100 b) 225	c) 50	d) 300
621. In the ABO system of blood groups, if both antigens	s are present but no antibo	dy, the blood group of the
individual would be		
a) B b) O	c) AB	d) A
622. Barr body in mammals represents		
a) All the heterochromatin in female cells		
b) One of the two X-chromosomes in somatic cells	of females	
c) All the heterochromatin in male and female cells	5	
d) The Y-chromosome in somatic cells of male		
623. When a segment of a chromosome breaks and later	r rejoins after 180°rotation	,it is known as
a) Deletion	b) Duplication	
c) Inversion	d) Interstitial translocat	ion
624. Human skin colour is controlled by several gene pa	•	
pairs on different chromosomes and that for each p		
that codes for no melanin deposition and an incom		
_		
deposition. If a very dark skinned person marries a	very light skinned women	, what will be the chance that

their offspring will have very dark skin? d) 9/64 a) 0 b) 1/4 c) 5/8 625. If a cross is made between AA and aa, the nature of F_1 -progeny will be b) Genotypically Aa, phenotypically a a) Genotypically AA, phenotypically a c) Genotypically Aa, phenotypically A d) Genotypically aa, phenotypically A 626. In Barr body (sex-chromatin) of a normal female a) One of the X-chromosome of paternal side becomes inactive to form Barr body b) Y-chromosomes form Barr body c) Heterochromatin condense near centre of nucleus to form Barr body d) One of the X-chromosome of maternal side becomes inactive and form Barr body near nuclear membrane 627. In certain plant species, red flower colour is incompletely dominant to white flower colour (the heterozygote is pink) and tall stems are completely dominant to dwarf stem. If a tall pink plant (TtRr) is crossed with a tall white plant (TTrr), which one of the following type of plants would be produced in the offsprings? a) Tall pink and tall white b) Dwarf pink and tall red c) Dwarf red and tall pink d) Tall pink and dwarf white 628. Which is true about meiotic cell division? I. Meosis only occurs in diploid organism without any exception II. RNA is replicated during S-phase III. Chromatids of a chromosome separate during anaphase-I IV. Only sperms are produce by this process a) I and III b) I and II c) Only I d) III and IV 629. Work of Beadle and Tatum on Neurospora crassa proved that a) Replication of DNA is semi-conservative b) Viruses have genetic material c) Every gene is responsible for specific enzymes d) Plant cells are totipotent 630. Which of the following pairs of chromosomal mutation are most likely to occur when homologous chromosomes are under going synapsis? a) Deletion and inversion b) Duplication and translocation c) Deletion and duplication d) Inversion and translocation 631. Down's syndrome is an example of a) Anueploidy b) Polyteny c) Polyploidy d) Monoploidy 632. Mendel's works were read out the a) Natural History Society in Russia b) Natural History Society in America c) Natural History Society in Brunn d) Natural History Society in Germany 633. Genes of which of the following disorder are present exclusively on the X-chromosome in humans or concerned with a) Baldness b) Red-green colour blindness c) Facial hair/moustaches in males d) Night blindness 634. In a given plant, red colour (R) of fruit is dominant over white fruit (r); and tallness (T) is dominant over dwarfness (t). If a plant with genotype RRTt is crossed with a plant of genotype rrtt, what will be the percentage of tall plants with red fruits in the next generation? a) 100% b) 25% c) 50% d) 75% 635. The figure depicits

Possibility I Possibility II		
Possibility I Two pair of homologous chromosomes		
Spindle fibres Anaphase-I Anaphase-I		
(Meiosis-I) (Meiosis-I)		
Pole		
Anaphase-II Anaphase-II (Meiosis-II) (Meiosis-II)		
a) Linkage	b) Independent assortme	ent
c) Law of dominance	d) Equational division	
636. Pick out the correct statements.		
I.Haemophilia is a sex-linked recessive disease		
II.Down's syndrome is due to aneuploidy III.Phenylketonuria is an autosomal dominant gene	dicordor	
IV.Phenylketonuria is an autosomal recessive gene		
V.Sickle cell anaemia is an X-linked recessive gene		
a) I , III and V are correct	b) I and III are correct	
c) II and V are correct	d) I , II and IV are correct	t
637. Allelic sequence variations, where more than one va	ariant (allele) at a locus in a	a human population with a
frequency greater than 0.01, is referred to as		
a) Incomplete dominance	b) Multiple allelism	
c) SNP	d) DNA polymorphism	
638. Sex chromosomes of a female bird are represented	-	1) 77147
a) X0 b) XX 639. How many types of gametes may be produced by ge	c) XY $D/d \cdot E/a \cdot E/f^2$	d) ZW
a) 27 b) 8	c) 3	d) 6
640. If a colourblind women marries a normal visioned	,	uj o
a) All normal visioned	b) One half normal and c	ne half colourblind
c) Three fourth colourblind and one fourth normal		
641. Genic balance theory of sex determination, stated b	y C B Bridges, is related to	
a) Drosophila melanogaster	b) rumex	
c) Snapdragon	d) None of the above	
642. In human beings, 45 chromosomes/single X/XO abi		
a) Down's syndrome b) Klinefelter syndrome		d) Edward's syndrome
643. When a cluster of genes show linkage behaviour the	•	
a) Do not show independent assortmentc) Do not show a chromosome map	b) Induce cell divisiond) Show recombination (during moiocic
644. Colour blindness is a failure to discriminate betwee	-	auting meiosis
a) Red and blue b) Red and green	c) Red and black	d) Red and white
645. Linkage group in <i>E</i> . <i>coli</i> is/are	ej neu una blach	aj nou una vinco
a) 4 b) 2	c) 1	d) 5
646. Linked Cross Over		
Gene pair Value (COV)		
T and U 25		

T and V 5				
V and U 30				
U and W 10				
V and W 20				
COV are given for linked gene pair. Find out their	sequence in the chromoso	me		
a) VTWU b) TVWU	c) BTWVU	d) VWTU		
647. The tendency of offsprings to resemble their pare				
a) Variation b) Heredity	c) Inheritance	d) Resemblance		
648. In case of incomplete dominance, what will be th				
a) 1:2:1 b) 3:1	c) 1:1:1:1	d) 9 : 3 : 3 :1		
649. The major reason for the success of Mendelian ex				
a) Garden pea was true breeding	b) Garden pea was cro	ss breeding		
c) Garden pea was heterozygous	d) Garden pea was not	-		
650. Which of the following is best suited for codomin				
a) Both of recessive b) Both of dominant	c) One is recessive	d) One is dominant		
651. ABO blood group system is given by		uj one is dominant		
a) Landsteiner b) Wallace	c) de Vries	d) I amarah		
	,	d) Lamarck		
652. Which of the following is generally used for induc		ants?		
a) Alpha particles	b) X-rays			
c) UV (260nm)	d) Gamma rays (from	cobalt 60)		
653. Genetic recombination is due to				
a) Fertilization and meiosis	b) Mitosis and meiosis			
c) Fertilization and mitosis	d) None of the above			
654. Identify the type of inheritance in the given diagr	am			
Q L				
$\diamond \diamond \circ \Box$				
a) Dominant X-linked	b) Recessive X-linked			
c) Dominant Y-linked	d) Cytoplasmic or mite	ochondrial inheritance		
655. Linkage gene do not shows				
a) Independent assortment	b) 9 : 3 : 3 : 1			
c) Segregation	d) All of the above			
656. Haploids are more suitable for mutation studies	-			
a) Haploids are reproductively more stable than	=	cuuse		
b) Mutagens penetrate in haploids more effective	-			
c) Haploids are more abundant in nature than di				
d) All mutations whether dominant or recessive				
-				
657. Mendel's work remain unrecognized for long tim	e uue to			
I. Communication was not easy II. Concept of factors which did not blend was not accepted				
III. Use of mathematics to explain biological prob	-			
IV. He could not provide any physical proof for th				
Choose the right combination				
a) I and II b) II and III	c) III and IV	d) All of these		
658. Ratio of progeny, when a red coloured heterozyg	ote is crossed with a white	coloured plant in which red		
colour is dominant to white colour				
a) 3 : 1 b) 1 : 1	c) 1:2:1	d) 9 : 3 : 3 : 1		
659. Mendel self-pollinated the F_2 -plant and found the	-	generate dwarf plant inB		
andC generations. He concluded that the gen				
0	-			

Choose the correct option for A, B, C and D			
a) A-dwarf, B-F ₃ , C-F ₄ , D-homozygous	b) A-dwarf, B-F ₃ , C-F ₄ , D-	heterogygous	
c) A-tall, B-F ₅ , C-F ₆ , D-homozygous	d) A-tall, B-F ₅ , C-F ₆ , D- he	terogygous	
660. The possibility of erythroblastosis foetalis occurring			
a) The baby is Rh^+ and mother Rh^-	b) The baby and mother a	re Rh ⁺	
c) The baby and mother are Rh^-	d) The baby is Rh^{-} and mo	other Rh ⁺	
661. I. Enborn error of metabolism			
II. Homozygous recessive autosomal alleles on chron	nosomes 12 causes absence	e of the specific enzyme	
III. A specific amino acid do not changes into tyrosin	е		
IV. Accumulation of phenylpyruvic acid and other de	rivatives leading to mental	retardation	
The above facts refer to			
a) Muscular dystrophy	b) Phenylketonuria		
c) Turner's syndrome	d) Down's syndrome		
662. How many phenotype and genotypes are possible in	ABO blood group systems	?	
a) Four, five b) Four, six	c) Four, seven	d) Three, four	
663. Lack of independent assortment of two genes-A and	B in fruit fly- <i>Drosophila</i> is	due to	
a) Repulsion b) Recombination	c) Linkage	d) Crossing over	
664. Mendel was successful in discovering the principles	of inheritance as		
a) He took pea plants for his experiments	b) He did not encounter li	nkage between the genes	
	for the characters he co	onsidered	
c) He had an in-depth knowledge on hybridization	d) He was a famous math	ematician	
665. The common point of attachment of all the arms of p	olytene chromosome, is kn	own as	
a) Centromere b) Chromomere	c) Chromocentre	d) centrosomes	
666. Choose the correct option for allotetraploid			
a) AABB b) AAAA	c) AAABB	d) BBBB	
667. Mutation is more common when it is present in			
a) Recessive condition	b) Dominant condition		
c) Constant in population	d) None of these		
668. Allelism refers to			
a) genic interactions controlling a character	b) Multiple genes control	•	
c) Expression of many characters by a single gene	d) Alternative forms of a g		
669. Which one pair of parents is most likely get a child, v	vho would suffer from haei	nolytic disease of new	
born?			
a) Rh^+ mother and Rh^- father	b) ^{Rh⁻} mother and Rh ⁻ fa	ather	
c) Rh ⁺ mother and Rh ⁺ father	d) $^{\rm Rh^-}$ mother and $^{\rm Rh^+}$ fa	ather	
670. Mendel performed test cross to know the			
a) Genotype of F_1 b) Genotype of F_2	c) Genotype of F_3	d) Genotype of F ₄	
671. Change in single base pair			
a) May not change the phenotype	b) Quickly changed the pl	ienotype	
c) Change the natural process	d) None of the above		
672. Find out the correct statement.	-		
a) Monosomy and nullisomy are the two types of eu	ploidy		
b) Polyploidy is more common in animals than in pla	ints		
c) Polyploids occur due to the failure in complete se	paration of sets of chromos	somes	
d) 2 <i>n</i> -1 condition results in trisomy			
673. In phenylketonuria, the phenylalanine gets converte	d to		
a) Acetic acid b) Phenyl acetic acid	c) Phenyl pyruvic acid	d) Pyruvic acid	
674. Which one of the following is a genetically transmitte	ed character?		

a) Colourblindness	b) Hydrocephalus	c) Haemophilia	d) All of these					
675. Identify the correct cho	pice for given symbols (A an	nd B)						
	\supset							
A B a) A-consenguineous n	acting R mating	b) A-mating; B-mating	botwoon relatives					
, ,	U	, , ,	, between relatives					
c) A-mating; B-conseng		d) Both (b) and (c)						
676. F_1 -hybrid is intermedia	ate between the two parent	=						
a) Codominance	_	b) Dominance						
c) Blending inheritance		d) Incomplete domina	ince					
677. Multiple phenotype see	en in							
a) Pleiotropy		b) Incomplete domina						
c) Multiple allelism		d) Polygenic inheritan						
678. After a mutation at a ge	enetic locus character of an		the change in					
a) Protein structure		b) DNA replication						
c) Protein synthesis pa		d) RNA transcription j	pattern					
679. In XX and XY type of se								
a) Homogametic	b) Heterogametic	c) Both (a) and (b)	d) Isogametic					
680. Dihybrid ratio of test c	=							
a) F ₁ hybrid produces f	our different progenies	b) F ₁ hybrid produces	two different progenies					
c) Parents produce two	o different progenies	d) None of the above						
681. A homozygous sweet p	ea plant with blue flowers	(RR) and long pollen (R_0)	R_0) is crossed with a					
homozygous plant hav	ing red flowers (rr) and rou	und pollen $(r_0 r_0)$. The res	sultant <i>F</i> ₁ hybrid is test crossed					
Which of the following	genotype does not appear	in its progeny?						
a) $\frac{1}{4}$	b) ¹	a) 1	d) $\frac{3}{16}$					
$\frac{a}{4}$	b) $\frac{1}{8}$	c) $\frac{1}{16}$	$\frac{1}{16}$					
682. Mendel's findings were	e rediscovered by							
a) De Vries	b) Correns	c) Tschermark	d) All of these					
683. The salivary gland chro	omosomes in the dipteran l	arvae are useful in gene m	apping because					
a) These are much long	ger in size	b) These are easy to st	tain					
c) These are fused		d) They have endored	uplicated chromosomes					
684. Percentage of recessive	e phenotype in a cross betw	veen PP and Pp, when P is	dominant, p recessive					
a) 25%	b) 50%	c) 35%	d) 100%					
685. Genes are made up of	,	,						
a) Histones	b) Hybrocarbons	c) Polynucleotides	d) Lipoproteins					
686. The diagram indicates	-))	-))	.,					
	ПП							
	9999							
Yy Rr	yy rr							
	4) _							
$ \square \qquad \bigcirc \qquad yR (1/4) \qquad \land \qquad $								
y y y y y y y y y y								
YR (1/4)	$\langle \land \rangle$							
	\backslash							
Yy Rr $(1/4)$ yy rr $(1/4)$	yy Rr (1/4) Yy rr (1/4)							
$\longleftarrow Parental \longrightarrow$	← Recombinant →							
a) Test cross of monoh	-	b) Test cross of dihyb						
c) Back cross of dihybi	rd	d) Back cross of mono	hybrid					

687. Type of substitution takes pla	co in ciclelo, coll anaomi	a ia			
a) Acidic amino acid to an neu		b) Glutamic acid by valine			
c) GUG to GAG	iti ai allillo aciu	-			
688. In the hexaploid wheat, the ha	poloid (n) and basis (v)	d) All of the above	270		
_	n=21 and $x=21$	c) $n=21$ and $x=14$			
		c) $n=21$ and $x=14$	d) <i>n</i> =21 and <i>x</i> =7		
689. Persons who are colourblind	can not distinguish	h) Vallour and white colou	и		
a) Red and green colour		b) Yellow and white colour	1		
c) Black and white colour	fam. datamination (h.	d) Yellow and blue colour			
690. Haploid-diploid mechanism o					
	Wasps	c) Ants	d) All of these		
691. Sickle-cell anaemia happens o		which	n is affected.		
Fill the correct option for A an	IU B	h) A shromosomal, D s sh	a la		
a) A-point; B- β -chain		b) A-chromosomal; B- α -ch	lain		
c) A-allele; B- α -chain	a is inharitad hu	d) A-non-allele; B-chain			
692. The gene of sickle cell anaemi	-	a) Cara alamana a ann a a	d)		
, , , , , , , , , , , , , , , , , , ,	Bone cells	c) Sex chromosomes	d) autosomes		
693. A character, which is express	-	a) Cadaminant	d)istatia		
	Recessive	c) Codominant	d) epistatic		
694. The first definite proof of mut	-				
	Hooker	c) Lister	d) Leeuwenhoek		
695. If the genotype of an individu					
	Heterozygous	c) Monoallelic	d) Uniallelic		
696. The nucleoprotein structures					
	Telomeres	c) Centromeres	d) Satellites		
697. In polytene chromosomes dan	rk bands are visible. I h	=			
a) Protein particles		b) Chromomeres on chrom	nonemata		
c) Nucleosomes	1	d) None of the above			
698. Chances of segregation of alle					
	35%	c) 50%	d) 75%		
699. In <i>Drosophila</i> , gene for white		sponsible for depigmentation	ion of body parts. Thus , a		
gene that controls several pho			م الم الم		
, , , , , , , , , , , , , , , , , , , ,	Epistatic gene	c) Hypostatic gene	d) Pleiotropic gene		
700. Hypertrichosis is an example	of which inneritance?	h) In complete con linked			
a) Holandric		b) Incomplete sex-linked			
c) Sex –influenced	allouing is	d) Sex –limited			
701. The mutagenic agent among f a) Ethyl methane b)	Ethylene	c) 2, 4-D	d) IAA		
702. The most important example	5	=	uj IAA		
	Night blindness	c) Down's syndrome	d) Sickle-cell anaemia		
703. When tall and dwarf plants ar	•	, ,			
	tt and tt	c) Tt and Tt	d) TT and Tt		
704. Failure of segregation of chro					
which as called	illatiu uuring teli uivisi	on cycle results in the gain			
	Hypopolyploidy	c) Hyperpolyploidy	d) Polyploidy		
705. Genes are present on	πγρομοιγμισιαγ		uj i olypioluy		
_	Lamellae	c) Plasma membrane	d) mesosomos		
706. Out of 7 contrasting trait pair		-	d) mesosomes		
	8 and 6	-			
		c) 6 and 8	d) 5 and 9		
707. Example of environmental de	-		d) All of those		
, , , , , , , , , , , , , , , , , , , ,	Turtles	c) <i>Bonelia</i>	d) All of these		
708. Dominant allele are expressed	u III				

a) Second generation	b) Homozygous conditi	on
c) Heterozygous condition	d) Both (b) and (c)	
709. If the ratio between X-chromosomes and com		en the individual will be
a) Female b) Superfemale	c) Male	d) Supermale
710. When a tall plant with rounded seeds (TTRR)	2	
F_1 -generation consists of tall plants with roun	-	
produce?		
a) One b) Three	c) Four	d)
711. The leaf colour of certain plants is controlled h	,	•
green. You have a plant with orange leaves, bu		
If you cross your unknown plant with one of the	_	
determine your unknown's genotype. With wh		
a) GG b) Gg	c) Gg	d) Either of parents
712. Which of the following discoveries resulted in	, .	
a) Recombination of linked genes		
b) Genetic engineering		
c) X-rays induce sex-linked recessive lethal m	utations	
d) Cytoplasmic inheritance		
713. A boy has a normal brother and a colourblind	sister. What is true about his p	arents?
a) His father was normal but mother was colo		
c) Both father and mother were colourblind	d) Both father and mot	
714. By seeing the ratio of F_1 and F_2 -generation Me	-	
unchanged over successive generation and cal		
a) Alleles b) Genes	c) Chromosomes	d) Factors
715. Extranuclear inheritance is a consequence of p	•	
a) Mitochondria and chloroplasts	b) Endoplasmic recticu	lum and mitochondria
c) Ribosomes and chloroplast	d) Lysosomes and ribo	
716. The F_2 genotypic ratio of monohybride cross is		
a) 0% b) 25%	c) 50%	d) 100%
717. Colour blindness is due to defect in		2
a) Cones b) Rods	c) Rods and cones	d) Rhodopsin
718. In F_2 -generation, quantitative inheritance 1 : 4	4 : 6 : 4 : 1 is obtained instead o	
a) 9 : 3 : 3 : 1 b) 8 : 6 : 4 : 1	c) 7:4:1:4	d) 6 : 6 : 4 : 7
719. Leaf colour in <i>Mirabilis jalapa</i> is an example o	of	
a) Non-Mendelian inheritance	b) Mendelian inheritan	се
c) Chemical inheritance	d) Both (b) and (c)	
720. I. Trisomy of sex (X) chromosome		
II. XXY+44		
III. 21st trisomy		
IV. Sterile male		
V. Gynaecomastia		
Choose the correct option for Klinefelter's syn	drome	
a) I, II, III and IV b) I, II, IV and V	c) II, III, IV and V	d) I, III, IV and V
721. Consider the following statement regarding A	BO blood group in human	
I. It is controlled by multiple allele		
II. It shows codominance		
III. Codominance can be manifested phenotyp	ically in human	
IV. It follows the Mendel law of inheritance		
Which of the following statements (s) are corr	rect?	
a) Only I is correct	b) I and II are correct	
c) II and III are correct	d) IV and II are correct	

722. Brachydactyly is due to a) Dominant gene on the autosome b) Recessive gene on the autosome c) Dominant gene on the sex chromosome d) None of the above 723. Which of the following chromosomal formulation is responsible for the expression of meta-male character in Drosophila? a) 2A+3X b) 3A+3X c) 4A+3X d) 3A+XY 724. When there are more than two allele controlling the same character. These are called a) Many alleles b) Polyalleles c) Multiple alleles d) All of these 725. Monohybrid cross deals with b) Two character a) One character c) Three characters d) Four characters 726. X-chromosomes of female, in a case of sex-linked inheritance, can be passed on to a) Only female progeny b) Only male progeny c) Only in grand daughter d) Male and female progeny 727. Identify the type of mutation in given diagram DNA AACTGAT CC Gene mutation a) Inversion b) Insertion c) Deletion d) Substitution 728. The recessive parental trait is expressed without any blending in the F_2 -generation, we can infer. That F_1 plants produce gamete by the process of ...A... and allele of parental pair separate ...B... from each other and only one gamete is transmitted a gamete. Here A and B are a) A-mitosis; B-aggregate b) A-meiosis; B-segregate d) A-mitosis; B-segregate c) A-meiosis; B-aggregate 729. If a cross between two individuals produces offspringe with 50% dominant character (A) and 50% recessive character (a), then the genotypes of parents are a) Sex linked alleles b) Asexually reproducing forms c) Sexually interbreeding forms d) Diploid homozygous forms 730. The similar and dissimilar sex chromosomes of females and males are described as a) Hormomorphic b) Heteromorphic c) Both (a) and (b) d) Isomorphic 731. Starch synthesis gene in pea plant in heterozygous condition produces starch grain of intermediate size. This shows a) Complete dominance b) Incomplete dominance c) Codominant d) Dominant 732. Select the correct bases of DNA, RNA and amino acid of beta chain resulting in sickle cell anaemia. Amino Acid DNA RNA a) CTC/GAG GUG Glutamic acid b) CAC/GAG Valine GUG c) CAC/GTC GAG Valine d) CTC/GAG GUG Valine

PRINCIPLES OF INHERITANCE AND VARIATION

	: ANSWER KEY :														
1)	b	2)	а	3)	С		a 17		b	174)	b	175)	а	176)	а
-) 5)	c	6)	d	-) 7)	a	-		77)	b	178)	c	179)	d	180)	b
9)	а	10)	b	11)	d			B1)	а	182)	b	183)	d	184)	b
13)	d	14)	С	15)	а		a 18	85)	b	186)	b	187)	а	188)	а
17)	С	18)	а	19)	d		c 18	89)	а	190)	а	191)	С	192)	d
21)	с	22)	а	23)	а	24)	d 19	93)	С	194)	С	195)	d	196)	С
25)	а	26)	b	27)	С	28)	a 19	97)	d	198)	а	199)	b	200)	а
29)	С	30)	С	31)	С	32)	c 20	01)	b	202)	а	203)	d	204)	С
33)	а	34)	С	35)	С	36)	b 20	05)	а	206)	d	207)	d	208)	b
37)	b	38)	d	39)	а	40)	d 20	09)	а	210)	а	211)	С	212)	d
41)	d	42)	а	43)	С	44)	a 21	13)	а	214)	d	215)	а	216)	С
45)	b	46)	b	47)	b	48)	a 21	17)	а	218)	b	219)	а	220)	b
49)	d	50)	С	51)	а	52)	a 22	21)	С	222)	d	223)	а	224)	а
53)	а	54)	d	55)	b	56)	c 22	25)	а	226)	b	227)	b	228)	а
57)	а	58)	а	59)	а	60)	a 22	29)	а	230)	С	231)	d	232)	С
61)	d	62)	b	63)	С	64)	d 23	33)	d	234)	d	235)	d	236)	а
65)	С	66)	С	67)	b	68)	d 23	37)	b	238)	С	239)	b	240)	а
69)	С	70)	d	71)	С	72)	d 24	41)	а	242)	а	243)	а	244)	b
73)	d	74)	С	75)	С	76)	d 24	45)	а	246)	d	247)	а	248)	а
77)	а	78)	а	79)	а	80)	c 24	49)	d	250)	d	251)	b	252)	а
81)	С	82)	b	83)	b	84)	a 25	53)	b	254)	а	255)	b	256)	b
85)	d	86)	а	87)	а	88)	c 25	57)	b	258)	а	259)	а	260)	d
89)	С	90)	С	91)	а	92)	d 26	51)	b	262)	а	263)	d	264)	С
93)	а	94)	а	95)	d	96)	a 26	65)	b	266)	d	267)	b	268)	b
97)	d	98)	С	99)	С	100)	c 26	69)	b	270)	а	271)	С	272)	b
101)	С	102)	b	103)	С	104)	c 27	73)	b	274)	а	275)	b	276)	d
105)	С	106)	а	107)	а	108)	b 27	77)	b	278)	b	279)	b	280)	b
109)	d	110)	а	111)	С	,		81)	а	282)	а	283)	а	284)	а
113)	а	114)	d	115)	d	-	a 28	-	b	286)	С	287)	d	288)	d
117)	b	118)	С	119)	а	-	d 28	-	а	290)	а	291)	b	292)	С
121)	а	122)	а	123)	b	,	b 29	-	b	294)	а	295)	а	296)	b
125)	С	126)	а	127)	С	,	b 29	,	С	298)	С	299)	а	300)	b
129)	d	130)	С	131)	а	-	c 30	-	b	302)	d	303)	d	304)	b
133)	b	134)	d	135)	d	-	b 3(-	b	306)	b	307)	С	308)	С
137)	С	138)	d	139)	С	-	d 30	-	b	310)	a	311)	b	312)	b
141)	a	142)	b	143)	а		c 31	-	а	314)	d	315)	а	316)	d
145)	b	146)	b	147)	а	-	a 31	-	C	318)	a	319)	C	320)	b
149)	С	150)	d	151)	а	-	a 32	-	b	322)	d	323)	b	324)	b
153)	С	154)	d	155)	а	-	a 32	-	b	326)	С	327)	d	328)	а
157)	а	158)	С	159)	a	-	c 32	-	а	330)	С	331)	a	332)	а
161)	С	162)	С	163)	d	-	c 33	-	а	334)	a	335)	b	336)	а
165)	С	166)	С	167)	С	-	b 33	-	а	338)	d	339)	d	340)	а
169)	С	170)	С	171)	С	172)	c 34	¥1)	С	342)	b	343)	а	344)	а

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345)	а	346)	d	347)	b	348)	b	549)	d	550)	С	551)	d	552)	а
349)	b	350)	d	351)	b	352)	a	553)	b	554)	С	555)	b	556)	d
353)	а	354)	С	355)	d	356)	a	557)	С	558)	а	559)	b	560)	b
357)	а	358)	b	359)	С	360)	С	561)	а	562)	d	563)	С	564)	d
361)	С	362)	b	363)	d	364)	b	565)	а	566)	а	567)	d	568)	а
365)	С	366)	С	367)	а	368)	а	569)	d	570)	d	571)	b	572)	d
369)	d	370)	а	371)	а	372)	а	573)	С	574)	b	575)	b	576)	b
373)	b	374)	а	375)	d	376)	d	577)	d	578)	С	579)	b	580)	d
377)	а	378)	а	379)	b		b	581)	С	582)	b	583)	С	584)	d
381)	а	382)	а	383)	а	,	b	585)	а	586)	а	587)	d	588)	а
385)	С	386)	С	387)	а	,	b	589)	d	590)	С	591)	d	592)	С
389)	b	390)	b	391)	d	,	a	593)	а	594)	b	595)	b	596)	а
393)	С	394)	a	395)	а	,	С	597)	С	598)	b	599)	d	600)	С
397)	b	398)	b	399)	C	2		601)	d	602)	d	603)	C	604)	С
401)	а	402)	b	403)	b	2		605)	a	606)	b	607)	b	608)	С
405)	a	406)	а	407)	b	2		609)	b	610)	a	611)	а	612)	а
409)	b	410)	C	411)	а	-		613)	а	614)	b	615)	С	616)	С
413)	d	414)	b	415)	а	2		617)	С	618)	a	619)	С	620)	а
417)	а	418)	а	419)	a	,		621)	С	622)	b	623)	С	624) (20)	а
421) 425)	а	422)	С	423)	b	2		625)	С	626)	d	627)	а	628)	С
425) 420)	C	426) 420)	C h	427) 421)	b			629)	C h	630) (24)	C	631) (25)	a h	632) (2()	C d
429) 422)	a	430) 424)	b	431) 435)	b d	-		633) (27)	b d	634) (29)	C d	635) (20)	b h	636) (40)	d d
433) 437)	C C	434) 429)	C	435) 420)	d	2		637) 641)	d	638) 642)	d	639) 642)	b	640) 644)	d h
437) 441)	a d	438) 442)	a d	439) 443)	C b	2		641) 645)	a c	642) 646)	C	643) 647)	a b	644) 648)	b
441) 445)	d C	442) 446)	d d	443) 447)	b b			649)	с а	650)	a b	651)		652)	a d
449)	с а	440) 450)	u C	451)	b	2		653)		654)	d	655)	a d	656)	u d
453)	a b	450) 454)	с b	451) 455)	a	-		657)	a d	658)	u b	659)	u a	660)	u a
457)	c	458)	b	459)	a C		c	661)	b	662)	b	663)	a C	664)	a b
461)	b	462)	c	463)	c	2		665)	C	666)	a	667)	b	668)	d
465)	d	466)	d	467)	a	-		669)	d	670)	b	671)	a		c
469)	b	470)	d	471)	d	-		673)	c	674)	d	675)	d	676)	d
473)	c	474)	a	475)	b			677)	d	678)	a	679)	b	680)	a
477)	С	, 478)	d	479)	b	-		681)	a	682)	d	683)	d	684)	b
481)	а	482)	а	483)	b			685)	С	686)	b	687)	d	688)	d
485)	а	486)	d	487)	b	-		689)	b	690)	d	691)	а	692)	d
489)	d	490)	d	491)	d	,		693)	а	694)	а	695)	а	696)	b
493)	с	494)	а	495)	а	496)	с	697)	b	698)	С	699)	d	700)	а
497)	а	498)	а	499)	а	500)	с	701)	а	702)	d	703)	а	704)	а
501)	а	502)	а	503)	b	504)	a	705)	а	706)	а	707)	d	708)	d
505)	а	506)	а	507)	b	508)	a	709)	С	710)	d	711)	b	712)	С
509)	d	510)	b	511)	d	512)	a	713)	b	714)	d	715)	а	716)	С
513)	а	514)	С	515)	b	516)	a	717)	а	718)	а	719)	а	720)	b
517)	С	518)	а	519)	b	520)	С	721)	b	722)	С	723)	b	724)	С
521)	d	522)	С	523)	b	524)	a	725)	а	726)	d	727)	b	728)	b
525)	С	526)	d	527)	а	528)	b	729)	b	730)	С	731)	b	732)	b
529)	С	530)	С	531)	d	532)	d								
533)	С	534)	С	535)	а	536)	b								
537)	b	538)	а	539)	С	540)	b								
541)	а	542)	а	543)	а		d								
545)	а	546)	С	547)	а	548)	С								

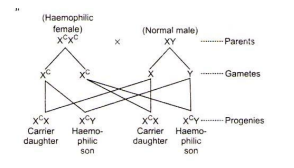
PRINCIPLES OF INHERITANCE AND VARIATION

BIOLOGY

: HINTS AND SOLUTIONS :

1 **(b)**

Haemophilia is a recessive X-linked disease. A female with defected single X-chromosome is normal but, carrier of disease, and male with defected single X-chromosome is haemophilic.



2 **(a)**

Genetic disorder may be grouped into two categories

(i) **Mendelian Disorders** These genetic disorder are mainly caused by alternation and mutation in the single gene. They are transmitted to offsprings following the principle of inheritance. Mendelian disorder can be dominant or recessive. *e. g.*, haemophilia, colour blindness, sickle-cell anaemia, cystic fibrosis, phenylketonuria, thalassaemia.

(ii) **Chromosomal Disorders** Chromosomal disorder are caused due to excess, absence, or abnormal arrangement of one or more chromosome, *e.g.*, Turner's syndrome, Down's syndrome, etc

3 **(c)**

The term gene was coined by Johanssen.

4 **(a)**

A dihybrid cross involves two pairs of contrasting characters, *e.g.*, yellow round seeded plant and wrinkled. Green seedes plant(both pure lines) homozygous. When a dihybrid cross is made between two pure line of homolzygous parents, then the F_1 generation shows hybrids with dominant phenotypic effect. When F_1 heterozygous plants are self-ferilized to produce F_2 generation, four types of combinations are obtained of which two are similar to parental combination and other two are new combinations. The phenotypic dihybrid ratio of these four combinations in F_2 generation comes out to be 9:3:3:1, while the genotypic dihybrid ratio is 1:2:2:4:1:2:1.

5 **(c)**

Chromosome is made up of DNA and histone proteins.

6 **(d)**

Baldness is not a sex-limited trait. Balaness is a sex influenced trait.

Linkage is an exception to the principle of independent assortment in heredity.

Galactosemia is a hereditary disease that is caused by the lack of a liver enzyme required to digest galactose.

Small population size results in random genetic drift in population.

7 (a)

The F_1 offsprings of pure tall and pure dwarf are heterozygoous tall, which on selfing produces 1 : 1 ratio of breeding tall to breeding dwarf.

8 **(b)**

Exposure of 'X' rays enhance the frequency of crossing over

9 **(a)**

The genotype of trihybrid would be AaBbCc. Eight different types of gametes ABC, ABc, AbC, Abc, aBC, aBc, abC, abc would be formed. The number of zygotes would be $8^2 = 64$.

10 **(b)**

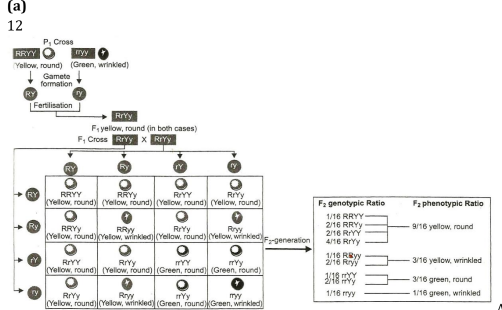
The genetic composition of an organism, *i.e.*, the combination of all alleles possessed by an organism is called genotype

11 **(d)**

In polyploidy there are more than one set of chromosomes is presen't in an organisms. It only

happens when cytokinesis doesn't take place in proper way

12 **(a)**



A dihybrid cross in pea

plant between yellow round (smooth) seeded and green wrinkled seeded plant. The cross proves the principle of independent assortment

13 **(d)**

When the F_1 -hybrid (Rr Yy) of a dihybrid cross is test crossed (crossed with double recessive parent rryy), the F_2 -offspring appear in the phenotypic and genotypic ratio of 1:1:1:1confirming that F_1 -offspring was heterozygous in both the traits. It is a cross between RrYy × rryy.

14 **(c)**

Allelomorphs or simply called allele represents a pair of contrasting characters

15 **(a)**

Harmful mutation does not get elimated from the gene pool because most of the harmful mutations are recessive and they carried by heterozygous condition in the individual. If they (mutation) are dominant then they easily get eliminated by the death of an organism

16 **(a)**

A-Common, B-Rare

17 **(c)**

Mendel died in 1884 long before his work came to recognized. It was in 1900 when three worker independently rediscovered the principles of heredity already worked out by Mendel. They were Hugo de Vries of Holland, Carl Correns of Germany and Eric Tiron and Tschermark of Austria

18 **(a)**

In gynandromorphs, some cells of body contain

XX and some cells XY genotype.

19 **(d)**

Post Mendelian Discoveries

Gene interaction is the influence of alleles and nonalleles ion the normal phenotypic expression of genes. It is two types, intragenic (allelic) and intergenic (nonallelic). In the intragenic interaction the two allels (present on the same gene locus on the two homologous chromosome) of a gene interact in such a way as to produce a phenotypic expression different from typical dominant-recessive phenotype, e.g., incomplete dominance, codominance, multiple alleles. In intergenic or non-allelic interaction, two or more independent gene present on the same or different chromosomes interact to produce different expression, e.g., epistasis, duplicate genes, complementary genes, supplementary genes, lethal genes, inhibitory genes, etc.

20 **(c)**

Intermediate inheritance is incomplete dominance in which dominant factor of a heterozygote does not completely mask the expression of recessive allele. In incomplete dominance, genotypic and phenotypic ratio remain the same and is 1 : 2 : 1.

21 **(c)**

Green pod colour is dominant. 7 dominant traits, 7 recessive traits total 14 traits

or 7 oppossing pairs of traits

Characters	Dominant Traits	Recessive Traits
Seed shape	Round	Wrinkled
Seed colour	Yellow	Green
Flower	Violet	White
colour		
Pod shape	Full	Constricted
Pod colour	Green	Yellow
Flower	Axial	Terminal
position		
Stem	Tail	Dwarf
height		

22 (a)

 ${\rm I}^{\rm A}\,{\rm I}^{\rm B}$ are the dominant form of I gene, I is recessive form

23 (a)

A-statistical analysis; B-mathematical logic

24 (d)

ZW and ZZ and ZOZZ.

ZW and ZZ Type of Sex Determination This mechanism operates in certain insects (butterflies and moths) and in vertebrates (fishes, reptiles and birds). The male has two homomorphic sex chromosomes (ZZ) and is homogametic and the female has two heteromorphic sex chromosomes (ZW) and is heterogametic. There are thus two

types of eggs with Z and with W and only one type of sperms. i.e., each with Z

	A + Z	A + O
A + Z	AA + ZZ	AA + ZO
A + Z	AA + ZZ	AA + ZO
	Males	Females

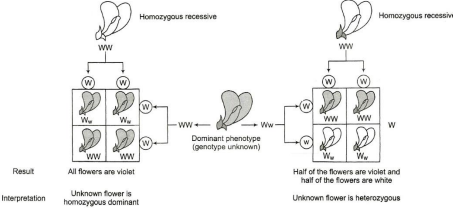
WZ-ZZ types of sex determination

25 (a)

If a character is expressed equally in the homozygous and heterozygous conditions, it is called **dominant** and the other character is said to be recessive. In given question, 'Tall' character is dominant over 'dwarf', hence, the cross shows dominance and segregation of traits.

26 **(b)**

The best method to determine homozygosity and heterozygosity of an individual is back cross. Crossing of F₁ (dominant phenotype) to any one of the parent called back cross and when phenotype of crossing parent is recessive than this is called test cross. The progeny of such cross can easily be analysed to predict the genotype of test organism



27 (c)

Presence of one Barr body indicates the person under investigation is a normal female.

28 (a)

A-Male, B-Female, C-Gametes.

XY and XY type sex determination seen in many insect and mammals including humans. Males have X and Y chromosome along with autosome

and females have pair of 'X' chromosome along with autosome Parents Phenotypes Male Female

Genotypes 44A + XY 44A + XXGametes 22A + X 22A + Y22A + A22A + X22A+X 22A+X

Children 22A + X 44A + XX 44 A + XY Female

22A + Y 44 A + XY 44 A + XY

Male

Sex ratio Female : Male = 1 : 1

In plants The flowering plants are mostly bisexual and lack sex chromosomes. The unisexual flowering plants tent to have XX-XY type of sex chromosomal mechanism for sex determination. The female plants are XX and male plants are XY.

XX and XO Type of Sex Determination

Found in insect like grasshopper, cockroaches and bugs. Males have only X sex-chromosome and autosomes, female have pair of X-chromosome and autosome

Parents	Phenotypes	Male	Female
	Genotypes	AA + XO	AA + XX
Gametes	Α	+ X, A + O	A + X, A + Y

 F_1 -generation

	A + X	A + X			
	AA + XO				
A + O	AA + XO	AA + XO			
	Genotypes				

XX-XO type of sex determination

29 **(c)**

Female is sterile.

Disorders	Autosomal/Sex Linked	Symptoms	Effects
Down's syndrome	Autosomal aneuploidy (trisomy, +21)	Mongolian eyefold (epicanthus), open mouth, protruded tongue, projected lower lip, many loops on finger tip, palm crease	Retarded mental development IQ (below 40)
Turner's syndrome	Sex chromosomal monosomy 44 + XO	Short stature females (<5'), webbed neck, body hair absent menstrual cycle absent. Sparse pubic hair, underdeveloped breasts narrow lips puffy fingers	Sterile hearing problem
Klinefelter's syndrome	Sex chromosomal aneuploidy (tri/tetrasomy of X chromosome) 44+XXY 44+XXY	The males are tall with long legs, testes small, sparse body hair, Barr body present, breast enlargement	Gynaecomast ia azospermia sterile

Some Examples of Aneuploidy

(i) Down's syndrome-21 trisomy

Symptoms

- (a) Short statured with small round head
- (b) Partially open mouth with protruding furrowed tongue
- (c) Palm is broad with characteristic palm crease
- (d) Slow mental development

(ii) Turner's syndrome

Cause Absence of one of the X-chromosomes, resulting in the karyotype 44+X0

Symptoms

- (a) Sterile female with rudimentary ovaries
- (b) Shield-shaped thorax
- (c) Webbed neck
- (d) Poor development of breasts
- (e) Short stature, small uterus, puffy fingers

(iii) Klinefelter's syndrome

Cause Presence of an additional copy of X-chromosome resulting in the karyotype 44+XXY **Symptoms**

- (a) Sex of the individual is masculine but possess feminine characters
- (b) Gynaecomastia, *i.e.*, development of breasts
- (c) Poor beard growth and often sterile
- (d) Feminine pitched voice

30 **(c)**

The environmental stress (as pesticides) does not cause the direct changes in genome, instead, it simply selects rather persisting mutations, which result in phenotypes that are better adapted to the new environment (*e.g.*, certain pesticides).

31 **(c)**

A **Lampbrush chromosomes** is made up of two homologous chromosomes held at several places by chiasmata. The chromosomes are found in oocytes of many invertebrates and all vertebrates except some mammals. Lampbrush chromosomes are found during the extended diplotene phase of first meiotic division.

32 **(c)**

A gene consists of a polynucleotide sequence that encodes a functional polypeptide or RNA sequence.

33 **(a)**

XY and XY type sex determination seen in many insect and mammals including humans. Males have X and Y chromosome along with autosome and females have pair of 'X' chromosome along with autosome

Parents Phenotypes Male Female

Genotypes 44A + XY

44A + XX			
Gametes		22A + X 22A+	-Y
22A + A2	22A + X		
		22A+ X	22A+X
Children	22A + X	44A + XX	44 A + XY
Female			
	22A + Y	44 A + XY	44 A + XY

Male

Sex ratio Female : Male = 1 : 1

In plants The flowering plants are mostly bisexual and lack sex chromosomes. The unisexual flowering plants tent to have XX-XY type of sex chromosomal mechanism for sex determination. The female plants are XX and male plants are XY.

XX and XO Type of Sex Determination

Found in insect like grasshopper, cockroaches and bugs. Males have only X sex-chromosome and autosomes, female have pair of X-chromosome and autosome

Parents	Phenotypes	Male	Female
	Genotypes	AA + XO	AA + XX
Gametes	s A	+X, A + 0	A + X, A + Y
F ₁ -gene	ration		
A	$+ X \qquad A + X$		

	A + X	A + X	
A + X	AA + XO	AA + XO	
A + O	AA + XO	AA + XO	
	Genotypes		

XX-XO type of sex determination

34 **(c)**

Frameshift mutations are the mutations caused by

insertion (*i.e.*, addition) or deletion of one or more nitrogen bases in the DNA or RNA. This type of mutation alters the nucleotide sequences in all the genes and hence, the genetic code is changed totally, fro the point of mutation which results in the change in biochemical behaviour of the genes. **Base pair substitution mutations** involve **substitution of a aitrogen base** by another base or by some derivative of nitrogen base.

35 **(c)**

According to law of segregation, the heredity character in the form of alleles segregate from each other during gamete formation, *i.e*, each gamete carry only one allele of each gene. This is also called law of purity of gametes. When tall and dwarf plants are crossed only tall plants are produced in F_1 generation. By selfing of these F_1 plants tall and dwarf plants produced in 3:1ratio.

36 **(b)**

Mendel's law are able to predict accurately the pattern of inheritance for a situation in which alleles shows the complete dominance. Effect of environment, other alleles did not explained by the Mendel. Mendel did not know about the polygenic traits also

37 **(b)**

Blood group-O has no antigens but A and B antibodies.

38 **(d)**

Colour blindness disease was detected by **Wilson** in 1910.

39 **(a)**

In the dihybrid cross between RRYY and rryy parents, the number of RrYy genotypes in F_2 -generation will be four.

40 **(d)**

Allelism refers to presence of alternative forms of a gene at a given locus. Alleles or allelomorphs are the two contrasting aspects of the same character present at a locus of homologous pair of chromosomes. Now –a-days, the same aspect in duplicate (TT or tt) of a character is also considered an allele.

41 **(d)**

The women with albinic father has gene for

albinism. When this women marries with albinic men, they produce normal and albinic in 1 : 1 ratio.

42 **(a)**

ZO and ZZ type of sex determination. This mechanism occurs in certain buttlerfiles and moths. The female is heterogametic and produces two types of eggs half with Z and half without Zchromosome. The males have homomorphic sex chromosomes and is homogametic. It forms only one kind of sperms, each with Z-chromosome

ParentsPhenotypesMaleFemaleGenotypesAA + ZZAA + ZOGametesA+Z, A+ZA+Z, A+OF1-generation F_1 -generation

$$\begin{array}{c|c} A+Z & A+O \\ \hline A+Z & AA+ZO \\ A+Z & AA+ZZ & AA+ZO \\ \hline Males & Females \end{array}$$

ZO-ZZ type of sex determination

43 **(c)**

Mendel's law of independent assortment states that,"the alleles of different genes segregate independently of each other during meiosis".

44 **(a)**

XX and XO chromosome.

XY and XY type sex determination seen in many insect and mammals including humans. Males have X and Y chromosome along with autosome and females have pair of 'X' chromosome along with autosome

Parents Phenotypes Male Female

Genotypes 44A + XY 44A + XX **Gametes** 22A + X 22A + Y 22A + A22A + X **Children** 22A + X 44A + XX 44A + XYFemale 22A + Y 44A + XY 44A + XYMale

Sex ratio Female : Male = 1 : 1

In plants The flowering plants are mostly bisexual and lack sex chromosomes. The unisexual flowering plants tent to have XX-XY type of sex chromosomal mechanism for sex determination. The female plants are XX and male plants are XY. **XX and XO Type of Sex Determination**

Found in insect like grasshopper, cockroaches and

bugs. Males have only X sex-chromosome and autosomes, female have pair of X-chromosome and autosome

Genotypes AA + XO

Male

Female

AA + XX

A + X, A + O A + X, A + Y

Gametes

Parents

 F_1 -generation

 $\begin{array}{c|c} A + X & A + X \\ A + X & AA + XO & AA + XO \\ A + O & AA + XO & AA + XO \\ \hline Genotypes \end{array}$

XX-XO type of sex determination

Phenotypes

45 **(b)**

A-Chromatid, B-Allele pair, C-Genetic composition

46 **(b)**

The position of centromere determines the shape of chromosome.

47 **(b)**

After schooling Mendel joined Augustinian monastery of St. Thomas at Brunn (then in Austria now Brunn in Czechoslovakia) in 1843 at the age of 21. At the age of 25 (1847), he was made a prist in that monastery

48 **(a)**

It was TH Morgan who clearly proved and define linkage on the basis of the breeding experiments in fruitfly. In 1911, Morgan and Castle proposed 'chromosomal' theory of linkage'

49 **(d)**

Francis Galton(1885) gave the term eugenics. Eugenics is the improvement of human race by the application of principles of genetics. The other meaning of eugenics is 'science of being well born'.

50 **(c)**

The **test cross** involves the crossing of F_1 hybrid with a double recessive genotypic parent. By test cross, the heterozygocity and homozygocity of the organism can be tested.

51 **(a)**

Tetraploid endosperm is obtained, when a diploid female and tetraploid male plants are crossed.

52 **(a)**

Colour Blindness

(i) It is a sex-linked recessive disorder

(ii) It results in defect in either red or and green cone cells of eye resulting in failure to discriminate between red and green colour

(iii) The gene for colour blindness is present on X-

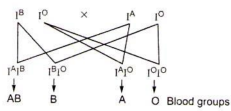
chromosome

(iv) It is observed more in males (X^cY) because of presence of only one X-chromosome as compared to two chromosomes in famales

53 **(a)**

When the F_1 -hybrid is crossed with recessive parent, both phenotypes appear in progeny and this is called test cross. It gives 1 : 1 ratio in monohybrid cross and 1 : 1 : 1 : 1 ratio in dihybrid cross.

54 (d)



Thus, the genotype of parents will be $BO \times AO$.

55 **(b)**

Symbol in pedigree chart represents still death 56 **(c)**

Histones are basic proteins found in the eukaryotic chromosomes. These are rich in basic amino acids lysine and arginine. There are basically five types of histones, *i.e.*,

 H_1 , H_2A , H_2B , H_3 and H_4 which have been studied in almost all eukaryotic cells.

57 **(a)**

In monosomic condition, one chromosome is missing from the somatic chromosome complement. It is denoted by 2n-1. When somatic cells of an organism contain three copies of one chromosome, the condition is known as trisomy. It is denoted by 2n+1. Therefore, monosomic trisomy is represented as 2n-1+1.

58 **(a)**

Baldness is common in humans. Hereditary baldness is carried by a dominant autosomal gene. It develops only in men and never in women.

59 **(a)**

Thalassaemia

(i) It is an autosome-linked recessive disesase(ii) It occurs due to either mutation or deletion resulting in reduced rate of synthesis of one of globin chains of haemoglobin

(iii) Anaemia is the characteristic of this disease

(iv) Thalassaemia is classified into two types

- 1. α -thalassaemia Production of α -globin chain is affected. It is controlled by the closely linked genes HBA1 and HBA2 on chromosome 16. It occurs due to mutation or deletion of one or more of the four genes.
- β-thalassaemia Production of β-globin chain is affected. It occurs due to mutation of one or both HBB genes on chromosome 11

60 **(a)**

The term 'mutation' was introduced by Hugo de Vries in 1901 and his mutation theory of evolution called mutation theory of evolution. Mutation is new sudden inheritable change in organism due to permanent change in their genotype

61 **(d)**

Because in sex linked inheritance the chance of girl or female to be affected is almost nill. Generally, the females are carriers and in heterozygous condition

62 **(b)**

Non-disjunction is the condition in which the separation of chromosome doesn't take place during cell division. In 44+XY non-disjunction there is non-separation of XY gene is there, which leads to the formation of sperm having genotypes, 22+XY and 22

63 **(c)**

Linkage prevents independent assortment.

64 **(d)**

Sex influenced trait.

Finalization of sex at the time of fertilization is known as sex determination. All sex linked character show criss-cross inheritance and firstly it was studied and discovered by TH Morgan (1910). *Sex related trait may be divided into three types*

(i) Sex Linked Traits They are those traits the determining genes of which are found on the sex chromosomes. All the sex-linked traits present on a sex chromosome are inherited together
(ii) Sex Limited Traits They are autosomal traits which are expressed in a particular sex in response to sex hormones although their genes also occur in the other sex, *e.g.*, milk secretion in mammalian females, pattern baldness in males. The gene for baldness behaves as an autosomal

dominant in males and autosomal recessive in females

(iii) **Sex Influenced Traits** The traits are not due to particular genes but are by products of sex hormones, *e.g.*, low pitched voice, beard moustaches. In males, pattern baldness is related to both autosomal genes as well as excessive secretion of testosterone

5 **(c)**

Since in an individual only two alleles can be present, multiple alleles can be found only when population studies are made

66 **(c)**

Biological concept of species says that only the members of a species can breed freely in nature to produce fertile offsprings. The plant tobacco (*Nicotiana*) has two different species, *Nicotiana tobaccum* and *Nicotiana sylvestris*. These two species cannot reproduce freely.

67 **(b)**

In duplication there is increase in size of genes by duplication of it segment of a chromosome. Mainly seen in case of plants. But in deletion there is loss of genes or segment of chromosome

68 **(d)**

The types of gametes produced by a plant depend upon the number of hetrozygous pair.

Number of types of gametes= 2^n

N=Number of heterozygous pair

 $2^1 = 2$

The gametes are-ABC and AbC.

69 **(c)**

In **trisomic condition**, diploid organism have extra chromosome represented by the chromosomal formula2n + 1. One of the pairs of chromosomes has an extra member, so that a trivalent may be formed during meiotic prophase, *e.g.*, **Down's syndrome** (45+XX or 45+XY), Klinefelter's syndrome (44 + XXY).

In **monosomic**, diploid organism has one chromosome of a single pair missing with genomic formula 2n - 1. Monosomics can from two kind of gametes, (*n*) and (*n*-1),

e.g., Turner's syndrome (44 + X).

70 **(d)**

When a normal man marries a normal woman, whose father was colourblind then their 50% sons are colourblind (50% sons normal) and all the daughters are phenotypically normal(carrier woman also are phenotypically normal).

The female parent is carrier as it receives a defective X-chromosome from her father.

71 **(c)**

Testosterone in male secreted by Leydig cells. In the male testosterone is essential for development of secondary sexual character and related to baldness also

72 **(d)**

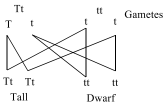
The disorder cystic fibrosis, sickle cell anaemia, colour blindness and haemophilia are caused due to the abnormality in gene (Mendelian factor). Turner's syndrome is due to chromosomal abnormality. It is characterized by 44 autosomes and only one X-chromosome.

73 **(d)**

Haemophilia is a X-linked disorder not Y-linked.

74 **(c)**

(Homozygous tall) × (Homozygous dwarf)



Phenotypic ratio 1 : 1 (50% tall, 50% dwarf) Genotypic ratio 1 : 1

75 **(c)**

 γ – rays generally induce mutations.

76 **(d)**

Since genotypes/phenotypes of both parents are same but only sources of gametes are reversed, these crosses are called **reciprocal crosses**.

77 **(a)**

Man has only one X-chromosome that is inherited to his daughter. Therefore, a hereditary disease, which is X-chromosomal linked, is never passed on from father to son.

78 **(a)**

Bateson gave the coupling and repulsion hypothesis for linkage and crossing over.

Similar genes remain together they don't go for crossing over. Bateson called them coupling gones.

While on the other hand dissimilar gene segregate crossing over takes place. Bateson called them repulsion parents gene

79 **(a)**

The blood group type in human provides an example of multiple allelism (*i.e.*, presence of more than two allele for one gene). There are three A, B and O blood groups allele usually given the symbolI^A, I^B and Iⁱ. I^A and I^B are codominant to each other but both are dominant to Iⁱ. The offspring of parents having I^A and I¹ allele with be I^AIⁱ having blood group-A.

A-I^AIⁱ

B-I^BIⁱ

AB-I^AI^B

0-IⁱIⁱ

80 **(c)**

A person having 45 chromosomes instead of 46 due to lack of Y-chromosome is suffering from **Turner's syndrome.**

81 **(c)**

Opposite phenomena.

Strength of linkage between two genes inversely proposed to the distance of two gene means if two genes are closely placed then they have high percentage of linkage and if they placed for then there is low percentage of linkage. Linkage and recombination are two opposite phenomena. In linkage two genes remains united and in recombination the two get apart due to

crossing over during gametogenesis (meiosis) **(b)**

Johanssen (1909) proposed the term gene. Genes are made up of DNA, *i.e.*, a DNA segment associated with proteins, which can be copied in the form of RNA and is responsible for hereditary characters. Genes have full control over protein synthesis.

83 **(b)**

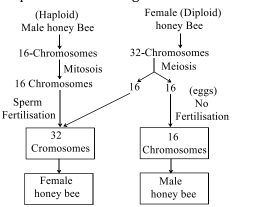
82

2*n* (diploid).

Haploid diploid mechanism of sex determination (haplodiploidy).

Hymenopterous insect such as bees, wasps ants show unique phenomena in which an unfertilized egg develops into male and female develops from fertilized egg.

In honeybee, the quality of food determines whether a diploid larva will become a fertile queen or a sterile worker female. A larva fed on royal jelly a secretion from the mouth of mussing workers grows into a queen, whereas a larva fed on pollen and nectar grows into a worker bee



84 (a)

For the given case, the imprints need to be used only on plates with streptomycin.

85 **(d)**

Phenylketonuria (PKU) is homozygous recessive autosomal disorder associated with metabolism and the gene for it is present on chromosome-12. **Huntington's disease (Huntington's chorea)** is a dominant autosomal disorder due to an allele on short arm of chromosome-4. **Sickle cell anaemia** is due to codominant autosomal allele Hb^S present on chromosome-11.

86 **(a)**

The genes are arranged in a linear fashion on the chromosome.

87 **(a)**

According to cis - trans effect of **Lewis**, when two genetic loci produce identical phenotypes in *cis* and *trans* position, they are considered to be pseudoalleles and phenomenon as **pseudoallelism**.

88 **(c)**

Factor-II	– Prothrombin
Factor –III	– Thromboplastin
Factor –VIII	– Antihaemophilic globulin

Factor – XII – Hageman factor

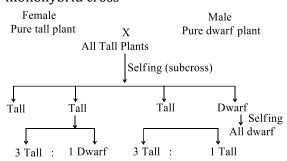
89 **(c)**

The longest chromosome is seen in *Trillium* (30μ) .

90 **(c)**

F₁.

Mendel cross-pollinated a pure tall pea plant (100-120 cm hight) and a pure dwarf pea plant. (only 22 to 44 cm hight). He called them parental generation, expressed now-a-days by symbol P. This hybridization popularly called as monohybrid cross



This three generations of pea plants after crossing a pure tall plant with a dwarf one. The plants of F_1 -generation are all tall, of F_2 -generation three tall and one dwarf. One third of the tall plants are pure, while the remaining behave as hybrids F_1 -generation Seeds collected from the parental generation called first filial generation or F_1 generation

F₂-generation F_1 -plants pollinated among them self (self breeding or inbreeding) and seed produced by F_1 -plants called F_2 -generation. They were in ratio 3:1 (three tall and one dwarf).

F₃-generation Mendel allowed F_2 -plant to form seed by self-pollination called F_3 -generation. Mendel observed that tall and dwarf plant behave differently

(i) Dwarf plant produced dwarf plant on selfpollinated

(ii) In tall plants one third plants breed true so they were pure

(iii) Other two third plant behave like parents and give tall to dwarf plants 3 : 1 indicate that their parents have dwarf genes also

91 **(a)**

A-Linked; B-Unlinked gene

92 **(d)**

Allelic sequence variation has traditionally been described as a DNA polymorphism if more than one variant (allele) at a locus occurs in human population with a frequency greater than 0.01. In simple terms, if an inheritable mutation is observed in a population at high frequency, it is referred to as DNA polymorphism.

93 **(a)**

94

A-Extremely, B-Carrier, C-Haemophilia (a)

The Rh factor causes erythroblastosis foetalis, when a woman who is Rh⁻ marries a man, who is Rh⁺, their first child will be safe (which is Rh⁺) but during pregnancy some blood of foetus and mother mixes due to which the mother develops antibodies against her foetus antigen which is Rh⁺.

95 **(d)**

Landsteiner divided human population into four groups based on the presence of antigens found in their RBCs. Each group represented a blood group. Thus, there are four types of blood groups A, B, AB and O. Blood group-O does not contain any antigen on RBCs, hence can be given to any person, that's why, this blood group is called universal donor.

96 **(a)**

Alleles or allelomorphs are alternative forms of the same gene, *e.g.*, for height of plant 'T' and 't'. Homologous chromosomes are a pair of chromosomes having similar genes, which control the same characters.

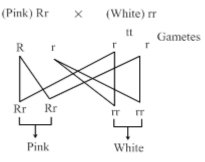
97 **(d)**

Telomeres are ends of chromosome, that have repetitive DNA sequences and are stable and resistant to exonuclease digestion hence, essential for chromosome stability.

98 **(c)**

Intermediate inheritance also called the incomplete dominance. In that inheritance the phenotypic and genotypic ratio are same. The classical examples are = four O' clock plant and snapdragon.

Mirabilis jalapa shows incomplete dominance



The ratio of pink and white flower will be 1 : 1 (c)

Mendelian disorder may be dominant or recessive 100 **(c)**

In African population, sickle cell anaemia provides immunity against malaria.

101 **(c)**

99

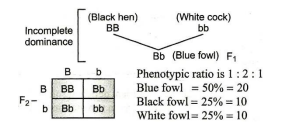
Aneuploidy is the variations in individual chromosomal number. Actually, loss or gain of individual chromosomes upsets the balance and, hence normal development is not possible.

103 (c)

Black feathered hen = BB

White feathered cock = bb

Blue feathered fowl = Bb



104 **(c)**

Histones are special type of basic protein associated with DNA and form chromosome. RNA, protein, carbohydrate, fat, doesn't find in chromosomes

105 **(c)**

In pedigree

Square represents male blackened square or circle represents affected individual. Horizontal line represents-parents The study of inheritance of genetic traits in several generations of a human family in the form of a family tree diagram is called **pedigree analysis**.

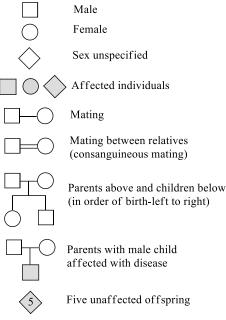
Advantages

(i) It helps in genetic counselling to avoid disorders

(ii) It shows the origin of a trait and flow of a trait

in a family

(iii) It is important to know the possibility of a recessive allele that can cause genetic disorders like colour blindness, haemophilia, etc. *Signosed in the pedigree are*



106 (a)

In the given pedigree chart only males are affected. So, it can be easily inferred that the given trait is connected to Y-chromosome. The genes, which are present on the Y-chromosome are called holoandric genes

107 **(a)**

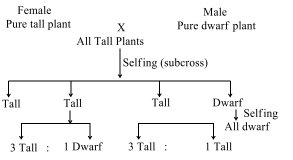
In the gametogenesis meiosis occur. The diploid chromosome become haploid, so the probability of side is sperm lacking one recessive autosomal allele and holandric gene is half

108 **(b)**

 $\rm F_3$ -generation obtained by selfing of $\rm F_2$ -generation.

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109 **(d)**

W Bateson and **R** C Punnett observed complementary gene interaction for flower colour in sweet pea (*Lathyrus odoratus*). In complementary interaction, two separate pairs of genes interact to produce the phenotype in such a way that neither of the dominant genes is expressive unless the other one is present. In F_2 generation, complementary genes produce a ratio of 9 : 7.

110 **(a)**

Occasionally a single gene product may produce more than one effect. For example starch synthesis in pea seeds is controlled by one gene. It has two allele (B and b). Starch is synthesized effectively by BB and have bigger grains. In contrast bb homozygous have lesser efficiency in starch synthesis and produce smaller grains

111 (c)

In *Drosophila*, sex is determined by the ratio number of X-chromosomes to the set of autosomes.

112 (c)

The genotypic and phenotypic ratio of **1** : **2** : **1** with **red**, **pink** and **white** flowers are produced in *Mirabilis jalapa*, when red flowered plants (RR) are crossed with white flowered (rr). It occurs

due to allelic gene interactions, called, **incomplete dominance**. In which, both of the allelomorphic genes will have partial or incomplete dominance and F_1 -hybrid will show mixture of characters of two parents.

113 **(a)**

Meiosis is an important stage in sexual reproduction. During meiosis, genetic recombination occurs as a result of crossing over.

114 **(d)**

Blood group of children may be A, B, AB and O.

115 **(d)**

The genes, which are present very far from each other tend to get unlinked and they arethe most chances for crossing over

116 **(a)**

Given diagram depicits the sex linked inheritance in given options haemophilia is the sex-linked character

117 **(b)**

Incomplete dominance or blending inheritance is the phenomenon, in which the two genes of allelomorphic pair are not related as dominant or recessive but each of them expresses itself partially, thus the F_1 hybrids exhibit a mixture or blending of characters of both the parents. In F_2 generation, the phenotypic ratio obtained is 1:2:1.

118 **(c)**

Haemophilia is a disease, which is caused due to lack of blood clotting factor. It appears only in human male which can be transferred to their grandson through his carrier daughter.

119 **(a)**

In the given option only codominance does not obey Mendel's laws.

The phenomenon of expression of both the alleles in heterozygote is called codominance. As the result the phenotype is different from both homozygous genotype.

Examples

- 1. Blood group is the good example codominance

any kind of sugar.

- 3. I^A, I^B are dominant alleles where as i is recessive alleles
- 4. Since, there are three different allele, there are six different combination of these three alleles are possible and four phenotypes (A, B, AB and O)

Genetic Basis of Blood Groups in Human Population

Allele	Allele	Genotype	Blood
from	from	of	Types of
Parent	Parent	Offspring	Offspring
1	2		
IA	IA	I ^A I ^A	А
IA	IB	IAIB	AB
IA	i	I ^A i	А
IB	IA	IAIB	AB
IB	IB	I ^B I ^B	В
IB	i	I ^B i	В
i	i	ii	0

When I^A and I^B are present together they both express their own types of sugars this is because of co-dominance. ABO blood grouping also provides a good example of multiple alleles.

Here, you can see that there are more than two, *i.e.,* three alleles governing the same character. Since, in an individual only two alleles can be present multiple alleles can be found only when population studies care made. Dominance is not an autonomous features of a gene. It depends on much on the gene product

120 **(d)**

11th.

Thalassaemia

(i) It is an autosome-linked recessive disesase(ii) It occurs due to either mutation or deletion resulting in reduced rate of synthesis of one of globin chains of haemoglobin

(iii) Anaemia is the characteristic of this disease(iv) Thalassaemia is classified into two types

- 5. α -thalassaemia Production of α -globin chain is affected. It is controlled by the closely linked genes HBA1 and HBA2 on chromosome 16. It occurs due to mutation or deletion of one or more of the four genes.
- β-thalassaemia Production of β-globin chain is affected. It occurs due to mutation of one or both HBB genes on chromosome 11

122 **(a)**

TH Morgan.

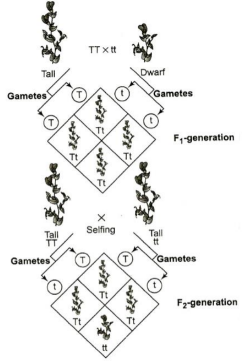
Father of experimental genetics is TH Morgan. He is also called the fly man of genetics because of selecting fruit fly (Drosophila melanogaster) as research material in experimental genetics

123 **(b)**

Mendel gave the laws of inheritance, which

124 (b)

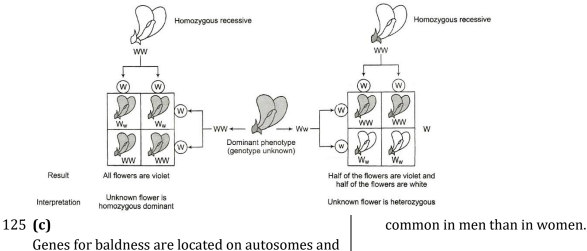
The genotypic ratio can be find out of a dominant phenotype by test cross or by simply Punnett square.



Phenotypic ratio Tall : Dwarf Genotypic ratio TT : Tt : tt 1:2:1

A Punnett square used to understand to typical monhybrid cross conducted by Mendal between truebreeding tall plants and true-breeding dwarf plants

Crossing of F₁ (dominant phenotype) to any one of the parent called back cross and when phenotype of crossing parent is recessive than this is called test cross. The progeny of such cross can easily be analysed to predict the genotype of test organism



influenced by androgens, thus, this is more

provides the mechanism that explains the pattern of inheritance



The number of characters studied by Mendel was

seven which were present on four chromosomes. Three contrasting characters with respect to pea pod are (i) pod shape,(ii) pod colour, (iii)pod position.

127 (c)

Chlorinated hydrocarbons are mutagen pollutants which can cause mutation in gene.

128 **(b)**

Both genes and chromosomes (Mendelian factors) whether dominant or recessive are transmitted from generation to generation in the pure or unaltered form. It is also called law of purity of gametes

129 **(d)**

The study of inheritance of genetic traits in several generations of a human family in the form of a family tree diagram is called **pedigree analysis**.

Advantages

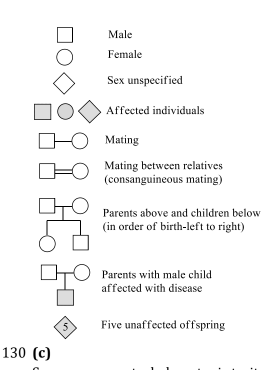
(i) It helps in genetic counselling to avoid disorders

(ii) It shows the origin of a trait and flow of a trait in a family

(iii) It is important to know the possibility of a recessive allele that can cause genetic disorders like colour blindness, haemophilia, etc. *Signosed in the pedigree are*

131 (a)

Absence of one X-chromosome (44 with XO).



Some genes control phenotypic traits and at the same time they also influence the viability of the individuals. The influence of these genes on viability is such that it may cause death of individual carrying them. Such genes are called **lethal genes**.

Disorders	Autosomal/Sex Linked	Symptoms	Effects
Down's syndrome	Autosomal aneuploidy (trisomy, +21)	Mongolian eyefold (epicanthus), open mouth, protruded tongue, projected lower lip, many loops on finger tip, palm crease	Retarded mental development IQ (below 40)
Turner's syndrome	Sex chromosomal monosomy 44 + XO	Short stature females (<5'), webbed neck, body hair absent menstrual cycle absent. Sparse pubic hair, underdeveloped breasts narrow lips puffy fingers	Sterile hearing problem
Klinefelter's syndrome	Sex chromosomal aneuploidy (tri/tetrasomy	The males are tall with long legs, testes small, sparse body hair, Barr	Gynaecomast ia azospermia sterile

of X chromosome)	body present, breast enlargement	
44+XXY	bi east emaigement	
44+XXXY		

Some Examples of Aneuploidy

(i) Down's syndrome-21 trisomy

Symptoms

- (a) Short statured with small round head
- (b) Partially open mouth with protruding furrowed tongue
- (c) Palm is broad with characteristic palm crease
- (d) Slow mental development

(ii) Turner's syndrome

Cause Absence of one of the X-chromosomes, resulting in the karyotype 44+X0

Symptoms

- (a) Sterile female with rudimentary ovaries
- (b) Shield-shaped thorax
- (c) Webbed neck
- (d) Poor development of breasts
- (e) Short stature, small uterus, puffy fingers

(iii) Klinefelter's syndrome

Cause Presence of an additional copy of X-chromosome resulting in the karyotype 44+XXY

Symptoms

- (a) Sex of the individual is masculine but possess feminine characters
- (b) Gynaecomastia, *i.e.*, development of breasts
- (c) Poor beard growth and often sterile
- (d) Feminine pitched voice

132 **(c)**

The recessive genes located on X-chromosome in humans are always expressed in males because a female may be homozygous or heterozygous, while male is always hemizygous (*i.e.*, only one allele is present).

133 **(b)**

Strength of linkage between two genes inversely proposed to the distance of two gene means if two genes are closely placed then they have high percentage of linkage and if they placed for then there is low percentage of linkage.

Linkage and recombination are two opposite phenomena. In linkage two genes remains united and in recombination the two get apart due to crossing over during gametogenesis (meiosis)

134 **(d)**

Fruitfly is excellent model for genetics because (i) Life cycle is very short (14 days)

(ii) Can be feed on simple synthesis medium

(iii) Single mating produces large number of progeny(iv) Clear differentiation of sexes

(v) Variation can be seen simply by hand lens or simple microscope

(vi) They (fruitfly) are easy to handle

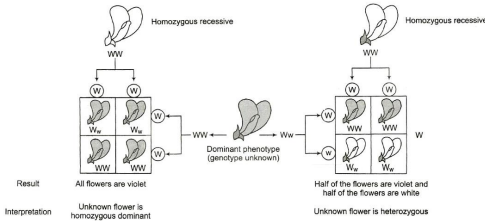
136 **(b)**

If both parents have blood group-AB then the possible blood groups of children are A, B and AB.

137 **(c)**

Test cross is a cross in which the dominant F_1 -plant crosses with the homozygous recessive parents plant. Crossing of F_1 (dominant phenotype) to any one of the parent called back cross and when phenotype of crossing parent is recessive than this is called test cross. The progeny of such cross can easily be analysed

to predict the genotype of test organism

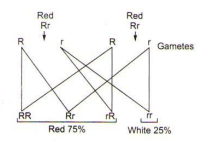


138 (d)

The ability of a gene to have multiple effects is known as **pleiotropy** and this phenomenon pleiotropism. The basis of pleiotropy is the interrelationship between the metabolic pathways that may contribute towards different phenotypes.

139 **(c)**

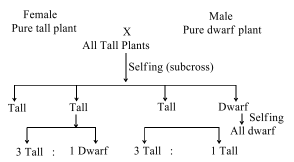
When two pea plants having red (dominant) coloured flowers with unknown genotype are crossed, the 75% red and 25% white flowered plants will be produced by following the law of Mendel. This is possible only if the parents are heterozygous.



141 **(a)**

 $\frac{1}{4}$ th and $\frac{3}{4}$ th.

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142 **(b)**

Night blindness is nutritional deficiency disease generally happens due to deficiency of vitamin-A

143 **(a)**

The genotypic ratio of monohybrid cross is 1 : 2 : 1, whereas the phenotypic ratio is 3 : 1.

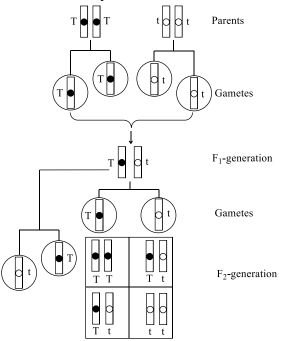
9:3:3:1 is the phenotypic ratio of dihybrid

cross (the cross made to study the inheritance of two pairs of factors or alleles of two genes).

144 **(c)**

Chromosomal Theory of Inheritance

Walter Sutton and Theodore Boveri noted that the behavior of chromosomes was parallel to the behaviour of genes and used chromosome movement to explain Mendel's laws.



Law of segregation interpreted on the basis of genes or factors (solid and hollow) situated on two homologous chromosomes.

Comparison between the Behaviour of

Chromosomes and Genes

Chromosomes	Genes
Segregate at the	Segregate of
time of gamete	gamete
formation such	formation and
that only one of	only one of
each pair is	each pair is
transmitted to a	transmitted to
gamete	a gamete
Independent	One pair
pairs segregate	segregates
independently	independently
of each other	of another pair

Sutton and Boveri argued that the pairing and separation of a pair of chromosomes would lead to the segregation of a pair of factors they carried. Sutton united the knowledge of chromosomal segregation with Mendelian principles and called it the chromosomal theory of inheritance. Experimental verification of chromosomal theory of inheritance was given by Thomas Hunt Morgan. Morgan worked with tiny fruitfly (*Drosphila*

melanogaster)

145 **(b)**

The females have **homozygous** XX sex chromosomes, while males have **heterozygous** XY-chromosome. Y-chromosome is shorter than X-chromosome.

146 **(b)**

The **Down's syndrome** (Mongolian idiocy) arises due to **trisomy** of **21st chromosome**, *i.e.*, total 47 chromosomes will present in such person. The main features are mental deficiency, short stature, round face, flaccid muscles, protruding tongue, etc.

147 **(a)**

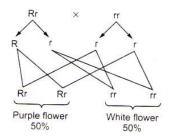
ZW-ZZ system of sex determination occurs in certain insects (gypsy moth) and vertebrates such as fishes, reptiles and birds and plants such as *Fragaris elatior*.

148 **(a)**

The Sudden and heritable change in the genetic make up of an individual is called **mutation**. The term mutation was introduced by Hugo de Vries.

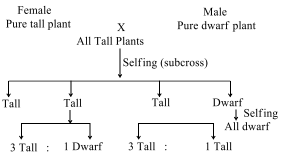
149 **(c)**

According to law of purity of gametes, when the gametes are formed, they carry only one allele of the gene considered.



150 **(d)**

The recessive trait shown by F_2 and F_3 -generation both but firstly it was observed in F_2 -generation. Mendel cross-pollinated a pure tall pea plant (100-120 cm hight) and a pure dwarf pea plant. (only 22 to 44 cm hight). He called them parental generation, expressed now-a-days by symbol P. This hybridization popularly called as monohybrid cross



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F₂-generation F_1 -plants pollinated among them self (self breeding or inbreeding) and seed produced by F_1 -plants called F_2 -generation. They were in ratio 3:1 (three tall and one dwarf).

 $F_3\mathchar`-generation$ Mendel allowed $F_2\mathchar`-plant$ to form seed by self-pollination called $F_3\mathchar`-generation.$ Mendel observed that tall and dwarf plant behave differently

(i) Dwarf plant produced dwarf plant on self-pollinated

(ii) In tall plants one third plants breed true so they were pure

(iii) Other two third plant behave like parents and give tall to dwarf plants 3 : 1 indicate that their parents have dwarf genes also

151 **(a)**

A-A; B-Genetic disorders

152 **(a)**

Heredity (L. *Hereditas* – Heirship or inheritance) is the transmission of genetically based characters from parents to their offspring.

The process by which characters are transferred from one generation to the next generation is called inheritance

153 **(c)**

Huntington's chorea is a fatal disease of man. It is characterized by uncontrolled jerking of body and progressive degeneration of central nervous system. The mean age for the onset of these symptoms is between 35 to 40. This disease is caused by an autosomal dominant gene.

154 **(d)**

Movement of chromosomes towards poles

requires centromere.

155 **(a)**

Klinefelter's syndrome is represented by 44 autosomes + XXY.

157 **(a)**

Man has only one X-chromosomes that is inherted to has daughter. Therefore, a heredftary disease, which is X-chromosomal linked, is never passed on from father to son.

159 **(a)**

Chromosomal theory of linkage states that (i) Linked gene present on same chromosome (ii) They lie in linear sequence in chromosome (iii) There is tendency to maintain the parental combination

(iv) Strength of linkage between two gene is inversely proposal to the distance of two gene and *vice-versa*

160 **(c)**

In the diploid organism (plants and animals) the chromosome or DNA number becomes double just before the cell division

161 **(c)**

Genetics is the branch of biology which deals with the inheritance and variations

162 **(c)**

Balbiani first observed these chromosomes in the salivary glands of midge *Chironomus* in 1881. These polytene chromosomes are considered as somatic giant tubules and also reported from fat bodies some other dipterans also, *e.g., Drosophila, Chironomus, Sciaca, Rhyncosciara,* etc.

163 **(d)**

Sir Archibald Edward Garrod was an English physician, who pioneered the field of inborn errors of metabolism. He was born on November 25, 1857, in London and died on March 28, 1936, in Cambridge.

164 **(c)**

Punnett square is a table, in which all possible combinations of gametes and progeny are displayed in a grid structure.

165 **(c)**

Erythroblastosis foetalis is a haemolytic disease of newborn children. Erythroblastosis foetalis can occur when father is Rh positive and mother is Rh negative. An Rh negative woman can be sensitized when she bears an Rh⁺ child and Rh⁺ children may have erythroblastosis.

166 **(c)**

When F_1 hybrid is crossed with its recessive parents, it is called as **test cross**. By test cross, the herterozygosity and homozygosity of the organism can be tested. The test cross ratio in monohybrid cross is 1 : 1 and in dihybrid cross, ratio wii be 1 : 1 : 1 : 1.

167 **(c)**

Inversion involves a reverse order of genes in a part of chromosome.

168 **(b)**

The gamete mother cells (2n) are called **meiocytes**, which undergo meiosis to form gametes (n). The chromosome number in the melocytes(2n) of housefly is 12.

169 **(c)**

Genes which codes for a pair a contrasting traits is called alleles. They are slightly different forms of the same gene, *e*. *g*., TT, tt, tT

170 **(c)**

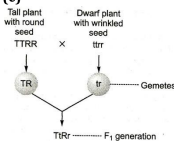
Haemophilia and colour blindness are both Xlinked recessive diseases. The gene for both is found on X-chromosome only.

Albinism, Sickle-cell anaemia and thalassemia are autosomal diseases.

171 **(c)**

Mendel described the inheritance of recessive and dominant genes. Phenylketonuria (PKU) is an autosomal recessive mutation of gene on chromosome-12.

172 **(c)**



Thus, there is no dwarf plant with wrinkled seeds in $\mathrm{F}_1\text{-}\mathrm{generation}.$

coined by **Bateson** in 1906. Genetics is the study of principles and mechanism of heredity and variations.

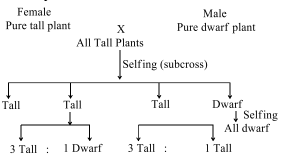
174 **(b)**

In human beings, 46 chromosomes are found, in which only one pair XY takes part in sex determination. These are known as **sex chromosomes** or **allosomes**, rest 22 pairs are known as **autosomes**.

175 **(a)**

Mendel obtained the recessive character in F_2 by self pollinating the F_1 -plants.

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173 **(b)** The term genetics (*Gk. Genesis*=descent) was

176 **(a)**

Criss-cross Inheritance It is a type of sex-linked inheritance, where a parent passes the traits to the grand child of the same sex through offspring of the opposite sex, that is, father passes the traits to grandson through his daughter (diagynic), while the mother transfers traits to her grand daughter through her son (dia-andric). It was first studied by Morgan (1910) in case of eye colour in *Drosophila*. Criss-cross inheritance is applicable to most sex-linked disorders in humans, *e. g.*, red green colour blindness, haemophilia

177 **(b)**

Heterozygous.

The diploid condition in which the alleles at a given locus are identical is called homozygous or pulls. In homozygous condition, organism have two similar genes or alleles for a particular character in homologous pair of chromosomes, *e. g.*, TT or tt.

Organisms containing two different alleles or individual containing both dominant and recessive genes of an allele pair, *e.g.*, Tt is known as heterozygous or hybrid

178 (c)

Mendel conducted artificial pollination/cross pollination using true breeding pea lines A true breeding line is one that having undergone continuous self pollination, shows stable trait inheritance and expression for several generation

179 **(d)**

Morgan and his group found that when genes were grouped on the same chromosome, some genes were very tightly linked (showed very low recombination), while others were loosely linked (showed higher recombination).

180 **(b)**

Polygene results in quantitative inheritance, which is characterized by occurrence of intermediate forms between the parental type. In case of crossing between AABBCC (dark colour) and aabbcc (light colour), in F_2 -generation seven phenotypes will obtain with ratio 1 : 6 : 15 : 20 : 15 : 6 : 1. The total number of progeny is 64, out of which only two will be likely resemble with either parents. Hence, their proportion in F_2 -generation would be 3.12, *i.e.*, less than 5%

181 **(a)**

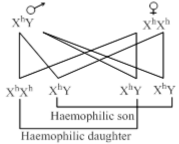
In cancer cells there is uncontrolled cell division. In them chromosomal abbreviation is commonly found

182 **(b)**

The given case is the example of **codominance**.

183 **(d)**

Males and female are haemophilic definately. If their father and mother both are haemophilic



184 **(b)**

Polyploidy is the phenomenon, which leads to increase in the number of chromosomes thus, increasing in the number of genes. Due to cumulative effect of genes, new characters appear, which results into formation of new species.

185 **(b)**

Mendel is called father of genetics. There are three laws of Mendel in respect of inheritance:

- 7. Law of dominance
- 8. Law of segregation or Law of purity of gametes or Law of splitting of hybrids.
- 9. Law of independent assortment

186 **(b)**

Test cross is a cross between F_1 hybrid with its recessive parent.

187 **(a)**

Homogametic.

XY and XY type sex determination seen in many insect and mammals including humans. Males have X and Y chromosome along with autosome and females have pair of 'X' chromosome along with autosome

Parents Phenotypes Male Female

Genotypes 44A + XY 44A + XX **Gametes** 22A + X 22A + Y 22A + A22A + X22A + X 22A + X Children 22A + X 44A + XX 44A + XYFemale 22A + Y 44A + XY 44A + XY

Male

Sex ratio Female : Male = 1 : 1

In plants The flowering plants are mostly bisexual and lack sex chromosomes. The unisexual flowering plants tent to have XX-XY type of sex chromosomal mechanism for sex determination. The female plants are XX and male plants are XY.

XX and XO Type of Sex Determination

Found in insect like grasshopper, cockroaches and bugs. Males have only X sex-chromosome and autosomes, female have pair of X-chromosome and autosome

Parents	Phenotypes	Male	Female
	Genotypes	AA + XO	AA + XX
Gametes	Α	+ X A + 0	$A + X \cdot A + Y$

F_1 -generation

 $\begin{array}{c|c} A+X & A+X \\ \hline A+X & AA+XO & AA+XO \\ A+O & AA+XO & AA+XO \\ \hline & Genotypes \end{array}$

XX-XO type of sex determination

In most of cases the female produce similar sex chromosome called homomorphic. In most of cases the male produce dissimilar sex chromosome called hetermorphic

188 (a)

In birds, usually female is designated as ZW, being heterogametic and male is designated as ZZ being homogametic.

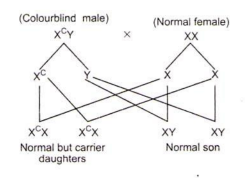
189 (a)

A cross of round yellow seeds (both dominant) and green wrinkled seed (both recessive) plants produced 9: 3: 3: 1 ratio of plants (phenotypic)in F_2 generation. The ratio of parental to recombinant is 10: 6 here because the 9 and 1 are of parental type and 3 & 3 are recombinant.

190 **(a)**

In genetics, a test cross, first introduced by **Gregor Johann Mendel**, is used to determine weather an individual exhibiting a dominant trait is homozygous or heterozygous for that traits. More simply, test cross determines the genotype of an individual with a dominant phenotype. The test cross is defind as being a type of back cross between the recessive homozygote parents and F_1 generation.

191 **(c)**



So, all sons in the progeny will be normal.

192 **(d)**

When a tall pea plant (TT) is crossed with dwarf plant (tt), the F_1 progeny shows all plants hybrid tall and on selfing of F_1 progeny, the F_2 generation shows both tall and dwarf plant in the ratio 3 : 1. Out of three tall plants, one is pure tall (TT) and two are hybrid tall (Tt).

193 **(c)**

A-Two, B-Chromosomal, C-Mutation

194 **(c)**

A-Heterozygous, B-Unaffected, C-Carrier

196 **(c)**

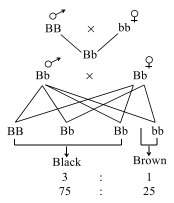
Linkage group will be equal to haploid number of chromosomes. *Pisum sativum* has seven pairs of chromosomes, therefore linkage group is also **seven**.

197 **(d)**

When a cross (dihybrid) is made between plants bearing round yellow (RRYY) and wrinkled green (rryy) seeds, all the plants in F_1 -generation are with yellow round seeds (showing the genotype RrYy).

198 **(a)**

Black colour is dominant over the recessive so by cross it is easily infered that 75% of the offspring are black and 25% are brown



199 **(b)**

There are only very few characters, which are present on the Y-chromosome of male. Like hypertrichosis. Given pedigree analysis is the example of Y-linked inheritance because all male progeny is affected

200 (a)

Haemophilia.

Genetic or chromosomal symbol used for person who is having sickle-cell anaemia Ps – Hb^s Hb^s. Sickle-cell Anaemia

Sickle-cell Anaemia

(i) It is an autosome-linked recessive trait

(ii) The disease is controlled by a single pair of allele Hb^{s} and Hb^{s}

(iii) Only the homozygous individuals for Hb^s, *i.e.*, Hb^sHb^s show the diseased phenotype

(iv) The heterozygous individuals are carriers $({\rm Hb}^{\rm A}{\rm Hb}^{\rm S})$

(v) Due to point mutation, glutamic acid (Glu) is replaced by valine (Val) at the sixth position of β -globin chain of haemoglobin molecule

(vi) A single base substitution at sixth codon of the beta globulin gene from GAG to GUG. GAG code for glutamic acid and GUG code for valine.
(vii) Hb^S behaves as normal haemoglobin except under the oxygen stress where erythrocytes lose their circular shape and become sickle-shaped. As a result, the cells cannot pass through narrow capillaries. Blood capillaries are clogged and thus, affect blood supply to different organs

201 **(b)**

Grasshopper is an example of XO type of sex determination in which the males have only one X-chromosome besides the autosomes, whereas females have a pair of X-chromosomes .

202 **(a)**

Tr Rr (heterozygous tall and pink) ↓ (self crossed) Tt Rr × Tt Rr Gametes TR Tr tR tr

TR	TTRR	TTRr	TrRR	TrRr
	(Red)	(Pink)	(Pink)	(Pink)
Tr	TTR	TTrr	TrRr	Ttrr
	(Pink)		(Pink)	
tR	TrRR	TrRr	ttRR	ttrR
				(Pink)
tr	TrRr	Ttrr	ttRr	ttrr
			(Pink)	

 $1/16 \mathrm{TTRR}$

2/16 TTRr (2/16 TtRR / 4/16 TtRr) 1/16 TTrr 2/16 Ttrr } 2/16 Ttrr } 2/16 ttRR 2/16 ttRR } 3/16 - 25% 3/16 - 50% 1/16 ttrr 1/16 - 50%

203 **(d)**

Chimera is an individual which has in its body cells of two or more genorypes *i.e.*, pletiotroic mutations. Chimeric individuals produced by transfections arise when some cells of an embryo become stably transfected.

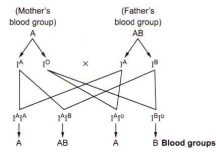
204 **(c)**

Mendel selected 14 pairs of true bruding pea plant varieties for his experiment

205 (a)

Syndrome stands for the group of symptoms, which indicates to a particular disease

206 (d)



Hence, parents with blood group-A and AB will not produce offsprings with blood group-O.

207 (d)

Nephrogenic diabetes is due to genetic deficiency of ADH-receptor linked to X-chromosome.

208 **(b)**

Mendel got only parental phenotype in the offspring. He didn't get only intermediate result. So, he could not formulated the blending theory of inheritance or observed linkage and crossing over

209 (a)

Mutations are large discontinuous sudden

heritable change in the genotype. Mutation are generally **recessive** in nature.

210 (a)

Cri-du-chat syndrome (Cat –cry syndrome) was discovered by **Lejeune** in 1963 and is due to the deletion of a large part of the small arm of the 5th autosomes. Deletion is a type of mutation in which a segment is removed from chromosomes or DNA molecules.

211 **(c)**

A-Human; B-Quantitative

212 **(d)**

When a pair of contrasting characters are crossed with each together then F_1 -generation has only one type of character. This expressed character is known as **dominant** character, while the character, which could not express in F_1 generation is known as **recessive** character. In pea plants, tallness, round seed, yellow seed, purple flower, green pod, inflated pod and axial flower are dominant over dwarfness, wrinkled seed, green seed, white flower, yellow pod, constricted pod and terminal flower, respectively.

213 **(a)**

Genes for cytoplasmic male sterility in plants are located in mitochondrial genome.

214 **(d)**

DNA **transposition** is the process, which involves the movement of DNA elements from one site in the genome to the other. It is mediated by transposase enzymes. These short segments of DNA (DNA elements) with remarkable capacity to move from one location in a chromosome to another, are called **transposons** or **jumping genes** or transposable elements or mobile genetic elements. These were first discovered by **Barbara McClintock** in maize (*Zea mays*) for which she got the Nobel Prize for physiology and medicine.

215 **(a)**

The phenomenon of expression of both the alleles in heterozygote is called codominance. As the result the phenotype is different from both homozygous genotype.

Examples

- 10. Blood group is the good example codominance
- 11. ABO blood groups are controlled by gene

I. The gene (l) has three allele I^A, I^B, i, I^A, I^B produce slightly different form of sugar while i does not produce any kind of sugar.

- 12. I^A, I^B are dominant alleles where as i is recessive alleles
- Since, there are three different allele, there are six different combination of these three alleles are possible and four phenotypes (A, B, AB and O)

Genetic Basis of Blood Groups in Human Population

Allele	Allele	Genotype	Blood
from	from	of	Types of
Parent	Parent	Offspring	Offspring
1	2		
IA	IA	I ^A I ^A	А
IA	IB	IAIB	AB
IA	i	I ^A i	А
IB	IA	IAIB	AB
IB	IB	I ^B I ^B	В
IB	i	I ^B i	В
i	i	ii	0

When I^A and I^B are present together they both express their own types of sugars this is because of co-dominance. ABO blood grouping also provides a good example of multiple alleles.

Here, you can see that there are more than two, *i.e.*, three alleles governing the same character. Since, in an individual only two alleles can be present multiple alleles can be found only when population studies care made. Dominance is not an autonomous features of a gene. It depends on much on the gene product

216 **(c)**

If pedigree initiated from male it is called proposity. It pedigree initiated from female it is called propista. So, individual from which a pedigree initiated could be proband or propositus

217 **(a)**

Hugo de Vries used *Oenothera lamarckana* for his mutation experiment.

218 **(b)**

Phenylketonuria is due to deficiency of liver enzyme phenylalanine hydroxylase.

219 **(a)**

Haemophilia is a sex-linked character (X-linked recessive trait). It is a rare human blood disorder, in which, blood clotting is deficient, resulting in

severe bleeding internally and externally. The condition is due to lack of fibrin in the blood and is controlled by two closely linked genes on the blood and is controlled by two closely linked genes on the X-chromosome that are responsible for the production of different clotting factors.

220 **(b)**

A cross between two individuals for studing inheritance of two characters is known as dihybrid cross.the phenotypic ratio in F_2 generation of a dihybrid cross is 9 :3 : 3 : 1, therefore, the maximum number of different phenotypes available are four.

221 **(c)**

Nullisomy is an aneuploid condition, in which a pair of homologous chromosomes is deficient and represented as 2*n*-2. **Monosomy** is an aneuploid condition, in which a chromosomes is deficient from its homologue and represented at 2*n*-1.

222 (d)

For the given case, the disease is sex-linked recessive.

223 (a)

Colour blindness and haemophilia are diseases caused by X-linked recessive genes. Colour blindness involving faulty perception of red and green light and follows an X-linked pattern of inheritance.

224 (a)

Principle or law of independent assortment has been called the second law of Mendelism by Correns

225 (a)

Mendel's hybridization strategy was use of two plants differing in a pair of contrasting character in case of monohybrid cross and two pair of contrasting characters in dihybrid cross

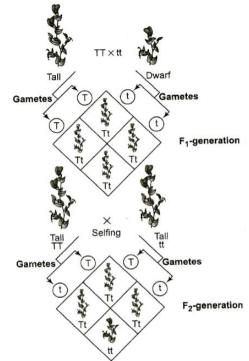
226 **(b)**

A-heterozygous, B-TT or tt, C-Heterozygous

- 227 **(b)**
- 229 **(a)**
 - 4.

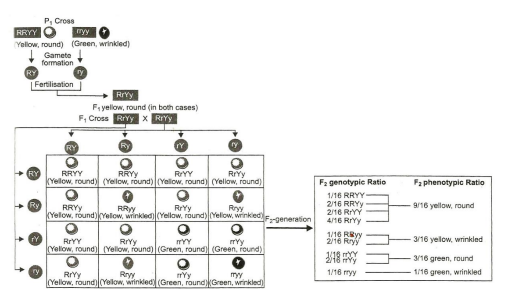
Drumsticks of Barr body are the sex chromatin present in the neutrophils (polymorphonuclear leucocyte) of 3 to 5 % cells in females and are absent in males.

228 (a)



Phenotypic ratio Tall : Dwarf Genotypic ratio TT : Tt : tt 1 : 2 : 1

A Punnett square used to understand to typical monhybrid cross conducted by Mendal between true-breeding tall plants and true-breeding dwarf plants



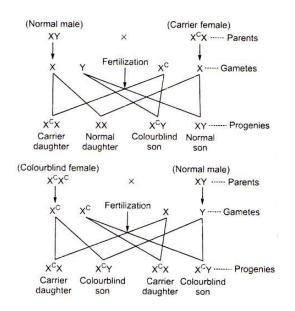
A dihybrid cross in pea plant between yellow round (smooth) seeded and green wrinkled seeded plant. The cross proves the principle of independent assortment

230 **(c)**

In the given pedigree chart, squares are representing males and circles representing females. In F_1 generation, 2-male are diseased and in next generation only male is diseased. This shows the inheriatnce of a Y-linked disease.

231 (d)

Colour blindness is a disease, in which a person is unable to differentiate between red and green colour. The gene for this disease is located on the X-chromosome. So, if a colourblind woman marries a normal man, it will produce all the sons colourblind (X^cY). In case of a carrier woman, the probability of a colourblind and normal son is 50 : 50



232 (c)

Sex Chromatin in Interphase Nuclei Barr and Bertram (1949) found that interphase nuclei of

human females stained with orcein possess small distinct chromatin body called **sex chromatin**, **Barr body** or **X-chromatin**

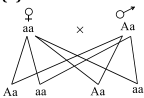
Barr body is found attached to nuclear envelops in oral mucosa, any where in the nucleus in nerve cells and as **drumstick** or small rod at one side of nucleus in neutrophil or polymorphonuclear leucocytes (Davidson and Smith) Barr body is produced due to partial inactivation of one X-chromosome and development of facultative heterochromatin in it. Any of the two X-chromosomes can become heterochromatic. It begins in the late blastocyst stage (roughly 16 day of embryonic life). Partial inactivation of one Xchromosomes in females is called **Dosage compensation**

233 **(d)**

As given in the chart the AB blood group percentage is 5% in India.

So, it can be said that any individual, selected at random from sample population has 1 in 20 chance of being blood group AB

234 **(d)**



The pedigree given in question is the most probable autosomal disease

235 **(d)**

The study of inheritance of genetic traits in several generations of a human family in the form of a family tree diagram is called **pedigree**

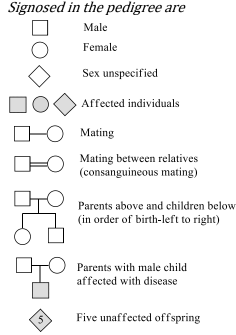
analysis.

Advantages

(i) It helps in genetic counselling to avoid disorders

(ii) It shows the origin of a trait and flow of a trait in a family

(iii) It is important to know the possibility of a recessive allele that can cause genetic disorders like colour blindness, haemophilia, etc.



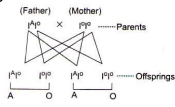
236 **(a)**

The non-ionizing radiations of visible light (UV rays) have been found to produce mutagens in culture media, which induced mutations in irradiated organism placed in these media for few hours. These rays cause hydration of cytosine and thymine. In humans, it causes DNA damage producing thymine dimers, however, generally it does not create any serious problem due to DNA repair system.

237 **(b)**

Nucleotide sequence is also called the base-pair sequence. In frame-shift mutation or base pair substitution the nucleotide sequence get changed

238 **(c)**



So, if a woman with 'O' blood group has a child with blood group-O claims a man with blood group-A as father, then the genotype of claimed person should be I^AI^O.

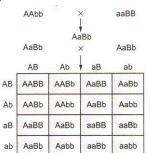
240 **(a)**

Mendel carried out hybridization experiments on garden pea for 7 years from 1856-1863

241 **(a)**

Cross between two genetically unlike individuals is called **hybridization**.

242 (a)



The phenotypic ratio will be 9:3:3:1.

Dihybrid test cross gives 1 : 1 : 1 : 1 ratio.

Law of segregation gives $1:2:1\ genotypic$ ratio in F_2 -generation.

243 (a)

Some Examples of Aneuploidy

(i) Down's syndrome-21 trisomy

Symptoms

(a) Short statured with small round head

(b) Partially open mouth with protruding

furrowed tongue

(c) Palm is broad with characteristic palm crease

- (d) Slow mental development
- (ii) Turner's syndrome

Cause Absence of one of the X-chromosomes,

resulting in the karyotype 44+X0

Symptoms

(a) Sterile female with rudimentary ovaries

- (b) Shield-shaped thorax
- (c) Webbed neck
- (d) Poor development of breasts
- (e) Short stature, small uterus, puffy fingers
- (iii) Klinefelter's syndrome

Cause Presence of an additional copy of Xchromosome resulting in the karyotype 44+XXY

Symptoms

(a) Sex of the individual is masculine but possess feminine characters

(b) Gynaecomastia, *i.e.*, development of breasts

(c) Poor beard growth and often sterile

(d) Feminine pitched voice

244 **(b)**

A recessive allele is not weaker than the dominant allele. It (recessive allele) does not shows its effect (in the presence of dominant allele) because of modified or different enzymes. A recessive allele make its gene product even when paired with the dominant allele. It is not necessary that dominant allele always better (in the case of dominant disease)

245 (a)

Given pedigree analysis indicates the transmission of autosomal recessive trait from parents to their offsprings

246 (d)

Phenotype is the observable characteristics or the total appearance of an organism. It is determined by its genes, the dominance relationships between the alleles and by the interaction during development between its genetic constitution (genotype) and the environment.

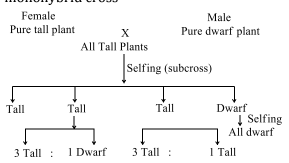
247 **(a)**

Cretinism is caused by deficiency of thyroid hormones in infants.

248 **(a)**

Tall.

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This three generations of pea plants after crossing a pure tall plant with a dwarf one. The plants of F_1 -generation are all tall, of F_2 -generation three tall and one dwarf. One third of the tall plants are 251 **(b)**

12.

pure, while the remaining behave as hybrids F_1 -generation Seeds collected from the parental generation called first filial generation or F_1 -generation

F₂-generation F_1 -plants pollinated among them self (self breeding or inbreeding) and seed produced by F_1 -plants called F_2 -generation. They were in ratio 3:1 (three tall and one dwarf). **F₃-generation** Mendel allowed F_2 -plant to form

seed by self-pollination called F_3 -generation. Mendel observed that tall and dwarf plant behave differently

(i) Dwarf plant produced dwarf plant on self-pollinated

(ii) In tall plants one third plants breed true so they were pure

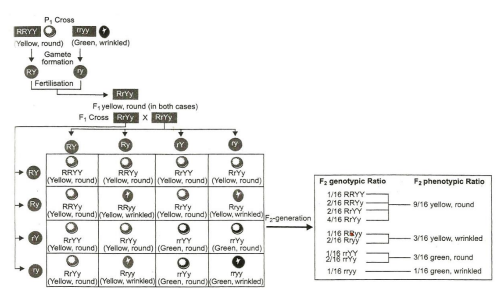
(iii) Other two third plant behave like parents and give tall to dwarf plants 3 : 1 indicate that their parents have dwarf genes also

249 **(d)**

The persons suffering from Klinefelter's syndrome have normal external genitelia but internally testes are absent or reduced. Sparse body hair and gynecomastia (female like breast development) is also seen.

250 **(d)**

Dominant lethal gene kills the organism.



A dihybrid cross in pea plant between yellow round (smooth) seeded and green wrinkled seeded plant. The cross proves the principle of independent assortment

252 (a)

A-Linkage; B-Crossing over

253 (b)

The actual distance between two genes is said to be equivalent to the percentage of crossing over between these two genes. Since the two genes lie at the ends of the chromosome, there are 100% chances of their segregation during crossing over.

254 (a)

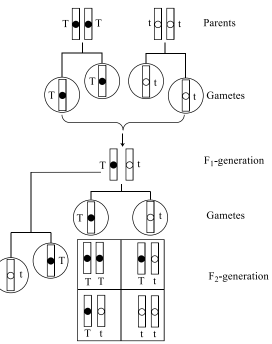
In **trisomic** (2n + 1) condition, organism has one extra chromosome, e.g., Klinefelter's syndrome (22 pair + XXY).

255 **(b)**

TH Morgan.

Chromosomal Theory of Inheritance

Walter Sutton and Theodore Boveri noted that the behavior of chromosomes was parallel to the behaviour of genes and used chromosome movement to explain Mendel's laws.



Law of segregation interpreted on the basis of genes or factors (solid and hollow) situated on two homologous chromosomes.

Comparison between the Behaviour of
Chromosomes and Genes

Chromosomos	Comos
Chromosomes	Genes
Segregate at the	Segregate of
time of gamete	gamete
formation such	formation and
that only one of	only one of
each pair is	each pair is
transmitted to a	transmitted to
gamete	a gamete
Independent	One pair
pairs segregate	segregates
independently	independently
of each other	of another pair

Sutton and Boveri argued that the pairing and separation of a pair of chromosomes would lead

to the segregation of a pair of factors they carried. Sutton united the knowledge of chromosomal segregation with Mendelian principles and called it the chromosomal theory of inheritance. Experimental verification of chromosomal theory of inheritance was given by Thomas Hunt Morgan. Morgan worked with tiny fruitfly (*Drosphila melanogaster*)

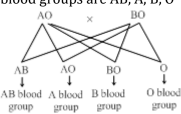
256 **(b)**

Rr and red because the R is dominant r so, the F_1 -hybrid will be red

257 **(b)**

There are two possibility for blood group A = AA and AO.

There are two possibility for blood group B = BB or BO cross between AO, BO. Give four types of blood groups are AB, A, B, O



258 **(a)**

Principle or law of inheritance were enunciated by Mendel. There are four principals or laws of inheritance based on monohybrid and polyhybrid cross.

(i) One gene inheritance

- 1. Principle of paired factors
- 2. Law of dominance
- 3. Principle of law of segregation
- (ii) Inheritance of two genes
- 4. Principles or law of independent assortment

259 (a)

Mendel's laws explained by the meiosis (gametogenesis). Like law of independent assortment, law of segregation, etc.

260 (d)

Mendel's discoveries concerning genetic inheritance were generally did not accepted by scientific community at that time. Mendel did not discovered linkage and blending inheritance

261 **(b)**

Deletion is a chromosomal aberration in which

there is loss of a chromosomal segment.

262 **(a)**

There are three symbols for the carrier (heterozygous condition). $\odot \otimes$ and \bigcirc Generally, the carriers are females so there is rounded structure

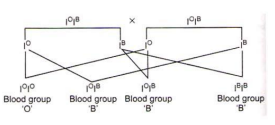
263 **(d)**

Turner's syndrome is a disorder caused due to the absence of one of the X-chromosome, *i.e.*, 45 with XO (44 autosomes + XO). Such females are sterile as ovaries are rudimentary besides other features including lack of other secondary sexual characters.

264 **(c)**

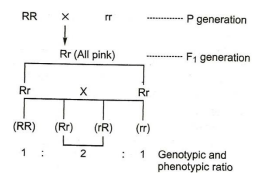
Genotype of a person with blood group-B may be I^BI^B or I^OI^B, person with genotype I^BI^B can not produce offsprings with blood group-O in any case but if the person's genotype is I^OI^B, then its offsprings may have blood group-O.

eg,



265 **(b)**

Incomplete dominance or **blending inheritance** was first seen in *Mirabilis jalapa*(4'0 clock plant). Here, when red flowers are crossed with white flowers variety the F_1 -hybrid is pink and F_2 ratio is 1 red : 2 pink : 1 white.



266 **(d)**

In **incomplete** (partial or intermediate) **dominance**, the effect of dominant allele is diluted or modified, so that the phenotypic expression of the concerned trait in a hybrid is distinguishable from both parental type. Consequently, both phenotypic and genotypic ratios in F_2 -generation are 1:2:1.

267 **(b)**

Cytoplasmic inheritance always shows maternal characters.

268 **(b)**

Type of gamete participating in selfing of members in monohybrid cross is of two types.

269 **(b)**

GAG code for glutamic acid in haemoglobin *m*RNA replaced by GUG code which code for valine in haemophillic haemoglobin *m*RNA

270 **(a)**

Linkage is the inheritance of certain genes as a group because they are parts of the same chromosome. Linked genes do not show independent assortmant. Linkage was first suspected and theorized in 1903 by Sutton and Boveri.

271 **(c)**

If a character is transmitted from father to his sons and then to grandson only, it means it is located on Y-chromosome (inheritance of Ylinked genes).

272 **(b)**

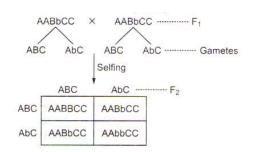
The term 'mutation' was coined by Hugo de Vries (1901). He also proposed mutation theory of evolution. The mutational theory of evolution published in 1903. Hugo de Vries worked on *Oenothera lamarckiana* (evening primorse). Out of a population of 54343 plants, de Vries observed 834 mutation and concluded that the primary force of evolution is mutation

273 **(b)**

'**Jumping genes'** or **movable genetic elements** were discovered by '**Barbara McClintock'** (1902-92) in maize. These 'controlling elements' could move from one location to another on the chromosome.

274 **(a)**

Since AABbCC contains only one heterozygous allelic pair, 'Bb', the cross would behave as monohybrid cross leading to phenotypic ratio in F_2 -generation.



Phenotypic ratio is 3 : 1

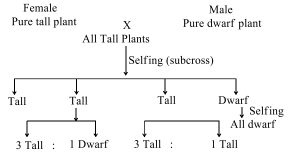
275 **(b)**

Dominant character.

Law or Principle of Dominance

In heterozygous individuals or hybrids a character is represented by two contrasting factors called alleles or allelomorphs. Out of the two contrasting alleles. Only one is able to express its effect in the individual. It is called dominant factor or dominant allele.

The other which does not show its effect in the heterozygous individual is called recessive factor or recessive allele. The cross between the pure tall and pure dwarf gives all progeny (F_1) tall.



The character shown by $F_{1}\xspace$ called dominant character

276 **(d)**

Sutton and **Boveri** proposed chromosomal theory of inheritance. This theory believes that chromosomes are vehicles of hereditary information possess mendelian factors segregate and assort independently during transmission from one generation to the next.

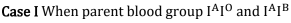
277 **(b)**

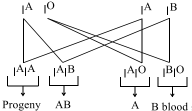
In sex-linkage, the speciality is criss-cross inheritance. Criss cross inheritance is a type of sex linked inheritance where a parent passes the traits to the grand child of the same sex through offspring of the opposite sex, that is father passes the traits to grandson through his daughter while the mother transfers traits to her grand daughter through her son, criss cross inheritance establish the relationship between gene and the sex chromosomes.

278 **(b)**

For the blood group A, there will be two type of genotype $-I^AI^A$, I^AI^D .

For the blood group AB there will be only one genotype $-I^AI^B$.





Case II When parent blood group are $I^A I^A$ and $I^A I^D$

279 **(b)**

Nullisomic=2n - 2

Monosomic= 2n - 1

Trisomic= 2n + 1

Haploid= n

280 **(b)**

Phenylketonuria (Folling; 1934). It is an inborn, autosomal, recessive metabolic disorder in which the homozygous recessive individual lacks the enzyme **phenylalanine hydroxylase** needed to change phenylalanine (amino acid) to tyrosine (amino acid) in liver. It results in hyperphenylalanine

281 (a)

Trisomy : An individual having one extra chromosome of a set (2n + 1).

282 **(a)**

Only two types of gametes are formed by genotype rrYy

	r	r	
у	ry	ry	
Y	rY	rY	

283 **(a)**

1906.

Genetics word is derived from the Greek word genesis, which stands for descent. Term genetics was introduced by Bateson in 1906 branch of Biology that deals with the study of heredity and variations Mutant gene that gives slightly modified phenotypes are called heteroalleles.

285 **(b)**

Colour blindness and haemophilia are well known examples of sex-linked diseases.

286 **(c)**

Gamma radiations are ionizing radiations and are physical mutagens. They are used as mutagens in such materials, where nucleus is deep seated, *e.g.*, seeds, stem, cutting, etc. Sharbati Sonora variety of wheat has been developed by gamma radiations on 'Sonora 64' variety (Mexican dwarf wheat variety).

287 (d)

Albinism is a genetic disorder.

288 (d)

Mendel's paper 'experiments on plant hybridisation' was published in the 'Proceeding of Brunn Natural Science Society' in 1805

289 **(a)**

If O gamete (no X or Y) fuses with X gamete, the resulting XO zygote will survive and form a sterile female. This situation is called **Turner's syndrome**.

290 **(a)**

Milk secreting and baldness both trait belongs to the sex limited trait.

Finalization of sex at the time of fertilization is known as sex determination. All sex linked character show criss-cross inheritance and firstly it was studied and discovered by TH Morgan (1910). *Sex related trait may be divided into three types*

(i) **Sex Linked Traits** They are those traits the determining genes of which are found on the sex chromosomes. All the sex-linked traits present on a sex chromosome are inherited together

(ii) **Sex Limited Traits** They are autosomal traits which are expressed in a particular sex in response to sex hormones although their genes also occur in the other sex, *e.g.*, milk secretion in mammalian females, pattern baldness in males. The gene for baldness behaves as an autosomal dominant in males and autosomal recessive in females

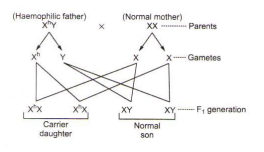
(iii) **Sex Influenced Traits** The traits are not due to particular genes but are by products of sex hormones, *e.g.*, low pitched voice, beard

284 **(a)**

moustaches. In males, pattern baldness is related to both autosomal genes as well as excessive secretion of testosterone

291 **(b)**

Haemophilia is an X-linked disease, it is transmitted from mother to son.



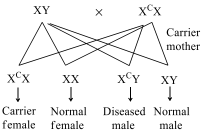
292 **(c)**

ABO blood group in human beings are as follow (i) codominant

- (ii) dominant-recessive
- (iii) multiple allele

294 **(a)**

Haemophilia is also called Bleeder's disease. It is a sex linked recessive disorder. It transmit from carrier female to male progeny



295 **(a)**

The important sources of variation are mutations and recombinations (crossing over).

296 **(b)**

Gregor Johann Mendel (1822-1884) is known as the father of genetics because he was the first to demonstrate the mechanism of transmission of characters from one generation to the other

297 **(c)**

Holandric gene occurs on the Y-chromosome only. It is inherited only by the male line and is a recessive gene that always expresses.

298 **(c)**

When a gene pair (allele) contains two different genes in an organism then the organism is

301 (b)

Disorders	Autosomal/Sex Linked	Symptoms	Effects
Down's	Autosomal	Mongolian eyefold	Retarded

considered as **heterozygous** for the particular character.

299 **(a)**

Finalization of sex at the time of fertilization is known as sex determination. All sex linked character show criss-cross inheritance and firstly it was studied and discovered by TH Morgan (1910). *Sex related trait may be divided into three types*

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300 **(b)**

The number of Barr bodies is one less than the number of X-chromosomes present in an individual, *e. g.*, 1 for normal XX, 2 for XXXY

syndrome	aneuploidy (trisomy, +21)	(epicanthus), open mouth, protruded tongue, projected lower lip, many loops on finger tip, palm crease	mental development IQ (below 40)
Turner's syndrome	Sex chromosomal monosomy 44 + XO	Short stature females (<5'), webbed neck, body hair absent menstrual cycle absent. Sparse pubic hair, underdeveloped breasts narrow lips puffy fingers	Sterile hearing problem
Klinefelter's syndrome	Sex chromosomal aneuploidy (tri/tetrasomy of X chromosome) 44+XXY 44+XXY	The males are tall with long legs, testes small, sparse body hair, Barr body present, breast enlargement	Gynaecomast ia azospermia sterile

302 (d)

Sex influenced traits are not due to particular genes but are by-products of sex hormones, *e.g.*, low pitched voice, beard, moustaches and baldness in human. The gene for balaness behaves as an autosomal dommant in males and autosomal recessive in females.

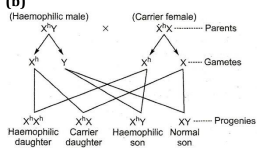
303 **(d)**

The numerous strands of polytene chromosomes are produced but to repeated replication of the paired chromosomes without any cell division (endoduplication) so, that the number of strands in a chromosomes doubles after every round of DNA replication.

304 **(b)**

50% of the sons would be colourblind.

306 **(b)**



So, half the daughters are haemophilic, whereas other half are carrier.

307 (c)

Genetic counselling is the giving of information and advice about the risk of genetic diseases like colour blindness, haemophilia, albinism and outcomes.Genetic screening is a part of genetic counselling, which includes parental diagnosis (like amniocentesis),carrier diagnosis and predictive diagnosis.

308 **(c)**

Dissimilar sex chromosome condition are(i) XY and XY(ii) XX and XO(iii) ZW and ZZ(iv) ZO and ZZ

309 **(b)**

Down's syndrome is due to the trisomy of 21^{st} pair of autosome. Therefore, a Down's syndrome patient has 47 chromosomes, *i.e.*, 45 A + XX/XY.

310 **(a)**

Mutation is the ultimate source of variation. It is said by Hugo de Vries. Mitosis and linkage does not lead to the variation in genetic material

311 **(b)**

External morphology or appearance or descriptive term of an genotype is called phenotype

312 **(b)**

Down's syndrome (Mongolian idiocy) is a congenital disorder caused by trisomy of

chromosome-21, often by non disjunction.

313 (a)

Sex influenced trait.

Finalization of sex at the time of fertilization is known as sex determination. All sex linked character show criss-cross inheritance and firstly it was studied and discovered by TH Morgan (1910). *Sex related trait may be divided into three types*

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314 **(d)**

Chromosomal Disorders These genetic disorders are caused due to absence or excess or abnormal arrangement of one or more chromosomes. These are **non-heritable** and pedigree analysis of a family does not help in tracing the pattern of inheritance of such chromosomal disorders. These are of two types abnormalities due to aneuploidy and aberrations either autosome or in sex chromosomes

315 (a)

Alfred Sturtevant.

Linkage and Recombination

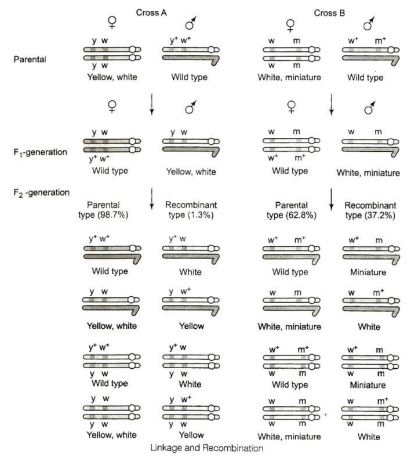
Morgan carried out several dihybrid crosses in *Drosophila* to study genes that were sex linked. The crosses were similar to the dihybrid crosses carried out by Mendel in peas.

He observed that when two genes were grouped on the same chromosome, same genes are highly linked of associated and show low recombination.

When the genes are clearly present linked they show higher recombination.

Morgan attributed this due to the physical association or linkage of the two genes and coined the term linkage to describe this physical association of genes on a chromosome and the term recombination to describe the generation of non-parental gene combinations.

His student Alfred Sturtevant used the frequency of recombination between gene pairs on the same chromosome as a measure of the distance between genes and 'mapped' their position on the chromosome. Morgan hybridized yellow bodied and white eyed females with brown bodied and red eyed male (wild type) (cross-A) and inter-crossed their F_1 -progeny



316 (d)

Hugo de Vries conducted his experiments on evening primrose and postulated the mutation theory. As per this theory, new species were formed by sudden appearance of variations (mutations) rather than continuous variations.

317 (c)

Many chemicals such as acenaphthene, chloral hydrate, colchicine, sulphanil amide, mercury chloride, hexachlorcyclohexane, etc, are used to induce polyploidy. Polyethyl glycol is used in protoplast fusion.

318 **(a)**

Linkage chromosome always present on the chromosome, which represents the same trait called homologous chromosomes

319 **(c)**

According to the genic balance theory of sex determination given by **Bridges**, ratio between the number of X-chromosomes and number of complete sets of autosomes will determine the sex. According to this theory, if the X/A ratio is 1.5, then organism will be **super female**.

320 **(b)**

Emasculation is necessary for the controlled or artificial pollination. Removal of male part

(anther) called emasculation

321 **(b)**

The genes, which remain confined to differential region of Y-chromosome are called **holandric genes**. These genes are not expressed in females Y-linked holandric genes are transmitted directly from father to son.

322 (d)

The given pedigree analysis shows inhertancs of a recessive sex-linked disease like haemophilia.

323 **(b)**

The word **mutation** was first described by **Hugo de Vries** in 1900. Mutation is a sudden heritable change in the characteristics of anorganism. These may result due to the change in genes (DNA or RNA), enzymes, change in chromosome that involves several genes or a change in the plasmogene. But mutation cannot change the **environment.**

324 **(b)**

It is the female reproductive cell, which usually carries more cytoplasm and cytoplasmic organelles than the male cell and hence, naturally would be expected to influence Non-Mendelian traits. Since in the present case, the male parent (not female) had mutation in mitochondria, there are negligible chances of the mutation being inherited.

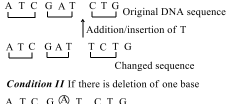
325 **(b)**

The principle of independent assortment states that members of one pair of factors assorted independently of member of other gene pair. Mendel studied the characters which were located on four different chromosome. It is not applicable for genes located on same chromosome, *i.e.*, linked genes.

326 **(c)**

Deletion and insertion of one base leads to entire change of DNA base pair sequence.

DNA base pair sequence is called reading frame *Condition I* If there is insertion of one base



327 **(d)**

The more cross over between A and C than A and B will be possible only when B is present in between A and C. So by taking this consideration the sequence would be $A \rightarrow B \rightarrow C$. This gene sequence also fulfil the other statements also (I, II, IV)

328 (a)

3:1

Law of Independent Assortment

Mendel also worked with and crossed pea plants that different in two characters/factor.

He crossed between pea plant that has seeds with yellow colour and round shape and other that has seeds of green colour and wrinkled shaped.

Ratio appeared as 9:3:3:1 such ratio appeared for several character that Mendel studied

9/16 = Yellow round 3/16 = Yellow wrinkled 3/16 = Green yellow 1/16 = Green wrinkled Based on such observation Mendel concluded second law of inheritance called law of independent assortment.

According to this principle or law the two factors of each character assort or separate independent

of the factors of other characters at the time of gamete formation and get randomly re-arranged in the offspring producing both parental and new combination of traits.

Thus, the phenotypic ratio of a dihybrid cross is 9 : 3 : 3 : 1. The occurrence of four types of plants (more than parental types) in the F_2 generation of dihybrid cross shows that the factors of each of the two characters assort independent of the others as if the other pair of factors are not present. It can also be proved by studying the individual characters of seed colour and seed texture sepraterly.

Seed colour Yellow (9+3 = 12) : Green (3+1 = 4) or 3 : 1

Seed Texture Round (9+3 = 12) : Wrinkled (3+1 = 4) or 3 : 1

The result of each character similar to monohybrid ratio. Generally, the dihybrid cross used to study the independent assortment of genes

329 **(a)**

AB blood group person cannot donate blood to others but can receive blood from all blood groups and hence, called **universal recipient**.

330 **(c)**

Down's syndrome or Mongolian idiocy is not related to sex chromosome X or Y. It is due to the trisomy of 21st chromosome, *i.e.*, total 47 chromosomes. The main features are mental deficiency, short stature, round face, flaccid muscles, small ears protruding tongue and epithelial folds over the eyes (to give Mongolian look).

331 **(a)**

Genes, (DNA) present in cytoplasm are known as cytoplasmic genes or plasma genes. These are responsible for cytoplasmic inheritance or extrachromosomal inheritance.

332 **(a)**

A-TT, B-dominant, C-Recessive

333 **(a)**

In polygenic inheritance, several sets of alleles may produce cumulative effect on the same character, *e.g.*, human height and skin colour.

335 **(b)**

When F_1 -hybrids exhibited a mixture of characters of two parents, the case is considered

as that of incomplete or blending inheritance. It simply means that two genes or allelomorphic pair are not related as dominant or recessive, but each of them expresses itself partially partially. For example, in 4 O'clock plant (*Mirabilis jalapa*),when plants with red flower (homozygous,RR) are crossed with plants having white flower (homozygous,rr),the herterozygous

 F_1 -hybrid (Rr) bear pink flower.

336 (a)

When a gene pair (allele) contains two different genes in an orgenism then the organism is considered as **heterozygous** for that particular character. Gene pair 'Rr' shows the heterozygous condition.

338 **(d)**

Test cross, crossing of F_1 -progeny to the recessive parent is used to find the genotype of the progeny.

339 **(d)**

Amoeba cheris contains 500 chromosomes and *Ascaris megalocephala* contains only **two** chromosomes in each nucleus, *ie*, least number of chromosomes.

340 (a)

Allele	Allele	Genotype	Blood
from	from	of	Types of
Parent	Parent	Offspring	Offsprin
1	2		g
IA	IA	IAIA	А
IA	IB	IAIB	AB
IA	i	I ^B i	А
IB	IA	I ^A I ^A	AB
IB	IB	I ^B I ^B	В
IB	i	I ^B i	В
i	i	ii	0

341 (c)

Chromatin is the material, of which eukaryotic chromosomes are composed. Chemically, chromatin consists of nucleic acids, histone, and non-histone proteins.

342 **(b)**

As we can see in the diagram given in question there are only one trait considered. So, this diagram depict law of segregation explained on the behalf of gene in which genes or factors are indicated as solid and yellow situated on two different chromosome

343 (a)

The disease in the given pedigree chart is X-linked dominant because, it is inherited by female child from her father and none of the male child is affected (X-linked). As disease expressed in female inspite of two XX-chromosomes, it will be dominant.

344 **(a)**

Mendel did not observe linkage that was observed later by another scientists (T H Morgan).

345 **(a)**

A-Male, B-Female, C-Sex unspecified.

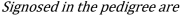
The study of inheritance of genetic traits in several generations of a human family in the form of a family tree diagram is called **pedigree analysis**.

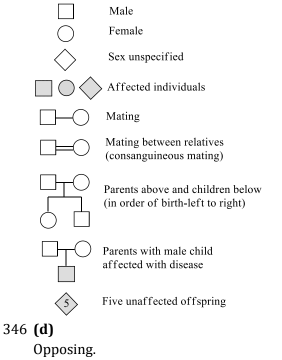
Advantages

(i) It helps in genetic counselling to avoid disorders

(ii) It shows the origin of a trait and flow of a trait in a family

(iii) It is important to know the possibility of a recessive allele that can cause genetic disorders like colour blindness, haemophilia, etc.





7 dominant traits, 7 recessive traits total 14 traits or 7 oppossing pairs of traits

Characters	Dominant Traits	Recessive Traits
Seed shape	Round	Wrinkled
Seed colour	Yellow	Green
Flower	Violet	White
colour		

Pod shape	Full	Constricted
Pod colour	Green	Yellow
Flower	Axial	Terminal
position		
Stem	Tail	Dwarf
height		

347 **(b)**

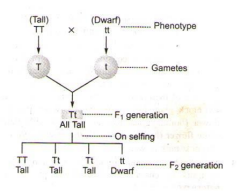
Autosomal recessive

348 **(b)**

Chromosomes (*Chromo* – coloured; some – body) So, the literal meaning of chromosome is the coloured body

349 **(b)**

Monohybrid cross involves single pair of contrasting traits or characters.



Phenotypic ratio 3 : 1

Genotypic ration 1:2:1

351 **(b)**

Selfing of F_1 hybrids in dihybrid cross gives 9:3:3: 1 ratio of progeny. The gametes produced by Aa, Bb are of four types (AB, aB, Ab, ab).

352 **(a)**

Haplopappus gracilis has lowest number of chromosomes out of the given options.

353 **(a)**

Scientist later discovered or found that the gene for seven characters that were took by Mendel in his experiment is present on four chromosome but Mendel was Lucky because seven gene were not linked

354 **(c)**

Drosophila have 3 pairs of autosome and one pair sex chromosome. Sex determination in *Drosophila* is exactly similar to the human beings, *i.e.*, female is homogenetic and male is heterogametic. In the given diagrams, 'A' belongs to the female *Drosophila* and 'B' belongs to the male *Drosophila* 1 indicates = X-chromosomes in female

Drosophila

2 indicates = Autosome

3 indicates = X-chromosomes in male *Drosophila* 4 indicates = Y-chromosome in male *Drosophila*

355 **(d)**

In domesticated fowls, the sex of progeny depends on the type of egg rather than sperm, as two different types of gametes are produced by females (ZW), while males have a same pair of sex chromosome (ZZ).

356 **(a)**

Given example is the example of inversion mutation in which the order of genes in a chromosome get inverted

357 **(a)**

Persons having Rh factor is called Rh⁺and without Rh factor Rh⁻. Rh⁺is **dominant** over Rh⁻.

358 **(b)**

Blood type 'O' has no igen but both types of antibodies 'a' and 'b'. The person with blood type 'O' is universal donor.

359 **(c)**

Law of independent assortment discovered by Mendel because all of his selected traits were present on the different chromosomes. None of his trait present on the same chromosome. If Mendel had studied 7 traits using plant of 12 chromosome instead of 14 then it is most probable that he would have not discovered the law of independent assortment

360 **(c)**

Thalassaemia have two major kinds α -thalassaemia and β -thalassaemia. According to defective gene in α or β -chain of haemoglobin.

Thalassaemia

(i) It is an autosome-linked recessive disesase(ii) It occurs due to either mutation or deletion resulting in reduced rate of synthesis of one of globin chains of haemoglobin

(iii) Anaemia is the characteristic of this disease

- (iv) Thalassaemia is classified into two types
- 5. α -thalassaemia Production of α -globin chain is affected. It is controlled by the closely linked genes HBA1 and HBA2 on chromosome 16. It occurs due to mutation or deletion of one or more of the four genes.

β-thalassaemia Production of β-globin chain is affected. It occurs due to mutation of one or both HBB genes on chromosome 11

361 **(c)**

In silkworm (insect), **female** has **XY** and **male** has **XX** type of sex chromosomes. This is essentially opposite to that of mammals.

362 **(b)**

The X-chromosome linked disease never passed on from father to son because the X-chromosome of father go to the daughter during inheritance.

363 **(d)**

Law of independent assortment does not applicable when the gene of different character occupy on the same homologous chromosome, *i.e.*, linked gene.

364 **(b)**

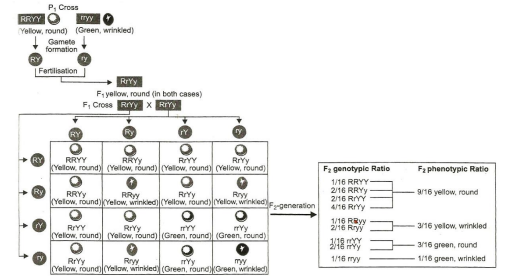
A-Meiosis; B-Allele pair

365 **(c)**

Genetic map is a diagram, which shows the relative position of genes on a chromosome. **Strutevant** in 1911 prepared the first genetic map of two chromosomes of fruitfly.

366 **(c)**

Haemophilia and red green colour blindness, both 370 (a)



are sex linked recessive gene on 'X' chromosome.

Body height is an example of polygenic inheritance.

Rhesus blood group is based on the presence or absence of

Rh-protein on the surface of RBC. Phenylketonuria (PKU) is a recessive autosomal variation.

367 **(a)**

Human's have 22 pairs of autosomes and one pair of sex chromosome

368 **(a)**

There are evidences that the gene for cytoplasmic male sterility particularly in maize reside in **mitochondria**.

369 **(d)**

In most cases if there is a change in genotype than it ultimately leads to change in phenotype also

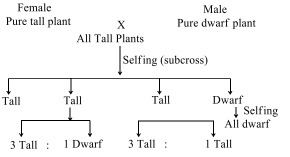
A dihybrid cross in pea plant between yellow round (smooth) seeded and green wrinkled seeded plant. The cross proves the principle of independent assortment

371 (a)

A cross in which parents differ in a single pair of contrasting character is called monohybrid cross. From the monohybrid cross the Mendel gave law of segregation, law of paired factor and law of dominance.

Law or Principle of Dominance In heterozygous individuals or hybrids a character is represented by two contrasting factors called alleles or allelomorphs. Out of the two contrasting alleles. Only one is able to express its effect in the individual. It is called dominant factor or dominant allele.

The other which does not show its effect in the heterozygous individual is called recessive factor or recessive allele. The cross between the pure tall and pure dwarf gives all progeny (F_1) tall.



The character shown by F_1 called dominant character

372 (a)

Genic Balance Theory of Sex The theory of genic balance given by Calvin Bridges (1926) states that instead of XY-chromosomes sex is determined by the genic balance or ratio between Xchromosomes and autosome genomes The theory is basically applicable to *Drosophila melanogaster* over, which bridges worked. He found that the genic ratio X IA of 1.0 produces fertile females whether the files have XX + 2A or XXX + 3A chromosome complement. A genic ration (X I A) of 0.5 forms a male fruitfly. This occurs in XY + 2A as well as XO + 2A

Chromosome	X/A Ratio	Sexual
Complement		Morphology
X X X + 2A	3/2 or 1.5	Metafemale
X X X + 3A	3/3 or 1.0	Female
X X + 2A	2/2 or 1.0	Female
X X + 3A	2/3 or 0.67	Inter sex
X X X + 4A	3/4 or 0.75	Inter sex
X O + 2A	1/2 or 0.5	Male
X Y +2A	1/2 or 0.5	Male
X Y +3A	1/3 or 0.33	Metamale

373 **(b)**

 $\rm Rr$ and red because the R is dominant r so, the $\rm F_{1^-}$ hybrid will be red

374 **(a)**

Grasshopper is an example of XO type of sex determination in which the male have only one Xchromosome besides the autosomes, whereas females have a pair of X-chromosomes (2A + XX).

375 **(d)**

9:3:3:1.

Law of Independent Assortment

Mendel also worked with and crossed pea plants that different in two characters/factor.

He crossed between pea plant that has seeds with yellow colour and round shape and other that has seeds of green colour and wrinkled shaped. Ratio appeared as 9:3:3:1 such ratio appeared for several character that Mendel studied 9/16 = Yellow round 3/16 = Yellow wrinkled 3/16 = Green yellow 1/16 = Green wrinkled Based on such observation Mendel concluded second law of inheritance called law of independent assortment.

According to this principle or law the two factors of each character assort or separate independent of the factors of other characters at the time of gamete formation and get randomly re-arranged in the offspring producing both parental and new combination of traits.

Thus, the phenotypic ratio of a dihybrid cross is 9 : 3 : 3 : 1. The occurrence of four types of plants (more than parental types) in the F_2 generation of dihybrid cross shows that the factors of each of the two characters assort independent of the others as if the other pair of factors are not present. It can also be proved by studying the individual characters of seed colour and seed texture sepraterly.

Seed colour Yellow (9+3 = 12) : Green (3+1 = 4) or 3 : 1

Seed Texture Round (9+3 = 12) : Wrinkled (3+1 = 4) or 3 : 1

The result of each character similar to monohybrid ratio. Generally, the dihybrid cross used to study the independent assortment of genes

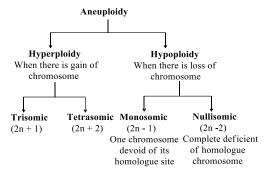
376 **(d)**

Trisomy stands for 2*n*+1.

Chromosomal disorders are caused due to excess, absence or abnormal arrangement of one or more chromosomes.

Sometimes the chromatids fail to segregate during cell division, resulting in gain or loss of a chromosome. This is called **aneuploidy** Aneuploidy is also called heteroploidy Aneuploidy is of two kinds

(i) Hyperploidy (ii) Hypoploidy



377 **(a)**

The genotype of Klinefelter syndrome is XXY. In this there is one extra X-chromosome. This extrachromosome comes when there is nondisjunction of X-chromosome in ova-or-sperm fuses with Y or X-chromosome of sperm or ova resulting XXY genotype

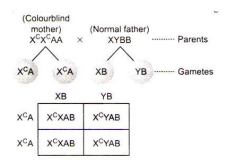
378 **(a)**

Colour blindness is caused by recessive sex-linked gene carried by X-chromosome. So, in male, one recessive gene is sufficient for its expression, *i.e.*, $(X^{C}Y)$ but female needs two recessive

gene $(X^{C}X^{C})$ for the expression of colour blindness.

Mother will colourblind with A blood group and father normal with blood group-B as-

So, all the sons will be colourblind with AB blood group.



379 **(b)**

Deletion and duplication occurs in homologous chromosomes usually during meiosis. In deletion, some part of chromosome is lost while in

383 **(a)**

Complete linkage.

Linkage and Recombination

Morgan carried out several dihybrid crosses in *Drosophila* to study genes that were sex linked. The crosses were similar to the dihybrid crosses carried out by Mendel in peas.

He observed that when two genes were grouped on the same chromosome, same genes are highly linked of associated and show low recombination.

When the genes are clearly present linked they show higher recombination.

duplication a piece of chromosome is copied next to an identical section, thus, increasing chromosome length.

380 **(b)**

The husband is heterozygous for Rh⁺so, it will (Rh rh) and wife is homozygous of Rh⁻ so, it will have genetic alleles (rh rh) :

So, 50% homozygous Rh⁻ children will be born.

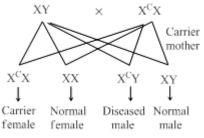
381 **(a)**

Mendel could not find out linkage because all of his experimental characters of pea were not linked. They were present far apart from each other

382 **(a)**

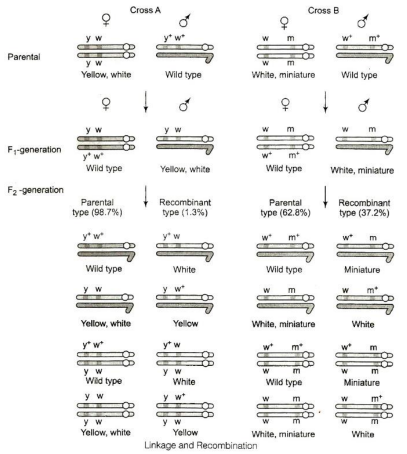
Bleeder's disease.

Haemophilia is also called Bleeder's disease. It is a sex linked recessive disorder. It transmit from carrier female to male progeny



Morgan attributed this due to the physical association or linkage of the two genes and coined the term linkage to describe this physical association of genes on a chromosome and the term recombination to describe the generation of non-parental gene combinations.

His student Alfred Sturtevant used the frequency of recombination between gene pairs on the same chromosome as a measure of the distance between genes and 'mapped' their position on the chromosome. Morgan hybridized yellow bodied and white eyed females with brown bodied and red eyed male (wild type) (cross-A) and inter-crossed their F_1 -progeny



In the incomplete linkage we get some recombinant progeny but in complete linkage the recombinant progeny percentage is very less as compared to incomplete linkage

384 **(b)**

With regard to transfusions of whole blood or packed red blood cells, individuals with 0 type negative blood are often called universal donors, and those with type AB positive blood are universal recipients, however, these terms are only generally true with respect to possible reactions of the recipients anti-A and anti-B antibodies to transferred red blood cells.

385 **(c)**

The individuals suffering from Klinefelter's syndrome disease show trisomy of X-chromosomes, *i.e.*, they contain 47 chromosomes in total with 22 autosomes (XXY).

386 **(c)**

Dominant and recessive were expressed or appeared together separately. This shows that

there is no mixing of characters means nonblending of character

387 **(a)**

Sex-limited and sex-linked genes are located on autosomes.

388 **(b)**

Types of gametes $= 2^n$

n=Number of heterozygotes considered, i.e., monohybrid cross(n=1), dihybrid cross (n=2), trihybrid cross (n=3) and so on. AA BB CC × aa bb cc is a trihybrid cross, therefore, different types of gametes are

$$= 2^{n}$$
$$= (2)^{3}$$
$$= 2 \times 2 \times 2 = 8$$

389	(b)	398	(b)
	The point mutations involves alterations in the		X =
	structure of gene by altering the structure of DNA.		Dro
	Point mutations are of two types- (i) base pair		sex
	substitution and (ii) frameshift mutation.		is e
200	(h)		is h
390	Type A blood group receive blood A and O type.		give
	Type A blobu group receive blobu A and o type.		Dro
391	(d)		1 in
	Pink colour flower is the intermediate character.		Dro
	It is obtained maximum by crossing of		2 in
	homozygous red and white flower		3 in
	Example (Red) (White)	200	4 in
	RR × rr	399	ln tl
	Rr (pink) – All progeny have pink colour		and
392			rela
	<i>Triticale</i> is the hybrid variety, which is obtain by		i cia
	crossing between wheat and rye.	400	(b)
	<i>'Triti</i> ' is for wheat and 'cale' is for rye together it is		The
	called <i>Triticale</i> . Its production value is higher than		que
202	both wheat (<i>Triticum</i>) and rye (<i>Secale</i>)	401	(a)
393	Mendel found that tallness, round shape of seed,		Gen
	yellow colour of seed, purple colour of flower,	402	ക
	green colour of pod,inflated nature of pod and	402	Ane
	axial position of flower were domianant over		chro
	dwarfness, wrinkled seed, green colour of seed,		CIII
	white colour of flower, yellow colour of pod ,	403	(b)
	constricted nature of pod and terminal position of		Gen
	flower.	101	(J)
		404	Allo
394			for
	Garden pea is the common name for <i>Pisum</i>	405	
	<i>sativum</i> which was the experimental material of	105	Cri-
205	Mendel's experiments		5p c
395			Leje
	Mutations can be induced by a number of agents		to a
	called mutagens.		syn
	Physical mutagen- UV rays, X-rays, cosmic rays,		lary
	high temperature.		-
		406	
	Chemical mutagens- 5 bromouracil, proflavin.		Dur

396 (c)

The observable characteristics of an organism is called phenotype. These are determined by its genes

397 **(b)**

Carrier organism refers to an individual, which carries a recessive gene that is not expressed.

1, 3 Y = 4.

sophila have 3 pairs of autosome and one pair chromosome. Sex determination in Drosophila xactly similar to the human beings, *i.e*., female omogenetic and male is heterogametic. In the en diagrams, 'A' belongs to the female sophila and 'B' belongs to the male Drosophila dicates = X-chromosomes in female sophila

dicates = Autosome

dicates = X-chromosomes in male Drosophila dicates = Y-chromosome in male *Drosophila*

he given case, embryo contains one Barr body one F-body, which corresponds to XXY ted to Klinefelter's syndrome.

chances of fifth child to be albino in previous stion would be one in four

otypes of the parents shall be I^Ai and I^Bi.

uploidy is the deletion or addition of few omosomes from the original genomes.

otypes of C and D are XX^c and XY respectively.

somes, heterosomes are the synonymous used sex chromosomes

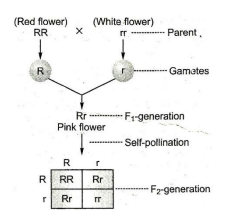
du-chat syndrome also known as chromosome deletion syndrome, 5p minus syndrome or eune's syndrome is a rare genetic disorder due missing part of chromosome-5. This droms is associated with malformation of the mx.

ing his experiments, Mendel called factors something unchanged that pass from parent to offspring through the gametes over successive generation

407 (b)

In case of incomplete dominance, when pink flowers of F₁-generation, are self-pollinated, they develop red (RR), pink (Rr) and white(rr) flowers in the ratio 1 : 2 : 1, respectively, genotypically as

well as phenotypically.



Genotypic ratio-RR : Rr : rr

1:2:1

Phenotypic ratio-Red : Pink : White

1:2:1

408 (d)

Genome of model organisms

Organism	Number of Bass Pair	Numbe r of Gene
Bacteriop	10	-
hage	thousand	
Escherichi	4.7 million	4,000
a coli		
Saccharo	12 million	6,000
myces		
cerevisiae		
Caenohab	97 million	18,000
ditis		
elegans		
Drosophil	180 million	13,00
а		
melanoga		
ster		
Human	3 million	30,000
Lily	106 billion	-

409 **(b)**

Albinism is caused by the absence of enzyme **tyrosinase**, which is necessary for the synthesis of melanin.

410 (c)

In human beings, ABO blood grouping is an example of dominance, codominance and multiple alleles.

411 (a)

In sickle-cell anaemia only one nucleotide

substitution takes place from T to A. Genetic or chromosomal symbol used for person who is having sickle-cell anaemia Ps – Hb^s Hb^s. **Sickle-cell Anaemia**

(i) It is an autosome-linked recessive trait
(ii) The disease is controlled by a single pair of allele Hb^s and Hb^s

(iii) Only the homozygous individuals for Hb^s, *i.e.*, Hb^sHb^s show the diseased phenotype
(iv) The heterozygous individuals are carriers
(Hb^AHb^S)

(v) Due to point mutation, glutamic acid (Glu) is replaced by valine (Val) at the sixth position of β globin chain of haemoglobin molecule (vi) A single base substitution at sixth codon of the beta globulin gene from GAG to GUG. GAG code for glutamic acid and GUG code for valine. (vii) Hb^S behaves as normal haemoglobin except under the oxygen stress where erythrocytes lose their circular shape and become sickle-shaped. As a result, the cells cannot pass through narrow capillaries. Blood capillaries are clogged and thus, affect blood supply to different organs

412 **(d)**

A cell or an organism having two copies of a single genome (with chromosome number 2x) is called **diploid.**

413 **(d)**

All the given statements are correct. Occasionally a single gene product may produce more than one effect. For example starch synthesis in pea seeds is controlled by one gene. It has two allele (B and b). Starch is synthesized effectively by BB and have bigger grains. In contrast bb homozygous have lesser efficiency in starch synthesis and produce smaller grains *In starch synthesis gene following condition to seen*

BB – rounded (due to more starch synthesis)
bb – wrinkled (due to less starch synthesis)
Bb – in between rounded of wrinkled size. It
produce starch of intermediate quantity between
BB to bb homozygous condition. So, it is
incomplete dominance.

Therefore, dominance is not an autonomous feature of a gene or the product that it has information for. It depends as much on the gene product and the production of a particular phenotype from this product as it does on the particular phenotype that we choose to examine, in case more than one phenotype is influenced by the same gene

414 **(b)**

When Rh⁻mother carries Rh⁺ foetus, in the first pregnancy no serious problem occurs because Rh⁺antigen arises in child's blood and the concentration of antibodies produced in mother's blood due to immunization by child's Rh⁺ antigen will be rather low.

415 **(a)**

Blood group-A has antigen-A and antibody-b.

416 **(b)**

Chromosomes are of two type, **autosomes** and **sex chromosomes** (allosomes). Y-chromosome is the sex chromosome.

417 **(a)**

Law of Independent Assortment

Mendel also worked with and crossed pea plants that different in two characters/factor.

He crossed between pea plant that has seeds with yellow colour and round shape and other that has seeds of green colour and wrinkled shaped.

Ratio appeared as 9 : 3 : 3 : 1 such ratio appeared for several character that Mendel studied

9/16 = Yellow round 3/16 = Yellow wrinkled 3/16 = Green yellow 1/16 = Green wrinkled Based on such observation Mendel concluded second law of inheritance called law of independent assortment.

According to this principle or law the two factors of each character assort or separate independent of the factors of other characters at the time of gamete formation and get randomly re-arranged in the offspring producing both parental and new combination of traits.

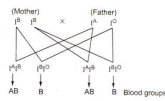
Thus, the phenotypic ratio of a dihybrid cross is 9:3:3:1. The occurrence of four types of plants (more than parental types) in the F_2 generation of dihybrid cross shows that the factors of each of the two characters assort independent of the others as if the other pair of factors are not present. It can also be proved by studying the individual characters of seed colour and seed texture sepraterly.

Seed Colour Yellow (9+3 = 12) : Green (3+1 = 4) or 3 : 1

Seed Texture Round (9+3 = 12) : Wrinkled (3+1 = 4) or 3 : 1

The result of each character similar to monohybrid ratio. Generally, the dihybrid cross used to study the independent assortment of genes

418 **(a)**



Thus, possible groups are :**AB** and **B**.

419 **(a)**

Recessive alleles influence the appearance of the phenotype only when they are in homozygous condition.

Mendel did not find the blending inheritance in F_2 -generation of a Mendelian monohybrid cross the tall and dwarf traits were identical to their parental types and doesn't show blending

420 **(c)**

The eggs are produced by meiosis, *i.e.*, reduction division. So, the egg contains one X-chromosome when released from ovary.

421 **(a)**

Variation.

Variation is the degree of difference in the progeny and between the progeny and the parents. The term variation is also used for a single difference in a trait

422 **(c)**

Pleiotropic gene is one which produces or controls more than one effects or characters. In other words, we can say that pleiotropic gene produces a major phenotypic trait and with that also influences some other phenotypic traits, *e.g.*, lethal genes, which are known to control the menifestation of some phenotypic trait alongwith affecting the viability of organism.

423 **(b)**

A-Dizygotic twins are the twins, which results from the fusion of two sperm with two ova. It is very rare in case of human beings.

B-Monozygotic twins are the twins, which results from the fusion of one sperm with one ova leads to zygote. This zygote later on divide and give rise to two or more zygote. In this cells of all progeny have the identical genome

425 **(c)**

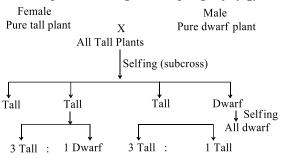
H J Muller was awarded Nobel Prize in 1946 for his discovery of the production of mutations by Xray irradiation.

426 **(c)**

Law or Principle of Dominance

In heterozygous individuals or hybrids a character is represented by two contrasting factors called alleles or allelomorphs. Out of the two contrasting alleles. Only one is able to express its effect in the individual. It is called dominant factor or dominant allele.

The other which does not show its effect in the heterozygous individual is called recessive factor or recessive allele. The cross between the pure tall and pure dwarf gives all progeny (F_1) tall.



The character shown by F_1 called dominant character

427 **(b)**

It is metaphase

428 **(b)**

When heritable alterations occur in a very small segment of DNA molecule, *i.e.*, a single nucleotide or nucleotide pair then this type of mutations are called **point mutations**.

429 **(a)**

Dihybrid genotype ratio. 1 : 2 : 1 : 2 : 4 : 2 : 1 : 2 : 1 total nine types of genotype and four types of phenotype

430 **(b)**

In Morgan's experiment on linkage, the percentage of white eyed, miniature-winged recombinants in F²generation is 37.2%. It is due to incomplete linkage, where two linked genes are sufficiently apart and the chances of their separation are quite good.

431 **(b)**

Dihybrid cross.

Law of Independent Assortment

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The result of each character similar to monohybrid ratio. Generally, the dihybrid cross used to study the independent assortment of genes

432 **(d)**

Dominant allele shows its effect in homozygous or heterozygous condition and recessive allele shows its effect only in homozygous condition. Given pedigree chart possible only when the male parent in heterozygous for hyperdactyle. If it is homozygous for hyperdactyle then its son would also be the hyperdactyle as well

433 **(c)**

Mutation happens by itself. It is the spontaneous phenomena

434 **(c)**

The **centromere** lies within a thinner segment of chromosome, the **primary constriction**.

435 (d)

Mutations are not acquired. They are selected by the nature. Gametes fuse with regard to which alleles they carry Mendel's knowledge background was mathematics and physics. In Vienna university he studied botany and physics. Later he became the teacher of physics and natural sciences.

Main reasons for Mendel's success were.

(i) Mendel took those traits, which were not linked

(ii) Mendel choose distinctive contrasting pairs(iii) Mendel took one or two character at a time

(iv) Pea plant is ideal for controlled breeding

(v) Mendel kept complete record of every cross(vi) Mendel used statistical method and law of probability (vi) Mendel used statistical method

437 **(a)**

Insect, grasshopper, cockroaches and bugs have XX and XO type of sex determination in which XO happens to be male and XX happens to be female

438 (a) Linkage and Recombination

and law of probability

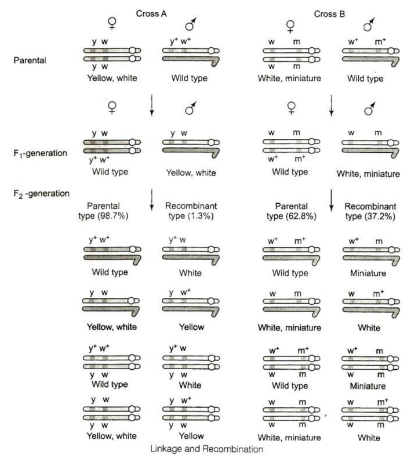
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439 (c)

In Mendel's dihybrid cross out of 16 progenies, 6 are recombinants so in case of 1280 progenies, the recombinants are

 $=\frac{1280\times 6}{16} = 480$ progenies.

440 **(b)**

A child of blood group-O cannot have parents of AB and O blood groups.

441 (d)

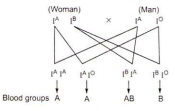
Rh factor was discovered by Landsteiner and Wiener (1940) in Rhesus

monkey (*Macaca rhesus*). It is found in man and rhesus monkey only. Erythroblastosis foetails occurs when the mother is Rh⁻, father is Rh⁺ and foetus is Rh⁺.

442 (d)

Monosomics (2n-1) one chromosome less then diploid set of somatic chromosome number.

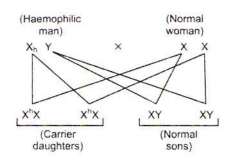
443 **(b)**



In this case, progeny with blood group-B is produced only when man is heterozygous, *i.e.*, I^AI^O.

444 (d)

The children of a haemophilic man and normal woman will be all normal. Neither sons nor daughters (only carrier) will be haemophilic



445 (c)

A set of three alleles present on chromosome number 9 is responsible for the four blood groups.

446 (d)

The gene for colour blindness is present on Xchromosome. If one X-chromosome of female have gene for colour blindness, this will be carrier (normal) but if a male have gene on Xchromosome it will be colourblind (since only one X-chromosome is present in male).

447 **(b)**

Cri-du–chat syndrome is caused by a conspiuous deletion in the short arm of 5th chromosome. These individuals asre severly impaired and their cat-like crying give the syndrome its name.

448 (c)

Inversion invoves a reverse order of genes in part of chromosome. ABCDEFGH has been shown in figure, where breaks occur between A and E. Reunion at broken ends may leads to inversion of the segment BCD into DCB. Indeletion, a section of chromosome is lost.

450 (c)

Ischihara chart is used to detect colour blindness.

451 **(b)**

The gene, which produces more than one phenotypic effect in an organism is called pleiotropic gene. It produces a major phenotypic traits and along with influences some other phenotypic traits, *e.g.*, sickle cell anaemia in man.

452 (a)

Persons with blood group-AB are called universal recipients because both antigens A and B are found in their blood and the two antibodies 'a' and 'b' are absent. Therefore, such persons can receive blood of all the blood groups.

453 **(b)**

Genes in the non-homologous region of Ychromosome pass directly from male to male. In man, the Y-linked or holandric genes such as ichthyosis, hystrix, gravis hypertrichosis (excessive development of hairs on pinna of ear) are transmitted directly from father to son.

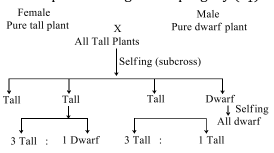
454 (b)

3:1 ratio in F_2 -generation explained by law of dominance Principle of law of dominance state that only dominant allele shows its effect evan in the heterozygous condition.

Law or Principle of Dominance

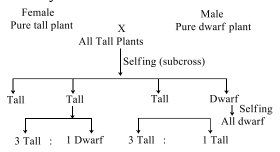
In heterozygous individuals or hybrids a character is represented by two contrasting factors called alleles or allelomorphs. Out of the two contrasting alleles. Only one is able to express its effect in the individual. It is called dominant factor or dominant allele.

The other which does not show its effect in the heterozygous individual is called recessive factor or recessive allele. The cross between the pure tall and pure dwarf gives all progeny (F_1) tall.



The character shown by F_1 called dominant character

Mendel cross-pollinated a pure tall pea plant (100-120 cm hight) and a pure dwarf pea plant. (only 22 to 44 cm hight). He called them parental generation, expressed now-a-days by symbol P. This hybridization popularly called as monohybrid cross



This three generations of pea plants after crossing a pure tall plant with a dwarf one. The plants of F_1 -generation are all tall, of F_2 -generation three tall and one dwarf. One third of the tall plants are pure, while the remaining behave as hybrids F_1 -generation Seeds collected from the parental generation called first filial generation or F_1 generation

F₂-generation F_1 -plants pollinated among them self (self breeding or inbreeding) and seed produced by F_1 -plants called F_2 -generation. They were in ratio 3:1 (three tall and one dwarf). **F₃-generation** Mendel allowed F_2 -plant to form seed by self-pollination called F_3 -generation. Mendel observed that tall and dwarf plant behave differently

(i) Dwarf plant produced dwarf plant on self-pollinated

(ii) In tall plants one third plants breed true so they were pure

(iii) Other two third plant behave like parents and give tall to dwarf plants 3 : 1 indicate that their parents have dwarf genes also

455 **(a)**

Rr and red because the R is dominant r so, the $F_1\mathchar`-$ hybrid will be red

 $\begin{array}{cccccc} A & B & C & D \\ RR & Rr & Rr & rr \\ \downarrow & & \downarrow & \downarrow \\ Red & Red & White \\ 1 & : & 2 & : & 1 \end{array}$

456 **(b)**

In dominant epistasis, a dominant gene(epistatic) masks the expression of another dominant or recessive gene (hypostatic). Such interactions give the modified F_2 ratio as 12:3:1.

457 **(c)**

Studies of human sex-linked trait shows that males are affected and females are carrier in most casese.

The recessive genes located on X-chromosome in humans are always expressed in male because human male is **hemizygous**. It is the condition in which even recessive genes get expressed when it is present on one chromosome because the another copy of chromosome have very less genes

458 **(b)**

A cross between homozygous recessive and hetrozygous plant is called test cross. It gives 1 : 1 ratio in monohybrid and 1 : 1 : 1 : 1 ratio in dihybride cross.

459 **(c)**

The experimental material garden pea used by Mendel is a self-fertilised crop and artificial pollination

460 **(c)**

W Bateson (1905) explained the lack of independent assortment in sweet pea and **T H Morgan** (1905) in *Drosophila* due to linkage.

461 **(b)**

Buffs and Kips are not the types of blood groups or blood factor.

462 **(c)**

463 (c)

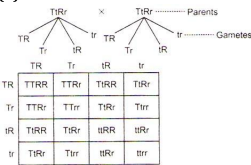
Environment can also influence the result of same genotype. Like, human skin colour. This is the example of incomplete dominance In ABO blood group system, inheritance of grouping is controlled by a single autosomal gene on chromosomal-9 with three major alleles A, B, an O (I^A , I^B and I^O). The ABO blood group system has at least 6 alleles. On the basis of presence or absence of antigens and antibodies four blood groups (phenotypes) have been differentiated-A, B, AB and O blood groups. The phenotype, genotype, antigen and antibody of blood group ABO.

Phen	Genotyp	Antige	Antibo
otype	e	n	dy
А	I ^a I ^o , I ^a I ^a	А	b
В	I ^b I ^o , I ^b I ^b	В	а
AB	I ^a I ^b	A, B	None
0	IoIo	None	a, b

464 (a)

The probability of genotype TTrr in F_2 generation of a dihybrid cross is $\frac{1}{16}$.





So, the number of genotypes of TtRr = 4

and TtRR = 2

466 **(d)**

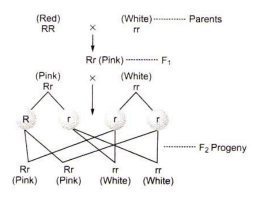
Blood group-O does not contain any antigen. Thus, ii does not produce any sugar polymer on the surface of the RBC.

467 **(a)**

Idiogram is the diagrammatic representation of the chromosomes of an individual.

468 **(b)**

Mirabilis jalapa shows incomplete dominance.



The ratio pink and white flowers will be 1 : 1.

469 **(b)**

The marriage between normal visioned man and colourblind woman will produce colourblind sons and carrier daughters.

470 (d)

In this pedigree chart of a family with five children, the parents are heterozygous.

471 **(d)**

Correct statements are

(i) Myotonic dystrophy is an autosomal dominant trait

(ii) Sickle-cell anaemia is autosomal recessive disease

(iii) Failure segregation result in chromosomal loss and gain. One daughter cell get one extrachromosome and other gets one less chromosome

(iv) Cystic fibrosis is Mendelian disorder

472 **(c)**

Haemophilia is X-linked recessive mutation thus, commonly seen in human males than in females.

473 **(c)**

Mendel 's work was rediscovered independently by three scientists, *i.e.*, **de Vries**(Dutch). **Carl Correns**(German) and **Tschermak**(Austrian).

474 **(a)**

 $\frac{1}{4}:\frac{1}{2}:\frac{1}{4}$ ratio of TT, Tt, tt can be depicted mathematically binomial expression as =

 $(ax + by)^2$.

Monohybrid cross can be denoted as 1:2:1(genetic) and the expression of binomial is also 1:2:1 so monohybrid cross can be represented in any of the given option (a or b or c) because their results are same

476 **(c)**

In the case of Klinefelter's syndrome, the male

possesses a Barr body, while in the case of Turner's syndrome, the Barr body is absent.

477 **(c)**

When a dominant gene and other recessive gene are present on opposite chromosomes of homologous pair, the heterozygotes are called as *trans* heterozygotes.

478 **(d)**

The superiority of the hybrids over either of the parents (dominant or recessive) is called hybrid vigour (G. Shull).Heterosis is equivalent of hybrid vigour.

479 **(b)**

As heterozygous alleles have two different types of chromosomes, *so they produce two types of genes*



480 **(b)**

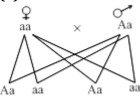
Ť

Prokaryotic nucleoid consists of DNA only; no histones associated with it.

481 **(a)**

In acrocentric chromosomes, centromere appears sub-terminal, *i.e.*, with a very small and a very long arm.

482 **(a)**



The pedigree given in question is the most probable autosomal disease

483 **(b)**

Sickle cell anaemia is caused by a change in a single base pair of DNA. It is a genetic disease reported from Negroes. In sickle cell haemoglobin the glutamic acid in β –chain is replaced by valine. The individuals of sickle cell anaemia are immune to malaria.

484 **(b)**

Directional selection operates on the range of phenotypes existing within the population and exerts selection pressure, which moves the mean phenotype towards one phenotypic extreme.

485 (a)

A-Homozygous; B-Heterozygous

486 **(d)**

The basic chromosome number of wheat is 7(x=7) and its hexaploid species contain $42(6 \times 7)$ chromosomes. Thus, it's monosomic (one chromosomes missing) contains 41(42-1) chromosomes. Haploid contains 21 (half to the 42) chromosomes. Nullisomic (one chromosome pair missing) contains 40(42-2) chromosomes and trisomic contain (one chromosome extra) 43(42+1) chromosomes.

487 **(b)**

In the incomplete linkage we get some recombinant progeny but in complete linkage the recombinant progeny percentage is very less as compared to incomplete linkage

488 (a)

Phenylalanine hydroxylase.

Phenylketonuria (Folling; 1934). It is an inborn, autosomal, recessive metabolic disorder in which the homozygous recessive individual lacks the enzyme **phenylalanine hydroxylase** needed to change phenylalanine (amino acid) to tyrosine (amino acid) in liver. It results in hyperphenylalanine

489 **(d)**

Monohybrid cross can be denoted as 1:2:1(genetic) and the expression of binomial is also 1:2:1 so monohybrid cross can be represented in any of the given option (a or b or c) because their results are same

490 **(d)**

Telomeres have unique structures, including short nucleotide sequences present as tandemly repeated units. Clusters of G residues in one strand and C residues in the other characterise telomeric DNA. Also, in some species the telomeres terminate with a single-stranded DNA (12-16 nucleotides long) rich in guanine.

491 **(d)**

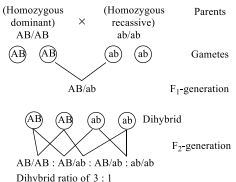
Heterozygous Round Yellow has genotype RrYy. On selfing, Round Green offsprings produced and are represented by RRyy and Rryy, genotypes only (R=Round; yy=Green).

492 **(d)**

Dihybrid linked gene cross (3 : 1)

Linked Gene The linked genes do not show independent assortment but remain together and are inherited *en*-block producing only parental type of progeny. They give a dihybrid ratio of 3 : 1 and a test cross ratio of 1 : 1

Dihybrid ratio of two linkage genes



493 **(c)**

Law of segregation states that heredity characters in the form of allele separate from each other in the formation of gametes. Half of the gametes carry one allele and the other half carry other allele.

494 **(a)**

Law of purity of gamete states characters recovered in F_2 -generation that was hiden F_1 . **Principle of Law of Segregation** This law is also called the purity of gametes. This law states that the two factors of a character present in individual keep their identity distinct separate at the time of gametogenesis (meiosis) or sporogenesis, factors get randomly distributed to different gametes and then get paired again in different offspring as per the principle of probability. The principal of segregation can be deduced in Punnett square

495 **(a)**

In *Melandrium*, sex is determined by X and Ychromosomes. X-chromosome is shorter than the Y-chromosome. If Y-chromosome is present, the individual shall be male and if it is absent, it will be female.

496 **(c)**

Mutated genes are mostly recessive, which will affect the new generations.

497 **(a)**

Genes at more than one locus are called as **polygenes**. Variation in these genes in a particular population have a combined effect upon a particular phenotypic character. **Multiple alleles** are the more than two different forms of a gene, present on the same locus.

498 **(a)**

Linkage was first suggested by Sutton and Boveri when they formulated the famous chromosomal theory of inheritance. Betason and Punnett (1906) while on working on sweet pea also noticed that some factors remain together and don't show recombination or segregation

499 (a)

X-linked genes are represented twice in female and once in male.

500 **(c)**

Colour blindness is due to a recessive gene carried on the X-chromosome and therefore men are more likely to show the defect although women may be carriers.

501 **(a)**

A-Female B-Meta female C-Male.

Genic Balance Theory of Sex The theory of genic balance given by Calvin Bridges (1926) states that instead of XY-chromosomes sex is determined by the genic balance or ratio between Xchromosomes and autosome genomes The theory is basically applicable to *Drosophila melanogaster* over, which bridges worked. He found that the genic ratio X IA of 1.0 produces fertile females whether the files have XX + 2A or XXX + 3A chromosome complement. A genic ration (X I A) of 0.5 forms a male fruitfly. This

occurs in XY + 2A as well as XO + 2A

Chromosome	X/A Ratio	Sexual
Complement		Morphology
X X X + 2A	3/2 or 1.5	Metafemale
X X X + 3A	3/3 or 1.0	Female
X X + 2A	2/2 or 1.0	Female
X X + 3A	2/3 or 0.67	Inter sex
X X X + 4A	3/4 or 0.75	Inter sex
X O + 2A	1/2 or 0.5	Male
X Y +2A	1/2 or 0.5	Male
X Y +3A	1/3 or 0.33	Metamale

502 (a)

The ABO blood group are controlled by I gene, which have three alleles $(\mathrm{I}^{A},\mathrm{I}^{B},\mathrm{I}^{O})$

503 **(b)**

Mendel studied seven different pair of contrasting characters, on the basis of studies on garden pea (*Pisum sativum*).

Dominant and recessive characters are as follows:

Character	Dominant	Recessive
The length	Tall	Dwarf
of stem		
The	Axial	Terminal
position of		
flower		
The colour	Green	Yellow
of the pod		
The shape	Inflated	Constrict
of the pod		ed
The shape	Round	Wrinkled
of the seed		
The colour	Coloured	White
of the seed		
coat		
The colour	Yellow	Green
of the		
cotyledon		

504 **(a)**

When an animal has both the characters of male and female, it is called **intersex**.

505 (a)

Mutation that takes place due to single base pair is called point mutation, *e. g.*, Sickle-cell anaemia

506 **(a)**

Due to hemizygous condition in male the most of recessive sex linked disease seen in males. Generally female are carrier. Occasionally they affected by sex linked disease

507 **(b)**

Law of independent assortment deduced by Mendel by performing dihybrid cross (9:3:3:1). Incomplete dominance was not deduced by Mendel

508 **(a)**

Genetic or chromosomal symbol used for person who is having sickle-cell anaemia Ps – Hb^s Hb^s. Sickle-cell Anaemia

Sickle-cell Anaemia

(i) It is an autosome-linked recessive trait
(ii) The disease is controlled by a single pair of allele Hb^s and Hb^s

(iii) Only the homozygous individuals for Hb^s, *i.e.*, Hb^sHb^s show the diseased phenotype

(iv) The heterozygous individuals are carriers $({\rm Hb}^{\rm A}{\rm Hb}^{\rm S})$

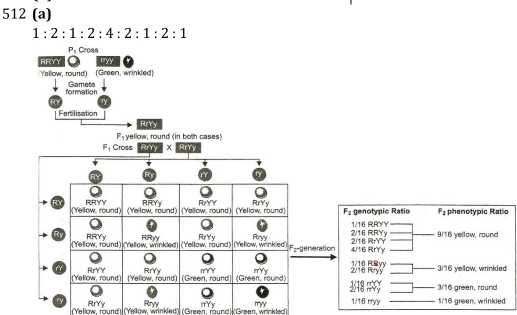
(v) Due to point mutation, glutamic acid (Glu) is replaced by valine (Val) at the sixth position of β -globin chain of haemoglobin molecule

(vi) A single base substitution at sixth codon of the beta globulin gene from GAG to GUG. GAG code for glutamic acid and GUG code for valine. (vii) Hb^S behaves as normal haemoglobin except under the oxygen stress where erythrocytes lose their circular shape and become sickle-shaped. As a result, the cells cannot pass through narrow capillaries. Blood capillaries are clogged and thus, affect blood supply to different organs

509 (d)

In human most characters are controlled by one gene but some characters like human skin colour is controlled by more than one gene and some characters like ABO blood group and human stairs colour are not inherited according to Mendel inheritance pattern





Foetal sex is determined by Barr body test.

511 **(d)**

The sex-linked allele or disease never pass from men to his sons because alleles of sex linked disease present on the sex chromosome-X not Y. This condition is also called hemizygous condition

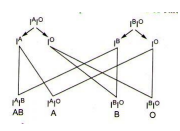
A dihybrid cross in pea plant between yellow round (smooth) seeded and green wrinkled seeded plant. The cross proves the principle of independent assortment

513 **(a)**

The law of segregation of characters is also called the law of purity of gametes because gametes have only one of the two alleles for each character.

514 **(c)**

In men, ABO blood group is best example of multiple allelism. The four children of blood group A, B, AB and O will be born if both the parents are heterozygous for A and B, *i.e.*, the genotype of one parent is I^AI^O and of other is I^BI^O.



Linkage and incomplete dominance were the post Mendelian discoveries.

Post Mendelian Discoveries

Gene interaction is the influence of alleles and nonalleles ion the normal phenotypic expression of genes. It is two types, intragenic (allelic) and intergenic (nonallelic). In the intragenic interaction the two allels (present on the same gene locus on the two homologous chromosome) of a gene interact in such a way as to produce a phenotypic expression different from typical dominant-recessive phenotype, *e.g.*, incomplete dominance, codominance, multiple alleles. In intergenic or non-allelic interaction, two or more independent gene present on the same or different chromosomes interact to produce different expression, e.g., epistasis, duplicate genes, complementary genes, supplementary genes, lethal genes, inhibitory genes, etc.

516 (a)

Barr body is, infact, an X-chromosome, which has become inactive or heterochromatic. It is present in the nuclei of the cells of females (not males).

517 **(c)**

A man can inherit his X-chromosome from his maternal grandfather only because the Xchromosome of the paternal grandfather goes to sister of his father.

519 **(b)**

Test cross is the cross of an individual with its homolozygous recessive parent. In other words, we can say that it is a specialized back cross of F_1 -hybrid with it's homozygous recessive parents in order to determine whether it is homozygous or heterozygous for a character.

520 **(c)**

Organism with more than two sets of chromosomes are known as **polyploids**. These may be triploid (3*n*), tetraploid (4*n*), and so on. Polyploidy may be of three types, *i.e.*, autopolyploidy, allopolyploidy and autoallopolyploidy.

522 (c)

L –shaped chromosomes are also called **sub – metacentric chromosomes**.

524 **(a)**

In the given case, the gene for disease is sexlinked dominant.

525 **(c)**

Polygenes show polygenic inheritance or quantitative inheritance. As genes are present on different chromosomes they will segregate independently during meiosis.

Number of individuals in $F_2 = 4^n$

n =

number of genes present

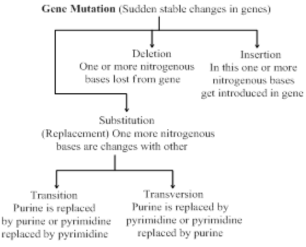
 $= 4^3 = 64$ individuals

$$1+6+2x+20+6+1=64$$

$$x = \frac{30}{2} = 15$$

526 (d)

Chromosomal mutation or change is the sudden inheritable change in the hereditary material of an organism. It caused due to the several ways like deletion, duplication, translocation, inversion, etc.



Main types of Mutation

Loss of Chromosome	Process
Loss of fragment of	Deletion
chromosome	
Inversion of fragment of	Inversion
chromosome	
Attachment of segment of one	Translocatio
chromosome to another	n
Repetition of a segment in a	
chromosome	Duplication

Gene Mutation	Process
Replacement of one base or	Substitution
more nucleotide	
Removal of one base or more	Deletion
nucleotide	
Addition of one base or more	Addition
nucleotide	

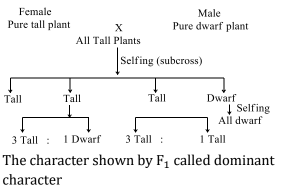
527 (a)

Law of dominance.

Law or Principle of Dominance

In heterozygous individuals or hybrids a character is represented by two contrasting factors called alleles or allelomorphs. Out of the two contrasting alleles. Only one is able to express its effect in the individual. It is called dominant factor or dominant allele.

The other which does not show its effect in the heterozygous individual is called recessive factor or recessive allele. The cross between the pure tall and pure dwarf gives all progeny (F_1) tall.



(dominant allele)

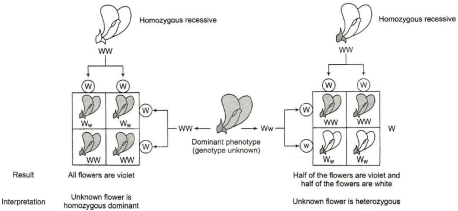
528 **(b)**

In mitosis cell division the chromosomal number

529 **(c)**

In monohybrid test cross the unknown dominant trait progeny crossed with recessive parent and the ratio between dominant and recessive comes out to be 1 : 1 only if the testing progeny was heterozygous dominant and if it was homozygous than all the progeny would have dominant character.

Crossing of F_1 (dominant phenotype) to any one of the parent called back cross and when phenotype of crossing parent is recessive than this is called test cross. The progeny of such cross can easily be analysed to predict the genotype of test organism



The cross of heterozygous dominant with its recessive parent is called test cross. The test cross gives 1:1 ratio in monohybrid condition, whereas 1:1:1:1 in dihybrid condition

530 **(c)**

Genetics word is derived from the Greek word genesis, which stands for descent. Term genetics was introduced by Bateson in 1906 branch of Biology that deals with the study of heredity and variations

531 **(d)**

Sickle cell anaemia is a biochemical disorder inherited as a recessive trait. In this disease, the haemoglobin differs in electrophoretic mobility and physiochemical properties from normal haemoglobin.

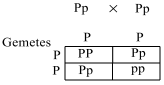
532 **(d)**

A mutated gene Hb^s produces sickle cell haemoglobin, in which the sixth amino acid, *ie*, glutamic acid in β -chain of normal haemoglobin is replaced by amino acid valine causing sickleshaped haemoglobin.

533 **(c)**

When F_1 hybrids exhibited a mixture on blending of characters of two parents, the case is considered as that of incomplete blending inheritance. It simply means that the two genes of allelomorphic pair are not related as dominant or recessive, but each them express itself partially.

534 **(c)**



Progeny 1, 2, 3 have the dominant trait and 4 progeny have recessive trait so the ratio between dominant of recessive progeny is 3 : 1 or

remain the same that's way it is called equatorial division. In meiosis cell division the chromosomal number remain the half of the original one that is way it is called reductional division percentage of dominant progeny is 75%

535 **(a)**

As homozygous have only one type of alleles, so they produce only one kind of gamete

536 **(b)**

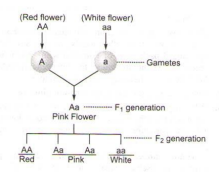
In higher plants, the gametes are formed by the mitotic division of microspores and megaspore. The microspores and megaspore are haploid since these are produced by the meiosis in microspore mother cell and megaspore mother cell respectively. The somatic cell has 40 chromosomes, *i.e.*, 2n = 40. Then the germ cell also have 40 chromosomes. The germ cells divide by meiosis and produce four haploid (*n*) gametes therefore, the chromosomes number will be 20.

537 **(b)**

Secondary oocyte is haploid, hence, it possess 22 + X-chromosomes.

538 **(a)**

Incomplete dominance is seen in *Antirrhinum* (snapdragon).



539 **(c)**

Test cross is the cross of F_1 with its recessive parent. It is used to observed that the F_1 is homozygous or heterozygous. It gives 1:1 ratio in monohybrid and 1:1:1:1 ratio in dihybrid cross.

540 **(b)**

In human beings, 46 chromosomes are found, in which only one pair takes part in sex determination. These are known as **sex chromosomes** or **allosomes**, rest 22 pairs are known as **autosomes**.

541 **(a)**

Euploidy is a normal state having balanced number of chromosome, that is an exact multiple of the haploid set, *e.g.*, if a haploid number is 5 then euploidy number would be 5, 10, 15, 20, etc.

542 **(a)**

In 1900 the Mendel's law were rediscovered. Mendel died in 1884 long before his work came to recognized. It was in 1900 when three worker independently rediscovered the principles of heredity already worked out by Mendel. They were Hugo de Vries of Holland, Carl Correns of Germany and Eric Tiron and Tschermark of Austria

543 **(a)**

Sickle cell anaemia, a hereditary disease is an example of **pleiotropy** (ability of a gene to have many effects).

544 **(d)**

The genes which individually have a small effect but collectively produce significant phenotypic expression are called polygenes. The inheritance of these genes is called polygenic inheritance, *e.g.*, skin colour in human.

545 **(a)**

In chromosome, heterochromatin remains condensed in secondary constriction.

546 **(c)**

Inheritance of ABO blood group shows **multiple** allelism.

547 **(a)**

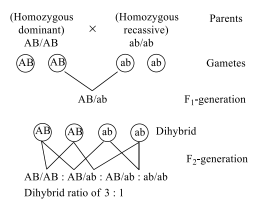
Neurospora complete their sexual life cycle in a few days and thus, make ideal organism for the study of laws of heredity. Penicillin, a potent antibiotic is obtained from *Penicillium notatum* and LSD is obtained from *Claviceps purpurea*.

548 **(c)**

Barr body (sex chromatin) is the densely staining mass that represents an inactivated X chromosomes found in nuclei of somatic cells of most female mammals. Number of Barr bodies is one less than the total X-chromosomes, therefore number of Barr bodies in XXXXY = 4 - 1 = 3.

549 **(d)**

Linked Gene The linked genes do not show independent assortment but remain together and are inherited *en*-block producing only parental type of progeny. They give a dihybrid ratio of 3 : 1 and a test cross ratio of 1 : 1 Dihybrid ratio of two linkage genes

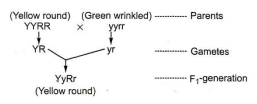


550 (c)

Colchicine is a poisonous chemical, isolated from seeds and bulbs of autumn crocus (*Colchicum autumnale*). It blocks spindle formation and thus, inhibits the movement of sister chromatids to the opposite poles. The resulting restitution nucleus includes all the chromatids. As a result, the chromosome number of the cell is doubled, which leads to polyploidy.

551 **(d)**

Mendel crossed a dominant homozygous yellow, round seeded plant with a recessive homozygous green and wrinkled seeded plant. The F_1 -offsprings produced are heterozygous yellow, round seeded plants.



552 **(a)**

Dominant and recessive these two words are commonly used for the describing alleles

553 **(b)**

Fruit fly (*Drosophila*) is used in genetic experiments. As polytene chromosomes, sex determination and sex linked inheritance have been studied in fruit fly.

554 **(c)**

Phenylketonuria is caused by the absence or deficiency of the enzyme phenylalanine hydroxylase, which results in the accumulation of phenylalanine in all body fluids.

555 **(b)**

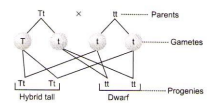
Balbiani rings are large RNA puffs reported in the salivary gland chromosomes of *Chironomus* insect during larval development.

556 **(d)**

Inheritance by multiple alleles causes a traits to exhibit more than ywo possible phenotypes.

557 **(c)**

Test cross is a cross between homozygous recessive parent and F_1 offsprings. The genotypic ratio as well as phenotypic ratio of such cross is 1 : 1.



Analysis of the given data in question shows almost 1 : 1 (94:89) ratio. Hence, the genotype of the two parents will be Tt and tt.

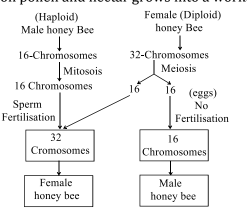
558 **(a)**

Haploid.

Haploid diploid mechanism of sex determination (haplodiploidy).

Hymenopterous insect such as bees, wasps ants show unique phenomena in which an unfertilized egg develops into male and female develops from fertilized egg.

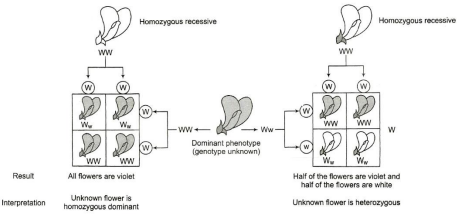
In honeybee, the quality of food determines whether a diploid larva will become a fertile queen or a sterile worker female. A larva fed on royal jelly a secretion from the mouth of mussing workers grows into a queen, whereas a larva fed on pollen and nectar grows into a worker bee



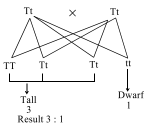
559 **(b)**

In monohybrid test cross the unknown dominant trait progeny crossed with recessive parent and the ratio between dominant and recessive comes out to be 1 : 1 only if the testing progeny was heterozygous dominant and if it was homozygous than all the progeny would have dominant character.

Crossing of F_1 (dominant phenotype) to any one of the parent called back cross and when phenotype of crossing parent is recessive than this is called test cross. The progeny of such cross can easily be analysed to predict the genotype of test organism



The cross of heterozygous dominant with its recessive parent is called test cross. The test cross gives 1:1 ratio in monohybrid condition, whereas 1:1:1:1 in dihybrid condition



560 **(b)**

When mutation is confined to only one substitution, deletion, insertion then this type of mutation is called point mutation. Sickle-cell anaemia is the example of point mutation in which the Glutamic acid (Glu) is replaced by valine (val) at the sixth position of β -globin chain of haemoglobin molecule

561 **(a)**

Mendel used letter symbols to denote factors. He used capital letters for dominant factors and small letters for recessive factor

562 **(d)**

Harmful mutation does not get eliminated from gene pool because mostly harmful mutation are recessive and carried by heterozygous individual

563 **(c)**

Father's blood group-B and mother's blood group-O will not be able to produce a offspring of blood group-A.

564 **(d)**

A-Sugar, B-I^A I^B, C-Sugar

565 (a)

Father of experimental genetics is TH Morgan. He

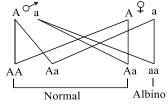
is also called the fly man of genetics because of selecting fruit fly (*Drosophila melanogaster*) as research material in experimental genetics

566 **(a)**

Thomas Hunt Morgan (the father of experimental genetics) selected fruitfly (*Drosophila melanogaster*) is also called Jackpot of genetics as experimental material

567 (d)

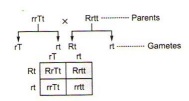
Albino is the recessive trait which comes only when there is homozygous condition. In the given problem the progenies are both albino and normal. This is possible only when their parents are heterozygous for normal colour



568 **(a)**

Linkage genes always arranged linearly on the homologous chromosome called linkage group

569 (d)



All the offsprings have different phenotypes.

Therefore, the phenotypic ratio obtained by crossing rrTt and Rrtt is 1 : 1 : 1 : 1.

570 (d)

Linked gene does not separate frequently. They remain together because linked gene lie very closely to each other

571 **(b)**

In translocation the segment of the one chromosome get attached to the other chromosome. Option A indicates inversion. Option 'b' indicates translocation. Option 'c' indicates deletion and option 'd' indicates duplication

572 **(d)**

Experimental verification of the chromosomal theory of inheritance was given by **Thomas Hunt Morgan** and his colleagues. This led to discovering the basis for variation that sexual reproduction produced.

573 **(c)**

A gene is said to be **epistatic**, when its presence suppresses the effect of a gene at another locus. Epistatic genes are sometimes called **inhibiting genes** because of their effect on other genes, which are described as **hypoststic**.

574 **(b)**

Jumping genes or mobile elements or transposons or transposable elements are DNA sequences that are able to move from one site to another. Transposons were discovered by **Barbara** McClintock, an American Geneticist, in a corn plant.

575 **(b)**

Originally, Mendel proposed two laws, **firstly law of segregation** and then law of independent assortment. Mendel coined the term dominant for any trait that express itself when present with the factors for the contrasting trait, and used the word recessive for any trait that is not expressed when present alongwith contrasting dominant trait in the hybrid.

576 **(b)**

Based on the ratio of F_2 and F_3 generation, Mendel proposed that something was being stably passed down F_1 and F_2 -generation) unchanged, from parent to offspring through the gametes, over successive generations.

He called these things as factors. Now we call them genes. Genes therefore, are the units of inheritance required to express a particular trait (d)

577 **(d)**

Finalization of sex at the time of fertilization is known as sex determination. All sex linked character show criss-cross inheritance and firstly it was studied and discovered by TH Morgan (1910). *Sex related trait may be divided into three types*

(i) **Sex Linked Traits** They are those traits the determining genes of which are found on the sex chromosomes. All the sex-linked traits present on a sex chromosome are inherited together (ii) **Sex Limited Traits** They are autosomal traits which are expressed in a particular sex in response to sex hormones although their genes also occur in the other sex, *e.g.*, milk secretion in mammalian females, pattern baldness in males. The gene for baldness behaves as an autosomal dominant in males and autosomal recessive in females

(iii) **Sex Influenced Traits** The traits are not due to particular genes but are by products of sex hormones, *e.g.*, low pitched voice, beard moustaches. In males, pattern baldness is related to both autosomal genes as well as excessive secretion of testosterone

578 **(c)**

Variation is the degree of difference in the progeny and between the progeny and the parents. The term variation is also used for a single difference in a trait

579 **(b)**

The more maternal influence can be expected in the cytoplasmic inheritances (*i.e.*, the inheritance of genes contained in the cytoplasm of a cell, rather than the nucleus). The reason is that of the female reproductive cell or the egg has large amount of cytoplasm containing many organelles which contain their own genes and can reproduce independently (*e.g.*, mitochondria and chloroplast) and which are consequently incorporated into the cytoplasm of all the cells of the embryo. The male reproductive cell (sperm and pollen) consists almost solely of a nucleus. Cytoplasmic organelles are thus, not inherited from the male parent. This is why, the cytoplasmic inheritance is also called **maternal inheritance**.

580 (d)

Mutagen may be physical or chemical agents, which causes change in DNA sequence. Like UVradiation acridne dye, etc

581 (c)

Linkage process was fully explained as a theory by **T H Morgan** (1911).

582 **(b)**

Blood group-AB has antigen A and B on RBCs but no antibodies in plasma. The alleles A and B are codominant.

583 **(c)**

A polyploid having two or more distinct genome usually produced by chromosome doubling of interspecific hybrids is called **allopolyploid** or **interspecific polyploid**.

584 (d)

Chromosomal theory of inheritance was proposed by Walter Sutton and T. Boveri in 1902. Sutton and Boveri made a correlation between Mendel's conclusion about genes and the behaviour of chromosome during mitosis and meiosis.

Polygenes show quantitative inheritance. Nilsson Ehle (1909) explained it in Kernel colour of wheat and Davenport (1910) for skin colour in humans.

585 **(a)**

Early agriculturists (8000-1000 BC) knew that cause of variation hidden in the process of sexual reproduction. However, our ancestors had no idea about the scientific basis of inheritance and variation

586 **(a)**

Punnett square was developed by British geneticist Reginald C. Punnett. It is graphical representation, to calculate the probability of all possible genotypes of offspring in genetic cross

587 **(d)**

ZW and ZZ Type of Sex Determination This mechanism operates in certain insects (butterflies and moths) and in vertebrates (fishes, reptiles and birds). The male has two homomorphic sex chromosomes (ZZ) and is homogametic and the female has two heteromorphic sex chromosomes (ZW) and is heterogametic. There are thus two types of eggs with Z and with W and only one type of sperms. *i.e.*, each with Z

	A + Z	A + W
A + Z	AA + ZZ	AA + ZW
A + Z	AA + ZZ	AA + ZW
	Males	Females

WZ-ZZ types of sex determination

588 **(a)**

Turner's syndrome is due to monosomy (2n - 1) and the chromosome constituent is 44 + XO = 45. The individual is female with under developed ovary, webbed neck. Kinefelter's syndrome is due to condition of 44+XXY=47.

The sex is male but have secondary sexual characters like female. Down's syndrome is due to trisomy of 21st chromosome.

589 **(d)**

According to Mendel' law of **independent assortment** two factors of each trait separate at random and independent of the factors of other traits at the time of meiosis (gametogenesis/sporogenesis) and get randomly as well as independently arranged in the offsprings.

590 **(c)**

A-Pair, B-Homozygous

591 **(d)**

Inversion : A piece of chromosomes is removed and rejoined in reverse orientation. It can be of two types:

- 7. **Pericentric Inversion** : The inverted segment does not contains centromere.
- 8. **Paracentric Inversion** : The inverted segment contains centromere.

592 **(c)**

Morgan and Castle.

It was TH Morgan who clearly proved and define linkage on the basis of the breeding experiments in fruitfly. In 1911, Morgan and Castle proposed 'chromosomal' theory of linkage'

593 **(a)**

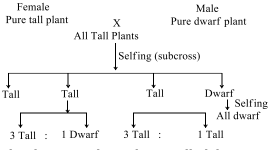
Dominant factor.

Law or Principle of Dominance

In heterozygous individuals or hybrids a character is represented by two contrasting

factors called alleles or allelomorphs. Out of the two contrasting alleles. Only one is able to express its effect in the individual. It is called dominant factor or dominant allele.

The other which does not show its effect in the heterozygous individual is called recessive factor or recessive allele. The cross between the pure tall and pure dwarf gives all progeny (F_1) tall.



The character shown by F_1 called dominant character

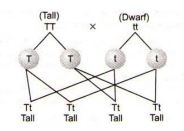
594 **(b)**

Human have 46 chromosomes out of which 22 pairs or 44 chromosomes are called **autosomes** and one pair of chromosomes (*i.e.*, XX in female and XY in male) are called **heterosomes** or **sex chromosomes**.

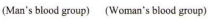
595 **(b)**

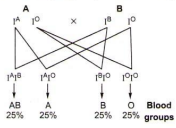
A tall plant was grown in nutrient deficient soil and remained dwarf; genetically, this plant has a genetic constitution of tall plant.

So, the cross between such plant and a dwarf plant will resulted into all hybrid tall plants.



596 **(a)**





There is 25% chances of first offspring having blood group-AB.

597 **(c)**

Paired factor or allele only applicable on the organism which are multicellular and diploid. Protista and Monera both are the unicellular

598 **(b)**

The allele which does not show its effect in heterozygous individual is called recessive factor or recessive allele. It shows its phenotype only in absence of dominant factor or dominant allele 599 (d)

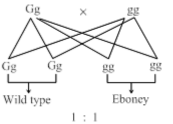
599 **(a)**

Human skin colour is the example of multiple gene inheritance. Human skin colour contributed by three separate genes. Environment also plays a significant role in determining human skin colour

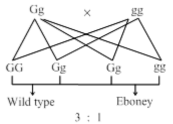
600 (c)

A-Clotting, B-Blood, C-Continuous 601 (d)

Condition I cross between Gg and gg



Condition II Cross between Gg and Gg



So, strain II and strain IV are heterogametic (Gg) 602 (d)

When Rh⁻ individual receive Rh⁺ blood, individual becomes **isoimmunized**.

603 **(c)**

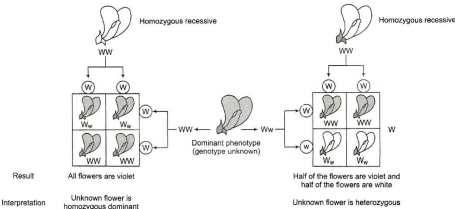
Incase of **transition**, purine base is replaced by another purine (*e.g.*, A by G) and pyrimidine is replaced by another pyrimidine (*e.g.*, C by T) and *vice versa*. In case of transversion, purine is replaced by a pyrimidine and *vice versa*.

604 **(c)**

Recessive characters are expressed when they are present in X-chromosome of male.

605 **(a)**

Crossing of F_1 (dominant phenotype) to any one of the parent called back cross and when phenotype of crossing parent is recessive than this is called test cross. The progeny of such cross can easily be analysed to predict the genotype of test organism



used to study the independent assortment of genes

607 **(b)**

Colour blindness is the sex-linked recessive disease in which the defective gene carried by the X-chromosome. So, if a person is colourblind then it is due to the defective gene present in the father and mother

608 **(c)**

When there is a loss of one chromosome from the homologous pair, this is called **monosomy** (2n-1) and when there is addition of one chromosome to the homologous pair, this called **trisomy** (2n+1).

609 **(b)**

Haemophilia, cystic fibrosis, thalassaemia. Sicklecell anaemia, colour blindness, phenylketonuria *Genetic disorder may be grouped into two categories*

(i) **Mendelian Disorders** These genetic disorder are mainly caused by alternation and mutation in the single gene. They are transmitted to offsprings following the principle of inheritance. Mendelian disorder can be dominant or recessive. *e. g.*, haemophilia, colour blindness, sickle-cell anaemia, cystic fibrosis, phenylketonuria, thalassaemia.

(ii) **Chromosomal Disorders** Chromosomal disorder are caused due to excess, absence, or abnormal arrangement of one or more chromosome, *e.g.*, Turner's syndrome, Down's syndrome, etc

610 **(a)**

16th.

Thalassaemia

(i) It is an autosome-linked recessive disesase

606 **(b)**

3:1

Law of Independent Assortment

Mendel also worked with and crossed pea plants that different in two characters/factor.

He crossed between pea plant that has seeds with yellow colour and round shape and other that has seeds of green colour and wrinkled shaped. Ratio appeared as 9 : 3 : 3 : 1 such ratio appeared for several character that Mendel studied

9/16 = Yellow round 3/16 = Yellow wrinkled 3/16 = Green yellow 1/16 = Green wrinkled Based on such observation Mendel concluded second law of inheritance called law of independent assortment.

According to this principle or law the two factors of each character assort or separate independent of the factors of other characters at the time of gamete formation and get randomly re-arranged in the offspring producing both parental and new combination of traits.

Thus, the phenotypic ratio of a dihybrid cross is 9: 3:3:1. The occurrence of four types of plants (more than parental types) in the F_2 generation of dihybrid cross shows that the factors of each of the two characters assort independent of the others as if the other pair of factors are not present. It can also be proved by studying the individual characters of seed colour and seed texture sepraterly.

Seed colour Yellow (9+3 = 12) : Green (3+1 = 4) or 3 : 1

Seed Texture Round (9+3 = 12) : Wrinkled (3+1 = 4) or 3 : 1

The result of each character similar to monohybrid ratio. Generally, the dihybrid cross

(ii) It occurs due to either mutation or deletion resulting in reduced rate of synthesis of one of globin chains of haemoglobin

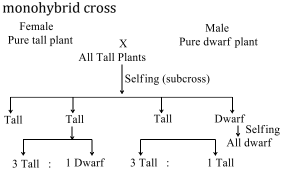
(iii) Anaemia is the characteristic of this disease

(iv) Thalassaemia is classified into two types

- 9. α -thalassaemia Production of α -globin chain is affected. It is controlled by the closely linked genes HBA1 and HBA2 on chromosome 16. It occurs due to mutation or deletion of one or more of the four genes.
- β-thalassaemia Production of β-globin chain is affected. It occurs due to mutation of one or both HBB genes on chromosome 11

611 **(a)**

Mendel cross-pollinated a pure tall pea plant (100-120 cm hight) and a pure dwarf pea plant. (only 22 to 44 cm hight). He called them parental generation, expressed now-a-days by symbol P. This hybridization popularly called as



This three generations of pea plants after crossing a pure tall plant with a dwarf one. The plants of F_1 -generation are all tall, of F_2 -generation three tall and one dwarf. One third of the tall plants are pure, while the remaining behave as hybrids F_1 -generation Seeds collected from the parental generation called first filial generation or F_1 generation

F₂-generation F_1 -plants pollinated among them self (self breeding or inbreeding) and seed produced by F_1 -plants called F_2 -generation. They were in ratio 3:1 (three tall and one dwarf).

F₃-generation Mendel allowed F_2 -plant to form seed by self-pollination called F_3 -generation. Mendel observed that tall and dwarf plant behave differently

(i) Dwarf plant produced dwarf plant on selfpollinated

(ii) In tall plants one third plants breed true so they were pure

(iii) Other two third plant behave like parents and

give tall to dwarf plants 3 : 1 indicate that their parents have dwarf genes also

613 **(a)**

An individual containing both dominant and recessive genes or traits or characters of a allelic pair is known as **heterozygous** or hybrid.

614 **(b)**

A-Less; B-More

615 **(c)**

In the cross of incomplete dominance the genotypic and phenotypic ratio both are same. Rr and red because the R is dominant r so, the F_1 -hybrid will be red

616 **(c)**

Thrombin facilitates the formation of the enzyme prothrombinase, which convert prothrombin into thrombin.

617 **(c)**

Mendel chose *Pisum sativum*(garden pea) to explain the laws of inheritance. His selection of garden pea was evidently not an accident, but the result of a long careful thought.

618 **(a)**

Blood groups are inherited from our biological parents in the same way as eye colour and other genetic traits. Within the ABO blood group system, the A and B genes are codominant, *i.e.*, these will be expressed whenever the gene is present. The O gene is silent and only expressed when neither A nor B is present.

619 **(c)**

Determination of sex of each child is an independent and exclusive event so the probability that whether the child will be a boy or a girl is 50% each in every case.

620 **(a)**

According to genotypic ratio of law of independent assortment for dihybrid cross, the RrTt genotype will be found in 100 plants out of 400 plants(as in 16 plants 4 having this type of genotype.

621 (c)

Blood Antigen Antibody

group	on RBC	in Serum
А	А	Anti-b
В	В	Anti-a
AB	A and B	None
0	—	Anti-a
		and Anti-
		b

Hence, blood group-AB has no antibodies in serum.

622 **(b)**

If there are more than one X-chromosome then one X-chromosome remains active, while the other one becomes inactive and condenses to form Barr body. Barr body is a sex chromatin particle. Barr bodies can be used as a sex marker always occur in numbers one less than the total number of X-chromosomes.

623 **(c)**

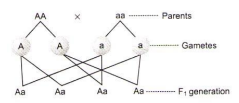
Inversions occur when there are two breaks in a chromosome and the intercalary segment reunites in a reverse order by rotating at 180°.

624 **(a)**

For the given case, there is no chance that the offsprings will have very dark skin.

625 **(c)**

If a cross is made between AA and aa, where **A** is dominant over **a**, then the nature of F_1 progeny will be **genotypically Aa** and **phenotypically A**. It means that the genotype of progeny will have both **A** and **a** but it will show the external appearance or character (s) regulated by gene A.



626 **(d)**

In body cells of a normal female, one of the Xchromosome become inactive and form Barr body near the nuclear membrane.

XY genotype has no Barr body.

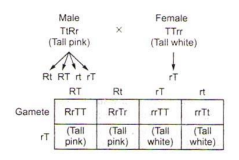
XX genotype has single Barr body.

XXX genotype have two Barr bodies.

XXXX genotype have three Barr bodies.

627 **(a)**

Offsprings would be tall pink and tall white (genotypic ratio 1 :1)



629 **(c)**

Beadle and **Tatum** conducted experiment on pink bread mould (*Neurospora cressa*)

And stated that each gene has the information to produce one enzyme. This concept was formulated as one gene-one enzyme hypothesis.

630 **(c)**

During synapsis, deletion and duplication occurs.

631 **(a)**

Aneuploidy is an abnormal number of chromosomes and is a type of chromosome abnormality. The presence of an extra chromosome 21 is found in Down's syndrome.

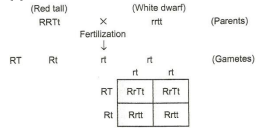
632 **(c)**

Mendel formulated his generalisation, which were read out at two meetings of Natural History Society of Brunn in 1855. His paper 'Experiments on Plant Hybridisation' was published in proceedings of Brunn Natural Science Society in 1866. Mendel died in 1884 without getting any recognition for his work

633 **(b)**

Red-green colour blindness or colour blindness is a genetic disorder in which eyes fail to distinguish red and green colours.

634 (c)



Phenotype of different plants is

11. All plants contain red fruits.

- 12. 50% plants are tall with red fruits.
- 13. 50% plants are dwarf with red fruits.

635 **(b)**

The process of gametogenesis or meiosis cell division is shown in the given diagram (in question). Which clearly indicates towards the Mendel's law of independent assortment. Because it says two factors of each character assort or separate independent of the factors of other characters at the time of gamete formation (gametogenesis) and get randomly re-arranged in the offspring producing both parental and new combination of traits

636 **(d)**

Haemophilia or Bleeder's disease is a sex linked (X-linked recessive) disease. Down's syndrome is caused due to trisomy of chromosome 21 *(i.e.,* hyperploidy type of aneuploidy). Phenylketonuria is an autosomal recessive gene disorder.

Sickle cell anaemia in human, is also inherited as an autosomal recessive disorder.

637 **(d)**

Allelic sequence variation has traditionally been described as a DNA polymorphism if more than one variant (allele) at a locus occurs in human population with a high frequency, it is referred to as DNA polymorphism.

638 **(d)**

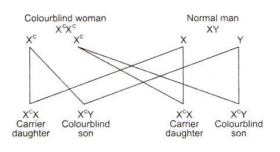
ZW-ZZ system of sex determination occurs in certain insects and vertebrates such as fishes, reptiles and birds. Here, the female sex has one Zchromosome and one W-chromosome.

639 **(b)**

Eight types of gametes will be produced.

640 **(d)**

In human colour blindness and haemophilia are X-linked disease.



All the sons are colourblind.

641 **(a)**

Bridges gave genic balance theory of sex determination, which is related to *Drosophila melanogaster*.According to him Xchromosome are carriers of genes for femaleness and autosomes are carriers of genes for maleness, so it is the ratio of X-chromosomes and autosomal sets which determine sex.

642 **(c)**

Turner's syndrome is a chromosomal disorder in which all or a part of one of the sex chromosomes is absent. Turners syndrome, having XO genotype are stelite females with poorly dereloped ovaries and underdeveloped breasts.

643 **(a)**

According to law of independent assortment of Mendel, alleles segregate randomly from each other, *i.e.*, all alleles separate from each other during gamete formation and are inherited independently from one anther. But genes, which are located on the same chromosomes (called linked genes) do not assort independently. Such type of genes are called **linked genes**, and this phenomenon is called as **linkage**.

644 **(b)**

Red and green.

Colour Blindness

(i) It is a sex-linked recessive disorder

(ii) It results in defect in either red or and green cone cells of eye resulting in failure to discriminate between red and green colour
(iii) The gene for colour blindness is present on X-chromosome

(iv) It is observed more in males (X^cY) because of presence of only one X-chromosome as compared to two chromosomes in famales

645 **(c)**

All those, which are located in the single chromosome set constitute a **linkage group**. *E. coli* contains a single linkage group.

646 **(a)**

The genes which have higher COV (Cross Over Value) are placed farthest and genes, which have lowest COV are placed close to each other. V and U have highest COV = 30 T and V have lowest COV = 5 After gathering the other COV the sequence of genes will be VTWU

647 **(b)**

Heredity.

Heredity (L. *Hereditas* – Heirship or inheritance) is the transmission of genetically based characters 655 (d) from parents to their offspring.

The process by which characters are transferred from one generation to the next generation is called inheritance

648 (a)

When the F_2 individuals are crossed with its pure recessive parent, the cross is called test cross. The result of it, is always 1 : 1 in monohybrid cross and 1:1:1:1 in hybrid cross.

649 (a)

Main reasons for Mendel's success were. (i) Mendel took those traits, which were not linked

(ii) Mendel choose distinctive contrasting pairs

(iii) Mendel took one or two character at a time

(iv) Pea plant is ideal for controlled breeding

(v) Mendel kept complete record of every cross (vi) Mendel used statistical method and law of probability

650 **(b)**

In codominance, both the genes of an allelomorphic pair express themselves equally in F_1 -hybrids. The ratio in F_2 -generation is 1:2:1, both genotypically as well as phenotypically, e.g., codominance of coat colour in cattle, and codominance of blood alleles in man.

651 (a)

Landsteiner recognized three blood groups, i.e. ,blood group-A (with antigen-A), blood group-B (with antigen-B) and blood group-O (without antigen).

652 (d)

Gamma rays are used to induce mutagenesis in crop plants.

653 (a)

Fertilization brings together the chromosomes of two mating types. Crossing over between these during meiosis leads to genetic recombination.

654 (d)

Cytoplasmic or mitochondrial inheritance is the inheritance in which the trait pass only from mother to all of their offspring. The genes of that inheritance present in the cytoplasm of ova that's way these genes goes to all of their offspring. As sperm have very less cytoplasm so this inheritance doesn't applicable for males

Linked gene don't show any accordance with Mendel's law because Mendel's law can be applied only on unlinked gene

656 (d)

Haploids plants can be produced in large number by anther and ovary cultures. Haploids may be useful for isolatio nof mutants, since, even recessive mutant alleles will be expressed in the mutagen treated generation itself.

657 (d)

Mendel's work remain unrecognized due to (i) Communication was not easy

(ii) Concept of factors which did not blend was not accepted

(iii) Using of mathematics to explain biological problem was unacceptable

(iv) He could not provide any physical proof for the existence of factors

658 (b)

The cross of heterozygous dominant with its recessive parent is called test cross. The test cross gives 1:1 ratio in monohybrid condition, whereas 1:1:1:1 in dihybrid condition.

659 (a)

A-Dwarf (F₂), B-F₃, C-F₄, D-Homozygous 660 (a)

> In first pregnancy, Rh⁺blood from the foetus enters the circulation of an Rh⁻mother, antibodies against the Rh antigen will be made.

661 **(b)**

Phenylketonuria (chromosomes 12) (i) It is inborn error of metabolism and is inherited as autosomal recessive trait (ii) The affected individual lacks an enzyme called phenylalanine hydroxylase that converts the amino acid phenylalanine into tyrosine in liver. (iii) Phenylalanine is accumulated and gets converted into phenylpyruvic acid and other derivatives. This effects the brain, resulting in mental disorder

662 **(b)**

Four, six.

The phenomenon of expression of both the alleles in heterozygote is called codominance. As the

result the phenotype is different from both homozygous genotype.

Examples

- 14. Blood group is the good example codominance
- ABO blood groups are controlled by gene
 I. The gene (l) has three allele
 I^A, I^B, i, I^A, I^B produce slightly different
 form of sugar while i does not produce
 any kind of sugar.
- 16. I^A, I^B are dominant alleles where as i is recessive alleles
- 17. Since, there are three different allele, there are six different combination of these three alleles are possible and four phenotypes (A, B, AB and O)

Genetic Basis of Blood Groups in Human Population

- op mano			
Allele	Allele	Genotype	Blood
from	from	of	Types of
Parent	Parent	Offspring	Offspring
1	2		
IA	IA	I ^A I ^A	А
IA	IB	IAIB	AB
IA	i	I ^A i	А
IB	IA	I ^A I ^B	AB
IB	IB	I ^B I ^B	В
IB	i	I ^B i	В
i	i	ii	0

When I^A and I^B are present together they both express their own types of sugars this is because of co-dominance. ABO blood grouping also provides a good example of multiple alleles.

Here, you can see that there are more than two, *i.e.*, three alleles governing the same character. Since, in an individual only two alleles can be present multiple alleles can be found only when population studies care made. Dominance is not an autonomous features of a gene. It depends on much on the gene product

663 **(c)**

W Bateson (1905) explained the lack of independent assortment in sweet pea and **T H Morgan** (1910) in *Drosophila* due to linkage. When genes closely present adhere or link together in a group and transmitted as a single unit, the phenomenon is called linkage. It stops the process of independent assortment. Incomplete linkage is broken down due to the crossing over.

664 **(b)**

Mendel was successful in discovering the principles of inheritance as he did not encounter linkage between genes for the characters he considered. One of his principles-independent assortment is applicable only if the genes are located on different non-homologous chromosome pairs.

665 **(c)**

The polytene chromosomes radiate as 5 long and 1 short arm from a deeply staining and more or less amorphous structure called **chromocentre**. This chromocentre is formed by the fusion of centromeric region of all the chromosomes and in males, entire Y-chromosomes.

666 **(a)**

Polyploid An organism or its karyotype having more than two genomes called polyploid *Polyploidy is three types*

(i) **Autopolyploidy** It is the type of polyploidy in which there is the numerical increase of same genome. *e. g.*, autotriploid (AAA), autotetraploid (AAAA) *e. g.*, maize, rice, gram

(ii) **Allopolyploidy** It has developed through hybridization between two species followed by doubling of chromosomes, *e. g.*, AABB *e. g.*, wheat, cotton, Nicotiana tobacum.
Raphanobrassica and Triticale recently allopolyploids

(iii) **Autoallopolyploidy** It is a type of alloplyploidy in which one genome is in more than diploid state commonly autoallopolyploids are hexaploids (AAAABB) *e.g., Helianthus tuberosis*

667 **(b)**

Matation is more common when it is present in dominant condition. The reason is that the dominant mutant gene can express in both homozygous and heterozygous conditions.

668 **(d)**

Allelism refers to presence of alternative forms of a gene at a given locus. Alleles or allelomorphs are the two contrasting aspects of the same character present at a locus of homologous pair of chromosomes. Now-a –days, the same aspect in duplicate (TT or tt) of a character is also considered an allele.

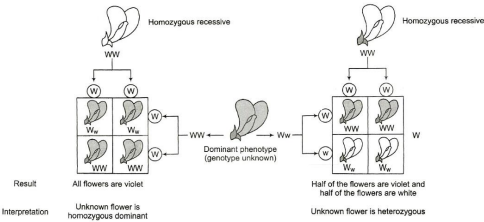
669 **(d)**

Rh factor was first reported by Landsteiner and Winer in rhesus monkey. When Rh⁺man marry with Rh⁻women the foetus will be Rh⁺. This cause the condition called haemolytic disease (erythroblastosis foetalis).

670 **(b)**

Genotype of F₁- plant

Crossing of F_1 (dominant phenotype) to any one of the parent called back cross and when phenotype of crossing parent is recessive than this is called test cross. The progeny of such cross can easily be analysed to predict the genotype of test organism



671 (a)

Chang in single base pair may not change in phenotype. The codon is triplet and degenerate.

672 (c)

Organisms with more than two sets of chromosomes are known as **polyploids**. It may be triploid with three sets of chromosomes (3*n*) or tetraploid with four sets of chromosomes (4*n*) and so on.

673 **(c)**

Due to absence of phenylalanine hydroxylase the phenylalanine changes in to phenyl pyruvic acid. Lack of this enzyme is due to autosomal recessive defective gene on chromosome number 12

674 (d)

All of the given disorders are genetically transferred.

675 (d)

Consanguineous mating, is the mating, which happens between the person's own relationship

676 **(d)**

When F_1 hybrid is intermediate between two parents, there is no perfect dominance of one character upon other this phenomenon is called **incomplete dominance** and their inheritance as blending inheritance. **Example** : 4 0' clock plant (*Mirabilis jalapa*).

678 **(a)**

After a mutation at a genetic locus, the character of an organism changes due to the change in **protein structure**.

679 **(b)**

Heterogametic.

XY and XY type sex determination seen in many insect and mammals including humans. Males have X and Y chromosome along with autosome and females have pair of 'X' chromosome along with autosome

Parents Phenotypes Male Female

(Genotypes	44A + XY	
44A + XX			
Gametes		22A + X 22A	+Y
22A + A2	22A + X		
		22A+ X	22A+X
Children	22A + X	44A + XX	44 A + XY
Female			
	22A + Y	44 A + XY	44 A + XY
Mala			

Male

Sex ratio Female : Male = 1 : 1

In plants The flowering plants are mostly bisexual and lack sex chromosomes. The unisexual flowering plants tent to have XX-XY type of sex

chromosomal mechanism for sex determination. The female plants are XX and male plants are XY.

XX and XO Type of Sex Determination

Found in insect like grasshopper, cockroaches and bugs. Males have only X sex-chromosome and autosomes, female have pair of X-chromosome and autosome

ParentsPhenotypesMaleFemaleGenotypesAA + XOAA + XXGametesA + X, A + OA + X, A + Y

F_1 -generation

	A + X	A + X	
A + X	AA + XO	AA + XO	
A + O	AA + XO	AA + XO	
	Genotypes		

XX-XO type of sex determination

In most of cases the female produce similar sex chromosome called homomorphic. In most of cases the male produce dissimilar sex chromosome called hetermorphic

680 (a)

Dihybrid cross is a cross involving two pairs of contrasting characters. A dihybrid test cross gives 1:1:1:1 ratio indicating that when F_1 hybrid is crossed with recessive parent the two pairs of factors segregate and assorting independently and produce four type of progenies.

682 **(d)**

In 1900 three workers independently rediscovered the principles of heredity already worked out by Mendel.

These workers were

(i) Hugo de Vries (Holland)

(ii) Carl Correns (Germany)

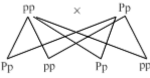
(iii) Erich von Tschermak (Austria)

683 **(d)**

The salivary gland chromosomes in the dipteran larvae have endoreduplicated chromosomes, which are useful in gene mapping.

684 **(b)**

Percentage of recessive phenotype = 50% (pp) percentage of dominant phenotype = 50% (Pp and PP)



685 **(c)**

A gene consist of a polynucleotide sequence that encodes a functional polypeptide or RNA

sequence

686 **(b)**

Test cross of dihybrid YyRr with double recessive yyrr gives four types of progeny 1 : 1 : 1 : 1 ratio due to presence of genes on separate chromosomes (unlinked genes)

687 **(d)**

In sickle-cell anaemia acidic amino acid (glutamic acid) is replaced by neutral amino acid (valine). It is caused because GUG (codes for glutamic acid) or placed by GAG (codes for valine)

688 **(d)**

The basic (*x*) numbers of chromosome of hexaploid wheat is 7 and haploid (*n*) number is 21.

689 **(b)**

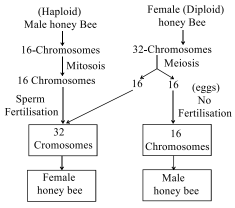
Persons who are colourblind cannot distinguish red and green colour. Colour due to absence of cone cells.

690 **(d)**

Haploid diploid mechanism of sex determination (haplodiploidy).

Hymenopterous insect such as bees, wasps ants show unique phenomena in which an unfertilized egg develops into male and female develops from fertilized egg.

In honeybee, the quality of food determines whether a diploid larva will become a fertile queen or a sterile worker female. A larva fed on royal jelly a secretion from the mouth of mussing workers grows into a queen, whereas a larva fed on pollen and nectar grows into a worker bee



691 **(a)**

A-Point; B-β-chain

692 **(d)**

Sickle cell anaemia is a genetic disorder reported from nigroes due to a molecular mutation of gene Hb^A on chromosome 11 (autosome), which produces the β -chain of haemoglobin. In sickle

cell anaemia, the sixth amino acid of haemoglobin (*i.e.*, glutamic acid) is replaced by valine.

693 **(a)**

In crossing between pure (homozygous) organisms for two contrasting characters, only one character of the pair appears in F_1 generation (hybrid), which is called **dominant**, while the other, which does not appear is called **recessive**.

694 **(a)**

Muller first gane the proof of mutagenic action of X-rays.

695 **(a)**

The diploid condition in which the alleles at a given locus are identical is called homozygous or pulls. In homozygous condition, organism have two similar genes or alleles for a particular character in homologous pair of chromosomes, *e. g.*, TT or tt.

Organisms containing two different alleles or individual containing both dominant and recessive genes of an allele pair, *e.g.*, Tt is known as heterozygous or hybrid

696 **(b)**

The tips of chromosomes are called **telomeres**. These show difference in structure and composition from rest of the chromosomes. Telomere has a unique property as it prevents the ends of chromosomes from sticking together alongwith it facilates attachment of chromosomes with nuclear envelope. Telomeres contain heterochromatin or repetitives DNA and bounded by specialized proteins.

697 **(b)**

Polytene chromosomes was reported by **Balbiani** (1881) from cells of salivary glands of insect larvae. Polytene chromosomes contain several dark stained regions called **bands** separated by lighter or less stained area called **interbands**. These dark bands are heterochromatin (genetically insert) area.

698 **(c)**

During meiosis or gametogenesis each chromosome/gene in allele has equal (50%) chances for separation. That is also called law of segregation

699 **(d)**

Pleiotropic gene (Gr. *Pleion* = *more*) is the gene

having the ability to show multiple effects. In*Drosophila*, a gene for white eye mutation is also responsible for depigmentation of body parts thus a gene controlling more than one phenotype is called as pleiotropic gene.

700 **(a)**

Hypertrichosis is the excessive growth of hair on body. It is an example of **holandric inheritance**. Genes responable for this are located on Y chromosomes only which are also known as holandric genes. Y-linked holandric genes are transmitted directly from father to son.

701 **(a)**

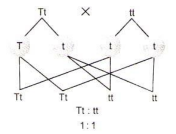
Ethyl methane is a mutagenic agent causing gene mutations.

702 **(d)**

Sickle-cell anaemia is caused due to base replacement leading to replacement of sixth amino acid in the β -chain. Glutamic acid is replaced by valine and causes sickle cell anaemia.

703 **(a)**

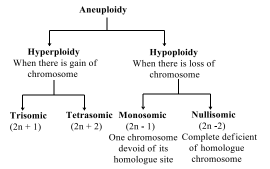
The cross between Tt and tt will show 1 : 1 ratio. Test cross is performed to know whether the parent is homozygous or heterozygous dominant.



704 **(a)**

Chromosomal disorders are caused due to excess, absence or abnormal arrangement of one or more chromosomes.

Sometimes the chromatids fail to segregate during cell division, resulting in gain or loss of a chromosome. This is called **aneuploidy** Aneuploidy is also called heteroploidy Aneuploidy is of two kinds (i) Hyperploidy (ii) Hypoploidy



705 **(a)**

Genes are segments of DNA. **Johanseen** (1909) described the gene as 'Unit of heredity' which can assigned to a particular character. **Morgan** and **Bridges** suggested that genes are located on chromosomes in linear fashion.

706 (a)

7 dominant traits, 7 recessive traits total 14 traits or 7 oppossing pairs of traits

Characters	Dominant	Recessive
	Traits	Traits
Seed shape	Round	Wrinkled
Seed colour	Yellow	Green
Flower	Violet	White
colour		
Pod shape	Full	Constricted
Pod colour	Green	Yellow
Flower	Axial	Terminal
position		
Stem	Tail	Dwarf
height		

707 (d)

Environmental Determination of Sex

In *Bonellia*, a marine worm, the swimming larva has no sex if it settles down alone, it develops into a large (2.5 cm) female. If it lands on or near an existing female, a chemical from female causes the larva to develop into a tiny (1.3 cm) male. In turtles, alligators, a temperature below 28°C produces more males, above 33°C produces more females and between 28° C – 33° C produces males and females in equal proportion

708 (d)

Dominant allele expressed in both homozygous and heterozygous condition.

or Principle of Dominance

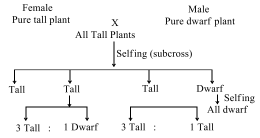
In heterozygous individuals or hybrids a character is represented by two contrasting factors called alleles or allelomorphs. Out of the two contrasting alleles. Only one is able to express

711 (b)

To know the genotype of dominant phenotype, we will cross that plant with the respective recessive phenotype. This is called test cross.

its effect in the individual. It is called dominant factor or dominant allele.

The other which does not show its effect in the heterozygous individual is called recessive factor or recessive allele. The cross between the pure tall and pure dwarf gives all progeny (F_1) tall.



The character shown by F_1 called dominant character

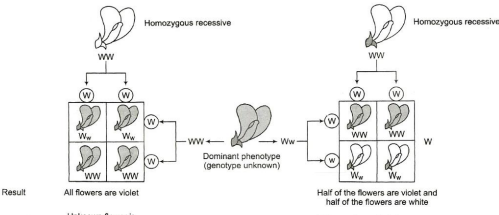
709 **(c)**

If the ratio between X-chromosomes and autosome is **0.5** then the individual will be **male** but if it is **1.0** then the individual will be **female**.

710 (d)

<u> </u>	
Column I	Column II
Metacentric	At the middle
Submetacentric	Slightly away
	from the middle
Acrocentric	Almost near the
	tip
Telocentric	At the tip

Crossing of F_1 (dominant phenotype) to any one of the parent called back cross and when phenotype of crossing parent is recessive than this is called test cross. The progeny of such cross can easily be analysed to predict the genotype of test organism



Interpretation Unknown flower is homozygous dominant

712 (c)

H J Muller was awarded **Nobel Prize** in 1946 for his discovery of the production of mutations by Xray irradiation.

713 **(b)**

If the father is colourblind (X^cY) and the mother is a carrier (X^cX), then their son will be normal(XY) and daughter will be colourblind (X^cX^c).

714 (d)

Based on the ratio of F_2 and F_3 generation, Mendel proposed that something was being stably passed down F_1 and F_2 -generation) unchanged, from parent to offspring through the gametes, over successive generations.

He called these things as factors. Now we call them genes. Genes therefore, are the units of inheritance required to express a particular trait

715 **(a)**

Extra –nuclear or extra-chromosomal or cytoplasmic or organellar inheritance is a consequence of presence of genes in mitochondrial and chloroplast DNA. Extrachromosomal units function either independently or in collaboration with nuclear genetic system.

717 (a)

720 **(b)**

Klinefelter's syndrome.

Disorders	Autosomal/Sex Linked	Symptoms	Effects
Down's syndrome	Autosomal aneuploidy (trisomy, +21)	Mongolian eyefold (epicanthus), open mouth, protruded tongue, projected	Retarded mental development IQ (below 40)

Colour blindness is a condition, in which, certain colours can not be distinguished due to a lack of one or more colour-absorbing pigments in the cone cells of the retina. Colour blindness is also occur due to recessive sex linked genes.

718 **(a)**

Unknown flower is heterozygous

In F_2 -generation, quantitative inheritance 1:4:6: 4 : 1 is obtained in a dihybrid cross instead of 9 : 3:3:1.

719 (a)

The inheritance due to the genes found in cytoplasm (mitochondria and chloroplast) is called **cytoplasmic inheritance** or **non-Mendelian inheritance**. The leaves of *Mirabilis jalapa* may be green, white or variegated. This is due to cytoplasmic inheritance.

Turner's syndrome	Sex chromosomal monosomy 44 + XO	lower lip, many loops on finger tip, palm crease Short stature females (<5'), webbed neck, body hair absent menstrual cycle absent. Sparse pubic hair, underdeveloped breasts narrow lips puffy fingers	Sterile hearing problem
Klinefelter's syndrome	Sex chromosomal aneuploidy (tri/tetrasomy of X chromosome) 44+XXY 44+XXY	The males are tall with long legs, testes small, sparse body hair, Barr body present, breast enlargement	Gynaecomast ia azospermia sterile

Some Examples of Aneuploidy

(i) Down's syndrome-21 trisomy

Symptoms

- (a) Short statured with small round head
- (b) Partially open mouth with protruding furrowed tongue
- (c) Palm is broad with characteristic palm crease
- (d) Slow mental development

(ii) Turner's syndrome

Cause Absence of one of the X-chromosomes, resulting in the karyotype 44+X0

Symptoms

- (a) Sterile female with rudimentary ovaries
- (b) Shield-shaped thorax
- (c) Webbed neck
- (d) Poor development of breasts
- (e) Short stature, small uterus, puffy fingers

(iii) Klinefelter's syndrome

Cause Presence of an additional copy of X-chromosome resulting in the karyotype 44+XXY

Symptoms

(a) Sex of the individual is masculine but possess feminine characters

- (b) Gynaecomastia, *i.e.*, development of breasts
- (c) Poor beard growth and often sterile
- (d) Feminine pitched voice

721 **(b)**

ABO blood group is an example of codominance because both allele express themself and is govern by multiple allele means it is controlled by more than two allele. ABO blood group system or A, AB, B and O blood group of human cannot judge by using physically the human individual

722 (c)

Brachydactyly is a disease characterized by small

sized finger and is due to dominant gene on the sex chromosome.

723 **(b)**

On mating female *Drosophila* to diploid males, their progeny consist of following types :

18. AAAXXX – Triploid female

19. AAXX	_	Diploid female
20. AXXY	_	Diploid female
21. AAAXX	_	Intersex
22. AAAXXY	_	Intersex
23. AAXY	_	Normal male
24. AAXXX	_	Superfemale
25. AAAXY	_	Super male or metamale.

724 **(c)**

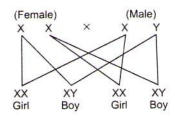
Human blood (ABO) is example of multiple alleles in which three alleles I^A, I^B, I^O governing the same trait

725 (a)

In monohybrid cross only one contrasting characters is taken like tallness and shortness, green-yellow

726 **(d)**

The X-chromosomes of females are represented in both the sexes as



Therefore, they are represented in both male and female progeny.

727 **(b)**

In the given diagram there is insertion of T in the given segment of gene so diagram depict insertion type of mutation

728 **(b)**

A-Meiosis; B-Segregate

729 **(b)**

Mendelian principles are based on sexual reproduction. Therefore, they are not applicable in case of asexually reproducing forms.

730 **(c)**

In most of cases the female produce similar sex chromosome called homomorphic. In most of

cases the male produce dissimilar sex chromosome called hetermorphic

731 **(b)**

In starch synthesis gene following condition to seen

BB - rounded (due to more starch synthesis)
bb - wrinkled (due to less starch synthesis)
Bb - in between rounded of wrinkled size. It
produce starch of intermediate quantity between
BB to bb homozygous condition. So, it is
incomplete dominance.

Therefore, dominance is not an autonomous feature of a gene or the product that it has information for. It depends as much on the gene product and the production of a particular phenotype from this product as it does on the particular phenotype that we choose to examine, in case more than one phenotype is influenced by the same gene

732 **(b)**

Sickle cell anaemia is a genetic disease reported from Negroes due to molecular mutation (missense mutation) of gene Hb^Aon chromosome 11, which produces the β -chain of mature haemoglobin. The mutated gene HB^s produces sickle cell haemoglobin. The sixth amino acid, in β -chain of normal haemoglobin is glutamic acid which is replaced by valine in sickle cell haemoglobin. The sequence of DNA in mutated β globin chain is CAC/GAG which on transcription results in codon GUG which produces valine in place of glutamic acid.