### Assessment Schedule - 2023

# **Biology: Demonstrate understanding of gene expression (91159)**

### **Assessment Criteria**

Achievement	Achievement with Merit	Achievement with Excellence
<ul> <li>Demonstrate understanding involves:</li> <li>defining, using annotated diagrams or models to explain, and giving characteristics of, or an account of, gene expression.</li> </ul>	Demonstrate in-depth understanding involves:  providing a reason as to how or why biological ideas and processes affect gene expression.	Demonstrate comprehensive understanding involves:  • linking biological ideas and processes about gene expression; explanations may involve justifying, relating, evaluating, comparing and contrasting, or analysing.

### **Cut Scores**

Not Achieved			Achievement with Excellence	
0 – 7	8 – 13	14 – 18	19 – 24	

### **Evidence**

Questio	n One					
Expected Coverage				Achievement	Achievement with Merit	Achieve. with Excellence
(a):  DNA Template Middle section of gene  mRNA Amino acid sequence	Normal gene sequence Point mutation 1 Point mutation 2 and m ONE		THREE boxes are correct and must include at least ONE mRNA and amino acid sequence/box.	All mRNA and amino acid sequences are correct.		
Both mut Substituti Point mut Point mut ATA char there are a usually has start / stop same ami be fully fi Point mut to STOP incorrectly	s for the same amino acide tation 2 changes A to T a station 1 is a silent mutation and to ATG, both triplet 20 amino acids and 64 triplet ave more than one codon p codons. Consequently, to acids, meaning it will functional.  Itation 2 will have an effect will result in the amino a stay/having the wrong shapits function will be comp	utations.  base is swapped in the I on where the change in the I on because of degeneracy is code for the same aminoplets / codons. This mean / triplet that codes for it. I the chain will be the same fold correctly / be the same ct on the final protein. The I on the	DNA base sequence.  the base sequences A to G  don.  of the code; even though of acid. This is because is that an amino acid will. There is no change to the elength and have the elength and so will still the change in amino acid ter and folding is shape is very different,	<ul> <li>Describes:</li> <li>a mutation</li> <li>point mutation 1 is a silent / same-sense mutation</li> <li>point mutation 2 codes for STOP / is a nonsense mutation.</li> <li>Identifies:</li> <li>both mutations as a substitution mutation OR describes a substitution mutation mutation</li> <li>that point mutation 1 will not change the final protein OR point mutation 2 will change the final protein</li> <li>that more than one codon can code for the same amino acid or mutation 1, due to redundancy / degeneracy.</li> </ul>	<ul> <li>Explains:</li> <li>point mutation 1 does not change the amino acid sequence therefore the protein can still fold into/have the correct shape/not affect function</li> <li>point mutation 2 does change the amino acid to a stop codon causing the amino acid sequence to be shorter (than normal) and fold incorrectly/protein does not function correctly</li> <li>silent mutations / degeneracy of the code (i.e. more codons than amino acids / redundancy in the code – 64 and 20)</li> <li>point mutation 1 will be the same length of gene / same number of amino acids because the start or stop</li> </ul>	Discusses, demonstrating comprehensive understanding of:  • why point mutation 1 will not affect the position of the start / stop codons (i.e. does not cause a change in length of amino acid sequence, and it will not change the amino acid, will not change folding / shape, will not impact protein folding / shape AND function; and makes links to silent mutations and degeneracy of the code  • how point mutation 2 causes a premature stop codon, (ATT codes for a stop), therefore changes the length of amino acids (shorter), affects folding / protein shape, and negatively affects function.

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Expected Coverage	Achievement	Achievement with Merit	Achieve. with Excellence
		codons remain unchanged / unaffected	
		• in both point mutations 1 and 2, the start codons are not affected; however, in point mutation 2, the stop codon is affected.	

Not Achieved		Achievement		Achievement with Merit		Achievement with Excellence	
N1	N2	А3	A4	M5	М6	<b>E</b> 7	E8
ONE evidence point at Achievement.	TWO evidence points at Achievement.	THREE evidence points at Achievement.	FOUR evidence points at Achievement.	THREE evidence points at Merit.	FOUR evidence points at Merit.	ONE evidence point at Excellence.	BOTH evidence points at Excellence.

**N0** = No response; no relevant evidence.

### **Question Two Expected Coverage Achievement with Excellence** Achievement **Achievement with Merit** Describes: An environmental factor (non-mutagenic) is a substance / factor **Explains: Discusses**, demonstrating that affects gene expression/phenotype but does not change the comprehensive understanding of • an environmental factor that • a metabolic pathway (as for A4 DNA base sequence / genotype. how: changes the phenotype but not the and includes the link to genes A mutagen is a substance that changes the DNA base sequence coding for enzymes catalysing only • UV light causes more enzyme 2 to genotype one reaction in the pathway) be made, so more melanin (i.e. permanently / genotype. • a valid example of an change in phenotype): therefore environmental factor, e.g. diet / An enzyme is a biological catalyst that speeds up a reaction. changes the phenotype but the temperature / rainfall (**not** UV A metabolic pathway is a series of enzyme-controlled, chemical explains this metabolic pathway in DNA remains unchanged / no radiation or sunlight) reactions where the product of one reaction is the substrate for the full (all genes, enzymes, substrates, change to genotype and products used in context next. • a mutagen • UV light causes the genotype to correctly) to show how melanin is In a metabolic pathway, one gene encodes one enzyme. Each enzyme • a metabolic pathway OR that an change / mutation in gene 2 produced can only catalyse one specific reaction due to its unique shape. enzyme is a biological catalyst dysfunctional / no enzyme 2 made • how UV causes increased levels / In a normal/functioning metabolic pathway, melanin is produced • that the genes in the metabolic so no melanin is made: therefore rates of enzyme 2 production, when gene 1 codes for enzyme 1, which converts phenylalanine pathway code for the enzymes the phenotype is changed because therefore more melanin is into tyrosine. Tyrosine is then the substrate in the next reaction • mutated DNA in people with of the change in genotype. produced where gene 2 codes for enzyme 2, which converts tyrosine into albinism means they are not • UV affects the genotype (mutated melanin. • how genes (genotype) + protected from UV light gene 2) so no melanin can be environment factors (UV light) Some UV light causes gene 2 to produce more of enzyme 2, which made; therefore, greater likelihood contribute to the phenotype then expresses more melanin in the skin. Therefore, environmental States that: of cancer when the underlying skin (melanin amount) factors (low UV light) + genotype = phenotype. Environmental factors • albinism is due to no melanin, is not protected by melanin and interact with the genes / genotype to express the phenotype type. • that the UV light enables the caused by build of tyrosine / less likelihood in individuals that genotypic potential to be reached Melanin is not produced in people with albinism if either or both phenylalanine (genes mutated) can produce melanin. because correct / enough melanin gene 1 and gene 2 has a mutation. A mutation to either gene would • phenotype / melanin production is made not produce the enzyme which cannot act on the phenylalanine and (not just skin colour) is a product • how / why melanin is not produced tyrosine substrates and convert into melanin. of both the environment and in people with albinism, using the People with albinism are more likely to get skin cancer because genotype metabolic pathway (gene 2 they do not have melanin pigmentation to protect the DNA in their OR mutated, enzyme 2 dysfunctional, skin cells. $G + E \rightarrow P$ (equation or in words). tyrosine not converted so no UV light can be both a non-mutagenic environmental factor and a melanin made) mutagen because some UV light causes gene 2 to code for more of • why people with albinism are more enzyme 2, which in turn converts more tyrosine into melanin, likely to get skin cancer, i.e. expressing increased pigment/skin colour. This allows the full because there is no melanin genetic potential of gene 2 to be expressed without changing DNA protecting the underlying skin / genotype. layers However, UV light can also cause DNA to permanently change • that excess UV can affect DNA and affect folding/shape/functioning of proteins produced and can and increase the rate of mutation, cause skin cancer. This changes the genotype of the individual

which will also change the phenotype.

e.g. linked to cancer.

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Not Achieved		Not Achieved Achievement		Achievement with Merit		Achievement with Excellence	
N1	N2	А3	<b>A</b> 4	M5	M6	<b>E</b> 7	E8
ONE evidence point at Achievement.	TWO evidence points at Achievement.	THREE evidence points at Achievement.	FOUR evidence points at Achievement.	THREE evidence points at Merit.	FOUR evidence points at Merit.	ONE evidence point at Excellence.	TWO evidence points at Excellence.

**N0** = No response; no relevant evidence.

### **Question Three**

Expected Coverage	Achievement	Achievement with Merit	Achievement with Excellence	
(a): 5'end 3'end  B  B  B  C  C  C  C  C  C  C  C  C  C	Labelled drawing showing correct anti-parallel DNA section, sugars, phosphates, and bases (one minor error, e.g. both strands same direction / single bonds between bases).	Labelled drawing showing correct anti-parallel DNA section, sugars, phosphates, and bases (no errors); must have double bonds between AT and triple bonds between GC.		
(b): Transcription: First step of protein synthesis. Translation: Second step of protein synthesis.	Names: • transcription AND translation in the correct order.			
(c):  Triplet: three consecutive nucleotide bases on the DNA strand.  Codon: the sequence of three consecutive nucleotides on the mRNA strand (that is complementary to a triplet).  Anticodon: three consecutive bases on a tRNA molecule (that is complementary to mRNA codon).  The base-pairing rule states that A can only bind with T (on the DNA) or U (on the RNA), T (DNA) or U (RNA) can only bind with A, and C can only bind with G.  The DNA holds the genetic code for the protein. The triplets on the DNA and the codons on the mRNA both code for amino acids.  In transcription, the triplets on the DNA need to be made into codons on the mRNA so that the code for the protein/gene can be taken out of the nucleus/into the cytoplasm/ribosome for translation. This must be done accurately so the mRNA carries the correct code/information for the sequence of amino acids for the protein.	Describes:  • a triplet  • a codon  • an anticodon  • the base-pairing rule (can come from diagram).  States that:  • protein is made of amino acids  • inaccurate transcription / translation leads to incorrect AA sequence / shape protein.	<ul> <li>Explains:</li> <li>C binds with G and T / U with A due to the number (2 and the 3) of hydrogen bonds</li> <li>C binds with G and A binds with T / U because of the size of the bases; double ring purines can only bind with single ring pyrimidines</li> <li>the triplets / codons determine the specific amino acids in the polypeptide chain</li> <li>triplets must be transcribed into codons in order for the code to leave the nucleus / go to ribosome</li> <li>the complementary nature of DNA and mRNA or of mRNA and tRNA</li> <li>accuracy is needed so the protein is folded correctly / has correct shape / functions correctly.</li> </ul>	Discusses, demonstrating comprehensive understanding that:  • links the number of hydrogen bonds that can form between AT and GC nucleotide bases with the reasoning that only certain combinations are possible; therefore, the complementary base-pairing rule ensures accurate transcription / translation of the protein  • links the size (either by names or number of loops) of the nucleotide bases with accurate transcription / translation because of complementary base pairing, e.g. given that single ring / small bases can only fit with double ring / larger bases, the base pairing rule ensures accurate transcription / translation  • links the accurate transfer of the	

Expected Coverage	Achievement	Achievement with Merit	Achievement with Excellence
In the ribosome, the mRNA is read in order to make the protein. Each codon codes for one amino acid, and the tRNA brings in the amino acids to the ribosome. Each tRNA carries one specific amino acid and has an anti-codon that complements the codon for this amino acid. In the ribosome, the codons are bonded to their complementary anti-codons on the tRNA with the correct amino acid. This ensures that the tRNA will put its specific amino acid into the correct sequence / order in the protein.			amino acids, i.e. correct protein synthesis for protein shape / function.
This accuracy is important in transcription to ensure that each triplet on the DNA is accurately transcribed to the correct codon so that the mRNA carries the correct code for the protein/information to make the protein. In addition, accuracy is important in translation so the tRNA puts the correct amino acid into the correct order in the polypeptide chain. The order of the amino acids determines the shape / folding of the protein/polypeptide, and the shape of the protein needs to be correct for the protein to function.			
Accurate transcription / translation is ensured by the base-pairing rule. The size of the bases determines which bases can bind together. A large (double ring / purines) base can complement only a small (single ring / pyrimidines) base. For example, A can bind only with T because A is large and T is small. Adenine and guanine are both large so can't fit together in either DNA or RNA. In addition, the placement of hydrogen bonds prevents other bonding combinations. A and T form the same number of hydrogen bonds together, and C and G form the same number of hydrogen bonds together. Adenine can't bind with cytosine because they have different numbers of hydrogen bonds and can't chemically fit together.			

Not Achieved		Achiev	/ement	Achievement with Merit		Achievement with Excellence	
N1	N2	А3	<b>A</b> 4	M5	M6	<b>E</b> 7	E8
ONE evidence point at Achievement.	TWO evidence points at Achievement.	THREE evidence points at Achievement.	FOUR evidence points at Achievement.	THREE evidence points at Merit.	FOUR evidence points at Merit.	ONE evidence point at Excellence.	TWO evidence points at Excellence.