**Q1.**The diagram below represents one process that occurs during protein synthesis.

 

(a)     Name the process shown.

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**(1)**

(b)     Identify the molecule labelled **Q**.

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**(1)**

(c)     In the diagram above, the first codon is AUG. Give the base sequence of:

the complementary DNA base sequence .....................................................

the missing anticodon ...................................................................................

**(2)**

The table below shows the base triplets that code for two amino acids.

|  |  |  |
| --- | --- | --- |
|   | **Amino acid** | **Encoding base triplet** |
|   | Aspartic acid | GAC, GAU |
|   | Proline | CCA, CCG, CCC, CCU |

(d)     Aspartic acid and proline are both amino acids. Describe how two amino acids differ from one another. You may use a diagram to help your description.

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**(1)**

(e)     Deletion of the sixth base (G) in the sequence shown in the diagram above would change the nature of the protein produced but substitution of the same base would not. Use the information in the table and your own knowledge to explain why.

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**(Extra space)** ................................................................................................

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**(3)**

**(Total 8 marks)**

**Q2.**CREB is a transcription factor in the mitochondria of neurones.

(a)     What is a **transcription factor**?

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**(2)**

(b)     CREB leads to the formation of a protein that removes electrons and protons from reduced NAD in the mitochondrion.

Huntington’s disease (HD) causes the death of neurones. People with HD produce a substance called huntingtin. Some scientists have suggested that binding of huntingtin to CREB may lead to the death of neurones.

Suggest how binding of huntingtin to CREB may lead to the death of neurones.

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*(Extra space)* .................................................................................................

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**(3)**

(c)     CREB is a protein synthesised in the cytoplasm of neurones. Transport of CREB from the cytoplasm into the matrix of a mitochondrion requires two carrier proteins.

Use your knowledge of the structure of a mitochondrion to explain why transport of CREB requires **two** carrier proteins.

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**(2)**

**(Total 7 marks)**

**Q3.**          The diagram shows part of the metabolic pathway involved in the clotting of blood in response to an injury.



Haemophilia is a condition in which blood fails to clot. This is usually because of a mutant allele of the gene for Factor VIII.

(a)     Explain how mutation could lead to faulty Factor VIII.

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**(2)**

(b)     Use information in the diagram to explain how faulty Factor VIII causes haemophilia.

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**(2)**

(c)     A boy had haemophilia caused by faulty Factor IX. When his blood was mixed with blood from a haemophiliac with faulty Factor VIII, the mixture clotted. Suggest an explanation for clotting of the mixture.

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**(2)**

**(Total 6 marks)**

**Q4.**          (a)     CFTR is a transmembrane regulator protein.  Its molecules have 1480 amino acids. People with cystic fibrosis produce defective CFTR protein which is missing one amino acid from its structure.

(i)      What is the minimum number of bases on DNA which would code for the normal CFTR protein? Explain your answer.

Number of bases ...............................

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**(2)**

(ii)     Which type of gene mutation produced the cystic fibrosis allele?
Explain your answer.

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**(2)**

(b)     The diagram shows part of the process of making normal and defective CFTR in a cell. A normal CFTR protein molecule has sugar molecules attached to it which make it functional.



Describe how the information on mRNA is translated into CFTR at the ribosome.

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**(4)**

**(Total 8 marks)**

**Q5.**          (a)     Complete the table to show **two** differences between the structure of DNA and RNA.

|  |  |
| --- | --- |
| **DNA** | **RNA** |
|  |  |
|  |  |

**(2)**

(b)     Explain how a gene codes for a protein.

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**(2)**

(c)     What are homologous chromosomes?

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**(2)**

**(Total 6 marks)**

**Q6.**          (a)     Complete the table to give **two** differences between DNA and RNA.

|  |  |  |
| --- | --- | --- |
| **Difference** | **DNA** | **RNA** |
| 1 |   |   |
| 2 |   |   |

**(2)**

(b)     Describe the part played by RNA in protein synthesis.

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*(Extra space)  ..............................................................................................*

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**(3)**

**(Total 5 marks)**

**Q7.**A species of flowering plant can have white, red or purple flowers. The colour of the flowers is controlled by two genes. Each gene is found on a different chromosome, and is responsible for one step in a biosynthetic pathway. The biosynthetic pathway is



Gene 1 has the dominant allele **A** and the recessive allele **a**. Gene 2 has the dominant allele **B** and the recessive allele **b**. In both cases, the dominant allele needs to be present for the production of the associated enzyme.

(a)     Explain how the two genes are involved in producing white, red or purple flowers.

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**(6)**

(b)     (i)      A homozygous red-flowered plant was crossed with a homozygous white-flowered plant. All the flowers of the offspring were purple. What was the genotype of

the red-flowered parent;

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the white-flowered parent?

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**(2)**

(ii)     The purple-flowered offspring were crossed. What phenotypic ratio would you expect in the next generation? Use a genetic diagram to explain your answer.

**(4)**

(c)     (i)      Genetically, there are different types of white-flowered plants of this species. Give their different genotypes.

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**(1)**

(ii)     You have samples of fresh petals from the two homozygous types of white flowers, and a pure sample of the red pigment, **K**. Explain, in outline, how you might distinguish the two types of petal from each other.

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**(2)**

**(Total 15 marks)**

**Q8.**          This question should be answered in continuous prose.
Quality of Written Communication will be assessed in the answer.

(i)      Starting with mRNA, describe how the process of translation leads to the production of a polypeptide.

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**(4)**

(ii)      Normal tomato plants have an enzyme that softens tomatoes as they ripen. Genetically engineered tomatoes ripen and soften more slowly. A gene was inserted which reduces the amount of softening enzyme produced.

The diagram shows matching parts of the base sequences for the mRNA produced by the gene for the softening enzyme and that produced by the inserted gene.

Softening gene mRNA                 …AAUCGGAAU…

Inserted gene mRNA                   …UUAGCCUUA…

Suggest how the inserted gene reduces the production of the softening enzyme.

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**(2)**

**(Total 6 marks)**

**Q9.**          (a)     Changes to the protein coat of the influenza virus cause antigenic variability. Explain how antigenic variability has caused some people to become infected more than once with influenza viruses.

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**(2)**

(b)     The drawings show the changes in a B lymphocyte after stimulation by specific antigens.



B lymphocyte before stimulation        B lymphocyte after stimulation

(i)      Describe the role of macrophages in stimulating B lymphocytes.

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**(1)**

**S**       (ii)     Explain how the changes shown in the drawings are related to the function of B lymphocytes.

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**(4)**

**(Total 7 marks)**

**Q10.**          Mitochondria contain the genes needed for the synthesis of the enzymes involved in the electron transport chain. One of these enzymes is cytochrome oxidase. If a mutation occurs during replication of the mitochondrial genes, functional cytochrome oxidase may not be produced.

**S**       Explain why mutation of a mitochondrial gene might result in no functional cytochrome oxidase being produced.

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**(Total 5 marks)**

**Q11.**          Lysozyme is an enzyme consisting of a single polypeptide chain of 129 amino acids.

(a)     What is the minimum number of nucleotide bases needed to code for this enzyme?

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**(1)**

(b)     The diagram shows the sequence of bases in a section of the mRNA strand used to synthesise this enzyme.

G   G   U   C   U   U   U   C   U   U   A   U   G   G   U   A   G   A   U   A   U

(i)      Give the DNA sequence which would be complementary to the first four bases in this section of mRNA.

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**(1)**

(ii)     How many different types of tRNA molecule would attach to the section of mRNA shown in the diagram?

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**(1)**

(c)     Give **two** factors which might increase the frequency at which a mutation in DNA occurs.

1 ...................................................................................................................

2 ...................................................................................................................

**(2)**

(d)     Two single base mutations occurred in the DNA coding for this section of mRNA. These mutations caused an alteration in the sequence of amino acids in the enzyme. The diagram shows the original and altered sequences of amino acids.

|  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- |
| Original amino acidsequence | Gly | Leu | Ser | Tyr | Gly | Arg | Tyr |
| Original mRNAbase sequence | GGU | CUU | UCU | UAU | GGU | AGA | UAU |

|  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- |
| Altered amino acidsequence | Gly | Leu | Tyr | Leu | Trp | Arg | Tyr |
| Altered mRNAbase sequence | GGU | CUU |   |   |   | AGA | UAU |

(i)      Use the mRNA codons provided in the table to complete the altered mRNA base sequence in the diagram.

|  |  |
| --- | --- |
| **Amino acid** | **mRNA codons which can be used** |
| Arg | AGA |
| Gly | GGU |
| Leu | CUU or UUA |
| Ser | UCU |
| Trp | UGG |
| Tyr | UAU or UAC |

**(1)**

(ii)     Use the information provided to determine the precise nature of the **two** single base mutations in the DNA.

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**(3)**

**(Total 9 marks)**

**Q12.**          (a)     Name **one** mutagenic agent.

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**(1)**

(b)     In flax plants the flowers are white, lilac or blue. The diagram shows the pathway by which the flower cells produce coloured pigments.



(i)      A deletion mutation occurs in gene 1. Describe how a deletion mutation alters the structure of a gene.

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**(2)**

(ii)     Describe and explain how the altered gene could result in flax plants with white-coloured flowers.

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**(4)**

(iii)     Electrophoresis was used to separate the enzymes involved in this pathway. When extracts of the differently coloured flax petals were analysed, four different patterns of bands were produced. In the table, only bands that contain functional enzymes are shown.

|  |  |
| --- | --- |
| **Result of electrophoresis** | **Colour of petal** |
|  | White |
|  |   |
|  |   |
|  |   |

Complete the table to give the colour of the petal from which each extract was taken.

**(2)**

**(Total 9 marks)**

**Q13.**          **S**       The diagram shows a single-celled organism called *Chlamydomonas*.



(a)     *Chlamydomonas* lives in fresh-water ponds. It uses its flagella to swim towards light of moderate intensity but away from very bright light. Using information in the diagram, explain the advantage of this behaviour.

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**(2)**

(b)     A *Chlamydomonas* cell has two flagella. These flagella contain a single sort of protein. A flagellum consists of a bundle of 242 filaments. Each filament consists of 7500 protein molecules. Each protein molecule contains 900 amino acid units.

(i)      What would be the minimum number of nucleotides in the coding region of the mRNA used to synthesise this protein?

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**(1)**

(ii)     In an investigation, a culture of *Chlamydomonas* was treated in a way that caused them to lose their flagella without any other damage to the cells. The flagella grew back to their original length in 60 minutes.

How many amino acid molecules would be incorporated into each growing flagellum per minute? Show your working.

Answer .........................................

**(2)**

(c)     The researchers investigated the rate at which the flagella grew in three different media.

1.      A medium containing actinomycin D, which prevents transcription by binding to the guanine in DNA

2.      A medium containing puromycin, which prevents translation by attaching to ribosomes

3.      A control medium

The results are shown in the graph.



(i)      Describe how the rate of growth was affected by puromycin.

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**(2)**

(ii)     The researchers concluded

1.       that the cells used mRNA that is already present in the cytoplasm for the regrowth of the flagella;

2.       that some of the regrowth uses protein molecules already present in the cell.

Explain the evidence for each of these conclusions.

1 ..........................................................................................................

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2 ..........................................................................................................

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**(4)**

**(Total 11 marks)**

**Q14.**          (a)     **Table 1** shows some of the events which take place in protein synthesis.

|  |  |  |
| --- | --- | --- |
|   | **A** | tRNA molecules bring specific amino acids to the mRNA molecule |
|   | **B** | mRNA nucleotides join with exposed DNA bases and form a molecule of mRNA |
|   | **C** | The two strands of a DNA molecule separate |
|   | **D** | Peptide bonds form between the amino acids |
|   | **E** | The mRNA molecule leaves the nucleus |
|   | **F** | A ribosome attaches to the mRNA molecule |

**Table 1**

(i)      Write the letters in the correct order to show the sequence of events during protein synthesis, starting with the earliest.

................  ................  ................  .................  .................  .................

**(2)**

(ii)     In which part of a cell does **C** take place?

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**(1)**

(iii)     Which of **A - F** are involved in translation?

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**(1)**

(b)     **Table 2** shows some mRNA codons and the amino acids for which they code.

|  |  |  |
| --- | --- | --- |
|   | **mRNA codon** | **Amino acid** |
|   | GUU | Valine |
|   | CUU | Leucine |
|   | GCC | Alanine |
|   | AUU | Isoleucine |
|   | ACC | Threonine |

**Table 2**

(i)      A tRNA molecule has the anticodon UAA.  Which amino acid does the tRNA molecule carry?

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**(1)**

(ii)     Give the DNA base sequence that codes for threonine.

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**(1)**

**(Total 6 marks)**

**Q15.**          (a)     **Figure 1** shows the exposed bases (anticodons) of two tRNA molecules involved in the synthesis of a protein.

**Figure 1**

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Complete the boxes to show the sequence of bases found along the corresponding section of the coding DNA strand.

**(2)**

(b)     Describe the role of tRNA in the process of translation.

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**(3)**

(c)     **Figure 2** shows the sequence of bases in a section of DNA coding for a polypeptide of seven amino acids.

**Figure 2**

TACAAGGTCGTCTTTGTCAAG

The polypeptide was hydrolysed. It contained four different amino acids. The number of each type obtained is shown in the table.

|  |  |
| --- | --- |
| **Amino acid** | **Number present** |
| Phe | 2 |
| Met | 1 |
| Lys | 1 |
| Gln | 3 |

Use the base sequence shown in **Figure 2** to work out the order of amino acids in the polypeptide. Write your answer in the table below.

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| Met |   |   |   |   |   |   |

**(2)**

**(Total 7 marks)**

**Q16.**          (a)     (i)      What is the role of RNA polymerase in transcription?

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**(1)**

(ii)     Name the organelle involved in translation.

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**(1)**

(b)     **Figure 1** shows some molecules involved in protein synthesis.

**Figure 1**

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Complete **Figure 1** to show

(i)      the bases on the DNA strand from which the mRNA was transcribed;

(ii)     the bases forming the anticodons of the tRNA molecules.

**(2)**

**Figure 2** shows the effects of two different mutations of the DNA on the base sequence of the mRNA. The table shows the mRNA codons for three amino acids.

**Figure 2**

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(c)     Name the type of mutation represented by mutation 1.

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**(1)**

(d)     Use the information in the table to

(i)      identify amino acid **X** in **Figure 1**;

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**(1)**

(ii)     explain how each mutation may affect the polypeptide for which this section of DNA is part of the code.

Mutation 1 ...........................................................................................

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.............................................................................................................

**(2)**

Mutation 2 ...........................................................................................

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**(2)**

**(Total 10 marks)**

**Q17.**          (a)     Give **two** factors, other than cost, that should be considered when selecting an antibiotic to treat a bacterial disease.

1 ...................................................................................................................

......................................................................................................................

2 ...................................................................................................................

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**(2)**

**S**       (b)     The table describes the effects of two antibiotics on bacteria.

|  |  |
| --- | --- |
| **Antibiotic** | **Effect** |
| Tetracycline | prevents tRNA binding |
| Chloramphenicol | prevents peptide bonds forming |

(i)      Explain how each of these antibiotics slows down the rate of growth of bacteria.

Tetracycline ........................................................................................

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Chloramphenicol .................................................................................

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**(4)**

(ii)     Suggest why tetracycline has no effect on human cells.

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**(1)**

**(Total 7 marks)**

**Q18.**          Read the following passage.

The sequence of bases in a molecule of DNA codes for proteins. Different sequences of bases
code for different proteins. The genetic code, however, is degenerate. Although the base
sequence AGT codes for serine, other sequences may also code for this same amino acid.
There are four base sequences which code for the amino acid glycine. These are CCA, CCC,

5    CCG and CCT. There are also four base sequences coding for the amino acid proline. These

are GGA, GGC, GGG and GGT.

Pieces of DNA which have a sequence where the same base is repeated many times are called
“slippery”. When “slippery” DNA is copied during replication, errors may occur in copying.
Individual bases may be copied more than once. This may give rise to differences in the

10  protein which is produced by the piece of DNA containing the errors.

          Use information in the passage and your own knowledge to answer the following questions.

(a)     Different sequences of bases code for different proteins (lines 1 – 2). Explain how.

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**(2)**

(b)     The base sequence AGT codes for serine (lines 2 – 3). Give the mRNA codon transcribed from this base sequence.

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**(2)**

(c)     Glycine-proline-proline is a series of amino acids found in a particular protein. Give the sequence of DNA bases for these three amino acids which contains the longest “slippery” sequence.

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**(2)**

(d)     (i)      Explain how copying bases more than once may give rise to a difference in the protein (lines 9 – 10).

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**(2)**

(ii)     At what stage in the cell cycle would these errors in copying DNA bases occur?

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**(1)**

(e)     Starting with mRNA in the nucleus of a cell, describe how a molecule of protein is synthesised.

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**(6)**

**(Total 15 marks)**

**Q19.**          (a)     Complete the table to show the differences between DNA, mRNA and tRNA.

|  |  |  |
| --- | --- | --- |
| **Type of nucleic acid** | **Hydrogen bonds present () or not present ()** | **Number of polynucleotide strands in molecule** |
| DNA |   |   |
| mRNA |   |   |
| tRNA |   |   |

**(2)**

(b)     The diagram shows the bases on one strand of a piece of DNA.



(i)      In the space below, give the sequence of bases on the pre-mRNA transcribed from this strand.

**(2)**

(ii)     In the space below, give the sequence of bases on the mRNA produced by splicing this piece of pre-mRNA.

**(1)**

**(Total 5 marks)**

**Q20.**          (a)     The table shows the mRNA codons for some amino acids.

|  |  |  |
| --- | --- | --- |
|   | **Codon** | **Amino acid** |
|   | CUA | Leucine |
|   | GUC | Valine |
|   | ACG | Threonine |
|   | UGC | Cysteine |
|   | GCU | Alanine |
|   | AGU | Serine |

(i)      Give the DNA sequence coding for cysteine.

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**(1)**

(ii)     Name the amino acid coded by the tRNA anticodon UCA.

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**(1)**

(b)     A particular gene is 562 base-pairs long. However, the resulting mRNA is only 441 nucleotides long. Explain this difference.

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**(1)**

(c)     Tetracycline binds to bacterial ribosomes. This is shown in the diagram.



Protein synthesis in bacteria is similar to that in eukaryotic cells. Explain how tetracycline stops protein synthesis.

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**(2)**

**(Total 5 marks)**

**Q21.**          In a hospital laboratory, a sterile Petri dish of nutrient agar was inoculated with bacteria from a patient with a throat infection. Four discs, each of which had been soaked in a different antibiotic, were placed on top of the bacteria. The dish was incubated at 37 °C. **Figure 1** shows the appearance of the dish after incubation.



**Figure 1**

(a)     Explain why there are clear zones around some of the discs containing antibiotic.

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**(2)**

(b)     It was suggested that ampicillin might be the best antibiotic to treat the patient’s throat infection. Give the evidence from the laboratory test to support this suggestion.

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**(1)**

(c)     Tetracycline binds to bacterial ribosomes. This is shown in **Figure 2**.



**Figure 2**

Tetracycline prevents bacterial growth by preventing protein synthesis. Give **two** other ways in which antibiotics can prevent bacterial growth.

1 ...................................................................................................................

......................................................................................................................

2 ...................................................................................................................

......................................................................................................................

**(2)**

**(Total 5 marks)**

**Q22.**          The table shows the sequence of bases on part of the coding strand of DNA.

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| Base sequence on coding strand of DNA | **C** | **G** | **T** | **T** | **A** | **C** |
| Base sequence of mRNA |   |   |   |   |   |   |

(a)     Complete the table to show the base sequence of the mRNA transcribed from this DNA strand.

**(2)**

(b)     A piece of mRNA is 660 nucleotides long but the DNA coding strand from which it was transcribed is 870 nucleotides long.

(i)      Explain this difference in the number of nucleotides.

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**(1)**

(ii)     What is the maximum number of amino acids in the protein translated from this piece of mRNA? Explain your answer.

Number of amino acids .......................................................................

Explanation .........................................................................................

.............................................................................................................

**(2)**

(c)     Complete the table to give **two** differences between the structure of mRNA and the structure of tRNA.

|  |  |
| --- | --- |
| **mRNA** | **tRNA** |
|   |   |
|   |   |

**(2)**

**(Total 7 marks)**

**Q23.**          The diagram shows part of a DNA molecule.



(a)     Name the **two** components of the part of the DNA molecule labelled **M**.

1 ...................................................................................................................

2 ...................................................................................................................

**(2)**

(b)     What is the maximum number of amino acids for which this piece of DNA could code?



**(1)**

(c)     Scientists calculated the percentage of different bases in the DNA from a species of bacterium. They found that 14% of the bases were guanine.

(i)      What percentage of the bases in this species of bacterium was cytosine?

Answer .......................................

**(1)**

(ii)     What percentage of the bases in this species of bacterium was adenine?

Answer .......................................

**(1)**

(d)     The scientists found that, in a second species of bacterium, 29% of the bases were guanine.

Explain the difference in the percentage of guanine bases in the two species of bacterium.

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**(2)**

**(Total 7 marks)**

 **Q24.**          **Figure 1** shows a short section of a DNA molecule.

**Figure 1**

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(a)     Name parts **R** and **Q**.

(i)      **R** ....................................................

(ii)     **Q** ....................................................

**(2)**

(b)     Name the bonds that join **A** and **B**.

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**(1)**

(c)     Ribonuclease is an enzyme. It is 127 amino acids long.

          What is the minimum number of DNA bases needed to code for ribonuclease?



**(1)**

(d)     **Figure 2** shows the sequence of DNA bases coding for seven amino acids in the enzyme ribonuclease.

**Figure 2**

**G  T  T  T  A  C  T  A  C  T  C  T  T  C  T  T  C  T  T  T  A**

The number of each type of amino acid coded for by this sequence of DNA bases is shown in the table.

|  |  |
| --- | --- |
| **Amino acid** | **Number present** |
| Arg | 3 |
| Met | 2 |
| Gln | 1 |
| Asn | 1 |

Use the table and **Figure 2** to work out the sequence of amino acids in this part of the enzyme. Write your answer in the boxes below.

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| Gln |   |   |   |   |   |   |

**(1)**

(e)     Explain how a change in a sequence of DNA bases could result in a non-functional enzyme.

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**(3)**

**(Total 8 marks)**

 **Q25.**          SCID is a severe inherited disease. People who are affected have no immunity. Doctors carried out a trial using gene therapy to treat children with SCID. The doctors who carried out the trial obtained stem cells from each child’s umbilical cord.

(a)     Give **two** characteristic features of stem cells.

1 ...................................................................................................................

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2 ...................................................................................................................

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**(2)**

The doctors mixed the stem cells with viruses. The viruses had been genetically modified to contain alleles of a gene producing full immunity. The doctors then injected this mixture into the child’s bone marrow.

The viruses that the doctors used had RNA as their genetic material. When these viruses infect cells, they pass their RNA and two viral enzymes into the host cells.

(b)     One of the viral enzymes makes a DNA copy of the virus RNA. Name this enzyme.

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**(1)**

The other viral enzyme is called integrase. Integrase inserts the DNA copy anywhere in the DNA of the host cell. It may even insert the DNA copy in one of the host cell’s genes.

(c)     (i)      The insertion of the DNA copy in one of the host cell’s genes may cause the cell to make a non-functional protein. Explain how.

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**(2)**

(ii)     Some of the children in the trial developed cancer. How might the insertion of the DNA have caused cancer?

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**(2)**

(d)     Five out of the 20 children in the trial developed cancer. Although the cancer was treated successfully, the doctors decided to stop the trial in its early stages. They then reviewed the situation and decided to continue. Do you agree with their decision to continue? Explain your answer.

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**(2)**

**(Total 9 marks)**

 **Q26.Figure 1** shows part of a gene that is being transcribed.

**Figure 1**

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(a)     Name enzyme **X**.

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**(1)**

(b)     (i)      Oestrogen is a hormone that affects transcription. It forms a complex with a receptor in the cytoplasm of target cells. Explain how an activated oestrogen receptor affects the target cell.

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**(2)**

(ii)     Oestrogen only affects target cells. Explain why oestrogen does not affect other cells in the body.

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**(1)**

(c)     Some breast tumours are stimulated to grow by oestrogen. Tamoxifen is used to treat these breast tumours. In the liver, tamoxifen is converted into an active substance called endoxifen. **Figure 2** shows a molecule of oestrogen and a molecule of endoxifen.

**Figure 2**

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Use **Figure 2** to suggest how endoxifen reduces the growth rate of these breast tumours.

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**(2)**

**(Total 6 marks)**

**Q27.**          The black mamba is a poisonous snake. Its poison contains a toxin.

The table shows the base sequence of mRNA that codes for the first two amino acids of this toxin.

|  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- |
|   | Base sequence of anticodon on tRNA |  |  |  |  |  |  |
|   | Base sequence of mRNA | **A** | **C** | **G** | **A** | **U** | **G** |
|   | Base sequence of DNA |  |  |  |  |  |  |

Complete the table to show

(a)     (i)      the base sequence of the anticodon on the first tRNA molecule that would bind to this mRNA sequence

**(1)**

(ii)     the base sequence of the DNA from which this mRNA was transcribed.

**(1)**

(b)     The length of the section of DNA that codes for the complete toxin is longer than the mRNA used for translation. Explain why.

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**(1)**

(c)     A mutation in the base sequence of the DNA that codes for the toxin would change the base sequence of the mRNA.

Explain how a change in the base sequence of the mRNA could lead to a change in the tertiary structure of the toxin.

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**(1)**

(d)     The black mamba’s toxin kills prey by preventing their breathing. It does this by inhibiting the enzyme acetylcholinesterase at neuromuscular junctions. Explain how this prevents breathing.

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**(3)**

**(Total 7 marks)**

**Q28.**          (a)     What name is used for the non-coding sections of a gene?

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**(1)**

**Figure 1** shows a DNA base sequence. It also shows the effect of two mutations on this base sequence. **Figure 2** shows DNA triplets that code for different amino acids.

**Figure 1**

|  |  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| Original DNA base sequence | A | T | T | G | G | C | G | T | G | T | C | T |
| Amino acid sequence |   |   |   |   |
| Mutation **1** DNA base sequence | A | T | T | G | G | A | G | T | G | T | C | T |
| Mutation **2** DNA base sequence | A | T | T | G | G | C | C | T | G | T | C | T |

**Figure 2**

|  |  |
| --- | --- |
| **DNA triplets** | **Amino acid** |
| GGT, GGC, GGA, GGG | Gly |
| GTT, GTA, GTG, GTC | Val |
| ATC, ATT, ATA | Ile |
| TCC, TCT, TCA, TCG | Ser |
| CTC, CTT, CTA, CTG | Leu |

(b)     Complete **Figure 1** to show the sequence of amino acids coded for by the original DNA base sequence.

**(1)**

(c)     Some gene mutations affect the amino acid sequence. Some mutations do not.
Use the information from **Figure 1** and **Figure 2** to explain

(i)      whether mutation **1** affects the amino acid sequence

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**(2)**

(ii)     how mutation **2** could lead to the formation of a non-functional enzyme.

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**(3)**

(d)     Gene mutations occur spontaneously.

(i)      During which part of the cell cycle are gene mutations most likely to occur?

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**(1)**

(ii)     Suggest an explanation for your answer.

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**(1)**

**(Total 9 marks)**

 **Q29.**          The diagram shows the life cycle of a fly.



When the larva is fully grown, it changes into a pupa. The pupa does not feed. In the pupa, the tissues that made up the body of the larva are broken down. New adult tissues are formed from substances obtained from these broken-down tissues and from substances that were stored in the body of the larva.

(a)     Hydrolysis and condensation are important in the formation of new adult proteins.
Explain how.

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**(2)**

(b)     Most of the protein stored in the body of a fly larva is a protein called calliphorin.
Explain why different adult proteins can be made using calliphorin.

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**(1)**

The table shows the mean concentration of RNA in fly pupae at different ages.

|  |  |  |
| --- | --- | --- |
|   | **Age of pupa as percentage of total time spent as a pupa** | **Mean concentration of RNA / μg per pupa** |
|   |     0 | 20 |
|   |   20 | 15 |
|   |   40 | 12 |
|   |   60 | 17 |
|   |   80 | 33 |
|   | 100 | 20 |

(c)     Describe how the concentration of RNA changes during the time spent as a pupa.

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**(2)**

(d)     (i)      Describe how you would expect the number of lysosomes in a pupa to change with the age of the pupa. Give a reason for your answer.

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**(2)**

(ii)     Suggest an explanation for the change in RNA concentration in the first 40% of the time spent as a pupa.

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**(2)**

(e)     Suggest an explanation for the change in RNA concentration between 60 and 80% of the time spent as a pupa.

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**(2)**

(f)      The graph shows changes in the activity of two respiratory enzymes in a fly pupa.

•        Enzyme **A** catalyses a reaction in the Krebs cycle

•        Enzyme **B** catalyses the formation of lactate from pyruvate



During the first 6 days as a pupa, the tracheae break down. New tracheae are formed after 6 days. Use this information to explain the change in activity of the two enzymes.

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**(4)**

**(Total 15 marks)**

 **Q30.**The body markings of cheetahs vary, in particular the pattern of bands on their tails. Cheetahs are solitary animals but the young stay with their mother until they are between 14 and 18 months old.

Scientists investigated the banding pattern on the tails of cheetahs living in the wild.

•        They drove a car alongside a walking cheetah and used binoculars to study the tail pattern.

•        They gave each cheetah a banding pattern score based on the width of the dark and light bands on the end of the tail.

•        They scored the width of the bands on the right and left side of the tail using a 5 point scale of width.

A typical pattern on the right side of one cheetah’s tail is shown in **Figure 1**.

**Figure 1**

|  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- |
|   | Band number | 1 | 2  3 | 4 | 5 | 6 | 7 |

 

|  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- |
|   | Band width score | 3 | 1  1 | 4 | 3 | 3 | 3 |

The scientists collected data from each cheetah on four separate occasions. **Figure 2** shows the data for one of the cheetahs.

**Figure 2**

|  |  |  |
| --- | --- | --- |
|   | **Side oftail** | **Mean band width score (± standard deviation)** |
|   | **Band 1** | **Band 2** | **Band 3** | **Band 4** | **Band 5** | **Band 6** | **Band 7** |
|   | Right | 3.00 (± 0.82) | 1.00 (± 0.00) | 1.00 (± 0.00) | 3.75 (± 0.50) | 2.75 (± 0.50) | 3.00 (± 0.00) | 3.00 (± 0.00) |
|   | Left | 3.75 (± 0.50) | 3.25 (± 0.50) | 2.00 (± 0.50) | 3.00 (± 0.00) | 2.00 (± 0.00) | 2.50 (± 0.50) | 3.00 (± 0.50) |

(a)     The scientists only used data from cheetahs which were fully grown. Suggest why.

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**(1)**

(b)     The scientists estimated the width of the bands on the same cheetah on four separate occasions. They did not always get the same score.

(i)      Give **two** pieces of evidence from **Figure 2** which show that the scientists sometimes obtained different scores for the same band.

1 ............................................................................................................

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2 ............................................................................................................

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**(2)**

(ii)     The method the scientists used resulted in them getting different scores for the same band. Suggest why.

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**(1)**

(c)     What is the evidence from **Figure 2** that the dark and light bands do **not** form rings of equal width around the tail?

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**(1)**

(d)     The scientists found the difference in banding pattern between

•        offspring in the same family

•        cheetahs chosen randomly.

Explain how scientists could use this information to show that some variation in tail banding was genetic.

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**(3)**

**(Total 8 marks)**

**Q31.**          The diagram shows part of a pre-mRNA molecule.



(a)     (i)      Name the **two** substances that make up part **X**.

................................................... and .................................................

**(1)**

(ii)     Give the sequence of bases on the DNA strand from which this pre-mRNA has been transcribed.

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**(1)**

(b)     (i)      Give one way in which the structure of an mRNA molecule is different from the structure of a tRNA molecule.

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**(1)**

(ii)     Explain the difference between pre-mRNA and mRNA.

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**(1)**

(c)     The table shows the percentage of different bases in two pre-mRNA molecules.
The molecules were transcribed from the DNA in different parts of a chromosome.

|  |  |  |
| --- | --- | --- |
|   | **Part of chromosome** | **Percentage of base** |
|   | **A** | **G** | **C** | **U** |
|   | Middle | 38 | 20 | 24 |   |
|   | End | 31 | 22 | 26 |   |

(i)      Complete the table by writing the percentage of uracil (U) in the appropriate boxes.

**(1)**

(ii)     Explain why the percentages of bases from the middle part of the chromosome and the end part are different.

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**(2)**

**(Total 7 marks)**

**Q32.**          The diagram shows a short sequence of DNA bases.

**T T T G T A T A C T A G T C T A C T T C G T T A A T A**

(a)     (i)      What is the maximum number of amino acids for which this sequence of DNA bases could code?



**(1)**

(ii)The number of amino acids coded for could be fewer than your answer to part (a)(i).

Give **one** reason why.

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**(1)**

(b)Explain how a change in the DNA base sequence for a protein may result in a change in the structure of the protein.

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**(3)**

(c)A piece of DNA consisted of 74 base pairs. The two strands of the DNA, strands **A** and **B**, were analysed to find the **number** of bases of each type that were present. Some of the results are shown in the table.

|  |  |  |
| --- | --- | --- |
|   |   | **Number of bases** |
|   |   | C | G | A | T |
|   | Strand **A** | 26 |   |   |   |
|   | Strand **B** | 19 |   | 9 |   |
|   |  |  |  |  |  |  |

Complete the table by writing in the missing values.

**(2)**

**(Total 7 marks)**

**Q33.**The diagram shows part of a DNA molecule.

 

(a)     (i)      DNA is a polymer. What is the evidence from the diagram that DNA is a polymer?

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**(1)**

(ii)     Name the parts of the diagram labelled **C**, **D** and **E**.

|  |  |  |
| --- | --- | --- |
|   | Part **C** | ....................................................................... |
|   | Part **D** | ....................................................................... |
|   | Part **E** | ....................................................................... |

**(3)**

(iii)    In a piece of DNA, 34% of the bases were thymine.

Complete the table to show the names and percentages of the other bases.

|  |  |  |
| --- | --- | --- |
|   | **Name of base** | **Percentage** |
|   | Thymine | 34 |
|   |   |   |
|   |   | 34 |
|   |   |   |

**(2)**

(b)     A polypeptide has 51 amino acids in its primary structure.

(i)      What is the minimum number of DNA bases required to code for the amino acids in this polypeptide?

|  |  |
| --- | --- |
|   |  |

**(1)**

(ii)     The gene for this polypeptide contains more than this number of bases.

Explain why

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**(1)**

**(Total 8 marks)**

**Q34.**(a)    The genetic code is described as being degenerate. What does this mean?

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**(1)**

(b)     What is a codon?

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**(2)**

(c)    (i)      What is the role of RNA polymerase during transcription?

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**(1)**

(ii)     mRNA can be converted to cDNA.

Name the enzyme used in this process.

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**(1)**

(d)     The diagram shows the base sequence on DNA where a restriction endonuclease cuts DNA.



Use evidence from the diagram to explain what is meant by a palindromic recognition sequence on DNA.

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**(1)**

**(Total 6 marks)**

**Q35.**The Amish are a group of people who live in America. This group was founded by 30 Swiss people, who moved to America many years ago. The Amish do not usually marry people from outside their own group.

One of the 30 Swiss founders had a genetic disorder called Ellis-van Creveld syndrome. People with this disorder have heart defects, are short and have extra fingers and toes. Ellis-van Creveld syndrome is caused by a faulty allele.

In America today, about 1 in 200 Amish people are born with Ellis-van Creveld syndrome. This disorder is very rare in people in America who are not Amish.

(a)     In America today, there are approximately 1250 Amish people who have Ellis-van Creveld syndrome. Use the information provided to calculate the current Amish population of America.

Amish population .....................................

**(1)**

(b)     The faulty allele that causes Ellis-van Creveld syndrome is the result of a mutation of a gene called *EVC.* This mutation leads to the production of a protein that has one amino acid missing.

(i)      Suggest how a mutation can lead to the production of a protein that has one amino acid missing.

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**(2)**

(ii)     Suggest how the production of a protein with one amino acid missing may lead to a genetic disorder such as Ellis-van Creveld syndrome.

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**(2)**

**(Total 5 marks)**

**Q36.**(a)     (i)      A mutation of a tumour suppressor gene can result in the formation of a tumour.

Explain how.

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**(2)**

(ii)     Not all mutations result in a change to the amino acid sequence of the encoded polypeptide.

Explain why.

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**(1)**

(b)     Some cancer cells have a receptor protein in their cell-surface membrane that binds to a hormone called **growth factor**. This stimulates the cancer cells to divide.

Scientists have produced a monoclonal antibody that stops this stimulation.

Use your knowledge of monoclonal antibodies to suggest how this antibody stops the growth of a tumour.

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**(3)**

**(Total 6 marks)**

**Q37.**(a)     Messenger RNA (mRNA) is used during translation to form polypeptides.
Describe how mRNA is produced in the nucleus of a cell.

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**(6)**

(b)     Describe the structure of proteins.

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**(5)**

(c)     Describe how proteins are digested in the human gut.

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**(4)**

**(Total 15 marks)**

**Q38.**(a)     Explain how the structure of DNA is related to its functions.

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**(6)**

Scientists investigated three genes, **C**, **D** and **E**, involved in controlling cell division.
They studied the effect of mutations in these genes on the risk of developing lung cancer.

The scientists analysed genes **C**, **D** and **E** from healthy people and people with lung cancer.

•        If a person had a normal allele for a gene, they used the symbol N.

•        If a person had two mutant alleles for a gene, they used the symbol M.

They used their data to calculate the risk of developing lung cancer for people with different combinations of N and M alleles of the genes. A risk value of 1.00 indicates no increased risk. The following table shows the scientists’ results.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
|   | **Gene C** | **Gene D** | **Gene E** | **Risk ofdevelopinglung cancer** |
|   | N | N | N | 1.00 |
|   | M | N | N | 1.30 |
|   | N | N | M | 1.78 |
|   | N | M | N | 1.45 |
|   | N = at least one copy of the normal allele is presentM = two copies of the mutant allele are present |

(b)     What do these data suggest about the relative importance of the mutant alleles of genes **C**, **D** and **E** on **increasing** the risk of developing lung cancer? Explain your answer.

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**(3)**

Chemotherapy is the use of a drug to treat cancer. The drug kills dividing cells.
The figure below shows the number of healthy cells and cancer cells in the blood of a patient receiving chemotherapy. The arrows labelled **F** to **I** show when the drug was given to the patient.

 
                                    Time / days

(c)     Calculate the rate at which healthy cells were killed between days 42 and 46.

.............. cells killed per unit volume of blood per day

**(1)**

(d)     Describe similarities and differences in the response of healthy cells and cancer cells to the drug between times **F** and **G**.

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**(3)**

(e)     More cancer cells could be destroyed if the drug was given more frequently.

Suggest why the drug was **not** given more frequently.

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**(2)**

**(Total 15 marks)**

**Q39.**(a)     (i)      Why is the genetic code described as being universal?

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**(1)**

(ii)     The genetic code uses four different DNA bases. What is the maximum number of different DNA triplets that can be made using these four bases?

 

**(1)**

Transcription of a gene produces pre-mRNA.

(b)     Name the process that removes base sequences from pre-mRNA to form mRNA.

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**(1)**

(c)     The figure below shows part of a pre-mRNA molecule. Geneticists identified two mutations that can affect this pre-mRNA, as shown in the figure.

|  |  |  |  |
| --- | --- | --- | --- |
|   | Base sequence codingfor amino acids | Base sequence removedfrom pre-mRNA | Base sequence codingfor amino acids |



|  |  |  |
| --- | --- | --- |
|   | **Mutation 1,single basedeletion** | **Mutation 2,single basesubstitution** |

(i)      **Mutation 1** leads to the production of a non-functional protein.

Explain why.

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*(Extra space)* ........................................................................................

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**(3)**

(ii)     What effect might **mutation 2** have on the protein produced?

Explain your answer.

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**(2)**

**(Total 8 marks)**

**Q40.**Read the following passage carefully.

|  |  |  |
| --- | --- | --- |
|   | A large and growing number of disorders are now known to be due to types of mitochondrial disease (MD). MD often affects skeletal muscles, causing muscle weakness. |   |
|   | We get our mitochondria from our mothers, via the fertilised egg cell. Fathers do not pass on mitochondria via their sperm. Some mitochondrial diseases are caused by mutations of mitochondrial genes inside the mitochondria.Most mitochondrial diseases are caused by mutations of genes in the cell nucleus that are involved in the functioning of mitochondria. These mutations of nuclear DNA produce recessive alleles. |  5 |
|   | One form of mitochondrial disease is caused by a mutation of a mitochondrial gene that codes for a tRNA. The mutation involves substitution of guanine for adenine in the DNA base sequence. This changes the anticodon on the tRNA.This results in the formation of a non-functional protein in the mitochondrion. | 10 |
|   | There are a number of ways to try to diagnose whether someone has a mitochondrial disease. One test involves measuring the concentration of lactate in a person’s blood after exercise. In someone with MD, the concentration is usually much higher than normal. If the lactate test suggests MD, a small amount of DNA can be extracted from mitochondria and DNA sequencing used to try to find a mutation. |  15 |

Use information in the passage and your own knowledge to answer the following questions.

(a)     Mitochondrial disease (MD) often causes muscle weakness (lines 1–3). Use your knowledge of respiration and muscle contraction to suggest explanations for this effect of MD.

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**(Extra space)** ................................................................................................

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**(3)**

Two couples, couple **A** and couple **B**, had one or more children affected by a mitochondrial disease. The type of mitochondrial disease was different for each couple.

None of the parents showed signs or symptoms of MD.

•        Couple **A** had four children who were all affected by an MD.

•        Couple **B** had four children and only one was affected by an MD.

(b)     Use the information in lines 5–9 and your knowledge of inheritance to suggest why:

•        all of couple **A**’s children had an MD

•        only one of couple **B**’s children had an MD.

Couple **A** ........................................................................................................

........................................................................................................................

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Couple **B** ........................................................................................................

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**(Extra space)** ................................................................................................

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**(4)**

(c)     Suggest how the change in the anticodon of a tRNA leads to MD (lines 10–13).

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**(Extra space)** ................................................................................................

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**(3)**

(d)     If someone has MD, the concentration of lactate in their blood after exercise is usually much higher than normal (lines 15–17). Suggest why.

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**(Extra space)** ................................................................................................

........................................................................................................................

........................................................................................................................

**(3)**

(e)     A small amount of DNA can be extracted from mitochondria and DNA sequencing used to try to find a mutation (lines 18–19).

From this sample:

•        how would enough DNA be obtained for sequencing?

•        how would sequencing allow the identification of a mutation?

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........................................................................................................................

........................................................................................................................

**(2)**

**(Total 15 marks)**

**M1.**(a)     Translation.

**1**

(b)     Transfer RNA / tRNA.

**1**

(c)     TAC;

UAC.

**2**

(d)     Have different R group.

*Accept in diagram*

**1**

(e)     1.      Substitution would result in CCA / CCC / CCU;

2.      (All) code for same amino acid / proline;

3.      Deletion would cause frame shift / change in all following codons / change next codon from UAC to ACC.

**3**

**[8]**

**M2.**(a)     1.      (Protein / molecule) that moves from cytoplasm to DNA;

*Accept ‘it’ as TF.*

*Accept moves into nucleus*

2.      (TF) binds to specific gene / genes / to specific part of / site on DNA / binds to promoter / RNA polymerase;

*Accept regulator / enhancer region*

3.      Leads to / blocks (pre)mRNA production / allows / blocks binding of RNA polymerase (to DNA) / allows RNA polymerase to work;

*Ignore translation unless context wrong*

*Max 1 if refer to oestrogen as a transcription factor*

**2 max**

(b)     1.      (Binding to CREB) prevents transcription / mRNA formation;

*Accept that lack of protein leaves NAD reduced*

2.      (Binding of huntingtin) prevents production / translation of protein (that removes electrons / protons from NAD);

3.      Fewer electrons to electron transport chain / electron transport chain slows / stops / stops / slower oxidative phosphorylation;

4.      Fewer protons for proton gradient;

5.      Not enough ATP produced / energy supplied to keep cells alive / anaerobic respiration not enough to keep cell alive;

*Accept neurones require ATP for active transport of ions*

*Ignore references to resting potential*

**3 max**

(c)     1.      Mitochondrion has two membranes / inner and outer membranes;

*Accept cristae for inner membrane*

2.      For each (different) membrane a (different) carrier required;

*Ignore reference to channel proteins*

**2**

**[7]**

**M3.**          (a)     mutation changes the amino acid sequence / primary structure of Factor VIII protein;
changes the tertiary structure / 3D shape;

**2**

(b)     (mutant) Factor VIII protein is non-functional / does not work with Factor IX;
so no conversion of Factor X to active form and pathway blocked;

**2**

(c)     boy’s blood contains (active) Factor VIII;
Factor VIII haemophiliac’s blood contains (active) Factor IX;
the mixture has both Factors and so the pathway can
complete / blood clots;

**2 max**

**[6]**

**M4.**          (a)     (i)      number of bases = 4440

*allow 4446 if they refer to start / stop*

each amino acid coded for by triplet / three bases
(so three times more bases than amino acids);

**2**

(ii)     deletion;
(deletion) of three bases;
because substitution / addition would change amino acid(s);

**2 max**

(b)     codon on mRNA;
specific / complementary base pairing with;
anti-codon on tRNA;
specific tRNA for each amino acid;
protein formed by condensation reactions /
peptide bonds formed;

**4 max**

**[8]**

**M5.**          (a)     Any two of:

|  |  |
| --- | --- |
| DNA | RNA |
| Large molecule | Smaller |
| Double stranded | Single stranded |
| Contains Thymine (T) | Contains Uracil (U) |
| Contains deoxyribose | Contains ribose |

**2 max**

(b)     Base sequence (on DNA/in gene);Determines sequence of amino acids;By determining base sequence on (messenger) RNA;Code is a triplet code/three base code for an amino acid;

**2 max**

(c)     Pairs of chromosomes/two chromosomes;With genes for same features / with same genes;At same loci / in same sequence;

*Accept same alleles*

**2 max**

**[6]**

**M6.**          (a)     Two suitable differences between DNA and RNA;

*1 mark per correct row to 2 max*

e.g.

DNA is double stranded, RNA is single stranded;DNA has thymine present, RNA has Uracil present;

*Accept T and U*

DNA is larger/heavier/longer, RNA is smaller/lighter/shorter;DNA has a deoxyribose sugar, RNA has a ribose sugar;DNA stays in the nucleus, RNA leaves the nucleus;

**2 max**

(b)     Three suitable examples;

e.g.

Carries coded information about the sequence of amino acids;Copied from DNA/gene;Code is in sequence of bases / triplet / three bases / a codon codes
for one amino acid;Moves out of nucleus/goes into cytoplasm;To ribosomes;

*Accept codons allow anticodons / tRNA to bind*

*Accept carries ‘start’ and ‘stop’ codes*

*Accept moves through ribosomes*

**3 max**

**[5]**

**M7.**(a)     (Gene 1) allele A makes enzyme converting J to K / colourless to red;

Allele a produces no / non-functional enzyme;
(Gene 2) allele B makes enzyme converting K to L / red to purple;
Allele b produces no / non-functional enzyme;
(“Recessive alleles produce no / non-functional enzyme” = 2)
White flowers result from genotype aa;
... regardless if B or b / even if aaB\_ ;
Colourless (substance) / J produces white;
Red flowers when A\_ bb / enzyme 1 only;
Purple flowers when A\_ B\_ / enzymes 1 and 2;

**6 max**

(b)     (i)      (1) (red parent) AAbb;

(2) (white parent) aaBB;

**2**

(ii)     F1 are AaBb;
F2 ratio of 9 : 3 : 4;
Purple : red : white;
Suitable working shown;

**4**

(c)     (i)      aabb, aaBb, and aaBB; (allow aabb & aaB\_)

**1**

(ii)     (Crush each type of white petal to make an extract, and) add some of the (red) pigment / K, to petal OR incubate with K;
(extract becoming) purple is identified as aaBB OR that staying red, after K is added, is aabb;

**2**

**[15]**

**M8.**          (i)      mRNA attaches to ribosome;
codon on mRNA;
binds to an anti-codon on tRNA;
each tRNA brings a specific amino acid;
sequence of codons / bases on mRNA determines order of amino acids;
formation of peptide bonds / amino acids joined by condensation
reactions;

**4 max**

(iii)     inserted gene / mRNA complementary to normal gene / mRNA;
binds to it to prevent protein synthesis / form double strand / prevents
mRNA binding to ribosomes;
will not stop all translation, some mRNA reaches ribosomes /
because not all mRNA is bound by inserted gene mRNA;

**2 max**

**[6]**

**M9.**          (a)     memory B / T cells do not recognise (new antigens);
antibodies previously produced are not effective
as shape not complementary to new antigen;

**2**

(b)     (i)      antigen in membrane presented to lymphocytes /
produce cytokinins;

**1**

(ii)     mitochondria provide (more) ATP / energy;
(more) RER / ribosomes synthesise proteins;
(more) Golgi body secretes / modifies or packages proteins /
produces glycoproteins;
(B lymphocytes) produces antibodies;

**4**

**[7]**

**M10.**          change in base / nucleotide (in DNA);
change in base sequence of mRNA / change in codons / idea of
frameshift following deletion or addition / incorrect tRNA / anticodon;
incorrect amino acids / different primary structure / fomation of new
stop codon;
different tertiary structure / different 3D structure / different
polypeptide / shortened polypeptide;
different shape of active site / no active site present;

**[5]**

**M11.**          (a)     387;

**1**

(b)     (i)      CCAG;

**1**

(ii)     5;

**1**

(c)     high energy radiation / X rays / ultraviolet light / gamma rays;
high energy particles / alpha particles / beta particles;
named chemical mutagens e.g. benzene / caffeine / pesticide /
mustard gas / tobacco tar / free radicals;
*(two named examples of any of the above = 2 marks)*length of time of exposure (to a mutagen);
dosage (of mutagen);

**2 max**

(d)     (i)      UAC UUA UGG;

**1**

(ii)     addition and deletion (of bases / nucleotides);
thymine added;
adenine deleted;
*(addition of thymine and deletion of adenine = 3 marks)
(allow addition of adenine (RNA) and deletion of uracil (RNA)
= 2 marks)*

**3**

**[9]**

**M12.**          (a)     high energy radiation / ionising particles;

named particles / α, β, γ;

colchicine;

x rays / cosmic rays;

uv (light);

carcinogen / named carcinogen;

mustard gas / phenols / tar (qualified);

**1 max**

(b)     (i)      removal of one or more bases / nucleotide;

frameshift / (from point of mutation) base sequence change;

**2**

(ii)     sequence of bases in mRNA would change;

(sequence of) amino acids different / different primary structure;

(active site / enzyme 1) changed tertiary shape / changed active
sites;

white pigment does not bind;

lilac pigment not produced / white pigment remains unchanged /

enzyme 1 does not function;

**4 max**

(iii)     blue and lilac; white;

|  |
| --- |
| *colour of petal* |
| *(white)* |
| blue |
| lilac; |
| white; |

**2**

**[9]**

**M13.**         (a)     chloroplast, so cell photosynthesises and moves to optimum / best light intensity for photosynthesis;
avoids damage due to bright light;

**2**

(b)     (i)      2700

**1**

(ii)      = 27 225 000 / 27 × 106 = *2 marks*

*(allow 1 mark for principle: )*

**2**

(c)     (i)      rate slightly slower / not affected in first 20 / 30 minutes / lower
peak than control;
then decreases / much lower (than control);

*(allow 1 mark for increase in first 20 / 30 minutes, then decreased, if not compared with control / normal)*

*(disqualify flagellum grows longer)*

**2**

(ii)     1.       actinomycin has no effect (on growth of flagella);
          even though mRNA production / transcription prevented;

*(accept references to ‘expt 1’)*

2.       (re)growth little affected by puromycin at first;
protein synthesis inhibited, so likely to be using proteins
present;

**4**

**[11]**

**M14.**          (a)     (i)      C → B → E → F → A → D

*Mark links: 5 correct = 2, 4 correct = 1, <4 correct = 0*

**2**

(iii)     nucleus;

**1**

(iii)     A, D, F;    *(ignore E if evident)*

**1**

(b)     (i)      Isoleucine;

**1**

(ii)     TGG;

**1**

**[6]**

**M15.**          (a)     AGC; TTC;

**2**

(b)     anticodon complementary to codon / reads message on mRNA;

specific amino acid;

carried / transferred (to ribosome);

correct sequence of amino acids along polypeptide;

**3 max**

(c)

|  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- |
| (Met) | Phe | Gln | Gln | Lys | Gln | Phe |

**2**

*(three / four / five correct 1 mark; six correct 2 marks)*

**[7]**

**M16.**          (a)     (i)      join / attach nucleotides, to form a strand / along backbone / phosphodiester bonds;

*(reject reference to H bonds, complementary base pairing)*

**1**

(ii)     ribosome / RER;

**1**

(b)     (i)      CGTTACCAA;

**1**

(ii)     CGU UAC CAA;

**1**

(c)     substitution;

**1**

(d)     (i)      alanine;

**1**

(ii)     (mutation 1)
no change(to sequence of amino acids);
codon for alanine / degenerate codon / same amino acid coded for;

**2**

(mutation 2)
(change in sequence) valine replaced by alanine / codon for alanine;
folding / shape / tertiary structure / position of bonds may change;

*(reject peptide bonds)*

**2**

**[10]**

**M17.**          (a)     side effects / allergic reactions / low toxicity to cells;
interaction with other drugs / effective in conditions of use / reasonably stable;
should only act on the problem bacteria / narrow spectrum;
how much resistance the bacteria have built up;

**2 max**

(b)     (i)      tetracycline
prevents tRNA binding to ribosomes / amino acid / mRNA;

**1**

amino acids not available / brought / picked up;

**1**

chloramphenicol
prevents amino acids being joined / prevents primary structure forming;

**1**

no enzymes / no structural proteins formed;

*(accept cell wall formation if qualified) (prevents protein synthesis gains one mark in either section, once only)*

**1**

(ii)     only prevents tRNA binding to 70S / prokaryotic / bacterial
ribosomes / human ribosomes are different sizes / shapes / structure;

**1**

**[7]**

**M18.**          (a)     Protein made of (chain of) amino acids;
Each amino acid has its own base / triplet code;

**2**

(b)     UCA = 2 marks
TCA – 1 mark;

**2**

(c)     CCG;
GGG GGG;

**2**

(d)     (i)      Changes base sequence;
Of later triplets / amino acid codes;

**2**

(ii)     S-phase / interphase;

**1**

(e)     1.      mRNA leaves (nucleus) through nuclear pore;

2.      To ribosome;

3.      tRNA molecules bring amino acids (to ribosome);

4.      Specific tRNA molecule for specific amino acid;

5.      Anticodon of tRNA corresponds / complementary to codon on mRNA;

6.      Peptide bonds form between amino acids;

7.      tRNA detaches and collects another amino acid;

8.      Ribosome moves along mRNA;

**max 6**

**[15]**

**M19.**          (a)

|  |  |  |
| --- | --- | --- |
| DNA |  | 2 |
| mRNA |  | 1 |
| tRNA |  | 1 |

*One mark for each correct column
Regard blank as incorrect in the context of this question
Accept numbers written out: two, one, one*

**2**

(b)     (i)      Marking principles
1 mark for complete piece transcribed;

*Correct answer
UGU CAU GAA UGC UAG*

1 mark for complementary bases from sequence transcribed;

*but allow 1 mark for complementary bases from section transcribed, providing all four bases are involved*

**2**

(ii)     Marking principle
1 mark for bases corresponding to exons taken from (b)(i)

*Correct answer
UGU UGC UAG
If sequence is incorrect in (b)(i), award mark if section is from exons. Ignore gaps.*

**1**

**[5]**

**M20.**          (a)     (i)      ACG;

**1**

(ii)     serine;

**1**

(b)     idea that DNA contains introns / mRNA is only exons / mRNA is “edited’;
*(allow junk / non-sense DNA)*

**1**

(c)     binds to / blocks codon / triplet on mRNA so anticodon / tRNA will not fit in / base-pair;
amino acids not delivered / joined;

*Accept translation will not occur for 1 mark*

**2**

**[5]**

**M21.**          (a)     antibiotic has diffused / spread / moved into agar;
killed / inhibited bacteria;

**2**

(b)     largest clear area / inhibition zone / killed the most bacteria;

**1**

(c)     disrupts cell wall / prevents cell wall synthesis;
stops DNA replication;

**2**

**[5]**

**M22.**          (a)     GCAAUG; ;

*Allow one mark if T instead of U, i.e. GCAATG*

**2**

(b)     (i)      DNA is edited / introns present in DNA;

*Allow reference to ‘junk’ or non-coding DNA*

**1**

(ii)     220; allow 218 or 219-allow 2

Three bases / nucleotides code for one amino acid;

Correct explanation for 218 or 219;

**2**

(c)     mRNA has no base-pairing, tRNA has base-pairing / mRNA linear,
tRNA cloverleaf shape; mRNA has no binding site for amino acids,
tRNA has; mRNA different for each gene / many kinds, only few / 20 / 64
kinds of tRNA; accept mRNA longer / larger / more nucleotides than tRNA

                **max 2**

**[7]**

**M23.**          (a)     Phosphate;

Deoxyribose;

***Q*** *Candidates must specify deoxyribose. This term is a specification requirement.
Ignore anything that is not incorrect.*

**2**

(b)     4;

**1**

(c)     (i)      14;

**1**

(ii)     36;

*If (c)(i) incorrect accept [50 – (c)(i)]*

**1**

(d)     Different genes;

Different (DNA) base sequences;

**2**

**[7]**

**M24.**          (a)     (i)      Deoxyribose;

*pentose / 5C sugar = neutral*

**1**

(ii)     Phosphate / Phosphoric acid;

*phosphorus / P = neutral*

**1**

(b)     Hydrogen (bonds);

**1**

(c)     381 / 384 / 387;

**1**

(d)     (Gln) Met Met Arg Arg Arg Asn;

**1**

(e)     Change in (sequence of) amino acids / primary structure;

Change in hydrogen / ionic / disulfide bonds leads to change in tertiary structure / active site (of enzyme);

Substrate cannot bind / no enzyme-substrate complexes form;

***Q*** *Reject = different amino acids are formed*

**3**

**[8]**

**M25.**          (a)     Will replace themselves / keep dividing / replicate;

Undifferentiated / can differentiate / develop into other cells / totipotent / multipotent / pluripotent;

*Accept tissues*

**2**

(b)     Reverse transcriptase;

*Allow phonetic spelling*

**1**

(c)     (i)      Alters base / nucleotide sequence / causes frame shift;

Different sequence of amino acids in polypeptide / protein / primary structure alters the tertiary structure;

*Accept any reference, such as adding bases, to changing the base sequence of the gene. Reject deletion / substitution.
Idea of sequence essential so not makes different amino acids.*

*Accept answers involving stop / start codons and effect on protein.*

**2**

(ii)     Affects tumour suppressor gene;

Inactivates (tumour suppressor) gene;

Rate of cell division increased / tumour cells continue to divide;

*Ignore answers relating to oncogenes. May gain third point.*

**2 max**

(d)     Yes
SCID patients unlikely to survive / quality of life poor unless treated;
Cancer that develops is treatable / only affects 25% / five children;

No
Risk of developing cancer is high / 25%;
Cancer may recur / may not be treated successfully in future / only short time scale so more may develop cancer;

*No mark for yes or no. Marks are for supporting argument based on biological reasoning.
Accept any points*

**2 max**

**[9]**

**M26.**(a)     RNA polymerase;

*DNA polymerase is incorrect
Ignore references to RNA dependent or DNA dependent
Allow phonetic spelling*

**1**

(b)     (i)      (Receptor / transcription factor) binds to promoter which stimulates RNA polymerase / enzyme X;

Transcribes gene / increase transcription;

**2**

(ii)     Other cells do not have the / oestrogen / ERα receptors;

*But do not accept receptors in general.*

**1**

(c)     Similar shape to oestrogen;

Binds receptor / prevents oestrogen binding;

Receptor not activated / will not attach to promoter / no transcription;

*Accept alternative
Complementary to oestrogen;
Binds to oestrogen;
Will not fit receptor;*

**2 max**

**[6]**

**M27.**         (a)     (i)     UGC;

**1**

(ii)     TGCTAC;

**1**

(b)     (DNA) contains introns / non-coding bases / mRNA only contains exons / coding bases;

*Assume that ‘it’ refers to DNA*

*Neutral: DNA contains introns and exons*

*Neutral: ‘splicing’*

*Neutral: pre-mRNA contains introns*

*Ignore refs. to start and stop codons*

**1**

(c)     Different primary structure / amino acid sequence / amino acid coded for;

*Reject: different amino acids produced / formed*

*Neutral: refs. to bonds*

**1**

(d)     1.      Acetylcholine not broken down / stays bound to receptor;

2.      Na+ ions (continue to) enter / (continued) depolarisation / Na+ channels (kept) open / action potentials / impulses fired (continuously);

3.      (Intercostal) muscles stay contracted / cannot relax;

*‘Muscles contract’ is not enough*

*Accept: diaphragm stays contracted / cannot relax*

**3**

**[7]**

**M28.**          (a)     Introns;

**1**

(b)     Ile Gly Val Ser;

**1**

(c)     (i)      Has no effect / same amino acid (sequence) / same
primary structure;

***Q*** *Reject same amino acid formed or produced.*

**1**

Glycine named as same amino acid;

**1**

*It still codes for glycine = two marks.*

(ii)     Leu replaces Val / change in amino acid (sequence) / primary structure;

Change in hydrogen / ionic bonds which alters tertiary structure / active site;

***Q*** *Different amino acid formed or produced negates first marking point.*

Substrate cannot bind / no longer complementary /
no enzyme-substrate complexes form;

*Active site changed must be clear for third marking point but does not need reference to shape.*

**3**

(d)     (i)      Interphase / S / synthesis (phase);

**1**

(ii)     DNA / gene replication / synthesis occurs / longest stage;

*Allow ‘genetic information’ = DNA.*

*Allow ‘copied’ or ‘formed’ = replication / synthesis*

**1**

**[9]**

**M29.**          (a)     1.      Hydrolysis breaks proteins / hydrolyses proteins / produces amino acids (from proteins);

2.      Protein synthesis involves condensation;

**2**

(b)     Amino acids (from calliphorin) can be joined in different sequences /
rearranged;

**1**

(c)     1.      Fall, rise and fall;

2.      Rise after 40 and fall after 80;

*Ignore concentration values.*

**2**

(d)     (i)      Fall / increase then fall;

Lysosomes associated with tissue breakdown;

**2**

(ii)     1.      Tissues / cells are being broken down;

2.      RNA is digested / hydrolysed / broken down;

3.      By enzymes from lysosomes;

4.      New proteins not made / no new RNA made;

**2 max**

(e)     1.      (RNA) associated with making protein;

2.      New / adult tissues are forming;

**2**

(f)      1.      In the first 6 days no / little oxygen supplied / with breakdown of tracheae, no / little oxygen supplied;

2.      (Without tracheae) respire anaerobically;

3.      Anaerobic respiration involves reactions catalysed by enzyme **B**  / conversion of pyruvate to lactate / involves lactate production;

4.      Enzyme **A** / Krebs cycle is part of aerobic respiration;

*Or, with emphasis on aerobic respiration:*

*1. Tracheae supply oxygen / after 6 days oxygen supplied;*

*2. (With tracheae) tissues can respire aerobically.*

**4**

**[15]**

**M30.**(a)     Banding pattern changes as cheetah gets older / difficult to judge as tail is short / fluffy;

**1**

(b)     (i)      Mean not (always) a whole number;
Standard deviation not (always) zero;

**2**

(ii)     Movement of tail / angle of sight / confused it with another band / subjective estimation;

*Accept reference to* ***Figure 1***

*E.g. Bands 2 and 3 have same thickness but look different*

**1**

(c)     Band width not the same on both sides of tail;

**1**

(d)     Offspring of the same family will be more similar genetically;
As have same mother (and father) / parent;
Expect to see more differences in randomly chosen cheetahs;

**3**

**[8]**

**M31.**          (a)     (i)      Phosphate and ribose;

*Accept in either order. Both correct for one mark.*

*For phosphate accept PO4 / Pi /  but not P.*

*Do not accept phosphorus.*

*Ignore references to pentose / sugar.*

**1**

(ii)     TAGGCA;

**1**

(b)     (i)      Does not contain hydrogen bonds / base pairs / contains
codons / does not contain anticodon / straight / not folded / no
amino acid binding site / longer;

*Assume that “it” refers to mRNA.*

*Do not accept double stranded.*

**1**

(ii)     (pre-mRNA) contains introns / mRNA contains only exons;

*Assume that “it” refers to pre-mRNA.*

*Accept non-coding as equivalent to intron.*

**1**

(c)     (i)

|  |  |
| --- | --- |
| **Part of chromosome** | **U** |
| Middle | 18 |
| End | 21 |

*One mark for both figures correct*

**1**

(ii)     1.      Have different (base) sequences / combinations of (bases);

2.      (Pre-mRNA) transcribed from different DNA / codes for different proteins;

**2**

**[7]**

**M32.**          (a)     (i)      9;

*Accept: nine*

**1**

(ii)     Introns / non-coding DNA / junk DNA;

Start / stop code / triplet;

*Neutral: Repeats.*

*Accept: ‘Introns and exons present’.*

*Reject: ‘Due to exons’.*

**1 max**

(b)     Change in amino acid / s / primary structure;

Change in hydrogen / ionic / disulfide bonds;

Alters tertiary structure;

*Reject: ‘Different amino acid is formed’ – negates first marking point.*

*Neutral: Reference to active site.*

**3**

(c)     Number of bases

|  |  |
| --- | --- |
|   | Number of bases |
| C | G | A | T |
| Strand A | 26 | **19** | **20** | **9** |
| Strand B | 19 | **26** | 9 | **20** |

Second column correct;

Columns three and four correct;

**2**

**[7]**

**M33.**(a)     (i)     Repeating units / nucleotides / monomer / molecules;

*Allow more than one, but reject two*

**1**

(ii)     1.      C = hydrogen bonds;

2.      D = deoxyribose;

*Ignore sugar*

3.      E = phosphate;

*Ignore phosphorus, Ignore molecule*

**3**

(iii)

|  |  |  |
| --- | --- | --- |
|   | **Name of base** | **Percentage** |
|   | Thymine | 34 |
|   | Cytosine / Guanine | 16 |
|   | Adenine | 34 |
|   | Cytosine / Guanine | 16 |

*Spelling must be correct to gain MP1*

*First mark = names correct*

*Second mark = % correct, with adenine as 34%*

**2**

(b)     (i)     153;

**1**

(ii)     Some regions of the gene are non-coding / introns / start / stop code / triplet / there are two DNA strands;

*Allow addition mutation*

*Ignore unqualified reference to mutation*

*Accept reference to introns and exons if given together*

*Ignore ‘junk’ DNA / multiple repeats*

**1**

**[8]**

**M34.**(a)     One / an amino acid (can be) coded for by more than one triplet;

*Accept codon for triplet*

*Accept description of triplet − three bases / nucleotides*

**1**

(b)     1.      Triplet / three bases on mRNA;

*1. Accept nucleotide for base*

*1. Accept DNA for mRNA*

*1. Ignore references to RNA unqualified*

2.      That code for an amino acid;

*2. Accept code for stop / start*

**2**

(c)     (i)       To join nucleotides together to form mRNA / premRNA / RNA;

*Reject forming base pairs*

*Accept checking and correcting mismatched base pairs*

**1**

(ii)     Reverse transcriptase;

*If they give two enzymes, no mark*

**1**

(d)     GGATCC same as CCTAGG in opposite direction;

*Accept reads same both ways / same forward and back*

*Neutral bases are the opposite of each other / reference to base pairs*

**1**

**[6]**

**M35.**(a)     250 000;

**1**

(b)     (i)      Loss of 3 bases / triplet = 2 marks;;

*‘Stop codon / code formed’ = 1 mark max unless related to the last amino acid*

Loss of base(s) = 1 mark;

*eg triplet for last amino acid is changed to a stop codon / code = 2 marks*

*3 bases / triplet forms an intron = 2 marks*

*Accept: descriptions for ‘intron’ eg non-coding DNA*

*‘Loss of codon’ = 2 marks*

**2**

(ii)     1.      Change in tertiary structure / active site;

*Neutral: change in 3D shape / structure*

2.      (So) faulty / non-functional protein / enzyme;

*Accept: reference to examples of loss of function eg fewer E-S complexes formed*

**2**

**[5]**

**M36.**(a)     (i)      1.      (Tumour suppressor) gene inactivated / not able to control / slow down cell division;

*Ignore: references to growth*

2.      Rate of cell division too fast / out of control.

*1 and 2 Accept: mitosis*

*1 and 2 Reject: meiosis*

**2**

(ii)     1.      (Genetic) code degenerate;

*Accept: codon for triplet*

*Accept description of degenerate code, e.g. another triplet codes for the same amino acid*

2.      Mutation in intron.

*Accept: mutation in non-coding DNA*

**1 max**

(b)     1.      Antibody has specific tertiary structure / binding site / variable region;

*Do not accept explanations involving undefined antigen*

2.      Complementary (shape / fit) to receptor protein / GF / binds to receptor protein / to GF;

*Ignore: same shape as receptor protein / GF*

3.      Prevents GF binding (to receptor).

**3**

**[6]**

**M37.**(a)      1.      Helicase;

2.      Breaks hydrogen bonds;

3.      Only one DNA strand acts as template;

4.      RNA nucleotides attracted to exposed bases;

5.      (Attraction) according to base pairing rule;

6.      RNA polymerase joins (RNA) nucleotides together;

7.      Pre-mRNA spliced to remove introns.

**6 max**

(b)     1.      Polymer of amino acids;

2.      Joined by peptide bonds;

3.      Formed by condensation;

4.      Primary structure is order of amino acids;

5.      Secondary structure is folding of polypeptide chain due to hydrogen bonding;

*Accept alpha helix / pleated sheet*

6.      Tertiary structure is 3-D folding due to hydrogen bonding and ionic / disulfide bonds;

7.      Quaternary structure is two or more polypeptide chains.

**5 max**

(c)     1.      Hydrolysis of peptide bonds;

2.      Endopeptidases break polypeptides into smaller peptide chains;

3.      Exopeptidases remove terminal amino acids;

4.      Dipeptidases hydrolyse / break down dipeptides into amino acids.

**4**

**[15]**

**M38.**(a)     1.      Sugar-phosphate (backbone) / double stranded / helix **so** provides strength / stability / protects bases / protects hydrogen bonds;

*Must be a direct link / obvious to get the mark*

*Neutral: reference to histones*

2.      Long / large molecule **so** can store lots of information;

3.      Helix / coiled **so** compact;

*Accept: can store in a small amount of space for ‘compact’*

4.      Base sequence allows information to be stored / base sequence codes for amino acids / protein;

*Accept: base sequence allows transcription*

5.      Double stranded **so** replication can occur semi-conservatively / strands can act as templates / complementary base pairing / A-T and G-C so accurate replication / identical copies can be made;

6.      (Weak) hydrogen bonds **for** replication / unzipping / strand separation / many hydrogen bonds **so** stable / strong;

*Accept: 'H-bonds' for ‘hydrogen bonds’*

**6**

(b)     1.      (Mutation) in **E** produces highest risk / 1.78;

2.      (Mutation) in **D** produces next highest risk / 1.45;

3.      (Mutation) in **C** produces least risk / 1.30;

*Must be stated directly and not implied*

***E*** *>* ***D*** *>* ***C*** *= 3 marks*

*Accept: values of 0.78, 0.45 and 0.30 for MP1, MP2 and MP3 respectively*

*If no mark is awarded, a principle mark can be given for the idea that all mutant alleles increase the risk*

**3**

(c)     **180**;

**1**

(d)     **(Similarities):**

1.      Same / similar pattern / both decrease, stay the same then increase;

2.      Number of cells stays the same for same length of time;

*Ignore: wrong days stated*

**(Differences):**

(Per unit volume of blood)

3.      Greater / faster decrease in number of healthy cells / more healthy cells killed / healthy cells killed faster;

*Accept: converse for cancer cells*

*Accept: greater percentage decrease in number of cancer cells / greater proportion of cancer cells killed*

4.      Greater / faster increase in number of healthy cells / more healthy cells replaced / divide / healthy cells replaced / divide faster;

*Accept: converse for cancer cells*

*For* ***differences****, statements made must be comparative*

**3 max**

(e)     1.      More / too many healthy cells killed;

2.      (So) will take time to replace / increase in number;

*Neutral: will take time to ‘repair’*

3.      Person may die / have side effects;

**2 max**

**[15]**

**M39.**(a)     (i)      (In all organisms / DNA,) the same triplet codes for the same amino acid;

*Accept codon / same three bases / nucleotides*

*Accept plurals if both triplets and amino acids*

*Reject triplets code for an amino acid*

*Reject reference to producing amino acid*

**1**

(ii)     64;

**1**

(b)     Splicing;

*Ignore deletion references*

*Accept RNA splicing*

**1**

(c)     (i)      1.      (Mutation) changes triplets / codons after that point / causes frame shift;

*Accept changes splicing site*

*Ignore changes in sequence of nucleotides / bases*

2.      Changes amino acid sequence (after this) / codes for different amino acids (after this);

*Accept changes primary structure*

*Reject changes amino acid formed / one amino acid changed*

3.      Affects hydrogen / ionic / sulfur bond (not peptide bond);

4.      Changes tertiary structure of protein (so non-functional);

*Neutral 3-D structure*

**3 max**

(ii)     1.      Intron non-coding (DNA) / only exons coding;

*Context is the* ***intron***

*Do not mix and match from alternatives*

*Neutral references to introns removed during splicing*

*1.and 2. Ignore ref. to code degenerate and get same / different amino acid in sequence*

2.      (So) not translated / no change in mRNA produced / no effect (on protein) / no effect on amino acid sequence;

*Accept does not code for amino acids*

***OR***

3.      Prevents / changes splicing;

4.      (So) faulty mRNA formed;

*Accept exons not joined together / introns not removed*

5.      Get different amino acid sequence;

**2 max**

**[8]**

**M40.**(a)      1.      Reduction in ATP production by aerobic respiration;

2.      Less force generated because fewer actin and myosin interactions in muscle;

3.      Fatigue caused by lactate from anaerobic respiration.

**3**

(b)     Couple **A**,

1.      Mutation in mitochondrial DNA / DNA of mitochondrion affected;

2.      All children got affected mitochondria from mother;

3.      (Probably mutation) during formation of mother’s ovary / eggs;

Couple **B**,

4.      Mutation in nuclear gene / DNA in nucleus affected;

5.      Parents heterozygous;

6.      Expect 1 in 4 homozygous affected.

**4 max**

(c)     1.      Change to tRNA leads to wrong amino acid being incorporated into protein;

2.      Tertiary structure (of protein) changed;

3.      Protein required for oxidative phosphorylation / the Krebs cycle, so less / no ATP made.

**3**

(d)     1.      Mitochondria / aerobic respiration not producing much / any ATP;

2.      (With MD) increased use of ATP supplied by increase in anaerobic respiration;

3.      More lactate produced and leaves muscle by (facilitated) diffusion.

**3**

(e)     1.      Enough DNA using PCR;

2.      Compare DNA sequence with ‘normal’ DNA.

**2**

**[15]**

**E2.**(a)    In this part, many students expressed themselves unsatisfactorily and there were a number of misconceptions; almost half failed to score. Some simply said that a transcription factor affects transcription. Others attempted to use oestrogen as an example of a transcription factor, rather than a hormone that binds to a receptor to form a transcription factor. Yet others failed to state that a transcription factor binds to a *specific* site (or sites) on DNA (however expressed). Only about 20% of students obtained both marks.

(b)     There were some very good, clear and concise answers to this part that obtained all three marks. All the points on the mark scheme were seen but perhaps the commonest observations were that binding of huntingtin to CREB stops production of the protein (that removes protons and electrons from reduced NAD). This stops / slows the electron transfer chain. This leads to not enough ATP being produced and the nerve cells die. A lot of students did not read the stem carefully and thought that CREB removes electrons and protons. Others made general references to respiration slowing but did not mention the electron transfer chain, or proton gradients, or ATP production.

(c)     It was pleasing in this part to find many students connecting the requirement for two carriers for CREB to the two membranes of a mitochondrion; just over 50% obtained both marks. Some students had problems expressing themselves and others seemed genuinely confused about the membranes of the mitochondrion; for example, some wrote about a membrane round the cytoplasm and then the cristae.

**E3.**          Quite a large number of candidates failed to perform well on this question because of poor use of language and terminology in parts (a) and (b).

(a)     Most candidates obtained the mark for the idea that the mutation alters the amino acid sequence in Factor VIII protein. Good candidates then related this to a change in the tertiary structure (or three-dimensional shape) of the protein. Poorer candidates made vague references to changes in the protein, or it not being able to work.

(b)     Most candidates scored one mark for the idea that the faulty Factor VIII leads to a failure of the activation of Factor X and blocking of the clotting pathway. Fewer candidates were able to give a reasonable description of Factor X not being activated, because non-functional Factor VIII cannot work with Factor IX. Weaker candidates misunderstood the diagram and thought that Factor IX and Factor VIII were used to make Factor X.

(c)     Only the best candidates understood that the blood from each haemophiliac contained the functional factor that the other lacked. Some who understood this only gained one mark because they simply stated that the mixture contained both factors. The examiners wanted a clear statement for the second mark that blood from the boy with faulty Factor IX contained working Factor VIII and the blood from the haemophiliac with faulty Factor VIII contained working Factor IX. A common misconception was that the mutations to the genes for Factor VIII and Factor IX would produce proteins that were able to interact, because they were both the products of mutation.

**E4.**          (a)     (i)      Most candidates calculated the number of bases correctly. Many obtained a second mark for explaining that three bases on DNA (a triplet) code for each amino acid. A significant number did not get the second mark, because they stated or implied that bases are amino acids.

(ii)     This proved very difficult for many candidates. Quite a large number obtained one mark for suggesting a deletion mutation but only the best candidates went on to suggest the deletion of a triplet of bases. Some candidates continued a misconception evident in (i) and stated that an amino acid was deleted. Credit was given to candidates who suggested another type of mutation affecting a terminal amino acid.

(b)     There were many good answers to this question and many candidates obtained all four marks. Surprisingly few candidates made clear reference to complementary base-pairing between codon and anti-codon.

**E5.**          In (a), nearly all candidates were able to describe two differences between the structure of DNA and RNA. The answers to (b) and (c) were polarised; either candidates knew the relevant material and scored well, or they produced very weak answers. In (b), weaker candidates used amino acids and nitrogenous bases, and gene and triplet completely interchangeably. Very few mentioned the base sequence on DNA being copied to (messenger) RNA. In (c), many candidates did not specify that homologous chromosomes form a pair and, of those who did, too many just wrote ‘they are identical’ with no mention of genes and positioning (loci).

**E6.**          In (a), over 80% of candidates correctly stated two differences between DNA and RNA, all alternatives in the mark scheme being seen frequently. A few candidates failed to score because they mismatched features across the two columns, e.g., giving a base for DNA opposite a sugar for RNA.

Part (b) discriminated well between candidates, with a fairly even spread of marks from three down to zero. Many candidates knew that RNA is copied from DNA, or that it moves from the nucleus to the ribosomes, but relatively few were able to give a clear or correct account of coding.

**E7.**This question involved epistasis and was based on the effect of two different enzymes in a biochemical pathway determining flower colour. Those candidates who were confident about genetics found little difficulty in gaining all 15 available marks in this question. In some cases, however, candidates answered in terms of monohybrid inheritance, even though two separate genes were clearly implicated. Some candidates gave answers in terms of multiple alleles, or even sex linkage, in which it appeared that they had confused the concepts of allele and gene.

In part (a), candidates could receive full credit by using a number of different approaches to explain how the two genes were involved in producing differently coloured flowers. Some candidates concentrated purely on the alleles and their effect on the two enzymes and the pigments produced, whereas others tackled the question only in terms of the flower colour produced by different genotypes. To gain all six marks available it was necessary for candidates to explain aspects of both.

In part (b)(i), the parents were Aabb (red-flowered) and aaBB (white-flowered). Good answers to part (b)(ii) showed the purple-flowered offspring as AaBb, and proceeded to derive the offspring 2 ratio of 9:3:4, purple: red: white, showing appropriate working. Examiners were surprised and pleased to find that it was not impossible for candidates who had achieved no marks in (b)(i), having offered answers that were totally wrong, to recover and gain full credit in (b)(ii).

In part (c)(i), three different genotypes, aaBB, aaBb, and aabb, were required for the single mark. A proportion of candidates failed to score because they omitted one of these. In part (c)(ii), candidates gained credit by suggesting the addition of the red pigment, K, to extracts of each homozygous type of white petal, going on to achieve full marks for explaining that the extract of the aaBB petal would contain enzyme 2 and that this would catalyse the conversion of K to L, turning the extract purple. In contrast, the other white petal from the aabb flowers would remain red after K had been added. The most commonly encountered wrong approach was for the petals somehow to be ‘crossed’.

Genetics questions, as a rule at this level, tend to produce extremes of either very low or very high marks, depending on candidates’ understanding of the topic. This question, in requiring quite different skills in its three main parts, proved an exception to this, with many intermediate scores being achieved as well. Where candidates understood the concept of dihybrid inheritance and associated enzymes, answers were very good indeed and showed an improvement in the General standard seen at Advanced level in recent years.

**E8.**          Many candidates who had done very well throughout the paper displayed surprising gaps in factual knowledge in this question, or expressed themselves very poorly.

(i)      This was well answered by the majority of candidates. Some failed to gain marks because they made reference to codons, anti-codons, messenger RNA and transfer RNA but made no attempt to say where the codons or anti-codons were found. A large number of candidates wrote about ‘amino acids’ being attached to tRNA rather than making the required point that each tRNA is specific to a certain amino acid. Weaker candidates were very confused in their terminology.

(iii)     This question was only accessible to the best candidates. Weaker candidates tended to answer along the lines that the inserted gene produced an ‘anti-enzyme’ that worked in opposition to the normal enzyme. Quite a number of candidates did note that the softening gene and inserted gene mRNAs are complementary and gained one mark. Very few then linked this in some way to the idea that they might bind to each other and this would reduce synthesis of the softening enzyme. It might have helped some candidates if they had noted that this was part (ii) of a question which started in (i) with a request to describe translation, given that parts of questions are often linked to point candidates in the right direction.

**E9.**          This question produced a wide range of marks and proved to be an effective discriminator.

(a)     Answers to this question were rather disappointing, often lacking the precise details expected at Advanced level. Although some candidates referred to ‘memory cells’, many did not specify that these are T or B cells (lymphocytes). Better candidates did mention antibodies but they often failed to explain that antibodies previously produced were ineffective or that it takes time to produce new effective antibodies following infection by a new strain of the influenza virus.

(b)     In part (c)(i), although many candidates appreciated that macrophages engulf pathogens, few candidates precisely described that the antigen is then displayed on the macrophage cell membrane. An alternative mark point credited was the role of macrophages in producing cytokinins which stimulate B lymphocytes. Part (c)(ii) was generally well answered with most candidates obtaining at least two marks. Many candidates explained that the mitochondria provide ATP and that the RER or ribosomes are involved in protein synthesis. Although some candidates then linked protein synthesis to antibody production, only the best candidates provided a correct function of the Golgi body in terms of packaging and/or secreting proteins or glycoproteins.

**E10.**          This question was often poorly answered with many candidates having insufficient synoptic knowledge to answer it. The most able candidates scored well, giving complete and accurate answers.

Although most candidates scored some marks, only a minority scored full marks by giving a full account explaining how a mutation would affect protein synthesis. There was a tendency for candidates to limit their account to the changes in amino acids and protein structure, making no reference to either mRNA or tRNA. Some candidates just gave an account of protein synthesis without any reference to the change that would occur. Many who did mention changes in the protein failed to describe the nature of the changes. There was considerable confusion between bases and amino acids in describing DNA and proteins. Many candidates also stated that amino acids are made in the process.

**E11.**          Most candidates were able to score at last two marks on this question. However, very few candidates obtained maximum marks, usually because of their responses in part (d)(ii).

(a)     The majority of candidates correctly gave 387 as the answer. However, a significant number of candidates divided the number of amino acids by three and gave an incorrect answer of 43.

(b)     Part (i) caused few difficulties with the majority of candidates correctly giving CCAG as the complementary DNA sequence. A few candidates incorrectly gave CCTG. Part (ii) proved to be far more discriminating with many candidates giving seven as an answer rather than five, having failed to realise that two of the codons appeared twice.

(c)     Most candidates were able to gain at least one mark by naming a specific mutagen. Unfortunately, a significant number of candidates were not specific enough, using vague references to ‘radiation’ and/or ‘chemicals’ which were not credited. Similarly, tobacco tar was credited but ‘smoking’ was not. Better candidates had no problem obtaining both marks by naming a chemical mutagen, e.g. benzene, and a named radiation source, e.g. ultraviolet rays. Answers referring to the length of time of exposure to a mutagen and to the dosage of mutagen were also credited.

(d)     This proved to be the most difficult question on the paper. Most candidates incorrectly used a substitution mutation when determining the altered mRNA base sequence in part (i), rather than an addition and deletion. Consequently they had difficulty obtaining any marks for part (ii). Although approximately a third of candidates provided the correct mRNA base sequence in part (i), few of them gained more than a single mark in part (ii). A common error was to refer to amino acid mutations rather than to base mutations. Many candidates in part (ii) referred to RNA nucleotides rather than to DNA nucleotides, which restricted their maximum mark.

**E12.**          This question produced a very wide spread of marks. Candidates frequently failed to gain marks through their inability to select appropriate information to answer the specific question asked. This particularly applied to part (b) (ii). Again, inaccurate use of terminology compromised the marks gained by many candidates.

(a)     The majority of candidates gained the mark, UV light being the most popular response. Vague reference to cigarettes or tar, without further qualification, did not gain credit.

(b)     In part (i), most candidates recognised the loss of a base and the frame shift occurring in consequence and gained both marks. Weaker candidates confused the change in base sequence with amino acid sequence, seemingly unaware of the distinction between the two. Very few candidates scored full marks in part (ii) and a substantial minority gained only one. The most common point to gain credit was reference to the enzyme.s inability to function. Weaker candidates wrote in general terms about enzyme function and did not specifically refer to enzyme 1 in the question. Again, as in section (i), some candidates confused the structure of a gene with the structure of a protein and gained no marks. The change to the mRNA was rarely mentioned and descriptions of alteration to tertiary shape were too often vague and imprecise to gain credit. In part (iii), any candidates interchanged the lilac and blue colour when completing the table. Errors also included ‘no pigment’ and ‘albino’ for the unlabelled white petal.

**E13.**          (a)     Although more demanding than the others, it was pleasing to find that many candidates performed well on this question, and that a considerable number gained at least 6 or 7 marks. In particular, an encouraging number managed to get to grips with part (c).

Surprisingly, this was rather poorly answered by many candidates. Often this was because they failed to use the diagram, and, for example, quite often the presence of the chloroplast was not related to photosynthesis. Very few suggested that the movements might position the cells in the optimum light intensity for photosynthesis. The commonest suggestion was that moving away from very bright light would avoid damage, more often to the eyespot than the chlorophyll. Some equated bright light with heat and suggested it would denature enzymes. Others were sidetracked into accounts of avoiding predators or referred to physiological processes that could not be seen in the diagram. Quite a few suggested that bright light was avoided so that the cells would not make too much starch or sugar and burst.

(b)     (i)      This proved more discriminating than expected, and only the better candidates simply multiplied the 900 amino acids by three.

(ii)     There was a good proportion of correct answers, and many more made a sufficiently sensible attempt to gain a mark for the principle. Some gave a figure for two flagella, rather than for each flagellum, and others made errors with the number of zeros or the indices.

(c)     (i)      A good proportion of candidates gained both marks for this part, but quite a large number failed to compare the results for puromycin with those for the control. The latter group simply described the curve for puromycin. Some weaker candidates misinterpreted the graph and stated that the flagella stopped growing longer after 20 minutes.

(ii)     This question was targeted at the more able candidates and the examiners were looking for a precise explanation of the evidence. It was, however, pleasing to note that large numbers of candidates were able to make the links between transcription and mRNA production and between translation and protein synthesis. Better candidates did point out that actinomycin D had no significant effect on regrowth of the flagella and that therefore the cells must be using existing mRNA. Fewer specified that the rate of regrowth declined after about 20 minutes in the presence of puromycin, so only some of the regrowth could be attributed to proteins already present in the cells.

**E14.**          **BYA2**

Most candidates were able to give the correct sequence in (a) (i). In (a) (ii), almost everybody gave the correct answer of nucleus, though there were some references to cytoplasm. There were fewer correct answers to (a) (iii), however, as many candidates assumed that only one stage had to be chosen. Part (b) was well answered by most. A small number of candidates gave the complementary sequence for the tRNA anticodon, UAA, rather than the name of an amino acid in (b) (i). The commonest error in (b) (ii) was to include uracil in the base sequence.

          **BYA3**

(a)     The sequence of events was familiar to most. One mistake tended to lead to others, however. Nearly everyone was able to identify the nucleus. Some appeared to think only one letter was required in part (iii), but many successfully produced all three.

(b)     This question was accessible to many although part (ii) offered more scope for errors and several did not work through all the steps needed.

**E15.**          The majority of candidates gained the marks for application of knowledge. Straightforward recall proved difficult for some, through lack of adequate preparation for the examination.

(a)     There was a very disappointing number of correct responses. The vast majority of candidates quoted ‘mRNA codons’ and gained no marks.

(b)     Candidates who had not thoroughly learnt the factual material necessary to understand protein synthesis gave confused accounts and failed to gain any marks. Many candidates gained two marks but common omissions were failure to mention the specific nature of the amino acid carried and also failure to complete the story by reference to the correct sequence on the completed polypeptide.

(c)     The vast majority of candidates gained both marks.

**E16.**          Most candidates were able to apply their knowledge and gained credit but poor expression marred the answers of the weaker candidates.

(a)     The role of RNA polymerase was not well known. There were very few answers worthy of credit. The majority of candidates described the role of RNA polymerase as catalysing complementary base pairing. Responses were often ambiguous and it was not clear if the enzyme was joining nucleotide to nucleotide along the backbone. In contrast, the vast majority could name the ribosome.

(b)     The majority of candidates gained both marks.

(c)     Well answered.

(d)     Most candidates recognised the amino acid alanine. Very few candidates scored full marks in part (ii) and a substantial minority gained only one. The most common point to gain credit was reference to mutation 1 having no effect. Weaker candidates described the change in the DNA triplet as causing a different amino acid to be made as the result of mutation 2 and then failed to relate description of the change to the polypeptide in terms of shape or tertiary structure.

**E17.**          Many candidates gained at least five marks in this question with weaker candidates scoring at least three.

(a)     Most candidates gained one mark. Cost was rarely mentioned. Answers were often vague and lacked precision.

(b)     (i)      Most identified protein synthesis. Candidates tended to just give the sequence of events rather than what would happen if they stopped. Weaker candidates just repeated the question. Some candidates were confused between transcription and translation and others wrote about DNA replication and mRNA production.

(ii)     There were many good answers but some just said ‘different sizes’.

**E18.**          **Unit 2**

(a)     Much confusion was shown here between bases, amino acids, DNA and protein. Few were able to give a clear account and many discussed translation, mRNA and tRNA and stumbled on markworthy points almost incidentally.

(b)     This usually scored two marks.

(c)     Although many produced a correct 9-letter code, several gave CCC rather than CCG, and some listed every triplet mentioned in the question.

(d)     This part of the question was not answered well. General accounts of mutation were often provided or changes other than addition of bases were given. Even those who were moving in the right direction failed to explain that the amino acid chain would only differ from this point onward. Many correctly identified the stage as interphase or the S phase but some offered translation or transcription.

(e)     Examiners commented on the precise use of language here even in papers where it was lacking elsewhere. Many started with transcription but got into their stride and gained 6 marks easily. Not all mentioned nuclear pores or the tRNA collecting more of the same amino acid. Several made creditable attempts to say that the amino acid was specific to the anticodon. Problems arose in the location of peptide bonds which were thought to join codons to ‘matching’ anticodons and a few failed to mention amino acids at all.

          **Unit 3**

In part (a), many candidates did not link the base code with different amino acids in the sequence. Weaker candidates were confused about DNA structure and referred to it containing amino acids. In (b), the correct base sequence was given by most candidates. In (c), many failed to give the correct answer, but produced a long list of triplet codes instead. In (d)(i), many did not link a change in the sequence of bases with a change in the amino acid sequence and, hence, a change in the protein. It was common to read that a change in base sequence gave a different protein without further detail. Most candidates gave the S-phase or interphase in (d)(ii) although some offered ‘transcription’ or ‘replication’. Part (e) was surprisingly badly done by many candidates. Many started with transcription, but when they reached the required part of the story, described the process erroneously. Many did not refer to mRNA leaving the nucleus via nuclear pores, nor to the fact that tRNA brings *specific* amino acids. The complementary codon/anticodon binding was well described, but many candidates described anticodons attaching to amino acids. Many referred to polypeptide bonds instead of peptide bonds. Weaker candidates confused protein synthesis with DNA replication.

**E19.**          (a)     This part of the question was often poorly answered. While errors in the first column were perhaps predictable, those not infrequently given in the second column suggested confusion between polynucleotide strands and bases or even chromosomes.

(b)     This question was marked in such a way that a candidate who made a single error was still able to gain some credit. The answers to both parts were generally sound although there were occasional errors involving giving the base sequence on the complementary DNA strand, or resulting from uncertainty over splicing.

**E20.**          (a)     (i)      Most correctly gave the coding sequence of ACG. The most common incorrect answer involved replacing A with T.

(ii)     Most identified serine correctly.

(b)     There was some good understanding of introns and many identified this correctly as the explanation for the different number of nucleotides. The main incorrect answer occurred when candidates concentrated solely on stop/start codons.

(c)     This was poorly answered by many candidates. Most had some understanding of what was happening but only the better candidates could express this in enough detail to achieve full marks.

Weak candidates failed to use terms like codon, anticodon or translation, and often simply restated the question. Many referred to tetracycline carrying a stop codon.

**E21.**          (a)     Almost all candidates scored one mark for the idea of bacteria being killed or inhibited but very few mentioned diffusion of the antibiotic. Candidates should note that two marks are rarely given for a single idea in a question of this type.

(b)     Some candidates failed to gain marks by casual references to large or larger clear areas instead of saying this was the largest.

(c)     Many candidates scored two marks but some, disregarding the question, wrote about protein synthesis or gave general statements about bactericidal or bacteriostatic compounds. Some were confused as to whether the cell wall or the cell membrane was disrupted.

**E22.**          **Unit 2**

          (a)     Two marks were scored regularly. Confusion between T and A does exist in a significant number however, giving rise to the incorrect responses of GCAAUG and GCUUTG.

(b)     In (i) the idea of editing of mRNA was well understood, although some did remove the exons and therefore retained the introns. Poor expression was in evidence here. A few thought that stop and start codons could account for the large difference in number of nucleotides. In (ii) some extremely good answers were seen. Once again many candidates failed to gain marks through poor quality of expression. Amino acids are not made from three bases and, unless qualified by referring to code, the mark was not allowed. Some believed there to be 290 amino acids, but for the correct reason. A significant number believed there to be two amino acids because they went back to the DNA sequence at the beginning of the question rather than the piece of mRNA 660 nucleotides long. This was, however, given credit.

(c)     The examiners were very surprised to find this was badly answered, even by many of the best candidates. Candidates do not appreciate the difference between structure and function. Equally they were unable to provide pairs of matching statements. Successful candidates discussed linear and clover-leafed structure or length of molecules. A major error in a significant number of responses was to imply that tRNA was only three nucleotides long. Amino acid binding sites were discussed in terms of carrying amino acids. It was obvious that many candidates mis-read the question and discussed differences between DNA and RNA, as they referred to the bases present in the molecules.

          **Unit 3**

(a)     This was a straightforward application question and most candidates gave the correct response.

(b)     Part (i) was reasonably well answered. The most common incorrect response was to explain the difference in length in terms of the start/stop codons. Responses to part (ii) showed candidates often chose to refer to the original sequence for part (a) rather than the section in part (b) to which the question actually referred. The examiners credited a correct response regardless of which section of mRNA was being used.

(c)     Only the most able candidates were able to give two structural differences. The vast majority of candidates gave a mixture of structural and functional differences.

**E23.**          (a)     Most of the more able candidates recognised that the feature labelled **M** in the diagram represented the sugar-phosphate backbone of the molecule and identified phosphate and deoxyribose as the relevant components. Others enjoyed less success and offered such suggestions as base, nucleotide or, even, hydrogen bond.

(b)     Although there were many candidates who identified the maximum number of amino acids coded by this piece of DNA as four, it was difficult to determine any pattern in the enormous range of incorrect responses. It was clear, however, that many candidates had little understanding of the concept of a triplet code.

(c)     Part (c) (i) was answered correctly by most candidates, but a substantial number were unable to make use of their responses to determine the percentage of adenine bases in part (c) (ii). The incorrect answer 86% featured frequently.

(d)     Better candidates were able to identify the principle involved here and suggest an explanation based on different base sequences coding for different proteins. This idea eluded many, however. Some clearly thought the question related to DNA hybridisation, while others attempted to derive answers from an uncertain understanding of ratios. A common problem arose from imprecise use of the term, genetic code. This should only be regarded as a base sequence coding for a specific amino acid. Answers that attempted to explain the observation described in the question in terms of changes in the genetic code of the bacteria were, therefore, clearly incorrect.

**E24.**          (a)     (i)      Most candidates correctly named part **R** as deoxyribose. Answers identifying part **R** as pentose or as a five carbon sugar were considered too imprecise due to the question clearly identifying the molecule as being DNA.

(ii)     Most candidates correctly named part **Q** as a phosphate group or as phosphoric acid. Unfortunately, some candidates incorrectly named parts **R** and **Q** the wrong way round.

(b)     Almost every candidate correctly stated ‘hydrogen bonds’.

(c)     Approximately fifty percent of candidates obtained this mark. Although there was a wide range of incorrect answers, the most common error was to divide, rather than multiply the number of amino acids by three.

(d)     Over 90 % of candidates were able to correctly work out the sequence of amino acids.

(e)     This question proved to be an effective discriminator. Most candidates gained at least one mark, often by mentioning a change in the sequence in amino acids. However, a significant number of candidates incorrectly referred to ‘different amino acids being formed’. Many of these candidates gained a second mark for describing that the active site or tertiary structure would be altered. Better candidates gained maximum marks either by linking this to enzyme-substrate complexes not being formed or to changes in hydrogen/disulfide bonds.

**E25.**          (a)     While most candidates recognised that stem cells are undifferentiated, many failed to point out that they were capable of replacing themselves. A reference to totipotency ensured this first mark, but a second could not be awarded for simply explaining what the term meant.

 (b)     The examiners were of the opinion that many candidates had encountered the word but had very little understanding of its meaning.

(c)     In part (i), only the better candidates recognised and were able to describe the relationship between the altered sequence of nucleotide bases and the consequent change in the sequence of amino acids affecting the tertiary structure of the protein. There were many general answers that offered little more information than that the “gene was disrupted” or that a “different amino acid was made.” Many candidates found part (ii) challenging and, although they were able to gain credit for a general statement relating to the uncontrolled division of cancer cells, they attributed this to insertion of the gene in the base sequence of either the tumour suppressor gene or into a proto-oncogene, thereby converting it into an oncogene.

(d)     Many candidates had clearly given careful thought to their answers and made effective use of the data provided. They usually came to the conclusion that 25% was either an unacceptably high cancer risk or that 75% of children were unaffected by cancer. Many took the view that, as the cancer could be treated, whereas SCID was likely to prove fatal, this was an acceptable trial. Answers that adopted a more general approach, describing the trial as unethical, accusing the scientists concerned of “playing God” or commenting on children being unable to make informed judgement did not gain credit.

**E26.**(a)     The answers to this question and to Question 6 (b) suggested that many candidates were uncertain as to the roles of various enzymes used in gene technology.

(b)     Although the majority of candidates clearly understood the basic idea of transcription, they tended to gain limited credit for part (i) of this question. This was largely because they failed to make effective use of the material with which they had been provided. There were few references either to binding to the promoter or to stimulation of the enzyme. In part (ii), most candidates recognised the specificity of the receptors but were not always able to address the question of why oestrogen does not affect other cells. There was much discussion of oestrogen binding to receptors, but relatively little about receptors being confined to the cells of target organs.

(c)     Most candidates recognised the molecular structures as being either complementary or similar and used this information to construct sensible suggestions about reduction in growth of breast tumours. The terminology used by many, however, suggested confusion with enzyme action.

**E27.**         (a)      (i)      Most students correctly gave the base sequence of the anticodon as **UGC**.

(ii)     Almost all students correctly gave the base sequence of DNA as **TGCTAC**. However, a minority of weaker students failed to read the stem of the question carefully enough. They replaced the thymine bases in this sequence with uracil.

(b)     Most students were aware that DNA contains introns or mRNA only contains exons. Weaker students were often let down by vague answers, such as ‘mRNA contains exons’ and ‘mRNA is spliced’.

(c)     It was widely appreciated that a change in the base sequence of mRNA could result in different amino acids being coded for or would produce a change in the primary structure of the toxin. A common response seen by a minority of weaker students was ‘different amino acids are produced’. This was not credited.

(d)     Most students gained at least one mark for stating that acetylcholine would not be broken down. However, the ability to tell the rest of the story correctly proved to be a good discriminator. Weaker students often thought that depolarisation would be prevented, resulting in the muscles not being able to contract. Some students attempted to explain this in terms of the inability to reabsorb acetylcholine into the presynaptic membrane. They thought that this would prevent any further release of acetylcholine. It was usually only better students who went on to describe the opening of sodium ion channels. Similarly, the importance of the ‘continuous’ aspect of muscle contraction was only appreciated by the best students.

**E28.**          (a)     Less than half the candidates correctly named introns as the non-coding sections of a gene.

(b)     The vast majority of candidates correctly identified the amino acid sequence.

(c)     (i)      Most candidates obtained at least one mark for stating that the amino acid sequence would not change. However, less than half the candidates gained the second mark by explaining that the new base triplet would still code for glycine.

(ii)     Most candidates gained at least one mark, often by mentioning a change in the sequence of amino acids. However, a significant number of candidates incorrectly referred to ‘different amino acids being formed’. Many candidates gained a second mark for explaining that the active site/ tertiary structure would be altered. The best candidates gained maximum marks either by linking this to enzyme-substrate complexes not being formed or to changes in hydrogen or ionic bonds.

(d)     (i)      Almost two thirds of candidates correctly identified the part of the cell cycle as being interphase or the synthesis stage. Anaphase was a common incorrect response.

(ii)     Most candidates obtained this mark, often by indicating that DNA replication occurs during interphase.

**E29.**          This question was intended to be synoptic and as such required a basic understanding of principles established in other units. There were some outstanding answers but it was also disappointing to note that there were many candidates who clearly had little idea of the functions of cell organelles or of the role of ribosomes and RNA in protein synthesis.

(a)     There were, perhaps inevitably, candidates who confused condensation and hydrolysis but most were able used the terms appropriately in the context of protein digestion and synthesis.

(b)     Those who understood protein structure usually gained credit, but almost two-thirds of all candidates made no progress here. While the most frequent problems stemmed from confusing amino acids with bases, others appeared uncertain that proteins could be digested.

(c)     Most, but by no means all, candidates identified the overall trend of decrease, increase, decrease but rather fewer supported this with data from the table relating to the age of the pupa. Where the age was quoted, it was not uncommon to see it given in days or years. A little common sense might have excluded the latter.

(d)     Answers to part (i) might have been better had more candidates distinguished between the roles of lysosomes and ribosomes. There were many responses associating an increase in lysosomes with increased protein synthesis towards the end of the time spent as a pupa.

Others linked lysosomes with disease and answered in terms of increased exposure to bacterial infection. A major misconception in the answers to part (ii) was that protein synthesis would decrease RNA concentration as it was “used up” in the process.

(e)     Although some of the candidates answering this part of the question were unable to identify the trend in the table, most recognised that tissue formation involved protein synthesis and hence the increase in RNA.

(f)      This question discriminated very effectively over the range of available marks but, at all levels of ability, candidates appeared to find difficulty with spelling the words aerobic and anaerobic. Examiners try to avoid being unnecessarily pedantic over the spelling of technical terms but the onus is on candidates to make their intentions clear, particularly when the words concerned are closely similar. A considerable number of candidates failed to equate tracheae with insect gas exchange and wrote of breathing and the lungs.

**E30.**(a)     There was widespread recognition that tail band width would be likely to change with age.

(b)     In part (a), many candidates lacked the mathematical understanding to appreciate that a mean which had a value with decimal places suggested that measurements of the same band must differ. Likewise, they did not appreciate that a standard deviation with a value other than zero indicated variation in the measurements of the same band. However in part (b), having read the description of the procedure, most recognised that viewing an animal's tail through binoculars from a moving vehicle was likely to give rise to inconsistent data.

(c)     Most candidates correctly used the data about the width of bands from the left and right sides of the tail as evidence that rings of equal width were not found.

(d)     The most frequently awarded mark was for showing an understanding that unrelated animals would be expected to show more variation than animals from the same family. It was less usual to find a link to the idea that members of one family are genetically closely related, or a reference to the animals’ parentage.

**E31.**          (a)     Most candidates named the two substances required in answer to part (i) correctly, although there were responses such as sugar and pentose that lacked the necessary precision. Part (ii) was answered correctly by most candidates.

(b)     In part (i), a few candidates attributed the properties of DNA in containing thymine being double stranded to one or other of the specified forms of RNA. Most, however, were able to explain that tRNA was folded and contained hydrogen bonds. Part (ii) was also answered well with only occasional confusion between exons and introns.

(c)     Although part (i) was answered well, less able candidates experienced considerable difficulty with part (ii). There was much confusion between chromosomes and genes and there were frequent references to stop codons being found at only the end of chromosomes. Equally worrying was the number who considered that as the base sequence on DNA was random, then the percentage of bases was also random.

**E32.**          (a)     (i)      Almost ninety percent of candidates were able to determine the maximum number of amino acids which could be coded for by the sequence of DNA bases provided.

(ii)     There was almost an equal split here between candidates who correctly referred to introns or stop/start codons and those candidates who incorrectly provided an explanation in terms of the code being degenerate.

(b)     This question proved to be a very effective discriminator. Most candidates gained at least one mark, often by mentioning a change in the sequence of amino acids. However, a significant number of candidates incorrectly referred to ‘different amino acids being formed’. Many candidates gained a second mark for explaining that the tertiary structure would be altered. Better candidates gained maximum marks either by linking this to changes in hydrogen/ionic/disulfide bonds. Candidates were not penalised for references to ‘active sites’ even though the question did not indicate that the protein was an enzyme.

(c)     Rather surprisingly, only half of the candidates gained marks on this question. Those that did gain credit usually obtained both marks by realising that it was important to match the number of complementary base sequences between strand A and strand B.

**E33.**(a)      (i)       Students were aware that polymers were made of many monomers, but in many cases went on to include descriptions in their answers that implied they did not understand what the monomers in this case were. A number of answers suggested that each strand was a monomer or that the monomers were amino acids.

(ii)      Most students knew the names of the parts of the diagram; the most common mistake was giving ‘sugar’ or ‘ribose’ instead of deoxyribose.

(iii)     In order to gain both marks, students had to show that they knew the names of the bases. This proved very revealing in that almost all knew the initial letters of the four bases but only a minority could write down the names correctly. About half were able to complete the simple calculation to give the percentage of the other three bases.

(b)     (i)       Those who failed to gain credit often did so because they were, apparently, of the opinion that one base coded for three amino acids.

(ii)      Introns, non-coding DNA, start and stop codes were all known to be non-coding DNA and, thus, adding to the length of the gene without contributing to the polypeptide. Some students also mentioned addition mutations or the fact that there are two strands. A minority of students incorrectly linked the degeneracy of the genetic code to the difference in number of bases.

**E34.**This question was intended to provide an accessible start to the paper, since it was almost entirely factual recall. In the event, it proved challenging for many students.

(a)     Just over forty percent failed to obtain the mark. Some students failed to make reference to triplets, or three bases and made statements along the lines of ‘Many bases code for the same amino acid.’ Others got the definition the wrong way round and said that one triplet codes for more than one amino acid. Quite a few got confused between the genetic code being degenerate and non-overlapping.

(b)     About thirty percent obtained both marks. Some students again failed to refer to triplets, or three bases, and some failed to say that a codon is on mRNA (we accepted DNA).

(c)     (i)      This question demonstrated that the majority of students think that RNA polymerase causes base pairing, rather than joining together nucleotides that have already base paired.

(ii)     About seventy-five percent of students correctly named the enzyme.

(d)     A similar percentage obtained the mark for an explanation of what a palindromic recognition sequence is.

**E35.**(a)     Nearly all students gave the correct answer of **250,000**.

(b)     (i)      One-third of students gained at least one mark. This question required students to apply the principle that three bases code for one amino acid to an unfamiliar context. However, other creditworthy approaches were used to explain why the faulty protein has one amino acid missing. This said, many students simply defined the term ‘mutation’ or repeated information given in the question stem. Consequently, there were many references to a *change* in the base sequence or amino acid sequence. Only the best responses mentioned a loss of bases. Students who took a different approach fell into one of two camps. Some suggested that a stop codon had formed for one mark. However, it was rare to see this related to the final amino acid of the protein. Similarly,others were clearly aware of introns but rarely mentioned that three bases may form an intron. Unfortunately, a minority of students provided a good response to (c) (ii) for this question part.

(ii)     One-third of students gained full marks. Many were aware that the protein produced could be faulty or non-functional. However, the ability to explain this in terms of a change in tertiary structure or active site discriminated well. Unfortunately, some students went no further than to state that the protein would have a different primary structure. This was given in the question stem and therefore not credited.

**E36.**(a)     (i)      The examiners wanted a statement that a mutation could make the gene inactive and that this would lead to uncontrolled, or very rapid, cell division. About half of students obtained both marks. Some students did not mention cell division but just stated that a tumour would grow; apparently taking ‘growth’ to mean cell division. The examiners did not accept these terms as equivalent. Some students got into long explanations of how a mutation could lead to a faulty protein and eventually got the first mark point for an inactive gene. Some of these failed to score because they wrote about mutations leading to the production of faulty amino acids.

(ii)     A large majority of students managed to convey the idea of the genetic code being degenerate.

(b)     Very few students obtained all three marks in this part. This was because they didn’t address the reference to ‘this antibody’ in the stem. The examiners were looking for an observation that ‘this antibody’ will have a specific tertiary structure, or binding site, or variable region. Some of those who did consider this aspect, failed to score because they referred to a specific ‘active site’. Many students obtained two marks for suggesting that the antibody binds either to the receptor (protein), or growth factor, and this prevents growth factor binding to its receptor.

**E38.**Parts (a), (b) and (d) proved to be good discriminators.

(a)     It was disappointing that only just below 40% of students scored at least half marks. This was mainly due to simply describing the structure of DNA, without explaining how these features relate to its functions. Some students wrote about DNA structure and function in different paragraphs. This made it unclear which feature went with which function, as no direct links had been made. In contrast, there were some truly excellent responses, which had clearly been well planned before putting pen to paper. The most common mark points awarded were for the sugar-phosphate backbone providing strength or protecting bases, the helix allowing the molecule to be compact, weak hydrogen bonds allowing strand separation or replication and the two strands acting as templates or allowing semi-conservative replication. Relatively few students linked complementary base pairing with accurate replication or the production of identical copies of DNA. Similarly, few students referred to DNA as a large molecule that can store lots of information, or the base sequence coding for amino acids. Weaker responses often mentioned this in the context of the genetic code being degenerate. Indeed, some students thought that the base sequence causes amino acids to be *produced*. The ability to convey that *many* hydrogen bonds provide stability was rarely seen. It was also unfortunate that a number of students wasted their time by writing about irrelevant topics such as the differences between prokaryotic and eukaryotic DNA and the role of histones. There were also some lengthy accounts of DNA replication, enzyme structure and the different levels of protein structure.

(b)     Many students scored at least two marks for stating that a mutation in gene **E** produces the highest risk and a mutation in gene **C** produces the lowest risk. However, only the best responses also referred to gene **D**. Students who did not mention any of the genes usually picked up one mark for noting that all of the mutant alleles increase the risk of lung cancer. Surprisingly, some thought that a mutation in gene **D** produces the highest risk.

(c)     Just fewer than 40% of students gave the correct answer of **180**.

(d)     Two-thirds of students scored at least two marks. Many were able to identify the decrease, plateau and increase for healthy cells and cancer cells. However, relatively few made reference to the plateau occurring for the same length of time. Students who failed to gain a mark for a similarity usually ignored the plateau. Most students spotted that a greater number of healthy cells were killed or that they experienced a faster decrease in number. Similarly, it was impressive to see that some used data from the graph to calculate that a greater *proportion* of cancer cells were killed. Many students also noted the faster increase in the number of healthy cells.

(e)     Half of students scored full marks. This was usually for mentioning that too many healthy cells would be killed, which could kill the patient or cause side effects. However, relatively few appreciated that it would take time to replace the healthy cells that had been killed.

**E39.**(a)     (i)      This part asked students why the genetic code is described as universal. Universal in this context means found in all organisms. A large percentage of students wrote that it is universal because it is found everywhere. Only a quarter of students made correct references to the triplet code used in DNA. Some had the correct idea but wrote things such as, ‘The same triplet codes for all amino acids’ and failed to score.

(ii)     50% of students gave the correct answer.

(b)     This part discriminated well, but with over 40% getting all three marks. Most stated or described the idea of a frame shift. However, some wrote that this changed the sequence of bases afterwards, rather than the sequence of codons. Another fairly common misconception was that mRNA leads to the synthesis, or formation, of amino acids.

(c)     This part proved more challenging and only about a third obtained both marks. Most correct answers revolved around the idea of introns being non-coding and thus not affecting an amino acid sequence. Students who failed to score often ignored the fact that the mutation was in an intron and wrote about possible effects of a substitution on amino acid sequences. In the figure, it clearly states that the intron is removed from pre-mRNA.