Assessment Schedule – 2014 Biology: Demonstrate understanding of gene expression (91159) Assessment Criteria

Achievement	Achievement with Merit	Achievement with Excellence
<i>Demonstrate understanding</i> involves defining, using annotated diagrams or models to explain, and giving characteristics of, or an account of, gene expression.	<i>Demonstrate in-depth understanding</i> 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. The explanation may involve justifying, relating, evaluating, comparing and contrasting, or analysing.

Evidence Statement

Q	Expected Coverage	Achievement	Merit	Excellence
ONE (a) (b)	 CUAGUA The purpose of transcription is described: mRNA transcribes the code for a polypeptide from the DNA. The purpose of transcription is explained: mRNA transcribes the code for a polypeptide from the DNA in the nucleus and carries it to the ribosomes / cytoplasm. So that the original DNA does not get damaged leaving the nucleus. The purpose of translation is described: to use mRNA to make a polypeptide / protein. The purpose of translation is explained: to use mRNA to make a polypeptide / protein. So that the protein can be used for cellular functions (or named example given e.g. to make an enzyme). The process of transcription is described: e.g. DNA unwinds and a single mRNA strand is made using U instead of T. The mRNA strand leaves the nucleus through a nuclear pore. The process of transcription is explained by giving a substantially correct sequence of steps: an enzyme (RNA polymerase) separates / unzips the DNA strand, exposing the gene / bases / nucleotides. Free nucleotides are match to the exposed bases on the template strand using the base pairing rule, A-U and G-C. Transcription is complete when (RNA polymerase reaches the terminator sequence) mRNA detaches and moves out of the nucleus into the cytoplasm and attaches to a ribosome in preparation for translation. 	 Correctly transcribes the mRNA strand. Describes the purpose of transcription. Describes the process of transcription Describes the purpose of translation Describes the process of translation Describes one similarity. Describes one difference. 	 Explains the purpose of transcription. Explains the purpose of translation. Explains the process of transcription by giving a substantially correct sequence of steps. Explains the process of translation by giving a substantially correct sequence of steps. 	 Discusses the process of transcription and translation, with links to the purposes of protein synthesis. Compares and contrasts similarities and differences between transcription and translation

bases on the mRNA codes for an amino acid . tRNA carries the amino acid to the ribosome and drops it off		
The process of translation is explained by giving a substantially correct sequence of steps: ribosomes move along the mRNA from the start codon until the stop		
codons is reached. Each sequence of 3 bases (codon) on the mRNA is read by the ribosome and matched to the complementary unpaired three base sequence		
(anticodon) on the tRNA. The specific amino acid attached to the tRNA is then added (peptide bond forms)		
Similarities between transcription and translation include:		
 both use complementary base pairing both have mRNA involved in the process both have a start and stop sequence 		
 both nave a start and stop sequence both are controlled by enzymes code on both read in sets off three bases 		
Differences between transcription and translation include:		
• transcription occurs in the nucleus and requires DNA. Translation occurs in the cytoplasm on a ribosome and involves tRNA and amino acids (<i>explanation for</i> <i>compare and contrast differences</i>)		
 transcription makes mRNA, translation reads mRNA / makes proteins 		
 transcription uses DNA as a template, translation uses mRNA as a template 		
• transcription involves the pairing of DNA and free RNA nucleotides, whereas translation involves pairing anticodon bases of tRNA and codon mRNA (<i>explanation for compare and contrast differences</i>).		
Compares and contrasts similarities and differences between transcription and translation. Eg both transcription and translation involve the use of templates but they are different. Transcription uses a DNA template whereas translation a mRNA template.		
• Transcription takes place in the nucleus and involves DNA and free nucleotides to make mRNA whereas translation occurs in the cytoplasm using ribosomes, tRNA and amino acids to read mRNA. So both processes involve mRNA.		
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NØ	N1	N2	A3	A4	M5	M6	E7	E8
No response; no relevant evidence.	Describes any ONE statement from Achievement.	Describes any TWO statements from Achievement.	Describes any THREE statements from Achievement.	Describes any FOUR statements from Achievement.	Explains any TWO statements from Merit.	Explains any THREE statements from Merit.	Discusses EITHER criterion for Excellence.	Discusses BOTH criteria for Excellence.

Q	Exp	ected Covera	ge		Achievem	ent	Merit		Excelle	nce
TWO	A m cont Genu enzy Genu the r Expl expr serie react enzy path mela diffe mela meta Expl the r by a code step. the e sens alwa Expl ches enzy mela the r by a code step. the e sens alwa Expl the r by a code step. the e sens alwa Expl ches enzy mela temp Expl ches enzy non- mela whit to f f e sens alwa Expl ches enzy mola temp Expl ches enzy mola temp Expl ches enzy non- mela whit to f f e sens alwa f e f e sens f e f e f e f e f e f e f e f e f e f	etabolic pathw rolled chemica e A codes for me converts t e B codes for a nelanin in the lains the metal ession of coat s of enzyme of tions. As enzy me is needed way. In the fir way, the enzyme is needed way. In the fir way, the enzyme, min, is needed abolic pathway lains the tyrosin titve, that step type occur. lains the darke emities: if gene tryrosinase functional at min is not pro- e. scusses how t rosinase enzyme siamese cats: ne abnormal for ork as a cataly mperatures. The natures (loses gment is not pro- coduction of min dy are coloured the step and the step antains its sha e cooler extreme breakdown of oduction of min dy are coloured	vay is a series al reactions . the tyrosinase yrosine into m an enzyme tha coat colour. polic pathway colour in Siar ontrolled cher mes are specif for each step i st step of the r me tryosinase substrate tyros for the express in the second 7. inase mutation way is indirect DNA as that ge ne needed to c the DNA muta- tase is temperatu- be produced a e Siamese cats colour of the A mutates so t is heat sensiti- the warmer ch- duced and the he temperature in the pathwas colour of the A mutates so t is heat sensiti- the warmer ch- duced and the he temperature is is because shape) and as roduced. The of pe and its acti- nities so is ablo f tyrosine witt- elanin- these ped.	of enzyme enzyme. This helanin. It expresses for the mese cats as a mical fic, a different in a metabolic produces in. A sion of the step of the at Each step in thy controlled one has the atalyse that ates so that ature by may not e Sianese cats that the re sensitive, t the cooler s extremities. Siamese cats that the re sensitive, t the cooler s extremities. Siamese cats that the re sensitive, t the cooler s extremities. Siamese cats that the re sensitive t the cooler s extremities. Siamese cats that the re sensitive a cat body the enzyme a result enzyme ive site only at le to catalyse h subsequent parts of the	 Describent metabol pathway Describent A. Describent Cooler temperate produce Describent Describent metabol produce Describent met	es a ic es gene es gene B. es that tures pigment. es that tures do luce	 Explains the metabolic pathway for expression o coat colour is Siamese cats Explains the effect of the tyrosinase mutation in t metabolic pathway for expression o coat colour is Siamese cats Explains the darker colour Siamese cat? extremities. Explains the white colour Siamese cat? chest area. 	the f n	 Discu geness contra metal for m how t envira influe colou cats. Discu tempo sensit enzyr the co Siama metal 	asses how and enzymes of the polic pathway elanin and the he comment cts to ence the coat r in Siamese asses how the erature ive tyrosinase me produces pat colour in ese cats in the polic pathway.
NØ	ð	N1	N2	A3	A4	M5	M6		E7	E8
No respon relevant evidence.	nse; no	Describes any ONE statement from Achievement.	Describes any TWO statements from Achievement.	Describes any THREE statements from Achievement.	Describes any FOUR statements from Achievement.	Explains any TWO statements from Merit.	Explains any THREE statements from Merit.	Discu ONE for E	usses criterion xcellence.	Discusses BOTH criteria for Excellence.

Q	Expected Coverage	Achievement	Merit	Excellence
THREE (a)	Normal DNA TAC CAC GTG GAC TGA GGA CTC AAC Normal amino acids Met Val His Leu Thr Pro Glu Leu	 Correct DNA sequence. 6 / 8 correct amino acid sequence. 	 All correct normal amino acid sequence. AND mutated amino acid 	Discusses how the insertion mutation affects the bases sequence and final protein compared to a substitution
(b)	Mutated DNA TAC CAC GTG GAC TGA GGA CAC AAC Mutated mRNA AUG GUG CAC CUG ACU CCU GUG UUG Mutated Amino Acid Met Val His Leu Thr Pro Val Leu	 Correct mutated DNA sequence. Correct mutated mRNA sequence. Correct mutated amino acid 	 sequence produces Val instead of Glu. Explains how a substitution mutation affects the sequence of bases. 	 Degeneracy of the code is unable to buffer the effect of the mutation as every codon and thus every amino acid from the
(c)	When A is substituted into the DNA sequence instead of T, it causes a totally new amino acid to be made and changes the function of the final protein. When A is substituted into the DNA sequence instead of T, it still has the right number of bases to produce a final protein. However, a new amino acid is included, and this will affect final protein shape and functioning.	 sequence. Required protein will not be made. 	 Explains how a substitution mutation affects the final protein. Explains how an insertion mutation into sequence affects bases. 	 changes. Therefore having a more profound effect on the organism. OR Relates degeneracy to the sickle cell example
	Substitution mutation involves the exchange of one base for another, hence a different codon may code for a different amino acid; final protein is still made, but may function incorrectly. If an insertion occurred, this would cause all bases to move along one during translation, causing totally new amino acids to be formed from the mutation onwards. This is called a frame shift and because the degeneracy of the code is unable to buffer this mutation, because more than one codon has changed, this will significantly affect the functioning of the protein. Because the final amino acid chain would not be able to fold into a shape required for functioning.		 Explains how an insertion mutation affects final protein. Explains degeneracy of the genetic code. 	

NØ	N1	N2	A3	A4	M5	M6	E7	E8
No response; no relevant evidence.	Describes any ONE statement from Achievement.	Describes any TWO statements from Achievement.	Describes any THREE statements from Achievement.	Describes any FOUR statements from Achievement.	Explains any THREE statements from Merit.	Explains any FOUR statements from Merit.	Discusses FIRST statement for Excellence.	Discusses FIRST and ONE OTHER statement for Excellence.

Cut Scores

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