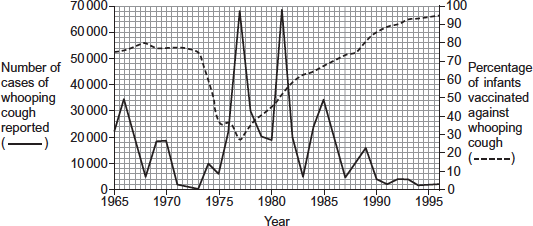
**Q1.**Whooping cough is a disease that affects some infants. Doctors collected data relating to whooping cough between 1965 and 1996.

They collected data for:

•        the number of cases of whooping cough reported

•        the percentage of infants vaccinated against whooping cough.

The graph shows the data collected by the doctors.



(a)     Suggest **two** reasons why the percentage of infants vaccinated decreased between 1973 and 1975.

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**(2)**

(b)     Between 1980 and 1990, there were three peaks in the number of reported cases of whooping cough. After 1981, the number of cases of whooping cough in each peak decreased.

Use the information from the graph to suggest why.

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**[Extra space]** ................................................................................................

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**(2)**

(c)     The percentage of the population vaccinated does **not** need to be 100% to be effective in preventing the spread of whooping cough.

Suggest why.

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**[Extra space]** ................................................................................................

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**(2)**

**(Total 6 marks)**

**Q2.**          Doctors use Zevalin to kill cancerous B-cells. Zevalin is a monoclonal antibody which has a highly radioactive substance called yttrium attached to it. The antibody binds to the surface of B-cells and the radioactivity kills the cells.

(a)     Only B-cells are killed by Zevalin.

Explain why.

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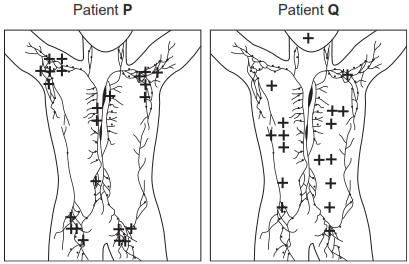
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**(2)**

The cancerous B-cells are found mainly in the lymphatic system of patients. Before treating any patient with Zevalin containing yttrium, doctors test the patient with a different form of Zevalin. This form has radioactive indium attached to the antibody instead of yttrium. The radioactivity from indium is strong enough for doctors to detect but not strong enough to kill a patient’s cells.

The diagram shows the lymphatic systems of two patients, **P** and **Q**, after being given Zevalin with indium.  The crosses (**+**) show where indium was detected.



(b)     The doctors decided they could treat Patient **P** with Zevalin containing yttrium but **not** Patient **Q**.  
Suggest why Patient **P** could be treated with Zevalin containing yttrium and Patient **Q** could not.

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**(3)**

(c)     Suggest **one** reason for the difference in distribution of the radioactivity detected in these patients.

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**(2)**

(d)     The antibody in Zevalin comes from mice. Patients are tested for antibodies against Zevalin before treatment for their cancer.  
Suggest why.

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**(2)**

**(Total 9 marks)**

**Q3.**          (a)     What is an antigen?

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**(2)**

(b)     A zookeeper was bitten by a snake. The bite contained venom which is a poison.  
He was given an injection of antivenom. This antivenom contained antibodies against this snake venom.

The antivenom did not give the zookeeper lasting protection against this snake venom. Explain why.

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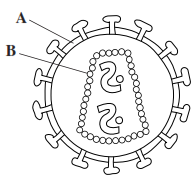
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**(2)**

**(Total 4 marks)**

**Q4.**          The diagram shows the human immunodeficiency virus (HIV).



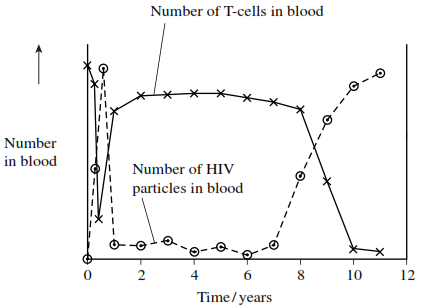
(a)     Name

**A** ..................................................................................................................

**B** ..................................................................................................................

**(2)**

The graph shows changes in the number of T-cells and HIV particles in the blood of a person following infection.



(b)     Explain why the number of HIV particles in the blood

(i)      rises during the first few months after infection

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**(2)**

(ii)     remains low between 1 and 7 years after infection.

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**(1)**

(c)     This person developed a large number of infections about 9 years after he first became infected with HIV. Using information from the graph, explain why.

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**(4)**

**(Total 9 marks)**

**Q5.**          Scientists have developed a new technique that can identify whether people smoke tobacco. Tobacco contains nicotine, which is broken down to cotinine. Cotinine is found in fingerprints. The new technique uses antibodies against cotinine.

(a)     These scientists injected laboratory mice with cotinine. Describe how this injection stimulates mice to produce antibodies against cotinine.

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**(4)**

(b)     The antibodies bind only to cotinine, and not to any other substance in the fingerprint.  
Explain why.

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**(2)**

**(Total 6 marks)**

**Q6.**          Read the following passage.

An anti-gal antibody is a type of antibody that helps to fight infections caused  
by bacteria. If a person has a bacterial infection, for example *Salmonella*, anti-gal  
antibodies bind to antigens on the surface of the *Salmonella*. Not all the  
anti-gal antibodies are used to fight the infection. Even after the infection, anti-gal  
antibodies remain in the blood.                                                                                                    5

Scientists have made adaptor molecules to try to use the anti-gal antibodies  
against viruses such as HIV. The adaptor molecules are proteins. Each adaptor  
molecule had a receptor site to which the HIV binds. This receptor site was  
similar to the receptor site on human cells to which the HIV binds. The  
adaptor molecule has another site to which an anti-gal antibody will bind.                               10

The scientists then investigated whether adding adaptor molecules and anti-gal  
antibodies can prevent HIV entering cells. They added adaptor molecules  
and anti-gal antibodies to a culture of human cells. They then added HIV  
to the culture. Their results showed that 90% of the virus particles failed  
to infect cells.                                                                                                                             15

The scientists are hoping to develop a different type of adaptor molecule to use against MRSA.

(a)     (i)      What is an antigen? (line 3)

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**(2)**

(ii)     Explain why antibodies against *Salmonella* do not normally bind to HIV.

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**(2)**

(iii)     Explain how the adaptor molecule allows anti-gal antibodies to associate with HIV.

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**(2)**

(b)     Describe how humans produce antibodies against a pathogen such as *Salmonella*.

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**(6)**

(c)     (i)      HIV infects some human cells, such as T-cells, but not others. Suggest why.

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**(2)**

(ii)     Antibiotics are **not** used to treat viral infections, such as HIV. Explain why.

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**(2)**

(d)     (i)      When HIV, anti-gal and the *adaptor molecule* were added to a culture of human cells, 90% of the virus did **not** infect human cells. (lines 12-15). Explain why.

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**(2)**

(ii)     Explain why a different type of adaptor molecule will have to be made to use against MRSA. (lines 16-17)

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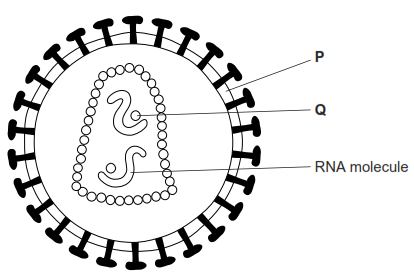
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**(2)**

**(Total 20 marks)**

**Q7.**          The diagram shows a human immunodeficiency virus (HIV).



(a)     (i)      Name structure **P** and enzyme **Q**.

Structure **P** ........................................................................................

Enzyme **Q** ........................................................................................

**(2)**

(ii)     What is the function of the RNA molecules in this virus?

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**(1)**

(b)     Describe how new viruses are produced after HIV has infected a T cell.

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**(3)**

**(Total 6 marks)**

**Q8.**          (a)     The MMR vaccine contains *attenuated* microorganisms.  
What is an *attenuated* microorganism?

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**(2)**

(b)     A child was given the MMR vaccine and was given a second dose of the vaccine as a booster later.

(i)      It took more than a week for antibodies to appear in the child’s blood after the first vaccination. Explain why.

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**(2)**

(ii)     The concentration of antibodies increased immediately after the second vaccination.  
Explain why.

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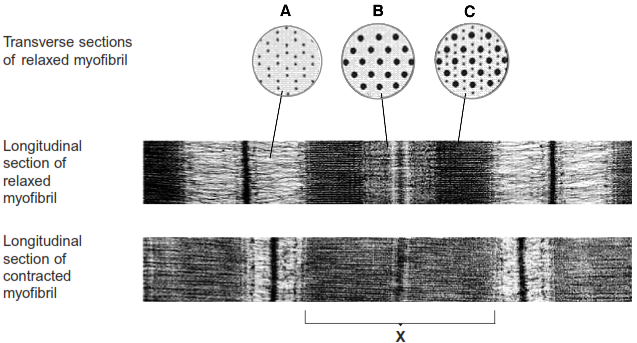
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**(2)**

**(Total 6 marks)**

**Q9.**          **Figure 1** shows sections through relaxed and contracted myofibrils of a skeletal muscle. The transverse sections are diagrams. The longitudinal sections are electron micrographs.

**Figure 1**

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(a)     (i)      The electron micrographs are magnified 40 000 times.  
Calculate the length of  band **X** in micrometres.  
Show your working.

Length of band **X** =..................................... µm

**(2)**

(ii)     Explain the difference in appearance between transverse sections **A** and **C** in **Figure 1**.

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**(1)**

(b)     Explain what leads to the differences in appearance between the relaxed myofibril and the contracted myofibril.

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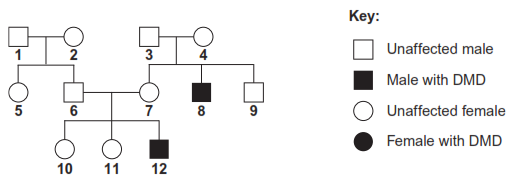
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**(4)**

(c)     Duchenne muscular dystrophy (DMD) is a condition caused by the recessive allele of a sex-linked gene. A couple have a son with DMD. They want to know the probability that they could produce another child with DMD. They consulted a genetic counsellor who produced a diagram showing the inheritance of DMD in this family.  
This is shown in **Figure 2**.

**Figure 2**

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The couple who sought genetic counselling are persons **6** and **7**.

(i)      Give the evidence to show that DMD is caused by a recessive allele.

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**(1)**

(ii)     Give the numbers of **two** people in **Figure 2** who are definitely carriers of muscular dystrophy.

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**(1)**

(iii)     Complete the genetic diagram to find the probability that the next child of couple **6** and **7** will be a son with muscular dystrophy. Use the following symbols:

**X**D= normal X chromosome  
**X**d= X chromosome carrying the allele for muscular dystrophy  
**Y** = normal Y chromosome

|  |  |  |
| --- | --- | --- |
|  | **6** | **7** |
| *Parental phenotypes* | Unaffected | Unaffected |
| *Parental genotypes* | *..............* | *..............* |
| *Gametes* | *..............* | *..............* |

*Offspring genotypes          .....................................................................*

*Offspring phenotypes        .....................................................................*

*Probability of having a son with DMD ...................................................*

**(4)**

(d)     DMD is caused by a deletion mutation in the gene for a muscle protein called dystrophin. A deletion is where part of the DNA sequence of a gene is lost. People in different families may inherit mutations in different regions of this gene.

Scientists isolated the dystrophin gene from DNA samples taken from children **10**, **11** and **12**. They cut the gene into fragments using an enzyme. The scientists then used two DNA probes to identify the presence or absence of two of these fragments, called **F** and **G**. This allowed them to find the number of copies of each fragment in the DNA of a single cell from each child.

The table shows their results.

|  |  |  |
| --- | --- | --- |
| **Child** | **Number of copies of gene fragment per cell** | |
| **F** | **G** |
| **10** (unaffected girl) | 2 | 1 |
| **11** (unaffected girl) | 2 | 2 |
| **12** (boy with DMD) | 1 | 0 |

(i)      The number of copies of gene fragments **F** and **G** shows that person **12** has DMD.  
Explain how.

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**(1)**

(ii)     The number of copies of gene fragments **F** and **G** shows that person **12** is male.  
Explain how.

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**(2)**

(iii)     The genetic counsellor examined the scientists' results. He concluded that person **10** is a carrier of DMD but her sister, **11**, is not.

Describe and explain the evidence for this in the table.

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**(3)**

(e)     Person **12** took part in a trial of a new technique to help people with DMD.

Doctors took muscle cells from person **12**’s father and grew them in tissue culture.

They suspended samples of the cultured cells in salt solution and injected them into a muscle in person **12**’s left leg. They injected an equal volume of salt solution into the corresponding muscle in his right leg. Person **12** was given drugs to suppress his immune system throughout the trial.

Four weeks later, the doctors removed a muscle sample from near the injection site in each leg. They treated these samples with fluorescent antibodies. These antibodies were specific for the polypeptide coded for by gene fragment **G** of the dystrophin gene.

The results are shown in the table.

|  |  |
| --- | --- |
| **Location and treatment** | **Percentage of muscle fibres labelled with antibody** |
| Left leg - injected with cultured cells suspended in salt solution | 6.8 |
| Right leg - injected with salt solution | 0.0 |

(i)      Why was it necessary to treat person **12** with drugs to suppress his immune system?

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**(1)**

(ii)     Explain why salt solution was injected into one leg and cultured cells suspended in salt solution into the other.

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**(1)**

(iii)     This technique is at an early stage in its development. The doctors suggested that further investigations need to be carried out to assess its usefulness for treating people with DMD.

Explain why they made this suggestion.

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**(4)**

**(Total 25 marks)**

**Q10.**          (a)     Describe how HIV is replicated after it has entered a human cell.

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**(4)**

(b)     The destruction of T-cells by HIV leads to the death of an infected person.  
Explain how.

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**(2)**

**(Total 6 marks)**

**Q11.**          **S**       A medical officer investigated the effectiveness of five different types of influenza vaccine. A total of 1350 people agreed to be vaccinated. The medical officer divided these into five groups. The number who suffered from influenza in the following year was recorded. The results are shown in the table.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
|  | **Number of people vaccinated** | | | |
| **Type of influenza vaccine** | **Suffered from influenza** | **Did not suffer from influenza** | **Total** | **Proportion suffering from influenza** |
| I | 43 | 237 | 280 | 0.15 |
| II | 52 | 198 | 250 | 0.21 |
| III | 25 | 245 | 270 | 0.09 |
| IV |  |  | 260 | 0.18 |
| V | 57 | 233 | 290 | 0.20 |

(a)     Complete the spaces in the table for the people vaccinated with type IV vaccine.

**(1)**

(b)     The medical officer used a statistical test to assess the effectiveness of the five different vaccines.

(i)      What would be the null hypothesis?

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**(1)**

(ii)     The statistical test gave a probability of less than 0.05. What conclusion can be drawn from this?

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**(1)**

(c)     It was suggested that the raw data showed that the type III vaccine was the most effective. Give **two** reasons why this conclusion may not be reliable.

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

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**(2)**

**(Total 5 marks)**

**Q12.**          (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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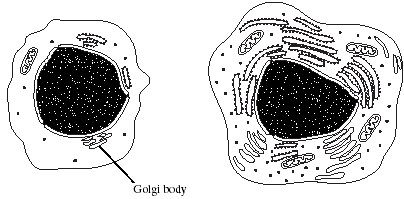
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**(2)**

(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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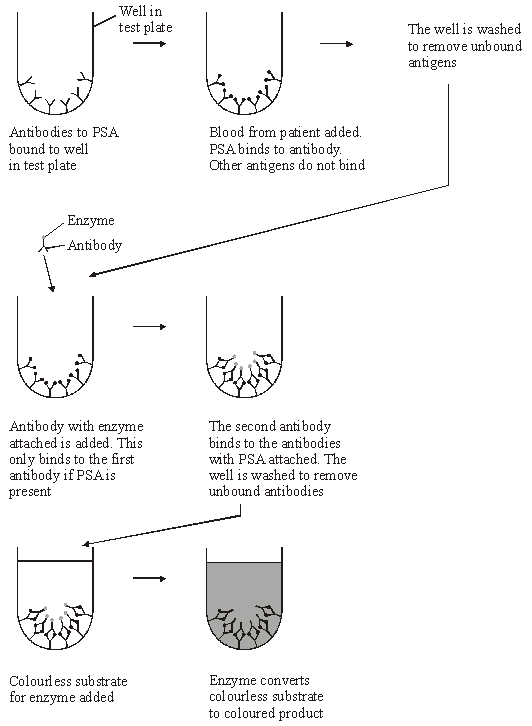
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**(4)**

**(Total 7 marks)**

**Q13.**          An antigen called PSA is present in the blood of men in the early stages of prostate cancer.

There is a blood test for PSA. The test uses monoclonal antibodies to PSA. The stages in the test are shown in the diagram.



(a)     (i)      What is an antigen?

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**(2)**

(ii)     What is a *monoclonal* antibody?

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**(2)**

(b)     (i)      Explain why this test detects prostate cancer, but not any other disease.

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**(2)**

(ii)     Explain why there will not be a colour change if the blood sample does not contain PSA.

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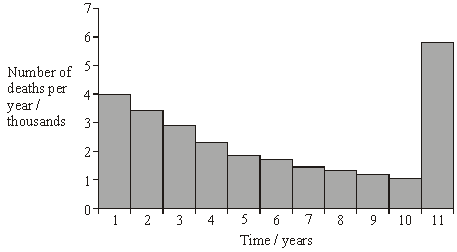
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**(2)**

**(Total 8 marks)**

**Q14.**          (a)     The graph shows the number of deaths from influenza per year in a developed country.



(i)      Suggest an explanation for the change in the number of deaths from influenza during the first 10 years.

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**(1)**

(ii)     Suggest an explanation for the large increase in the number of deaths from influenza in year 11.

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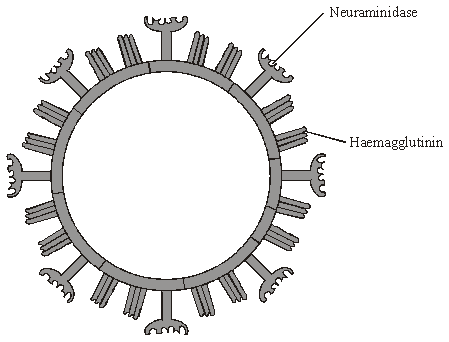
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**(2)**

(b)     The diagram shows some of the structures on the outside of an influenza virus.



Haemagglutinin and neuraminidase are protein molecules. Haemagglutinin binds to receptor molecules on the surface of epithelial cells in the breathing system.  
Neuraminidase is an enzyme which breaks down molecules in the surface membrane of epithelial cells and allows the viruses to be released from the cells.

(i)      Describe how T lymphocytes recognise and respond to the influenza virus.

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**(2)**

(ii)     Describe how B lymphocytes respond to the influenza virus.

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**(2)**

(c)     New drugs have recently become available for treating influenza. One type is a neuraminidase inhibitor. Explain how this type of drug would act as a treatment for influenza.

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**(2)**

**(Total 9 marks)**

**Q15.**          (a)     *Salmonella typhimurium* causes food poisoning in humans but not in other mammals. Explain why these bacteria attach to human cells but not to the cells of other mammals.

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**(2)**

**S**       (b)     Salmonella bacteria release toxins that cause the body temperature to rise. Although a small increase in body temperature can be beneficial, a large increase can cause serious harm.

Explain how a large increase in a person’s body temperature can cause harm.

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**(2)**

**S**       (c)     Some species of bacteria, which live in soil and decompose organic material, release exotoxins. Suggest how the release of exotoxins benefits the bacteria.

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**(1)**

(d)     Washing hands with anti-bacterial soap reduces the risk of transmission of the bacteria that cause food poisoning. Tea tree oil is a plant extract used in soaps. It is claimed to have anti-bacterial properties. Outline a method for investigating this claim.

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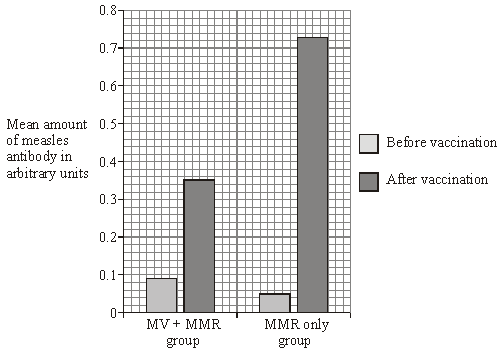
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**(4)**

**(Total 9 marks)**

**Q16.**          Measles is an infectious disease that can cause serious complications in children. In countries where measles is uncommon a combined measles, mumps and rubella vaccine (MMR) is given at 15 months. In a country where measles is common a single measles vaccine (MV) may be given at 9 months, followed by MMR at 15 months. In an investigation, the efficiency of the two vaccination programmes was compared in a country where measles is common. The amount of measles antibody in the blood of children before vaccination and after completing vaccination were measured. The graph shows the results. All difference are statistically significant.



(i)      What was the effect of vaccination in the MMR only group? Express your answer as the percentage increase in the amount of measles antibody in the MMR group after vaccination. Show your working.

Percentage increase ...................................... %

**(2)**

(ii)      The MV  +  MMR group had more measles antibodies in their blood before vaccination than the MMR only group. Suggest an explanation for this.

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**(1)**

**(Total 3 marks)**

**Q17.**          (a)     An antigen in a vaccine leads to the production of antibodies. Describe the part played by B lymphocytes in this process.

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**(4)**

**S**       (b)     Hepatitis B vaccine contains a viral antigen produced by genetically modified bacteria. Describe how the isolated gene that codes for a protein in the virus’s coat could be transferred to the bacterial cells.

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**(3)**

**(Total 7 marks)**

**Q18.**          (a)     Give **two** factors, other than cost, that should be considered when selecting an antibiotic to treat a bacterial disease.

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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)**

**Q19.** Read the following passage.

Malaria is a disease so deadly that it has devastated armies and destroyed great civilisations.  
It has been estimated that in the course of history malaria has been responsible for the death  
of one out of every two people who have ever lived. Even today, with all the advantages of  
modern technology, it is still responsible for some three million deaths a year.

5     The first half of the twentieth century was a time of hope for malarial control. The drugs

chloroquine and proguanil had just been discovered and there seemed a real possibility of a  
malaria-free world. Unfortunately, this honeymoon ended almost as soon as it had started,  
with the emergence of drug-resistant parasite populations. Scientists now accept that whatever  
new drug they come up with, it is likely to have a very limited effective life. As a result, they

10   are increasingly looking at combinations of drugs.

The approach to malaria control which holds the best hope is the production of a vaccine. One  
of these is being developed by a researcher in South America. His vaccine is based on a small  
synthetic polypeptide called SPf66 which is dissolved in a saline solution and given as an  
injection. A series of early trials on human volunteers produced confusing results. In one trial

15   the effectiveness of the vaccine was claimed to be 80% while, in others, the results were

statistically insignificant. Not only were the results inconclusive but the methods used were  
challenged by other scientists. In particular, the controls were considered inappropriate.

Another, possibly more promising, approach has been the development of a DNA-based  
vaccine. In theory, all that is required is to identify the DNA from the parasite which encodes

20   key antigens. Unfortunately, scientists have hit snags. Although they have succeeded in

sequencing the human genome, the genome of the malarial parasite has created major  
difficulties. This is partly because of the very high proportion of the bases adenine and  
thymine. In some places these two bases average 80%, and on chromosomes 2 and 3 nearly  
100% of the bases present are adenine and thymine. Because of this, it has proved impossible

25   to cut the relevant DNA with the commonly available restriction enzymes into pieces of a

suitable size for analysis.

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

(a)     Explain how a resistant parasite population is likely to arise and limit the life of any new anti-malarial drug (lines 8 - 9).

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**(3)**

(b)     A person has a 1 in 500 probability of being infected by a chloroquine-resistant strain of malarial parasite and a 1 in 500 probability of being infected by a proguanil-resistant strain. Use a calculation from these figures to explain why scientists are “increasingly looking at combinations of drugs” (lines 9 - 10).

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**(2)**

(c)     (i)      Explain why trials of the SPf66 vaccine needed a control.

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**(1)**

(ii)     The controls for the SPf66 vaccine trials were considered inappropriate (line 17).

Suggest how the control groups in these trials should have been treated.

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**(2)**

(d)     In some of the DNA of a malarial parasite, the proportion of adenine and thymine bases averages 80% (lines 22 - 23). In this DNA what percentage of the nucleotides would you expect to contain

(i)      phosphate; ..........................................................................................

(ii)     guanine? .............................................................................................

**(2)**

(e)     (i)      Use your knowledge of enzymes to explain why restriction enzymes only cut DNA at specific restriction sites.

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**(3)**

(ii)     Restriction enzymes that can cut the DNA of chromosomes 2 and 3 produce pieces that are too small for analysis. Explain why these restriction enzymes produce small DNA fragments.

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**(2)**

**(Total 15 marks)**

**Q20.** Read the following passage.

The life cycle of the malarial parasite consists of a number of stages. Some of these stages  
occur in humans and some occur in mosquitoes. At each stage, the parasite has different  
antigens on the surface of its cells. Attempts have been made to extract some of these antigens  
and use them to make vaccines to combat the disease. A trial has recently been carried out

5      with one of these vaccines. An injection of the vaccine was given to a group of people chosen  
at random at the start of the trial. Another injection was given 30 days later.

Blood samples were taken at regular intervals throughout the trial. After the first injection,  
the concentration of antibody in the blood rose slowly then fell quickly. After the second  
injection, the concentration rose quickly. It reached a maximum concentration of

10    approximately twice the concentration it reached after the first injection.

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

(a)     What is meant by *antigens* (line 3)?

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**(2)**

(b)     (i)      Use information from the passage to sketch a graph to show the effects of the two injections on the concentration of antibody in the blood.

**(3)**

(ii)     Suggest **one** reason why it was necessary to give two injections of the vaccine   
(line 6).

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**(1)**

(iii)     Although this vaccine is made from antigens from malarial parasites, it does not cause malaria. Explain why this vaccine does not cause malaria.

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**(2)**

(c)     The blood from those taking part in the trial was also examined under the microscope at the beginning of the trial. Explain how this would enable those who had malaria to be identified.

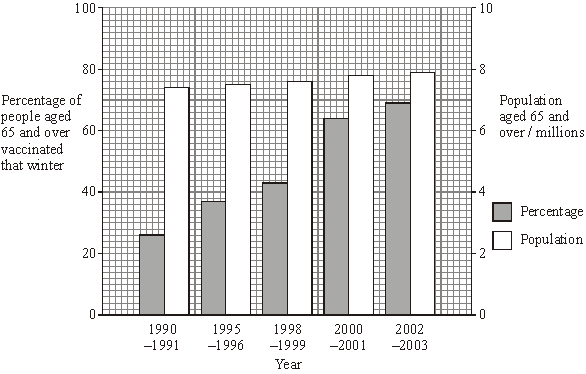
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**(1)**

**(Total 9 marks)**

**Q21.**          People considered ‘at risk’ are offered a vaccination against influenza each year. The bar chart shows the number of people in the UK population aged 65 and over and the percentage of those who were vaccinated against influenza each winter.



(a)     Suggest **one** reason to explain the change in the percentage of people aged 65 and over being vaccinated.

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**(1)**

(b)     (i)      Calculate the change in the total number of people aged 65 and over being vaccinated between 1990/91 and 2000/01. Show your working.

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

**(2)**

(ii)     A student suggested that some people aged 65 and over were being vaccinated every year. Explain how the information in the bar chart supports this suggestion.

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**(2)**

(iii)     Suggest why it is advisable for people to be vaccinated against influenza every year.

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**(2)**

(c)     An influenza virus consists of a protein coat surrounding nucleic acid. The influenza vaccine consists only of the protein coat of the virus. Explain how the influenza vaccine produces immunity in the body.

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**(2)**

**(Total 9 marks)**

**Q22.**          Some strains of the bacterium that causes gonorrhoea are resistant to antibiotics. This makes the disease difficult to treat. One way of testing the effectiveness of antibiotics is to use discs of paper soaked in antibiotic. These are placed in the centre of an agar plate covered by bacteria. A clear zone forms around the disc if the antibiotic is effective.

The table shows some results of an investigation into the effect of four different antibiotics on gonorrhoea bacteria.

|  |  |  |
| --- | --- | --- |
| **Antibiotic** | **Diameter of clear zone / mm** | **Minimum diameter of clear zone if antibiotic is effective / mm** |
| **A** | 47 | 52 |
| **B** | 30 | 28 |
| **C** | 22 | 40 |
| **D** | 33 | 34 |

(a)     Give **two** reasons why it would be important to use sterile techniques during this investigation.

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

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

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**(2)**

(b)     (i)      The antibiotic reached the bacteria by diffusion. Suggest why an effective antibiotic may produce only a small clear zone.

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**(1)**

(ii)     Which antibiotic used in the investigation would be most useful for treating gonorrhoea? Explain your answer.

Antibiotic ...............................................................

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

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**(2)**

**(Total 5 marks)**

**Q23.**          (a)     What is vaccination?

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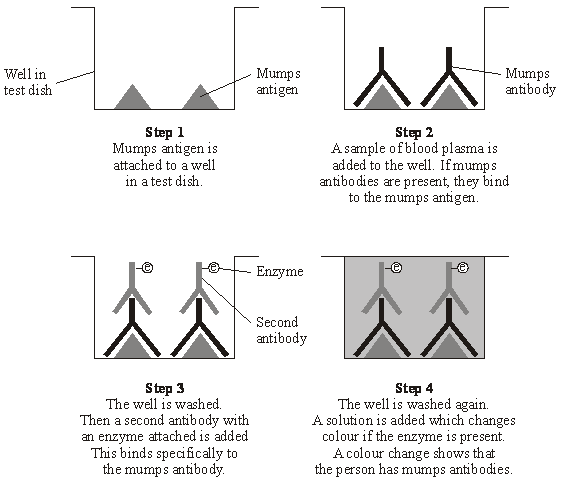
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**(2)**

(b)     A test has been developed to find out whether a person has antibodies against the mumps virus. The test is shown in the diagram.



(i)      Explain why this test will detect mumps antibodies, but not other antibodies in  
the blood.

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**(1)**

(ii)     Explain why it is important to wash the well at the start of **Step 4**.

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**(2)**

(iii)     Explain why there will be no colour change if mumps antibodies are not present in the blood.

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**(2)**

**(Total 7 marks)**

**Q24.**          Read the following passage.

Herpes viruses cause cold sores and, in some cases, genital warts. Scientists are well  
on the way to producing an antibody which will counteract herpes infection. This antibody works  
by sticking to the virus and blocking its entry into cells. It has proved very effective in animal  
tests.

5       One drawback with this approach, however, is that antibodies are at present produced using  
hamster ovary cells. This method is expensive and only produces limited amounts. A new  
technique is being developed to produce antibodies from plants. It involves introducing the  
DNA which codes for the required antibody into crop plants such as maize.

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

(a)     (i)      What is an antibody?

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**(2)**

(ii)     Describe how antibodies are produced in the body following a viral infection.

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**(6)**

(b)     Describe how the antibody gene could be isolated from an animal cell and introduced into a crop plant such as maize (lines 7-8).

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**(4)**

(c)     Taking a course of these antibodies from plants to treat a herpes infection would not produce long-term protection against disease. Explain why.

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**(2)**

(d)     Explain **one** advantage of using antibodies from plants to treat a disease, rather than antibodies produced in an experimental animal (lines 5-6).

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**(1)**

**(Total 15 marks)**

**Q25.**          The box jellyfish produces a poison (venom) which enters the blood when a person is stung. A person who has been stung can be treated with an injection of antivenom. This antivenom is produced by injecting small amounts of venom from box jellyfish into sheep, then extracting antibodies from the sheeps’ blood. These antibodies are then injected into the person who has been stung.

(a)     If a sheep is injected with the box jellyfish venom on more than one occasion a higher yield of antivenom is obtained. Explain why.

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**(2)**

(b)     Injecting antivenom does not give a person lasting protection against the venom of box jellyfish. Explain why.

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**(2)**

(c)     Suggest **one** possible problem in injecting people with antivenom made in this way.

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**(1)**

**(Total 5 marks)**

**Q26.**          A test has been developed to determine if a person is infected with variant CJD (vCJD), the human form of BSE (mad cow disease). The test detects the protein which causes vCJD in a urine sample.

The test kit contains the following components.



(a)     Complete the flow chart to describe how this test would be used.

|  |  |
| --- | --- |
|  | Urine sample is added to well in test plate |

**↓**

|  |  |
| --- | --- |
|  | Plate is washed to remove unbound vCJD protein |

**↓**

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| --- | --- |
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**(3)**

(b)     Explain why this test would detect vCJD, but not other antigens in the urine.

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**(2)**

**(Total 5 marks)**

**Q27.**          (a)     Phagocytes and lysosomes are involved in destroying microorganisms. Describe how.

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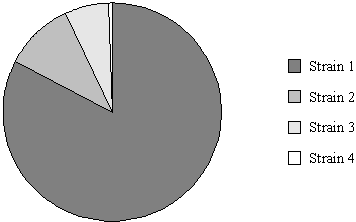
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**(3)**

(b)     The pie chart shows the proportions of people infected with four different strains of influenza virus early in 2004.



(i)      A person may develop influenza twice within a short time. Use information from the pie chart to explain why.

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**(2)**

(ii)     The information in the pie chart is valuable to companies who make influenza vaccines. Use your knowledge of antigens to explain why.

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**(2)**

**(Total 7 marks)**

**Q28.**          (a)     Describe how B-lymphocytes respond when they are stimulated by antigens.

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**(4)**

(b)     The table gives information about some components of a red blood cell.

|  |  |  |  |
| --- | --- | --- | --- |
| **Component** | Glycoprotein | Phospholipid | Haemoglobin |
| **Location in cell** | on outer surface of plasma membrane | within plasma membrane | in cytoplasm |

Suggest which component of an intact red blood cell is most likely to act as an antigen during a blood transfusion. Explain your answer.

Component ..................................................................................................

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

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**(2)**

**(Total 6 marks)**

**Q29.**          (a)     What is an antigen?

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**(2)**

(b)     Describe how B-lymphocytes respond when they are stimulated by antigens.

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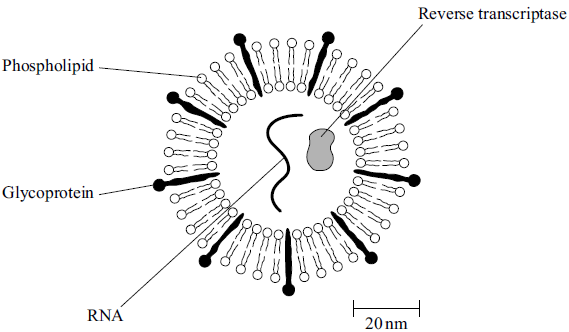
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**(4)**

(c)     The diagram shows some components of a human immunodeficiency virus (HIV).



(i)      Suggest which labelled component of the virus is most likely to act as an antigen. Give a reason for your answer.

Component .........................................................................................

Reason ...............................................................................................

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**(1)**

(ii)     A cell that HIV infects is 15 µm in diameter. Calculate how many times larger in diameter this cell is than an HIV particle. Show your working.

Answer ...................................... times larger

**(2)**

**(Total 9 marks)**

**Q30.**          Read the following passage.

|  |  |  |
| --- | --- | --- |
|  | 5        10 | *Campylobacter jejuni* is a bacterium. It is one of the commonest causes of diarrhoea in humans. The illness that it causes does not usually last very long and many sufferers do not even go to the doctor. The only treatment required is the use of oral rehydration solutions to replace the water lost by diarrhoea. In 1998, laboratory tests confirmed 60 000 cases of diarrhoea caused by this bacterium in the UK. The bacterium was more frequently found in males than in females with a ratio of 1.5 : 1.  In rare cases, the nervous system may be affected. Scientists are now beginning to understand the cause of this. Sugars in the antigens on the surface of the bacteria are identical to some of the sugars on the surface of nerve cells. Antibodies produced against the bacteria may therefore attack the body’s nerve cells. There can be serious problems if this leads to paralysis of the diaphragm. Breathing difficulties result and the patient may die. |

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

(a)     (i)      The number of cases of diarrhoea confirmed as being caused by *Campylobacter jejuni* in the UK in 1998 was 60 000 (lines 4–5). Explain why the true number of cases is thought to be more than this.

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**(1)**

(ii)     Calculate the number of cases of diarrhoea confirmed as being caused by *Campylobacter jejuni* in men in 1998.

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

**(1)**

(b)     Explain why antibodies produced against *Campylobacter jejuni* also attack nerve cells (lines 9 –10).

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**(3)**

(c)     Explain how paralysis of the diaphragm leads to breathing difficulties (line 11).

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**(2)**

**(Total 7 marks)**

**Q31.**          Read the following passage.

Several diseases are caused by inhaling asbestos fibres. Most of these  
diseases result from the build up of these tiny asbestos fibres in the lungs.

One of these diseases is asbestosis. The asbestos fibres are very small and  
enter the bronchioles and alveoli. They cause the destruction of phagocytes

5       and the surrounding lung tissue becomes scarred and fibrous. The fibrous  
tissue reduces the elasticity of the lungs and causes the alveolar walls  
to thicken. One of the main symptoms of asbestosis is shortness of breath  
caused by reduced gas exchange.

People with asbestosis are at a greater risk of developing lung cancer. The time

10      between exposure to asbestos and the occurrence of lung cancer is 20–30 years.

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

(a)     Destruction of phagocytes (lines 4–5) causes the lungs to be more susceptible to infections. Explain why.

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**(2)**

(b)     (i)      The reduced elasticity of the lungs (lines 6–7) causes breathing difficulty. Explain how.

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**(2)**

(ii)     Apart from reduced elasticity, explain how changes to the lung tissue reduce the efficiency of gas exchange.

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**(4)**

(c)     (i)      Doctors did not make the link between exposure to asbestos and an increased risk of developing lung cancer for many years. Use information in the passage to explain why.

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**(1)**

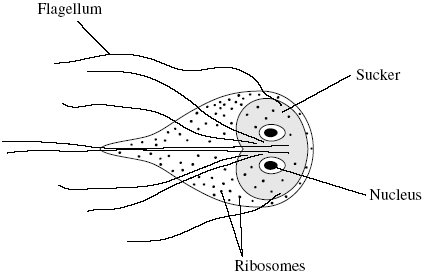
(ii)     Give **one** factor, other than asbestos, which increases the risk of developing lung cancer.

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**(1)**

**(Total 10 marks)**

**Q32.**          Giardiasis is an intestinal disease. It is caused by the microorganism *Giardia lamblia*. The drawing shows some of the structures present in *G. lamblia*.



(a)     Name **one** structure shown in the drawing which confirms that *G. lamblia* is a eukaryotic organism.

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**(1)**

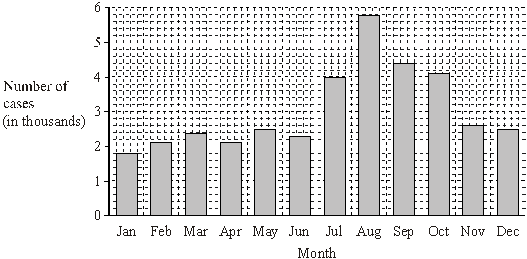
(b)     *G. lamblia* can attach itself with its sucker. Explain how this is an adaptation to living in the intestines.

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**(1)**

(c)     Giardiasis is one of the main causes of diarrhoea in the USA. It is usually transmitted by drinking contaminated water. The bar chart shows the number of cases of giardiasis in one state of the USA during one year.



(i)      Calculate the percentage increase in the number of cases of giardiasis from January to August. Show your working.

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

**(2)**

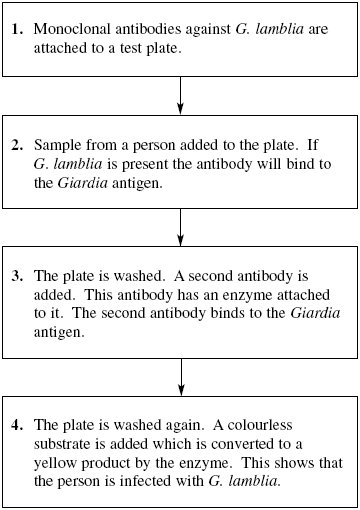
(ii)     Suggest **one** reason for the number of cases being highest in the late summer months.

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**(1)**

(d)     A test has been developed to find out whether a person is infected with *G. lamblia.* The test is shown in the flow chart.



(i)      Explain why the antibodies used in this test must be monoclonal antibodies.

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**(1)**

(ii)     Explain why the *Giardia* antigen binds to the antibody in step **2**.

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**(1)**

(iii)     The plate must be washed at the start of step **4**, otherwise a positive result could be obtained when the *Giardia* antigen is not present. Explain why a positive result could be obtained if the plate is not washed at the start of step **4**.

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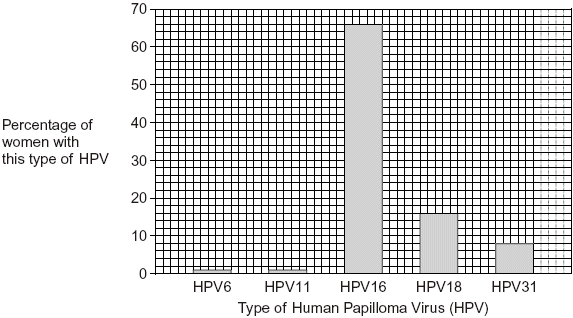
**(2)**

**(Total 9 marks)**

**Q33.**          Cervical cancer occurs in the neck of the uterus.

Scientists investigated the link between cervical cancer and infection with some types of Human Papilloma Virus (HPV).

The graph shows the frequency of five different types of HPV in women who had cervical cancer.



(a)     A local newspaper published an article about cervical cancer with the headline ‘HPV causes cervical cancer’.

Do the data shown in the graph support this claim? Explain your answer.

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**(3)**

(b)     Scientists have developed vaccines against HPV. One of the vaccines contains HPV antigens.

(i)      What is an HPV antigen?

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**(2)**

(ii)     A vaccine can be used to produce immunity to HPV. Describe how memory cells are important in this process.

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**(3)**

(c)     Some doctors suggested offering the vaccine to young men. Explain the advantage of vaccinating young men as well as young women.

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**(2)**

**(Total 10 marks)**

**Q34.**          (a)     Give **two** ways in which pathogens can cause disease when they enter the body of their host.

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

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

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**(2)**

(b)     Vaccines provide protection against disease. What is a vaccine?

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**(2)**

(c)     The only vaccine used against pulmonary tuberculosis is the BCG vaccine.  
Scientists have carried out trials on a ‘booster’ vaccine, MVA85A. This ‘booster’ vaccine is designed to increase the immune response to the BCG vaccine. One trial involved measuring the increase in the number of memory T cells in three groups of adult volunteers following different vaccination programmes.

•        Group **A** – injected with BCG

•        Group **B** – injected with MVA85A

•        Group **C** – injected with BCG and, two weeks later, injected with MVA85A

(i)      Suggest **two** factors the scientists should have considered when selecting adult volunteers for this trial.

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

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

**(2)**

(ii)     The adults in group C produced the greatest increase in the number of memory T cells. Suggest what this shows about the BCG and MVA85A vaccines.

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**(1)**

**(Total 7 marks)**

**Q35.**          Read the following passage.

Pathogens affect humans. They also affect farm animals. Once pathogens have  
entered the body of an animal they can cause disease. Vets sometimes have difficulty  
identifying the disease from which a particular animal is suffering. Until recently, they  
have had to take blood samples and send them to a laboratory. The laboratory carries

5       out tests on the sample.

New tests have been developed. Some of these new tests use monoclonal antibodies.  
Tests using monoclonal antibodies are fast, specific and allow vets to identify a disease  
while they are still on the farm.

Brucellosis is a disease of cattle. It is caused by bacteria. These bacteria can infect

10     people who drink milk or eat dairy products from infected cattle. A test using

monoclonal antibodies allows vets to identify cattle that are carriers. The carriers are  
cattle that carry the brucellosis bacteria but do not show any symptoms of the disease.

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

(a)     Other than bacteria, name **one** type of pathogen (line 1).

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**(1)**

(b)     Give **two** ways in which a pathogen may cause disease when it has entered the body (lines 1–2).

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

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

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**(2)**

(c)     Some new tests use monoclonal antibodies (lines 6–7).

(i)      Explain why these antibodies are referred to as monoclonal.

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**(1)**

(ii)     Tests using monoclonal antibodies are specific (line 7). Use your knowledge of protein structure to explain why.

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**(3)**

(d)     The tests using monoclonal antibodies allow vets to identify brucellosis while they are still on a farm. Explain the advantages of this.

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**(3)**

**(Total 10 marks)**

**Q36.**          Different cells in the body have different functions.

(a)     Some white blood cells are phagocytic. Describe how these phagocytic white blood cells destroy bacteria.

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**(4)**

(b)The epithelial cells that line the small intestine are adapted for the absorption of glucose. Explain how.

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**(6)**

**(Total 10 marks)**

**Q37.**In the early 1980s, before DNA analysis had been developed, scientists investigated the genetic variation of cheetahs living in captivity. They used skin grafts to do this. They carried out skin grafts on anaesthetised animals by

•        removing a small piece of skin from one animal. This animal was the recipient.

•        replacing the removed skin by a piece of skin taken from another animal. This animal was the donor.

•        attaching the new piece of skin with stitches.

A graft may be accepted by the recipient. It will be rejected if the recipient’s immune system recognises the antigens on the skin as foreign.

Scientists carried out skin grafts between cheetahs living in captivity and domestic cats.  
The table shows the data that they obtained.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
|  | **Recipient of skin graft** | **Donor of skin graft** | **Relationship** | **Time taken for the graft to be rejected / days** |
|  | Domestic cat 1 | Domestic cat 2 | Unrelated | 13 |
|  | Cheetah 1 | Domestic cat 3 | Unrelated | 12 |
|  | Cheetah 1 | Cheetah 2 | Sisters | No rejection after 52 days |
|  | Cheetah 3 | Cheetah 4 | Unrelated | 49 |
|  | Cheetah 5 | Cheetah 6 | Unrelated | No rejection after 78 days |
|  | Cheetah 7 | Cheetah 8 | Unrelated | No rejection after 41 days |
|  | Cheetah 9 | Cheetah 10 | Unrelated | No rejection after 24 days |
|  | Cheetah 11 | Cheetah 12 | Unrelated | No rejection after 14 days |
|  | Cheetah 13 | Cheetah 14 | Unrelated | No rejection after 44 days |

The scientists also grafted skin from one area to another on the same animal. These grafts were not rejected.

(a)     (i)      The scientists grafted skin from a domestic cat to a cheetah. Suggest why.

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**(1)**

(ii)     They also grafted skin from one area to another on the same animal. Explain why.

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**(1)**

(b)     (i)      Give **three** conclusions that you can make from the data in the table above about the time taken for rejection.

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**(3)**

(ii)     Give **one** reason why these conclusions may **not** be reliable.

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**(1)**

(iii)    There are proteins on the skin of cheetahs that act as antigens. What do the data in the table suggest about these cheetah antigens?

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**(1)**

(iv)    Antigens are proteins. Explain why a knowledge of antigens can show that animals are genetically similar.

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**(2)**

**(Total 9 marks)**

**Q38.**          Read the passage below.

Most cases of cervical cancer are caused by infection with Human Papilloma Virus  
(HPV). This virus can be spread by sexual contact. There are many types of HPV,  
each identified by a number. Most of these types are harmless but types 16 and 18  
are most likely to cause cervical cancer.

A vaccine made from HPV types 16 and 18 is offered to girls aged 12 to 13. Three

5       injections of the vaccine are given over six months. In clinical trials, the vaccine has   
proved very effective in protecting against HPV types 16 and 18. However, it will be   
many years before it can be shown that this vaccination programme has reduced   
cases of cervical cancer. Until then, smear tests will continue to be offered to

10      women, even if they have been vaccinated. A smear test allows abnormal cells

in the cervix to be identified so that they can be removed before cervical cancer   
develops.

The Department of Health has estimated that 80% of girls aged 12 to 13 need to be vaccinated to achieve herd immunity to HPV types 16 and 18. Herd immunity is where enough people have been vaccinated to reduce significantly the spread of HPV through the population.

          Use information from this passage and your own knowledge to answer the following

(a)     HPV vaccine is offered to girls aged 12 to 13 (line 5). Suggest why it is offered to this age group.

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**(1)**

(b)     The vaccine is made from HPV types 16 and 18 (line 5). Explain why this vaccine may **not** protect against other types of this virus.

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**(2)**

(c)     Three injections of the vaccine are given (lines 5 to 6). Use your knowledge of immunity to suggest why.

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**(2)**

(d)     It will be many years before it can be shown that this vaccination programme has reduced cases of cervical cancer (lines 7 to 9). Suggest **two** reasons why.

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**(2)**

(e)     Smear tests will continue to be offered to women, even if they have been vaccinated   
(lines 9 to 10). Suggest why women who have been vaccinated still need to be offered smear tests.

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**(1)**

(f)      Suggest **one** reason why vaccinating a large number of people would reduce significantly the spread of HPV through the population (lines 14 to 16).

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**(2)**

**(Total 10 marks)**

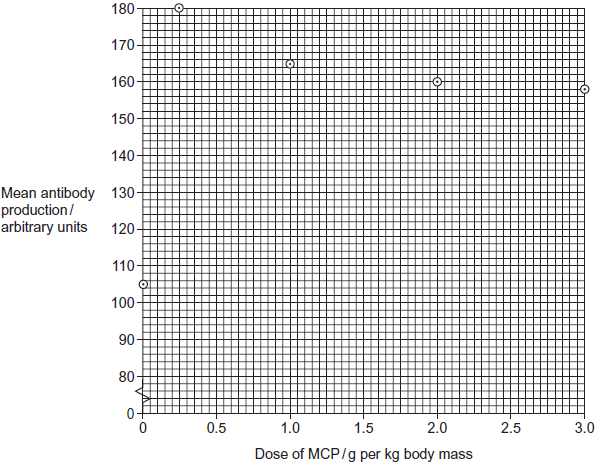
**Q39.**Scientists tested a claim that modified citrus pectin (MCP) increased the production of antibodies by the immune system.

•        They divided a large number of mice into five groups.

•        They gave the mice in each group a different amount of MCP in their food.

•        The scientists then stimulated antibody production in the mice. They did this by injecting them with a solution containing sheep red blood cells.

The results are shown in the graph.



(a)     The data obtained in this investigation have been plotted on a graph. How would you join the points? Give a reason for your answer.

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**(1)**

(b)     Use the graph to describe the effect of MCP on mean antibody production.

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**(2)**

(c)     Calculate the percentage increase in antibody production from when there was no MCP in the diet to when the dose is 1.0 g per kg.

Answer ....................................................%

**(2)**

(d)     The dose of MCP given to the mice was calculated in g per kg body mass. Explain why the dose was calculated per unit mass.

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**(1)**

(e)     Explain how antibodies were produced when the mice were injected with sheep red blood cells.

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**(3)**

(f)      A newspaper suggested that these data show that taking MCP will give people increased resistance to disease. With reference to the data give **two** reasons why this conclusion may **not** be valid.

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

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**(2)**

**(Total 11 marks)**

**Q40.**          (a)     *Clostridium difficile* is a bacterium that is present in the gut of up to 3% of healthy adults and 66% of healthy infants.

(i)*C. difficile* rarely causes problems, either in healthy adults or in infants. This is because its numbers are kept low by competition with harmless bacteria that normally live in the intestine.

Use this information to explain why some patients treated with antibiotics can be affected by *C. difficile*.

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**(2)**

(ii)     Suggest why older people are more likely to be affected by *C. difficile*.

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**(1)**

(b)The antibiotic methicillin inhibits the enzyme transpeptidase. This enzyme is used by some bacteria to join monomers together during cell wall formation. Methicillin has a similar structure to these monomers. Use this information to explain how methicillin inhibits the enzyme transpeptidase.

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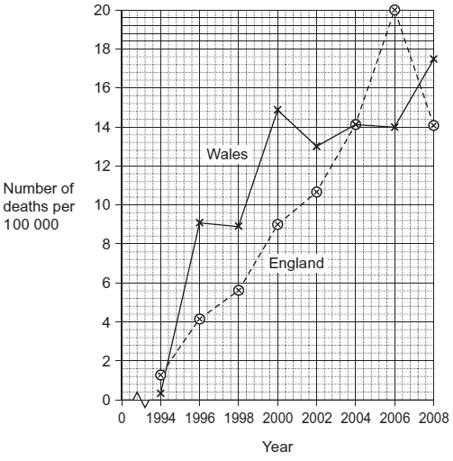
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**(2)**

(c)     MRSA is a variety of *Staphylococcus aureus*. It is difficult to treat infections caused by this bacterium because it is resistant to methicillin and to some other antibiotics. As a result, some patients who are already very ill may die if they become infected with MRSA. The graph shows the number of deaths in England and Wales between 1994 and 2008 caused by MRSA.



(i)It may be difficult to identify MRSA as the actual cause of death. Explain why.

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**(1)**

(ii)     Describe the change in the number of deaths caused by MRSA in England in the period shown in the graph.

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**(1)**

(iii)Calculate the percentage increase in the number of deaths caused by MRSA in Wales from 1996 to 2006. Show your working.

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

**(2)**

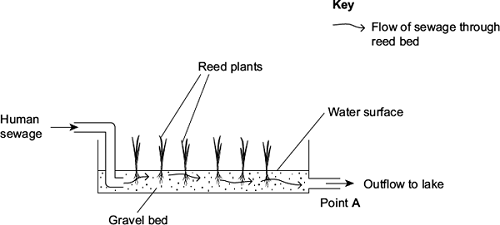
**(Total 9 marks)**

**Q41.**(a)     Name the process by which some bacteria oxidise ammonia to nitrate.

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**(1)**

Reeds are plants that grow with their roots under water. A reed bed contains a large number of growing reeds. Reed beds may be used to absorb nitrates produced when bacteria break down human sewage. The diagram shows a reed bed.



(b)     Reeds have hollow, air-filled tissue in their stems which supplies oxygen to their roots.  
Explain how this enables the roots to take up nitrogen-containing substances.

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**(2)**

(c)     (i)      There is an optimum rate at which human sewage should flow through the reed  
bed. If the flow of human sewage is too fast, the nitrate concentration at point **A** falls.  
Explain why.

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**(2)**

(ii)     An increase in nitrate concentration in the water entering the lake could affect algae and fish in the lake. Explain how.

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**(3)**

**(Total 8 marks)**

**Q42.**Vaccines protect people against disease. Explain how.

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**(Total 5 marks)**

**Q43.**          Read the following passage.

Chlamydia is a bacterium. Scientists have shown that infection with chlamydia can cause heart disease in humans. Infection with the bacterium can stimulate the formation of atheroma. This can lead to a heart attack.

Other scientists have been working with mice. These scientists have suggested that chlamydia may cause heart disease in a different way. They have found a protein on the surface of chlamydia cells which is similar to a protein in the heart muscle of mice. After an infection with chlamydia, cells of the immune system of the mice may attack their heart muscle cells and cause heart disease.

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

(a)(i)      Using information from the passage, explain what is meant by an antigen.

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**(2)**

(ii)     After an infection with chlamydia, cells of the immune system of the mice may attack the heart muscle cells (lines 7-8). Explain why.

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**(2)**

(b)Some scientists have suggested that people should be vaccinated to prevent infection by chlamydia. Evaluate this suggestion.

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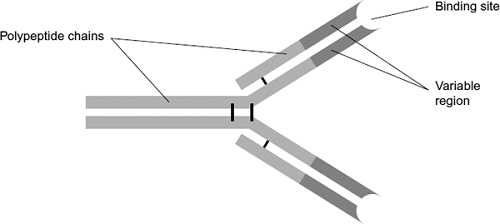
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**(3)**

**(Total 7 marks)**

**Q44.**The diagram shows an antibody molecule.



(a)     What is the evidence from the diagram that this antibody has a quaternary structure?

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**(1)**

(b)     Scientists use this antibody to detect an antigen on the bacterium that causes stomach ulcers. Explain why the antibody will only detect this antigen.

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**(3)**

**(Total 4 marks)**

**Q45.**Read the following passage.

|  |  |  |
| --- | --- | --- |
|  | Whooping cough is caused by the bacterium *Bordetella pertussis*. The first vaccines for whooping cough contained whole bacterial cells that had been heated for several minutes. Today, most vaccines only contain between one and three parts of the bacterial cells. People given whole-cell vaccines were more likely to develop harmful side effects than the people given the vaccines containing parts of the bacterial cells. Those given whole-cell vaccines produced a greater range of antibodies against the bacterium.  There have been suggestions that whooping cough vaccines may not work very well. These suggestions are due to recent reports of large 10 rises in the number of cases of whooping cough. Doctors who examined a group of patients with coughs diagnosed about 17% of them as having whooping cough. Scientists tested the blood of the same group of patients for antibodies against a toxin produced by *Bordetella pertussis*. They concluded that 4% of this group actually had whooping cough. | 5  10     15 |

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

(a)     (i)      People given whole-cell vaccines were more likely to develop harmful side effects than the people given the vaccines containing parts of the bacterial cells (lines 4–6).

Suggest reasons why.

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**(3)**

(ii)     People given whole-cell vaccines produced a greater range of antibodies against the bacterium than the people given the vaccines containing parts of the bacterial cells (lines 7–8).

Explain why.

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**(2)**

(b)     The scientists concluded from their test that 4% of patients with long-term coughs actually had whooping cough (line 15).

Explain how they used the results of their test to reach this conclusion.

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**(3)**

(c)     What does the scientists’ work suggest about reports of large rises in the number of cases of whooping cough (lines 10–11)?

Explain your answer.

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**(2)**

**(Total 10 marks)**

**Q46.**          (a)     What is a pathogen?

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**(1)**

(b)     When a pathogen enters the body it may be destroyed by phagocytosis.  
Describe how.

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**(4)**

(c)     When a pathogen causes an infection, plasma cells secrete antibodies which destroy this pathogen.

Explain why these antibodies are only effective against a specific pathogen.

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**(2)**

**(Total 7 marks)**

**Q47.**Read the following passage.

Microfold cells are found in the epithelium of the small intestine. Unlike other epithelial cells in the small intestine, microfold cells do not have adaptations for the absorption of food.

Microfold cells help to protect against pathogens that enter the intestine. They have receptor proteins on their cell-surface membranes that bind to antigens on the surface of pathogens.    5  
The microfold cells take up the antigens and transport them to cells of the immune system. Antibodies are then produced which give protection against the pathogen.

Scientists believe that it may be possible to develop vaccines that make use of microfold cells. These vaccines could be swallowed in tablet form.                                                                      10

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

(a)    (i)      Microfold cells have receptor proteins on their cell-surface membranes that bind to antigens (line 5). What is an antigen?

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**(1)**

(ii)     Microfold cells take up the antigens and transport them to cells of the immune system (lines 6-7). Antigens are not able to pass through the cell-surface membranes of other epithelial cells. Suggest **two** reasons why.

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**(2)**

(b)     Scientists believe that it may be possible to develop vaccines that make use of microfold cells (lines 9-10). Explain how this sort of vaccine would lead to a person developing immunity to a pathogen.

**(5)**

**(Total 8 marks)**

**Q48.**

|  |  |  |
| --- | --- | --- |
|  | The human immunodeficiency virus (HIV) leads to the development of acquired immunodeficiency syndrome (AIDS). Eventually, people with AIDS die because they are unable to produce an immune response to pathogens. |  |
|  | Scientists are trying to develop an effective vaccine to protect people against HIV. There are three main problems. HIV rapidly enters host cells. HIV causes the death of T cells that activate B cells. HIV shows a lot of antigenic variability. | 5 |
|  | Scientists have experimented with different types of vaccine for HIV. One type contains HIV in an inactivated form. A second type contains attenuated HIV which replicates in the body but does not kill host cells. A third type uses a different, non-pathogenic virus to carry genetic information from HIV into the person's cells. This makes the person's cells produce HIV proteins. So far, these types of vaccine have not been considered safe to use in a mass vaccination programme. | 10     15 |

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

(a)     People with AIDS die because they are unable to produce an immune response to pathogens (lines 2-4).

Explain why this leads to death.

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**(3)**

(b)     Explain why each of the following means that a vaccine might **not** be effective against HIV.

(i)      HIV rapidly enters host cells (lines 6-7).

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**(2)**

(ii)     HIV shows a lot of antigenic variability (lines 7-8).

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**(2)**

(c)     So far, these types of vaccine have not been considered safe to use in a mass vaccination programme (lines 14-15).

Suggest why they have **not** been considered safe.

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**(3)**

**(Total 10 marks)**

**Q49.**Nicotine is the addictive substance in tobacco. When nicotine reaches the brain, it binds to a specific protein. This causes the release of chemicals that give a feeling of reward to the smoker. This reward is part of the reason why people find it difficult to stop smoking.

Scientists have developed a vaccine against nicotine to help people stop smoking.  
They set up an investigation, which involved a large number of volunteers. Once a month for 5 months, one group of volunteers was given the vaccine and the other group was given a placebo.

At regular intervals, the scientists measured the concentration of antibodies to nicotine in the blood of each group of volunteers. They also calculated the percentage of volunteers who had stopped smoking from months 2 to 6 of the investigation.

(a)     (i)      In this investigation, neither the volunteers nor the scientists knew if a particular volunteer was receiving the vaccine or a placebo.

Suggest **two** reasons why this made the scientists’ results more reliable.

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

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

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**(2)**

(ii)     The scientists measured the concentration of nicotine in the blood of two volunteers who smoked the same number of cigarettes per day.

Suggest **two** reasons why the concentration of nicotine in the blood of these smokers might be different.

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**(2)**

(b)     (i)      Suggest how this vaccine could help people to stop smoking.

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**(3)**

(ii)     Some people have suggested that this vaccine should **not** be given free to smokers on the National Health Service (NHS). Evaluate this suggestion.

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**(3)**

The scientists measured the concentration of antibodies to nicotine in the blood of the volunteers for 12 months after the first vaccination. As a result of these measurements, they divided the volunteers who received the nicotine vaccine into three groups:

•        high antibody responders

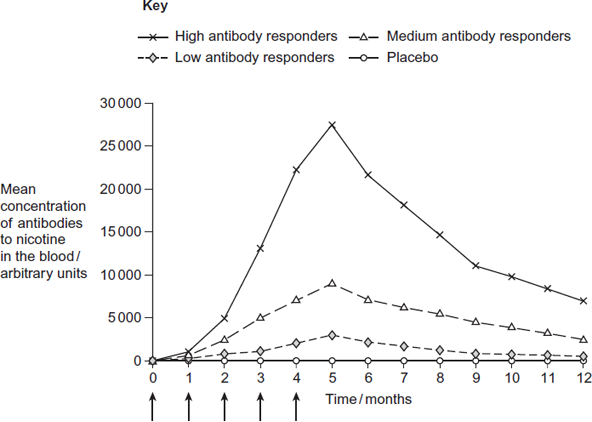
•        medium antibody responders

•        low antibody responders.

The figure below shows their results.

The scientists also recorded the number of volunteers who had stopped smoking from months 2 to 6 of the investigation.

The table below shows these results.

  
Month when vaccine or placebo was given

|  |  |  |
| --- | --- | --- |
|  | **Group** | **Percentage of volunteers who had stopped smoking from months 2 to 6 of the investigation** |
|  | High antibody responders | 56.6 |
|  | Low antibody responders | 38.1 |
|  | Medium antibody responders | 32.1 |
|  | Placebo | 31.3 |

(c)     A journalist reported that this vaccine is a major breakthrough in helping people to stop smoking. Do these data support this statement? Explain your answer.

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**(5)**

**(Total 15 marks)**

**Q50.**The figure below shows a test that has been developed to find out if a person has antibodies to the human immunodeficiency virus (HIV) antigen.

|  |  |  |
| --- | --- | --- |
|  | **Step 1** | HIV antigens are attached to a test well in a dish. |
|  |  |  |
|  | **Step 2** | A sample of blood plasma is added to the well. If HIV antibodies are present, they bind to the HIV antigen. |
|  |  |  |
|  | **Step 3** | The well is washed. A second antibody with an enzyme attached is then added. This binds specifically to the HIV antibody. |
|  |  |  |
|  | **Step 4** | The well is washed again. A yellow solution is added, which changes to blue if the enzyme is present. A blue colour shows that the person has HIV antibodies. |

(a)     This test only detects the presence of HIV antibodies. Give **two** reasons why it cannot be used to find out if a person has AIDS.

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**(2)**

(b)     The solution will remain yellow if a person is **not** infected with HIV. Explain why.

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**(2)**

(c)     A mother who was infected with HIV gave birth to a baby. The baby tested positive using this test. This does not prove the baby is infected with HIV.  
Explain why.

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**(2)**

(d)     A control well is set up every time this test is used. This is treated in exactly the same way as the test wells, except that blood plasma is replaced by a salt solution.

Use information from the figure above to suggest **two** purposes of the control well.

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**(2)**

**(Total 8 marks)**

**Q51.**(a)     When a vaccine is given to a person, it leads to the production of antibodies against a disease-causing organism. Describe how.

**(5)**

(b)     Describe the difference between active and passive immunity.

**(5)**

**(Total 10 marks)**

**Q52.**(a)     Describe how bacteria are destroyed by phagocytes.

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

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**(3)**

(b)     Give **two** structures a bacterial cell may have that a white blood cell does not have.

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

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**(2)**

**(Total 5 marks)**

**Q53.**Metastatic melanoma (MM) is a type of skin cancer. It is caused by a faulty receptor protein in cell-surface membranes. There have been no very effective treatments for this cancer.

Dacarbazine is a drug that has been used to treat MM because it appears to increase survival time for some people with MM.

Doctors investigated the use of a new drug, called ipilimumab, to treat MM. They compared the median survival time (ST) for two groups of patients treated for MM:

•        a control group of patients who had been treated with dacarbazine

•        a group of patients who had been treated with dacarbazine and ipilimumab.

The ST is how long a patient lives after diagnosis.

The doctors also recorded the percentage of patients showing a significant reduction in tumours with each treatment.

The total number of patients in the investigation was 502.

The table below shows the doctors’ results.

|  |  |  |  |
| --- | --- | --- | --- |
|  | **Treatment** | **Median survival time (ST) /  months** | **Percentage of patients  showing significant  reduction in tumours** |
|  | Dacarbazine | 9.1 | 10.3 |
|  | Dacarbazine and  ipilimumab | 11.2 | 15.2 |

(a)     The doctors compared median survival times for patients in each group.

How would you find the median survival time for a group of patients?

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**(2)**

(b)     In many trials of new drugs, a control group of patients is given a placebo that does not contain any drug.

The control group in this investigation had been treated with dacarbazine.  
Suggest why they had not been given a placebo.

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**(1)**

(c)     A journalist who read this investigation concluded that ipilimumab improved the treatment of MM.

Do the data in the table support this conclusion? Give reasons for your answer.

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

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**(4)**

(d)     MM is caused by a faulty receptor protein in cell-surface membranes.  
Cells in MM tumours can be destroyed by the immune system.

Suggest why they can be destroyed by the immune system.

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

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**(3)**

**(Total 10 marks)**

**Q54.**Read the following passage.

|  |  |  |
| --- | --- | --- |
|  | Low-density lipoprotein (LDL) is a substance found in blood. A high concentration of LDL in a person’s blood can increase the risk of atheroma formation. Liver cells have a receptor on their cell-surface membranes that LDL binds to. This leads to LDL entering the cell. A regulator protein, also found in blood, can bind to the same receptor as LDL. This prevents LDL entering the liver cell. People who have a high concentration of this regulator protein in their blood will have a high concentration of LDL in their blood. Scientists have made a monoclonal antibody that prevents this regulator protein working. They have suggested that these antibodies could be used to reduce the risk of coronary heart disease. | 5 |
|  | A trial was carried out on a small number of healthy volunteers, divided into two groups. The scientists injected one group with the monoclonal antibody in salt solution. The other group was a control group. They measured the concentration of LDL in the blood of each volunteer at the start and after 3 months. They found that the mean LDL concentration in the volunteers injected with the antibody was 64% lower than in the control group. | 10 |
|  | 15 |
|  | Use the information in the passage and your own knowledge to answer the following questions. | |

(a)     The scientists gave an injection to a mouse to make it produce the monoclonal antibody used in this investigation (line 7).

What should this injection have contained?

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**(1)**

(b)     LDL enters the liver cells (lines 3−4).

Using your knowledge of the structure of the cell-surface membrane, suggest how LDL enters the cell.

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**(2)**

(c)     Explain how the monoclonal antibody would prevent the regulator protein from working (lines 7−8).

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**(2)**

(d)     Describe how the control group should have been treated.

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**(2)**

**(Total 7 marks)**

**Q55.**Read the following passage.

|  |  |  |
| --- | --- | --- |
|  | Herpes simplex virus (HSV) infects nerve cells in the face, including some near the lips. Like many other viruses, HSV can remain inactive inside the body for years. When HSV becomes active, it causes cold sores around the mouth. |  |
|  | Human cells infected with a virus may undergo programmed cell death. While HSV is inactive inside the body, only one of its genes is transcribed. This gene is the latency-associated transcript (*LAT*) gene that prevents programmed cell death of an infected nerve cell. | 5 |
|  | Scientists have found that transcription of the *LAT* gene produces a microRNA. This microRNA binds to some of the nerve cell’s own mRNA molecules. These mRNA molecules are involved in programmed cell death of nerve cells. The scientists concluded that production of this microRNA allows HSV to remain in the body for years. | 10 |

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

(a)     HSV infects nerve cells in the face (line 1). Explain why it infects **only** nerve cells.

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

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**(3)**

(b)     HSV can remain inactive inside the body for years (lines 2–3). Explain why this virus can be described as **inactive**.

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**(2)**

(c)     Suggest **one** advantage of programmed cell death (line 4).

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**(1)**

(d)     The scientists concluded that production of this microRNA allows HSV to remain in the body for years (lines 10–12).

Explain how this microRNA allows HSV to remain in the body for years.

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

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**(4)**

**(Total 10 marks)**

**Q56.**Malaria is a disease caused by parasites belonging to the genus *Plasmodium*. Two species that cause malaria are *Plasmodium falciparum* and *Plasmodium vivax*.

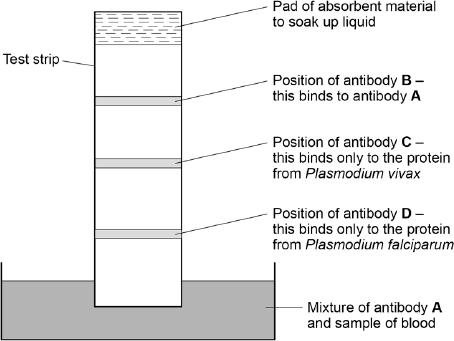
A test strip that uses monoclonal antibodies can be used to determine whether a person is infected by *Plasmodium*. It can also be used to find which species of *Plasmodium* they are infected by.

•        A sample of a person’s blood is mixed with a solution containing an antibody, **A**, that binds to a protein found in both species of *Plasmodium*. This antibody has a coloured dye attached.

•        A test strip is then put into the mixture. The mixture moves up the test strip by capillary action to an absorbent pad.

•        Three other antibodies, **B**, **C** and **D** are attached to the test strip. The position of these antibodies and what they bind to is shown in **Figure 1**.

**Figure 1**



(a)     Explain why antibody **A** attaches only to the protein found in species of *Plasmodium*.

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**(2)**

(b)     Antibody **B** is important if this test shows a person is not infected with *Plasmodium*.

Explain why antibody **B** is important.

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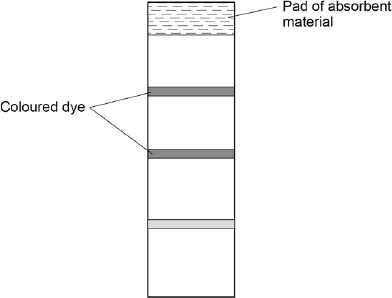
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**(2)**

(c)     One of these test strips was used to test a sample from a person thought to be infected with *Plasmodium*. **Figure 2** shows the result.

**Figure 2**



What can you conclude from this result?

Explain how you reached your conclusion.

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

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**(4)**

**(Total 8 marks)**

**Q57.**(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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**[Extra space]** ................................................................................................

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**(3)**

**(Total 6 marks)**

**Q58.**(a)     Give **two** ways in which pathogens can cause disease.

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

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

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**(2)**

(b)     Putting bee honey on a cut kills bacteria. Honey contains a high concentration of sugar.

Use your knowledge of water potential to suggest how putting honey on a cut kills bacteria.

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**[Extra space]** ................................................................................................

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**(3)**

**(Total 5 marks)**

**M1.**(a)     Any **two** from:

1.      (Decrease linked to) few(er) cases of whooping cough;

2.      (Decrease linked to) risk of / fear of side effects;

3.      Insufficient vaccine available / too expensive to produce / distribute.

*3. Too expensive unqualified is insufficient for mark*

**2 max**

(b)     1.      Vaccination rate increases;

2.      Fewer people to spread the disease / whooping cough / more people immune / fewer susceptible.

*2. Neutral − greater herd effect*

*2. Allow description of immune*

***Q*** *Reject ‘resistant’.*

**2**

(c)     1.      More people are immune / fewer people carry the pathogen;

*If neither point 1 or 2 awarded*

*Herd immunity = 1 mark*

*Unvaccinated does not mean infected*

*1.* ***Q*** *Do not accept disease for pathogen*

2.      So susceptible / unvaccinated people less likely to contact infected people.

**2**

**[6]**

**M2.**          (a)     Zevalin/antibody binds to specific receptor/cell surface protein/antigen;

(Only found) on B-cells;

**2**

(b)     Patient **P** treated with Zevalin/yttrium (no mark);

*Assume ‘Zevalin’ means ‘with yttrium’ unless they state  
otherwise*

Where indium/antibody (only) on lymphatic system/groin and armpits;

So only (cancerous) B-cells killed;

In patient **P** high concentration of radioactivity/antibodies high enough  
to kill cancer cells;

Patient **Q** ‒ radioactivity in places where other body cells could be killed/  
organs damaged/named example;

Could harm patient more than cancer;

Patient **Q** cancer has spread;

So too late to treat;

**3 max**

(c)     Patient **Q** ‒ (cancerous) B-cells outside of lymphatic system/metastasis;

So antibody bound in other parts of the body (as well);

Patient **Q** ‒ has different receptors/distribution of receptors compared  
to patient **P**;

Other body cells (than B-cells) have receptors for antibody;

**2 max**

(d)     Might be allergic to mouse antibody/protein;

(Mouse) antibody acts as an antigen;

Causes an immune response/antibody production;

Antibody destroys Zevalin;

Releases radioactivity into body/prevents activity against the cancer;

**2 max**

**[9]**

**M3.**          (a)     Protein / molecule/glycoprotein;On surface of cell/microorganism;Stimulates immune response/production of antibodies;

**2 max**

(b)     Zookeeper is not producing antibodies/passive immunity;No memory cells made;

ORAntivenom is an antigen/stimulates production of (anti-antivenom) antibodies;(Antivenom) destroyed by zookeeper’s own antibodies;

ORAntibody destroys antigen/venom;Before immune response/no immune response;

**2**

**[4]**

**M4.**         (a)     A = envelope/membrane/phospholipid (bilayer);B = capsid / nucleocapsid / capsomere / protein;

**2**

(i)      (HIV is) invading cells which make new viruses;Cells release viruses into blood;

**2**

(ii)     Virus remains dormant/exists as provirus/exists as DNA in host DNA;

*Accept virus stays in cells*

**1**

(c)     HIV destroys T cells;More (free) viruses produced leads to fall in T-cells;(So fewer) T-cells activate B-cells/memory cells;

Reduced/no antibody production;Immune system not working properly/inability to fight infection;Opportunistic infections;

**4 max**

**[9]**

**M5.**         (a)     Cotinine is an antigen;Antigen/cotinine binds to (specific) T-cell/activates T-cell;T-cell activates B-cells;Specific B cell becomes activated;(Specific) B cell divides/ clonal expansion;Forms (clone of) plasma cells;(Plasma) cell produces antibodies;

*Accept macrophage presents antigen for one mark*

*Ignore references to memory cells and secondary  
immune response*

**4 max**

(b)     Antibodies are proteins with tertiary structure/specific shape/binding sites;Antibodies specific shape for cotinine;Only cotinine fits;

*Do not credit active site*

**2**

**[6]**

**M6.**          (a)     (i)      Molecule/protein/glycoprotein;  
Stimulates immune response;  
(That causes) production of antibodies;

**2 max**

(ii)     Antigens on HIV are different (shape);  
So, antibody will not ‘fit’/not complementary (to antigen);  
Receptor sites on antibody specific to one antigen;

**2 max**

(iii)     (Has site with) same shape as salmonella antigen so binds  
to anti-gal antibodies;  
(Has site with) same shape as receptor molecule so that HIV will bind;  
Binds to both molecules;

**2 max**

(b)     Salmonella pathogen has specific antigen on surface;  
Salmonella pathogen engulfed by macrophage;  
T-cells activate B-cells;  
B-cell with complementary/specific receptor antibody activated/  
clonal selection;  
B-cells divide/form clone/clonal expansion;  
Plasma cells make antibodies;  
Specific to antigen/bind to salmonella bacterial antigen;

*Accept macrophage presents antigen to T/B cells;*

*Accept T-cells release factors;*

**6 max**

(c)     (i)      HIV binds to specific receptor;  
Only present on certain cells / T-cells;

**2**

(ii)     Antibiotics stop metabolism, viruses don’t have metabolism;  
Viruses hide in cells, antibiotics can’t reach;

Two suitable cell components antibiotics work against that  
viruses don’t have;  
e.g. some antibiotics work against ribosomes, that viruses don’t have

**2**

(d)     (i)      Adaptor molecule binds to HIV;  
(This) prevents the HIV binding to the receptor;  
Therefore few HIV available to infect cells;

**2 max**

(ii)     Would need to be complementary to MRSA (antigens);  
MRSA has different antigens;  
But would still need to have binding site for anti-gal;

**2 max**

**[20]**

**M7.**         (a)     (i)      P = membrane / lipid envelope / phospholipid bilayer;Q = reverse transcriptase;

*Accept (host) cell membrane;*

**2**

(ii)     Carries genetic information / to make DNA;

***Q*** *Do not accept ‘information’ on its ownAccept genes, alleles,to make (viral) protein;*

**1**

(b)     DNA copy made (of viral RNA);Inserted into host DNA / chromosomes;(Uses viral DNA to) make viral proteins/particles;Makes viral RNA;(Host) cell makes new viruses;“Budding off” / wrapped in cell membrane;

*Accept reverse transcriptase makes DNA for 2 marks in correct context;*

**3 max**

**[6]**

**M8.**          (a)     Microorganism alive/active;But does not cause symptoms of disease/Avirulent;

*Accept does not make you ill/harm*

**2**

(b)     (i)      (Takes time for) antigen to be recognised;

*Accept reference to presentation by macrophage  
for first marking point*

(Takes time for) T cells to be activated;

*Accept primary (immune) response*

B-cell activation/clonal selection/expansion;Plasma cells to make (specific) antibodies;Time for enough antibodies to measure;

**2 max**

(ii)     Memory cells (present);

*Accept secondary (immune) response*

Respond immediately / can produce antibodies immediately;

**2**

**[6]**

**M9.**          (a)     Correct answer: 1.25;

*Ignore working*

***OR*** (if wrong answer)

 / = 1 mark

*125 but wrong order of magnitude = 1 mark*

**2**

(ii)     **C** has myosin / thick (and actin / thin) filaments;

***OR***

**A** has only actin / thin (/ no myosin / no thick) filaments;

**1 max**

(b)     When contracted:

Thick & thin filaments/myosin & actin overlap more;

Interaction between myosin heads & actin / cross-links form;

Movement of myosin head;

Thin filaments / actin moved along thick filaments / myosin;

Movement of thin filaments / actin pulls Z-lines closer together;

Displacement of tropomyosin to allow interaction;

Role of Ca2+;

Role of ATP;

*Allow ref. to ‘sliding filament mechanism’ /  
described if no other marks awarded*

**4 max**

(c)     (i)      8 has DMD but 3 and 4 do not / 12 has DMD but 6 and 7  
do not / neither parent has the condition but their child has;

*Allow parents 3 and 4 give 8, parents 6 and 7 give 12*

**1**

(ii)     4 ***AND*** 7;

**1**

(iii)     Parental genotypes:  6 = **XDY** AND 7 = **XDXd**

***AND***

Gametes correct for candidate’s P genotypes ‒ e.g.

**X**Dand **Y** + **X**Dand  **X**d**;**

Offspring genotypes correctly derived from gametes e.g.

**X**D**X**D+ **X**D**X**d+ **X**D**Y** + **X**d**Y**;

Male offspring with MD correctly identified: **X**d**Y**;

Probability = 0.25 / correct for candidates offsprings genotypes;

*Accept ¼ / 1 in 4 / 1:3 / 25%*

*NOT ‘3:1’ / ‘1:4’*

**4**

(d)     (i)      No gene fragment **G**;

**1**

(ii)     Only one copy of gene fragment **F**;

Male has only one X-chromosome / is XY  
(c.f. female has two / is XX);

**2**

(iii)     10 has only one copy of gene fragment **G**;

10 has only one normal X-chromosome / has one abnormal /  
has only one normal allele / has one Xd / is XDXd / is heterozygous;

11 has two normal X-chromosomes / has 2 normal alleles /  
is XDXD / has not got Xd / has 2 copies of (F and) G;

**3**

(e)     (i)      To prevent rejection / prevent antibody production vs. injected cells /  
injected cells have (foreign) antigen (on surface);

**1**

(ii)     Shows effect of cells / not just effect of injection / not just effect of  
salt solution;

**1**

(iii)     Only one person tested so far ‒ need more to see if similar results /  
need more to see if reliable;

Need to assess if new (dystrophin positive) muscle fibres are  
functional / if muscle becomes functional;

Can’t tell how widespread effect is in the muscle / sample taken  
near injection site;

Need to test for harmful side effects;

Need to test if successful for other mutations of dystrophin gene;

Need to assess permanence / longevity of result/insufficient time  
allowed in investigation;

(In this patient) only small response / %;

Further sensible suggestion;

**4 max**

**[25]**

**M10.**          (a)     Reverse transcriptase;

*Accept integrase/description of action of*

Enzyme uses (HIV) RNA to make DNA (copy);

DNA joined to (host) cell’s DNA/chromosome;

DNA used to make HIV RNA (copies);

*Accept (HIV) DNA replicated when (T) cell divides*

And HIV capsid proteins/enzymes;

Made at (host) ribosomes;

Assembly of new virus particles;

Budding off from membrane (of host cell);

**4 max**

(b)     Not enough/no T-cells to activate B-cells/lead to antibody production/  
activate immune system;

*Accept death of T-cells weakens the immune system*

Person unable to fight /more prone to (opportunistic) infections/cancer;

*Accept diseases*

Example of infection/cancer;

*E.g. TB, pneumonia, cryptosporidium*

**2 max**

**[6]**

**M11.**          (a)     47 213;

**1**

(b)     (i)      there is no difference in the proportion / number of influenza cases  
between the 5 vaccines;

*(reject vaccinated versus no vaccinated)*

**1**

(ii)     significant difference in proportion / number of cases of influenza  
between the vaccines / the null hypothesis should be rejected;

**1**

(c)     sample size small;  
possible differences in exposure to infection;  
exposure to different strains / mutants;  
possible differences in existing immunity;  
possible differences in sex / age;  
possible differences in socio-economic status;

**2 max**

**[5]**

**M12.**          (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]**

**M13.**          (a)     (i)      protein / glycoprotein / glycolipid / polysaccharide / molecule;  
on surface / membrane (of cell);  
causes immune response / description / triggers antibody  
production;

**max 2**

(ii)     reference to hybrid cell from tumour / cancer and  
B-lymphocyte / hybridoma;  
antibodies all the same / from one type of plasma cell;  
specific to / complementary to / fits only one antigen;

**max 2**

(b)     (i)      antibodies specific / only binds to PSA;  
PSA only associated with prostate cancer / not with other  
diseases;

**2**

(ii)     antibody with enzyme only attaches if PSA present / washed  
away if no PSA;  
no colour change without enzyme;

**2**

**[8]**

**M14.**          (a)     (i)      fall in deaths due to rise in number of people with immunity / better care / targeting vaccination at vulnerable;

**1**

(ii)     mutation of virus / new strain;  
mutant form not recognised by memory cells (*allow antibodies*);

**2 max**

(b)     (i)      T lymphocyte receptors recognise shape of haemagglutinin /   
neuraminidase / viral antigen;  
clone (*once only*);  
destroy virus;

**2 max**

(ii)     clone (*once only*);  
produce antibodies;  
effect of antibody e.g. stimulation of phagocytosis /   
precipitation of toxins;

**2**

(c)     alter shape of active site of neuraminidase / block active site;  
virus unable to leave host cells;

**2**

**[9]**

**M15.**          (a)     bacteria have ligands / antigens / proteins / glycoproteins / polysaccharides (on membrane / wall);

**1**

complementary to receptors / fits / binds / attaches to specific receptor

**1**

(b)     enzymes denatured / tertiary / secondary structure altered / altered  
active sites / breaks hydrogen bonds;

**1**

prevents named chemical reactions / metabolic pathways;

**1**

(c)     inhibits / kills other bacteria / fungi / decomposers / reduces competition;

**1**

(d)     1       prepare a bacterial lawn / culture / sample;  
         *(accept mix bacteria with agar / medium)*2with oil and one with control / water / range of concentrations;  
3       appropriate method of standardising how sample applied,  
         e.g. discs / wells;  
4       appropriate measure of effectiveness / size / diameter of clear zone;  
5       the larger the zone the greater the effectiveness;  
6       use of aseptic technique;

*(ignore haemocytometer)*

**4 max**

**[9]**

**M16.**          (i)      1360 = 2 marks  
(general principle 0.68 ÷ 0.05 x 100 gains 1 mark)

**2**

(ii)      still have maternal antibodies;

**1**

**[3]**

**M17.**          (a)     1       macrophages present antigens to B lymphocytes;  
2       antigen binds to / is complementary to receptors on lymphocyte;  
3       binds to a specific lymphocyte;  
4       lymphocytes become competent / sensitised;  
5       (B) lymphocytes reproduce by mitosis / (B) lymphocytes cloned;  
6       plasma cells secrete antibodies;

**4 max**

(b)     1       restriction enzyme / endonuclease;  
2       to cut plasmid / to form sticky ends in plasmid;  
3       (use) ligase(to join) gene to plasmid;  
4       culture bacteria with (in medium containing) plasmids  
5       to allow uptake of plasmids / transformation;  
6       use of cold shock / chemical treatment (to enhance uptake) / heat  
         shock;  
*(ignore bullets / electroporation / microinjection)*

**3 max**

**[7]**

**M18.**          (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]**

**M19.**          (a)     Presence of resistant and non-resistant varieties / mutation produces resistant variety;  
Resistant ones survive / non-resistant ones killed by treatment;  
These will reproduce and produce more resistant parasites / pass on resistance allele;

**3**

(b)     Likelihood of being infected (by strain resistant to both drugs) is less;  
1/500 × 1/500/1/250 000;  
Drug has longer effective life;

**max 2**

(c)     (i)      As comparison / to show that nothing else in the treatment was responsible;

**1**

(ii)     Given injections of saline / injection without SPf66;  
(otherwise) treated the same as experimental group;

**2**

(d)     (i)      100%;

**1**

(ii)     10%;

**1**

(e)     (i)      Different lengths of DNA have different base sequences / cut at specific sequence;  
Results in different shape / different shape of active site;  
Therefore (specific sequence) will only fit active site of enzyme;

**3**

(ii)     Recognition sites contain only AT pairs;  
Which would occur very frequently;

**2**

**[15]**

**M20.**          (a)     molecule (on cell surface);  
that triggers immune response;

**2**

(b)     (i)      axes right way round and labelled;  
2nd peak drawn higher;  
steeper gradient on second rise;

**3**

(ii)     because one dose does not give a high enough level of  
antibody to be effective / because the antibody falls after a while;

**1**

(iii)     antigens are only single molecules / part of parasite;  
do not actually cause disease;

**2**

(c)     malaria sufferers would have parasites in red blood cells;

**1**

**[9]**

**M21.**          (a)     Publicity about vaccination / better health education / risks of ‘flu epidemics;

*(Accept: now free on NHS (though only since 2000) / better awareness / more commonly available)*

**1**

(b)     (i)      1990: 26% of 7.4million = 1.92million *and* 2000: 64% of 7.8 million = 4.99million;  
increase = 3.07 million;

**2**

*(Correct reading of all 4 figures from graph = 1)*

*(Correct answer but no ‘millions’ = 1)*

*(Correct method resulting from wrong graph reading = 1)*

(ii)     Over 50% of population being vaccinated;  
But only from 2000 onwards;  
*(Principle of more people being vaccinated each year = 1)*

**2**

(iii)     Different strain / type of virus each year / virus mutates;  
With different antigens;  
Influenza antibodies / memory cells (rapidly) destroyed / need replacing;

**max 2**

(c)     (Protein coat) carries antigens which stimulates B-cells / production of antibodies;  
Production of memory cells;

**2**

**[9]**

**M22.**          (a)     To prevent contamination of apparatus with other microorganisms / bacteria;  
To prevent personal contact with bacteria;  
To prevent release of bacteria into air;

**max 2**

(b)     (i)      Diffuses slowly;

**1**

(ii)     B;  
Produces inhibition zone greater than the minimum diameter;

**2**

**[5]**

**M23.**          (a)     Injection of antigens / toxoids;

(Antigen from) attenuated microorganism / non-virulent  
microorganisms / dead

microorganisms / isolated from microorganism;

Stimulates the formation of memory cells;

**max 2**

(b)     (i)      Antibodies are specific to mumps antigen;  
2nd antibodies specific to mumps antibody;

**1**

(ii)     Removes unbound 2nd antibodies;  
Otherwise enzyme may be present / may get colour change  
anyway / false positive;

**2**

(iii)     No antibodies to bind (to antigen);  
Therefore 2nd antibody (with the enzyme) won’t bind / no enzyme /   
enzyme-carrying antibody present  
(after washing in step 4);

**2**

**[7]**

**M24.**          (a)     (i)      protein / immunoglobulin;  
specific to antigen;  
idea of ‘fit’ / complementary shape;

**2 max**

(ii)     1. virus contains antigen;  
2. virus engulfed by phagocyte / macrophage;  
3. presents antigen to B-cell;  
4. memory cells / B-cell becomes activated;  
5. (divides to) form clones;  
6. by mitosis;  
7. plasma cells produce antibodies;  
8. antibodies specific to antigen;  
9. correct reference to T-cells / cytokines;

**6 max**

(b)     1. antibody gene located using gene probe;  
2. cut using restriction enzyme;  
3. at specific base pairs;  
4. leaving sticky ends / unpaired bases;  
5. cut maize / DNA / vector using same restriction enzyme;  
6. join using DNA ligase;  
7. introduce vector into maize / crop / recombinant DNA into maize;

**4 max**

(c)     passive / person is not making own antibodies / antibodies not replaced;  
memory cells not produced;

**2**

(d)     fewer ethical difficulties / less risk of infection;

**1**

**[15]**

**M25.**          (a)     Stimulates memory cells;

Secondary response, so antivenom / antibodies produced quicker;

**2**

(b)     Passive immunity; so no memory cells produced;

Antivenom breaks down / destroyed;

**2**

(c)     Could transfer disease / Allergy / Immune response to antibodies  
from animal;

**1**

**[5]**

**M26.**          (a)     add antibodies / enzyme;  
wash to remove unbound antibodies;  
add (colourless) solution;

*(mark correct responses sequentially)*

**3**

(b)     antibodies specific / shape only fits one antigen;  
other antigens different shape and would not bind to antibodies;

**2**

**[5]**

**M27.**          (a)     Phagocytes engulf pathogens / microorganisms;

Enclosed in a vacuole / vesicle / phagosome;

Lysosomes have enzymes;

That digest / hydrolyse molecules / proteins / lipids / microorganism;

**3 max**

(b)     (i)      Get another strain / there are different strains;

Therefore does not have memory cells against second strain;

***Q*** *The second marking point should only be awarded in the context of memory cells.*

**2**

(ii)     Vaccines only work against certain strains because the antigens they possess are different;

Enables company to target strain likely to be prevalent later / most common strain;

**2**

**[7]**

**M28.**          (a)     divide by mitosis / form clones;  
produce plasma cells;   
(plasma cells) make antibodies;  
(plasma cells) produce memory cells;

**4**

(b)     glycoprotein;  
different shape to body proteins / body phospholipids are the same /   
located on the outside of the cell / the haemoglobin is located  
inside the cell;

**2**

**[6]**

**M29.**          (a)     molecule / part of molecule / protein / glycoprotein / named molecule;  
that stimulates an immune response / eq;

**2**

(b)     divide by mitosis / form clones; produce plasma cells; (plasma cells)   
make antibodies;  
(plasma cells) produce memory cells;

**4**

(c)     (i)      glycoprotein AND  
different shape to body proteins / RNA and reverse transcriptase  
inside virus / phospholipids same as body’s / on the surface  
of the virus;

**1**

(ii)     187.5;;

*Accept 187 – 188*

*1 mark for HIV = 80nm;*

**2 max**

**[9]**

**M30.**          (a)     (i)      Many people do not go to the doctor;

**1**

(ii)     36000;

*No marks awarded for working here as calculation is very straightforward*

**1**

(b)     Same sugars / antigens on bacteria / nerve cells;

*Do not accept references to same shape as equivalent to complementary.*

Bind with antibody / form antigen-antibody complex;

*Reject react*

Have complementary shape / fit binding site;

*Reject active site*

**3**

(c)     Diaphragm will not move down / flatten / contract;

*Ignore references to breathing out*

Thoracic cavity / lung volume not increased so cannot breathe in;

**2**

**[7]**

**M31.**          (a)     Phagocytes engulf / ingest pathogens / microorganisms / bacteria / viruses;

Phagocytes destroy pathogens / microorganisms / bacteria / viruses;

Lung diseases are caused by pathogens / microorganisms / bacteria / viruses;

***Q*** *Allow description of process of engulfing*

**2 max**

(b)     (i)      Alveoli / lungs will not inflate / deflate fully / reduced lung capacity;

Breathing out particularly affected / no longer passive;

**2**

(ii)     Alveolar walls thicken;

Longer diffusion pathway;

Scarred / fibrous tissue;

Reduces surface area (for gaseous exchange);

***Q*** *Diffusion is essential for 2nd point and surface area for 4th point.*

**4**

(c)     (i)      Cancer develops 20 – 30 years after exposure (to asbestos);

**1**

(ii)     Smoking / air pollution / specified industrial source;

**1**

**[10]**

**M32.**          (a)     Nucleus;

**1**

(b)     Enables organism to remain in area (of food source) / prevent its removal;

***Q*** *‘To attach’ is not sufficient unless qualified*

**1**

(c)     (i)      Correct answer of 222(%);;

Incorrect answer that clearly identifies difference in number of cases as 5800 –1800 or 5.8 – 1.8;

*Correct answer gains two marks*

**2**

(ii)     More water-related activities / more ‘organisms’ with increased temperature;

***Q*** *Allow any reference to growth or replication of ‘organisms’. Do not penalise reference to bacteria.*

***Q*** *Do not allow increase in water consumption.*

**1**

(d)     (i)      All have same shape / only binds to *Giardia* / one type of / specific antigen;

**1**

(ii)     Has complementary (shape) / due to (specific) tertiary structure / variable region (of antibody);

***Q*** *Binds / fits not sufficient unless qualified;*

**1**

(iii)    Enzyme / second antibody would remain / is removed by washing;

Enzyme can react with substrate (when no antigen is present);

**2**

**[9]**

**M33.**          (a)     (yes):  
Many women (with cervical cancer) have HPV 16 (18 &31);

(no):  
Few women (with cervical cancer) have HPV 6 / 11;

(HPV infection does not mean causation because):  
Could be caused by another factor / example given / may be due to coincidence;

No control group / did not study HPV in healthy women / did not study all HPV types / having cancer may increase susceptibility to HPV / does not add up to 100% / not all women with cancer have HPV / individual may have more than one HPV type;

*Neutral: correlation between HPV (16) and cervical cancer*

*Reject: many women with HPV 16 (18 &31) have cervical cancer / not all women have cancer*

*Accept: figures from graph for ‘many’ and ‘few’*

*Accept: minor errors in reading HPV frequencies from graph*

*Reject: does not mean HPV vaccine causes cancer;*

*Neutral: refs. to sample size and factors that should have been kept constant*

**3 max**

(b)     (i)      Protein / glycoprotein / glycolipid / polysaccharide;

Causes immune response / antibody production;

*Accept: B / T cell production*

**2**

(ii)     Memory cells produced / remain / stored (from previous infection);

*Neutral: antibodies produced / remain*

(When individual) comes into contact with virus / antigen (again);

*Neutral: ‘cell’ instead of ‘virus’  
Reject: ‘bacteria’ once only*

Rapid / secondary / greater response / many or more antibodies produced;

*Accept: B cells / T cells*

Destroys virus / antigen before it can cause harm / symptoms / cancer;

*Reject: if destroys the virus / antigen in the vaccine before it can cause harm*

***Q*** *Do not allow ‘fights HPV’*

***Q*** *Do not allow ‘memory cells remember’*

**3 max**

(c)     HPV destroyed in males / prevents males being carriers of HPV;

*Neutral: prevents males catching HPV*

Prevents males passing on HPV (to unvaccinated females) / HPV  
may cause (other) cancers in males;

*Accept: reference to herd effect protecting the population*

**2**

**[10]**

**M34.**          (a)     Damage / destruction of cells / tissues;  
Production of toxins;

**2**

(b)     Contains antigen / proteins / dead / weakened microorganism / pathogen / virus / bacteria;  
Stimulates production of antibodies / plasma cells / memory cells;

***Q*** *Do not credit immune response unless qualified.*

**2**

(c)     (i)      Age;

Sex;

Ethnicity;

All healthy / not on other medication;

Not previously vaccinated / infected with TB;

***Q*** *Do not credit sample size.*

***Q*** *Allow any suitable reference to health not being affected for fourth marking point e.g. smoking, ‘depressed immune system’ etc.*

**2 max**

(ii)     Contain the same antigens;

**1**

**[7]**

**M35.**          (a)     Virus / fungus / protozoan;

*Neutral: named example*

**1**

(b)     Produces toxins;

*Neutral: infects / colonises / invades cells*

Damages cells / tissues / example given e.g. cell lysis;

**2**

(c)     (i)      (Antibodies) produced from a single clone of B cells / plasma cells;

*Accept: hybridoma cell line instead of B cell / plasma cell  
Reject: idea that antibodies are cloned*

***OR***

(Antibodies) produced from the same B cell / plasma cell;

**1**

(ii)     (Specific) primary structure / order of amino acids;

(Specific) tertiary / 3D structure;

(So) Only binds to / fits / complementary to one antigen;

*Reject: ‘active site’ for either point 2. or 3. only once*

**3**

(d)     (Rapid) treatment of carriers / infected cattle / disease;

*Neutral: reference to rapid identification of infected cattle*

Can isolate / cull carriers / infected cattle / infected (dairy) products not sold / consumed / tracked;

Reduces spread of disease / no need to kill / prevents the death of non-infected animals;

*Neutral: ethical arguments*

**3**

**[10]**

**M36.**          (a)     1.      Phagocyte attracted to bacteria by chemicals / recognise antigens on bacteria as foreign;

2.      Engulf / ingest bacteria;

3.      Bacteria in vacuole / vesicle;

4.      Lysosome fuses with / empties enzymes into vacuole;

5.      Bacteria digested / hydrolysed;

*1. Accept names chemical e.g. toxin*

*2. Allow description of engulfing*

*3. Accept: bacteria in phagosome*

*5. Neutral: Break down*

*5. Accept digestive enzymes destroy bacteria*

*5. Do not accept “destroy bacteria” as it is in question stem*

**4 max**

(b)     1.      Microvilli provide a large / increased surface area;

2.      Many mitochondria produce ATP / release or provide energy (for active transport);

3.      Carrier proteins for active transport;

4.      Channel / carrier proteins for facilitated diffusion;

5.      Co-transport of sodium (ions) and glucose or symport / carrier protein for sodium (ions) and glucose;

6.      Membrane-bound enzymes digest disaccharides / produce glucose;

*1. Reject villi on epithelial cells*

*1. Accept brush border*

*2. Accept large SA:vol ratio*

*3. Need idea of “lots”*

*4. Reject: energy produced*

*5. Accept Na+K+ pump*

*6. Neutral: Channel proteins*

*7. Accept named example*

**6**

**[10]**

**M37.**(a)     (i)      To show whether immune response occured / because cats are (genetically) related to cheetahs;

*Ignore reference to control.*

**1**

(ii)     To show that rejection did not normally occur / skin could (successfully) be grafted;

**1**

(b)     (i)      Rapid rejection between unrelated (domestic) cats / cats are **not** genetically similar;  
Rapid rejection between (domestic) cat and cheetah / cats and cheetahs are not genetically similar;  
Slow / no rejection in cheetahs / cheetahs are genetically similar;

**3**

(ii)     Sample size small;  
Time observed was short;

**1 max**

(iii)    Similar (antigens on all cheetahs);

*Accept same / not very different*

**1**

(iv)    Protein / antigen production determined by alleles / genes / base sequence on DNA;  
The more similar the proteins the more similar their alleles / genes / base sequence on DNA / the more they are genetically similar;

**2**

**[9]**

**M38.**          (a)     Girls are not sexually active / not likely to carry HPV / vaccine may not work if already infected / few girls sexually active (at this age);

*Neutral: girls are not sexually mature*

*Neutral: to provide better protection*

*Accept: provides immunity before sexually active*

*Neutral: girls are less likely to have ‘****it****’ as could mean the vaccine from the question stem*

**1**

(b)     Other (HPV) types have different antigens;

No memory cells for other types / memory cells not activated / antibodies cannot attach to antigen / correct antibodies not produced / antibodies are not complementary;

*Accept: refs. to antigenic variability*

*Accept: B cells for memory cells*

*Accept: memory cells cannot recognise antigen for ‘not activated’*

*Accept: examples of memory cell activation*

**2**

(c)     More antigen;

More memory cells;

So more antibodies produced / antibodies produced quicker (if infected);

*Accept: ‘many’ / ‘enough’ instead of ‘more’*

*Neutral: primary / secondary response*

*Accept: T cells / B cells / plasma cells instead of ‘antibodies’*

*Reject: the idea that vaccines contain antibodies*

***Q*** *Reject: antibodies ‘fight’ / ‘antibiotics’*

**2 max**

(d)     Cancer takes years to develop / develops later in life;

Takes time for females to become sexually active / females must become sexually active to obtain data;

Few people / only teenagers vaccinated;

*Neutral: will take time to vaccinate 80% of young girls*

*Accept: do not develop cancer instantly*

**2 max**

(e)     (Cervical cancer) can be caused by other types of HPV / other factors / example given;

***OR***

(Some) women may have been infected (with HPV) before receiving the vaccine;

***OR***

(As a precaution) in case vaccine does not work / a way of monitoring if the vaccine has worked;

*Accept: ‘caused by other types of HPV’ in the context of mutation*

*Neutral: to check for abnormal cells / that they are immune to the virus*

**1**

(f)      Virus cannot replicate / is destroyed / is not carried (in vaccinated people);

Non-vaccinated people more likely to contact vaccinated people;

*Neutral: ‘do not spread virus’ as in question stem*

*Must be in context of the individual and not the population as in question stem*

***Q*** *Do not allow ‘disease is destroyed’*

*Neutral: ‘herd effect’ as given in the question stem*

**2**

**[10]**

**M39.**(a)     Straight lines point to point as not possible to predict intermediate values / values between points;

**1**

(b)     Increases then levels / falls;  
Maximum antibody production 180 units / at dose of 0.25 g per kg;

**2**

(c)     Two marks for correct answer of 57.14 / 57.1;;  
One mark for incorrect answer in which candidate clearly divides difference in antibody production / 60 by 105;

**2**

(d)     Takes into account different masses of mice / allows comparison;

*Accept different weights of mice.  
Do not accept different size.*

**1**

(e)     Sheep red blood cells have antigens (on their surface);  
Antigens are proteins foreign to mice / are non-self;  
Stimulate B cells to produce antibodies;

**3**

(f)     Response only observed in mice;  
Disease organisms not investigated;  
Not all disease caused by pathogens / cured by antibodies;

*i.e. not tested on humans*

**2 max**

**[11]**

**M40.**          (a)     (i)      Antibiotics kill other bacteria / *Clostridium* is resistant;

Less / no competition so (*Clostridium*)  
reproduces / replicates / multiplies / increases in number;

*Reference to bacteria being ‘immune’ negates first marking point.*

*Reference to mitosis negates second marking point.*

**2**

(ii)     Immune system less effective / more likely to have other  
infections / been in hospital;

*Accept: ‘Weak / lower’ immune system’.*

**1**

(b)     Attaches to active site (of enzyme);  
(Methicillin) is a competitive inhibitor / prevents monomers / substrate  
attaching (to enzyme);

*‘Competes for active site’ = 2 marks.*

*Neutral: ‘Prevents monomers joining / attaching to each other’.*

*Allow one mark max for answers relating to non-competitive inhibitor changing active site / preventing substrate attaching.*

*Do not penalise Methicillin forms an enzyme / substrate complex.*

**2**

(c)     (i)      Have other illness / medical condition / ’weak’ immune system / disease / infection;

*Reject: Due to ‘other factors’, ‘are smokers’, ‘are obese’ unless related to disease or illness.*

**1**

(ii)     Increase up to 2006 / 20 (per 100 000) then decreases;

**1**

(iii)     Correct answer in range of 52 – 59.1% = two marks;

Incorrect answer but shows change as between 4.8 – 5.2 / shows  
correct subtraction giving this change e.g. 14 – 9 = one mark.

**2**

**[9]**

**M41.**          (a)     Nitrification;

*Accept nitrifying.*

*Do not accept nitrogen fixing.*

**1**

(b)     1.      Uptake (by roots) involves active transport;

*Reject all references to bacteria*

2.      Requires ATP / aerobic respiration;

**2**

(c)     (i)      1.      Not enough time / fast flow washes bacteria away;

*“Not enough time for bacteria to convert all the ammonia to nitrate” gains 2 marks*

2.      (Not all / less) ammonia converted to nitrate / less nitrification;

**2**

(ii)     1.      Algal bloom / increase in algae blocks light / plants / algae die;

2.      Decomposers / saprobionts / bacteria break down dead plant materials;

3.      Bacteria / decomposers / saprobionts use up oxygen in respiration / increase BOD causing fish to die;

*3. Accept alternatives such as microbes / saprophytes.*

**3**

**[8]**

**M42.**1.      Vaccines contain antigens / dead / weakened pathogens / antigens dead / weakened  
        pathogens are injected;

*Ignore references to T or B cells.*

2.      Memory cells made;

3.      On second exposure memory cells produce antibodies / become active / recognise pathogens;

*3. Idea of memory cells responding.*

4.      Rapidly produce antibodies / produces more antibodies;

*4. Production of antibodies must be qualified for mark. Underlined ideas essential.*

5.      Antibodies destroy pathogens;

*5. Accept bacteria / viruses etc but not disease*

**[5]**

**M43.**          (a)     (i)      Protein on (surface of) chlamydia;

That initiates an immune response (in mice) / causes antibody production;

*Neutral “foreign protein”*

*Do not accept glycoprotein.*

*2. Accept description of initiating immune response.*

**2**

(ii)     1.      Antibodies / memory cells against chlamydia (protein / antigen) are present;

2.      Protein on heart (muscle) similar to chlamydia protein / antigen so T cells / antibodies (attack heart muscle cells);

*2. Look for idea that both proteins are similar*

*2. Detail of what is attacking the heart muscle cells*

**2**

(b)     **FOR**

1.      Prevents / reduces heart disease / attacks;

2.      Cheaper to vaccinate than treat heart disease;

**AGAINST**

3.      Vaccination costly;

4.      Don’t know frequency of chlamydia infection;

5.      Research in mice might not be replicated in humans / humans might have a different protein;

6.      Vaccine could cause heart disease or immune response against heart (muscle);

*2 max for arguments against*

*Accept other valid answers*

**3 max**

**[7]**

**M44.**(a)     Has more than one / four polypeptide chains / made up of polypeptide chains;

**1**

(b)     1.      Antibody / variable region has specific amino acid sequence / primary structure;

2.      The shape / tertiary structure of the binding site is complementary to / fits / binds with these antigens;

*2. Do not accept active site for this point.*

3.      Forms complex between antigen and antibody;

**3**

**[4]**

**M45.**(a)     (i)      **(Whole-cell vaccine),**

*Accept converse statements for other vaccine*

*Reject references to the vaccine being alive or the disease reproducing etc*

1.      Heat(ing) supposed to kill bacteria;

2.      Some might be alive / active / viable;

*Accept active pathogens present*

3.      (If so) bacteria could reproduce;

4.      Bacterium makes or contains toxin;

5.      Toxin might not be affected / all destroyed by heat;

6.      Bacteria or toxins attacking / killing person’s cells;

**3 max**

(ii)     **(Whole-cell vaccine),**

*Ignore references to more / greater antigens unqualified. It is the variety of antigens that matters*

1.      (Contains) many different / greater range of antigens;

2.      Each antigen causes its own immune response / production of / has a specific (type of) antibody;

**2**

(b)     1.      Only patients who had whooping cough have toxin / antibody /   
immune response;

*Accept converse e.g. those without antibody had another disease*

2.      Toxin is an antigen and is (only) produced by this bacterium;

3.      Leading to presence of specific antibody / only 4% had this antibody / 13% did not have antibody;

**3**

(c)     1.      There may not be large rises;

2.      Might be the result of wrong diagnosis / reference to difference in figures / 13% diagnosed with whooping cough didn’t have it;

*Ignore reference to new strains or antigenic variability*

**2**

**[10]**

**M46.**          (a)     (Micro)organism that causes disease / harm to body / an immune response;

*Accept: named microorganism that causes disease*

*Allow infection*

**1**

(b)     1.      Phagocyte attracted by a substance / recognises (foreign) antigen;

*Accept named substance eg chemical / antigen*

2.      (Pathogen)engulfed / ingested;

*Accept: description*

3.      Enclosed in vacuole / vesicle / phagosome;

4.      (Vacuole) fuses / joins with lysosome;

5.      Lysosome contains enzymes;

*Accept named example of enzyme*

6.      Pathogen digested / molecules hydrolysed;

*Neutral: Destroyed*

**4 max**

(c)     1.      Antigens (on pathogen) are a specific shape / have specific tertiary / 3D structure;

*1 / 3 Structure alone is insufficient*

2.      Antibody fits / binds / is complementary to antigen / antibody-antigen complex forms;

*Reject - active site*

***OR***

3.      Antibodies are a specific shape / have specific tertiary / 3D structure;

4.      Antigens (on pathogen) fit / bind / are complementary to antibody / antibody-antigen complex forms;

**2**

**[7]**

**M47.**(a)     (i)      Substance that causes an immune response / production of antibodies;

*Ignore foreign / non-self*

**1**

(ii)     1.      Not lipid soluble;

2.      Too large (to diffuse through the membrane);

3.      Antigens do not have the complementary shape / cannot bind to receptor / channel / carrier proteins (in membranes of other epithelial cells);

**2 max**

(b)     1.      (Vaccine contains) antigen / attenuated / dead pathogen;

*1. Reject if in context of injection of vaccine*

2.       T-cells activate B-cells;

3.       B-cells divide / form clone / undergo mitosis;

4.      Plasma cells produce antibodies;

5.      Memory cells produced meaning more antibodies / antibodies produced faster in secondary response / on reinfection;

**5**

**[8]**

**M48.**(a)     1.      Infected by / susceptible to (other) pathogen(s) / named disease caused by a pathogen (from environment);

*Context is where immune system cannot prevent or stop these events*

*Allow attack / kill*

2.      Pathogen(s) reproduce / cause diease (in host);

*MPs not given in context of HIV*

3.      Damage cells / tissues / organs;

4.      Release toxins;

**3 max**

(b)     (i)      1.      (HIV enters cells) before antibodies can bind to / destroy it;

*Ignore SAFETY comments*

*1. and 2. Relate to antibodies*

2.      Antibodies cannot enter cells (to destroy HIV) / stay in blood;

***OR***

3.      (Enters cells) before (secondary) immune response caused / before memory cells have time to respond;

*3. and 4. Relate to virus*

4.      So no antibodies present (to attack HIV);

***OR***

5.      Vaccine taken up too quickly to cause immune response;

*5. and 6. Relate to vaccine*

6.      So no antibodies / memory cells formed;

**2 max**

(ii)     1.      Antigen (on HIV) changes;

*Accept mutates*

2.      (Specific) antibody / receptor no longer binds to (new) antigen;

*Ignore SAFETY comments*

***OR***

3.      Many different strains of HIV / many antigens present on HIV;

4.      Not possible to make a vaccine for all antigens / vaccine may not stimulate an antibody for a particular antigen;

**2 max**

(c)     3 suitable suggestions;;;

*QWC ignore reference to HIV cells*

E.g.

1.      Inactive virus may become active / viral transformation;

2.      Attenuated virus might become harmful;

3.      Non-pathogenic virus may mutate and harm cells;

4.      Genetic information / protein (from HIV) may harm cells;

5.      People (may) become / test HIV positive after vaccine used;

*Vaccinated people may develop disease from a different strain to that in the vaccine*

6.      This may affect their work / life;

*May continue high risk activities and develop or pass on HIV*

**3 max**

**[10]**

**M49.**(a)     (i)      1.      (Scientists) canߢt show bias / influence / may have a vested   
          interest / work for the company developing the vaccine;

*Relates to the scientists*

2.      (Volunteers) can’t show psychological / mental effects / ‘placebo effect’ / expectations;

*Relates to the volunteers*

*Accept: reduces the ‘Hawthorne effect’ / demand characteristics*

*Neutral: so they have no idea what they are taking*

**2**

(ii)     Any **two** suitable suggestions, eg

*Neutral: refs. to age and health*

1.      Amount of nicotine in cigarettes;

*Neutral: different types of cigarette / different ways / frequency of smoking*

2.      Amount inhaled / absorbed / time since last cigarette;

*Neutral: absorption by gut / digestion*

*Accept: absorption by mouth*

3.      (Different) amounts excreted / metabolism / rate of binding (of nicotine) to protein;

*Accept: broken down (differently)*

4.      (Different) blood volumes;

*Neutral: different body masses*

5.      Nicotine from passive smoking / other smokers / other sources;

6.      Some volunteers received the vaccine / placebo;

*Accept: some volunteers would have / would not have the antibodies*

**2 max**

(b)     (i)      1.      Antibodies to nicotine produced / antibodies bind to nicotine;

***Q*** *Reject: vaccine contains / produces antibodies*

***Q*** *Neutral: antibodies digest / kill / fight nicotine*

2.      (So) nicotine does not bind to protein / does not reach the brain;

***Q*** *Reject: any reference to ‘active site’*

*Neutral: idea that the antibodies bind to the protein*

3.      (So) cigarettes / smoking does not satisfy addiction / reward smokers / release (reward) chemicals;

**3**

(ii)     **(Agree):**

1.      People choose to smoke / know the risks;

2.      Should spend this money on education / preventing people from starting to smoke / treating other health problems / vaccines are expensive;

**(Disagree):**

3.      Unethical not to treat;

4.      Less money needed to treat the effects of smoking / cancer / smokers pay taxes so are entitled to treatment;

**3 max**

(c)     1.      High antibody responders have a high % to stop smoking / are more likely to stop smoking;

*‘People producing a high concentration of antibodies’ is equivalent to ‘high antibody responders’*

*Accept: reference to values from the table*

2.      Only a few may be high antibody responders / no numbers on how many are high / medium / low antibody responders;

*Neutral: not all people are high antibody responders*

3.      Percentage who stopped smoking is similar for placebo group and low / medium responders / some / % of placebo group (still) stopped smoking / placebo has the lowest value / % to stop smoking;

*Accept: reference to values from the table*

4.      Large sample size / double blind **so** reliable / representative;

5.      Antibody levels peak at / drop after 5 months / boosters may be needed at / after 5 months;

6.      May start smoking again after 5 / 6 months / do not know the percentage who stopped smoking after 5 / 6 months;

7.      Nicotine is not the only factor responsible for making people smoke;

*Must mention nicotine*

*Do not accept: correlation does not mean causation / could be due to other factors*

**5 max**

**[15]**

**M50.**(a)     (To diagnose AIDS, need to look for / at)

1.      (AIDS-related) symptoms;

2.      Number of helper T cells.

*Neutral: ‘only detects HIV antibodies’ as given in the question stem*

**2**

(b)     1.      HIV antibody is not present;

*Accept HIV antibodies will not bind (to antigen)*

2.      (So) second antibody / enzyme will not bind / is not present.

**2**

(c)     1.      Children receive (HIV) antibodies from their mothers / maternal antibodies;

2.      (So) solution will always turn blue / will always test positive (before 18 months).

*Allow 1 mark for the suggestion that the child does not produce antibodies yet so test may be negative*

**2**

(d)     (Shows that)

1.      Only the enzyme / nothing else is causing a colour change;

2.      Washing is effective / all unbound antibody is washed away.

**2**

**[8]**

**M51.**(a)     1.      Vaccine contains antigen from pathogen;

2.      Macrophage presents antigen on its surface;

3.      T cell with complementary receptor protein binds to antigen;

4.      T cell stimulates B cell;

5.      (With) complementary antibody on its surface;

6.      B cell secretes large amounts of antibody;

7.      B cell divides to form clone all secreting / producing same antibody.

**5 max**

(b)     1.      Active involves memory cells, passive does not;

2.      Active involves production of antibody by plasma cells / memory cells;

3.      Passive involves antibody introduced into body from outside / named source;

4.      Active long term, because antibody produced in response to antigen;

5.      Passive short term, because antibody (given) is broken down;

6.      Active (can) take time to develop / work, passive fast acting.

**5 max**

**[10]**

**M52.**(a)                         *QWC*

1.      (Phagocyte engulfs) to form vacuole / vesicle / phagosome;

*Accept surrounds bacteria with membrane*

2.      Lysosome empties contents into vacuole / vesicle / phagosome;

*Accept joins / fuses*

3.      (Releasing) enzymes that digest / hydrolyse bacteria;

*Ignore breakdown / destroy / lytic enzymes*

**3**

(b)     Two suitable structures;;

Examples,

1.      Cell wall;

2.      Capsule / slime layer;

3.      Circular DNA;

*Reject “circular chromosome”*

4.      Naked DNA / DNA without histones;

5.      Flagellum;

6.      Plasmid;

7.      Pilus;

8.      70s / smaller ribosomes;

9.      Mesosome;

**2 max**

**[5]**

**M53.**(a)     1.      Rank all STs in ascending order;

2.      Find value with same number (of people) above and below.

*Accept find middle value*

**2**

(b)     Not ethical to fail to treat cancer.

**1**

(c)     Yes since with ipilimumab:

1.      Median ST increased by 2.1 months;

2.      Percentage of patients showing reduction in tumours increased from 10.3% to 15.2%;

No because:

3.      No standard errors shown / no (Student) t- test / no statistical test carried out;

4.      (So) not able to tell if differences are (statistically) significant / due to chance (alone);

5.      Improvement might only be evident in some patients / no improvement in some patients;

6.      Quality of (extra) time alive not reported;

*If answers relate only to ‘Yes’ or ߢNo’, award 2 marks max*

**4 max**

(d)     1.      Faulty protein recognised as an antigen / as a ‘foreign’ protein;

2.      T cells will bind to faulty protein / to (this) ‘foreign’ protein;

3.      (Sensitised) T cells will stimulate clonal selection of B cells;

4.      (Resulting in) release of antibodies against faulty protein.

**3 max**

**[10]**

**M54.**(a)     Regulator protein.

*Accept regulator protein antigen*

*Reject regulator protein receptor*

*Ignore regular protein*

**1**

(b)     1.      Lipid soluble / hydrophobic

2.      Enters through (phospholipid) bilayer

***OR***

3.      (Protein part of) LDL attaches to receptor

4.      Goes through carrier / channel protein.

*4. Accept by facilitated diffusion or active transport*

*4. Reject active transport through channel protein*

**2**

(c)     Any **two** from:

1.      (Monoclonal antibody) has a specific tertiary structure / variable region / is complementary to regulator protein

*Do not award MP1 if reference to active site.*

2.      Binds to / forms complex with (regulator protein)

*“It” refers to monoclonal antibody in MP1 and MP2*

3.      (So regulator protein) would not fit / bind to the receptor / is not complementary to receptor

*3. Reject receptor on LDL*

**2 max**

(d)     1.      Injection with salt solution

*1. Accept inject placebo in salt solution*

2.      Otherwise treated the same.

**2**

**[7]**

**M55.**(a)     1.      Outside of virus has antigens / proteins;

2.      With complementary shape to receptor / protein in membrane of cells;

3.      (Receptor / protein) found only on membrane of nerve cells.

*Accept converse argument*

**3**

(b)     1.      No more (nerve) cells infected / no more cold sores form;

2.      (Because) virus is not replicating.

**2**

(c)     Prevents replication of virus.

**1**

(d)     MicroRNA binds to cell’s mRNA (no mark)

1.      (Binds) by specific base pairing;

2.      (So) prevents mRNA being read by ribosomes;

3.      (So) prevents translation / production of proteins;

4.      (Proteins) that cause cell death.

**4**

**[10]**

**M56.**(a)     1.      Antibody has tertiary structure;

2.      Complementary to binding site on protein.

**2**

(b)     1.      Prevents false negative results;

2.      (Since) shows antibody **A** has moved up strip / has not bound to any *Plasmodium* protein.

**2**

(c)     1.      Person is infected with *Plasmodium* / has malaria;

2.      Infected with (*Plasmodium*) *vivax*;

3.      Coloured dye where antibody **C** present;

4.      That only binds to protein from *vivax* / no reaction with antibody for *falciparum*.

*Person is infected with P. vivax / Plasmodium vivax = 2 marks (MP1 and MP2)*

**4**

**[8]**

**M57.**(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]**

**M58.**(a)     1.      (Releases) toxins;

2.      Kills cells / tissues.

*2. Accept any reference to cell / tissue damage*

*Ignore infecting / invading cells*

**2**

(b)     1.      Water potential in (bacterial) cells higher (than in honey) / water potential in honey lower (than in bacterial cells);

*Q candidates must express themselves clearly*

*1. Must be comparative e.g. high WP in cell and low WP in honey*

2.      Water leaves bacteria / cells by osmosis;

3.      (Loss of water) stops (metabolic) reactions.

*3. Needs a reason why lack of water kills the cell*

**3**

**[5]**

**E1.**(a)     Most students successfully used the graph to link the decrease in the percentage of infants vaccinated to fewer cases of whooping cough, with many correctly noting figures from the graph. Fewer students gave a second reason: the most common answer was the fear of side effects with many students linking the vaccine to potential side effects.

(b)     93% of the students scored one mark usually for correctly interpreting the graph and stating that the vaccination rate was increasing. Many then went on to discuss herd immunity in general terms rather than being specific and writing about more people being immune or fewer being susceptible as a result of the vaccination. Examiners expected students to use the correct terminology and students who wrote about ‘resistance’ to whooping cough did not gain credit.

(c)     Two-thirds of the students scored one mark here, for realising that herd immunity was involved, but very few explained clearly how it worked. Examiners were looking for the ideas that there were fewer people in the population in which the pathogen could survive, because many were immune, having been vaccinated, and secondly that contact between infected people and unvaccinated people was therefore less likely. Students confused the terminology with some assuming that all unvaccinated people were infected. Many incorrectly expressed the idea that infants did not need to be vaccinated because they had inherited immunity from their parents and a significant number simply restated the information in the question stem.

**E2.**          Part (a) proved difficult for many candidates and nearly two-thirds scored nought. Many candidates simply re-stated the information in the stem of the question that the antibody binds to B cells. Others got very confused between antibodies, antigens and cells and appeared to use all three interchangeably. Good answers included the deduction that the antibody must bind to an antigen found on the surface of B cells (only).

Part (b) discriminated across the whole range. Weaker candidates had a lot of trouble expressing themselves clearly and got confused between cancerous B cells and the normal functions and functioning of B cells. The best answers were in terms of the cancer having spread in patient Q and radioactivity doing damage to vital organs.

On reflection, part (c) was either too difficult, or not worded clearly enough. Very few candidates gave good responses to this question.

There were many good answers to (d). Many candidates obtained both marks for answers along the lines that existing antibodies would destroy the Zevalin before it became bound to cancerous B cells, thus making the treatment ineffective.

**E3.**          (a)     This question was not as well answered here as it has been on previous papers. Many candidates confused antigens with antibodies, and thought that they were produced by the body in response to an infection. While quite a large number of students gained one mark here, only the better candidates scored two.

(b)     Better candidates answered this well, but weaker candidates produced confused answers. Again, it was apparent that the immune system is not well understood by weaker candidates.

**E4.**          (a)     This was not well answered by most candidates. Many candidates offered suggestions such as ‘cell wall’ and ‘nucleus’.

(b)     In this question many candidates confined their answers to descriptions of the curve, instead of following the instruction to ‘explain’. Some candidates did relate the increase in particles in (i) to replication within T-cells, but it was unusual to find a reference to the virus remaining latent, inactive or dormant in (ii).

(c)     Answers here mainly focused on the body’s immune system being unable to ‘fight’ the infection. Weak candidates gave a simple description of the graph. Only the better candidates understood that T-cells were being destroyed as HIV particles were released from them, leaving the body exposed to opportunistic infections as the immune system was functioning less effectively.

**E5.**          (a)     It was unusual for candidates to score full marks here. Many thought that cotinine was a pathogen. The role of T-cells and B-cells was confused by many, and those who did have some understanding of the immune system frequently focused on the production of memory cells, which was not relevant to this question.

(b)     Many candidates scored a single mark here, but few gained both marks. The idea of complementary shapes and antigens fitting into receptor/binding sites on the antibody were not well known. A few incorrectly used the term ‘active site’, while many simply confined their answer to the statement that ‘antibodies are specific’.

**E6.**          (a)     In part (i) many vague answers were seen, usually about fighting cells and recognising self and foreign. Many described it as a ‘structure’ rather than a ‘molecule’ or a named molecule, and thought that it causes a disease. However, some good answers were also seen, which included the idea that an antigen produces an immune response.

Part (ii) was well understood by many, although there was some confusion with enzymes and active sites. A few candidates confused antibodies with antibiotics, explaining that antibiotics are not effective against viruses.

Part (iii) was left blank by many candidates, while many others did not understand and discussed induced fit. Where marks were awarded, it was usually for the idea that the adaptor molecule binds to both HIV and the anti-gal antibodies.

(b)     A surprising number of candidates scored no marks at all here, mainly because they wrote about memory cells and cloned antibodies. On the other hand, the better candidates scored 5 or 6 marks and gave comprehensive accounts. A few left the question blank, and some lost marks by confusing T and B cells.

(c)     Very few candidates scored marks in part (i) here at all. Common misconceptions were that the receptor was on the HIV or that antibodies were involved. In part (ii) some good answers were seen, usually for realising that antibiotics target metabolism, cell walls or protein synthesis, and explaining that viruses do not have these. A few said antibiotics only attack membrane bound organelles which was unfortunate. Many stuck with ‘’because they only work on bacteria’, ‘they would become resistant/immune’, ‘because viruses have RNA/ no antigens’. This was often a case of bad expression losing marks.

(d)     This question was very hard to mark. Many said the adaptor and anti-gal would bind to HIV so this was allowed but I looked for receptor for point 2, which was very rare. The idea of not many HIV available (point 3) was not clearly expressed. Many concentrated on the antigens of the HIV or thought the receptor was on the HIV. A lot had the virus being demolished by anti-gal and antibodies. Overall perhaps 15% of candidates gained 2, 40% gained 1 and the remainder 0 marks. Quite a few left the question blank.

In part (i) most candidates failed to score, usually because of poor expression. Very few referred to antigens. Hardly any candidates used the term ‘complementary’ though every other word e.g. active site / substrate was employed. Many talked about resistance, infection, attack or had the wrong name for parts.

Similarly in part (ii) poor expression lost marks. When the candidate used terms like ‘antigen’ and had the idea of shape and fit, both marks could be awarded.

**E7.**          (a)     Most candidates were able to identify structure **P** correctly in (i), although a few thought it was a capsule or a cell wall. Fewer candidates were successful in identifying **Q**, and despite being told that it is an enzyme, gave answers such as ‘ribosome’ or ‘nucleus’. In (ii), many candidates knew that this carries genetic information, but others simply guessed and gave answers such as ‘supplies energy’.

(b)     This question showed that there is widespread lack of understanding of how HIV affects infected cells. Many described the viral RNA pairing with a single strand of DNA to become double-stranded DNA. More worryingly, many candidates think that the helper-T cell becomes HIV once the HIV has invaded it.

**E8.**          (a)     This was very poorly known. Many candidates think that an attenuated microorganism is dead or inactive. Some who were aware that it is a weakened form of the microorganism went on to spoil their answer by describing it as ‘weakened or dead’. A few knew that the microorganism had been weakened, or repeatedly sub-cultured, but did not say that this means they do not cause the symptoms of the disease.

(b)     This section was well known by most candidates. In (i)most candidates could give an explanation for the time delay, such as it takes time for the B cells to become activated, or for plasma cells to produce antibodies. Part (ii) was also well known, though there were fewer right answers here. Nevertheless, the majority of candidates were able to make some relevant reference to memory cells.

**E9.**          (a)     Most candidates measured band X (the A-band in an electron micrograph of a myofibril) correctly. Many did not then understand that they had to divide this by the stated magnification. Among those who did, many had problems interconverting millimetres and micrometres and were often several orders of magnitude out. Only one quarter of candidates were entirely successful.

In part (ii), most candidates knew the correct distribution of actin and myosin filaments in the two distinct bands of the myofibril. One unusual, and erroneous, concept expressed by a number of candidates was that one part of the myofibril was contracted at the same time as the other part was relaxed.

(b)     Many candidates gave a full and clear account of the process of muscle contraction, including the roles of ATP, calcium ions, tropomyosin, the attachment of the myosin head to actin and its movement causing the actin filament to slide along the myosin. Weaker candidates just described how the appearance of the various bands changed when the myofibril contracted rather than offering the required explanation. Almost one-third of candidates scored full marks.

(c)     Using information from the pedigree diagram showing the inheritance of Duchenne muscular dystrophy (DMD) over three generations, almost two-thirds of candidates cited the production of a child with muscular dystrophy by unaffected parents as evidence for the condition being caused by a recessive allele. However, less than half the candidates were able to identify two carriers from the diagram.

In completion of the genetic diagram, common errors included switching the genders of the two parents, giving the male parent a genotype that would have resulted in him having muscular dystrophy, incomplete assignment of phenotypes to the offspring genotypes (both gender and having / not having DMD were important) and, having shown that 25% of the offspring would be expected to be male with DMD, to then halve this figure to 12.5 %. Additional, incorrect, answers on the probability line, e.g., ‘25% or 1 : 4’, failed to gain the mark. Despite this, almost one-third of candidates scored full marks in this section.

(d)     Just over half the candidates answered part (i) correctly, realising that the complete absence of one of the gene fragments indicated that the person would suffer from DMD. In part (ii), these candidates realised it was the single copy of the other gene fragment (compared with two copies in each of his sisters) that indicated the person concerned was male as he had just one X-chromosome while his sisters had two. Only about one-fifth of candidates were able to tell the complete story, although some two-thirds got half-way.

Part (iii) differentiated very well between candidates who gave varying degrees of appropriate detail in their answers. The most able noticed that one of the girls had two copies of one of the gene fragments while her sister, having but a single copy of this fragment, must have been the carrier as she would have had one normal X chromosome (hence being healthy herself) and one carrying the mutation responsible for DMD. Approximately one quarter scored full marks, although nearly two-thirds were able to make at least two of the three points required.

(e)     Far too many candidates failed to use appropriate terminology in part (i). There were no marks available for stating that the ‘immune system’ (given in the question) ‘fought against’ / ‘attacked’ the implanted cells. Terms such as *rejection*, *antibody* and *antigen* were required. Less than half the candidates used such terms.

Similarly, in part (ii), there was no mark available for merely stating that the injection with salt solution served as a ‘control’. The purpose of the control was required, e.g., so that the effect of the cells injected into the other leg became apparent, or to show it was not just the salt solution that had caused the effect in the other leg. Approximately half the candidates gave the appropriate detail.

In part (iii), there was plenty of scope for candidates to explain the limitations of the given investigation and to suggest appropriate further work that could be done. Candidates made general points about the limited sample size (i.e., just *one* individual), the short time period allowed to assess the effect of the treatment, or they made specific points relating to the given size of the response, the fact that success had so far been achieved only for this particular mutation, that only a measure of the *presence* of the appropriate type of muscle cells had been performed with no information about their ability to function, etc. The question differentiated very well amongst candidates who took varying amounts of care in selecting information, in assessing the reliability of the data and in applying their knowledge and understanding of how an investigation should be carried out in order to obtain reliable results and to draw valid conclusions. Although almost 90% of candidates were able to make at least one valid point, only 3% scored all 4 marks.

**E10.**          Some very good answers were seen to each part of this question. However, for a topic which has been so well covered by the media and educational programmes, and been a familiar part of A level for many years, it was disappointing to find many candidates scoring very poorly.

In (a), half of candidates failed to score any marks. The mark scheme only required candidates to know basic facts about replication of HIV. For example, the ideas that there is an enzyme that makes a DNA copy of the HIV RNA, this DNA is inserted into the host cell’s DNA and is used to make new HIV RNA and proteins. The statements in this sentence would have obtained all four marks.

Part (b) was better answered and the majority of candidates expressed the idea that people die from infections they are unable to suppress because of their compromised immune system.

**E11.**          Whilst a full range of marks was seen on this question, five marks were very rarely awarded. Most candidates’ powers of expression were not up to the task of explaining what they meant in part (b). Most candidates scored between one and three marks.

(a)     The majority of candidates obtained this mark.

(b)     Relatively few obtained the mark in part (i). Most candidates misunderstood the basic purpose of the study and wrote about vaccination (versus no vaccination) having no effect on the number of influenza cases. Many merely turned the stem of the question around and stated ‘there is no difference in effectiveness of the vaccines’. More candidates scored a mark in part (ii) by writing about the rejection of the null hypothesis.

(c)     This discriminated quite well. Good candidates usually obtained two marks, average candidates often failed to gain one mark for vague references along the lines of ‘it not being a fair test’, and weak candidates gave answers such as ‘there were a different number of people in each group’.

**E12.**          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.

**E13.**          (a)     Definitions of antigens and monoclonal antibodies indicated some confusion and there were many references to bacteria, foreign structures, one substance attacking another, or the body fighting off disease. Precise accounts were infrequent.

(b)     Again imprecise answers abounded, showing lack of understanding. In part (ii), candidates were required to explain that the enzyme had to be present and why it would not be. This needed extensive linguistic skills which were not always apparent. Some did make good use of the diagrams.

**E14.**          (a)     (i)      Build up of immunity in the community or advances in care were examples of acceptable responses. Again, ‘vaccination’, unqualified, was the most common answer and, again, this received no credit.

(ii)     Most candidates gained one mark for stating that a mutant form or new strain was involved. Relatively few went on to complete the explanation in terms of non-recognition by memory cells.

(b)     Weaker candidates continue to confuse the actions of T and B lymphocytes. However, even these candidates know that both types of cell clone in response to infection. References to phagocytosis by T lymphocytes were, unfortunately, quite common.

(c)     Few candidates read the information given and therefore answered in terms of viruses being unable to enter cells. Those who answered correctly in terms of leaving the cell could rarely explain inhibition, rather merely re-stating that it would occur.

**E15.**          (a)     Answers were generally good but some candidates confused the position of ligands and receptors. Some candidates did not know that humans are mammals.

(b)     Most candidates gained the mark for enzymes being deactivated but then failed to describe a specific relevant effect.

(c)     Some candidates appreciated the use of exotoxins. Incorrect responses suggested that they digested organic material as an aid to nutrition.

(d)     This question was well answered across the whole ability range. Most candidates had seen or done a similar experiment and described the method well.

**E16.**          (i)      Only a minority of candidates was able to perform a percentage increase calculation. The most common error was to divide the final mean amount (0.73) by the original amount (0.05), failing to subtract.

(ii)      Only the more able could suggest antibodies in the mother’s milk, or placental transfer of antibodies as methods. Weaker candidates frequently stated that the antibodies were inherited.

**E17.**          Most candidates gained at least four marks in this question.

(a)     The majority of answers lacked detail and a clear understanding of the correct sequence of events. Candidates had to select the facts needed to answer the question. Many gained a mark for replication by mitosis/cloning. All the other marking points were seen, with stronger candidates gaining all the marks. Few mentioned the importance of specific B lymphocytes, or differentiation resulting in plasma cells that release antibodies.

Antigen, antibody and receptor were often confused. Many mentioned memory cells and T cells.

Many candidates gained more than half the marks because there were six points on the mark scheme.

(b)     A large number of students wasted time describing how to isolate the gene. The gene had already been isolated. Marks were gained for references to restriction endonuclease, plasmid and ligase. Few suggested how to transfer plasmid into bacteria.

**E18.**          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’.

**E19.**          (a)     Many candidates understood the basic principles of natural selection underlying this part of the question and better answers related these to the development of resistance in malarial parasites. Responses, however, were frequently marred by imprecise use of terms. Thus malarial parasites were variously described as developing resistance, immunity or, in some cases, allergies to the drugs concerned, while resistance was described as taking place in bacteria, the disease or even in the human population.

(b)     Evidence from BYA5 suggests that many candidates understand that probabilities are combined by multiplication. However, they were unable to apply this principle to the example in this part of the question. The most frequent response was to add the two figures. The resulting value of 1/250 then proved difficult to explain, and the simple idea that the probability of being infected by a strain of malarial parasite resistant to both drugs was much lower eluded most.

(c)     The concept of a control proved surprisingly unfamiliar to most candidates and even the best seldom progressed beyond explaining that a control offered a standard against which to compare the effectiveness of the vaccine. This idea should have given rise, in part (ii), to injection with saline only in an otherwise identically treated control group. Answers ranged from those who clearly failed to appreciate the nature of a control and discussed issues which were largely ethical in nature, to responses which were in varying degrees incomplete. Such responses included making sure that both groups “lived in the same place” or “were the same age”, ignoring the fact that these were only part of a whole range of factors which should have been kept constant. Evidence from this question and from the coursework suggests that the issue of controls is one that needs to be addressed by centres.

(d)     Better candidates experienced little apparent difficulty in identifying the correct percentages here. Incorrect answers fell into no set pattern and most responses which could conceivably be given arose at least once.

(e)     Many candidates were obviously of the opinion that restriction enzymes function in a way that is totally different from other enzymes, and attempted to explain their specificity in part (i) in terms of base pairing. Others clearly understood the principles involved but neglected to relate their understanding of enzyme action to this particular question. A lack of precision characterised many of the answers to part (ii). Thus there were frequent references to adenine and thymine but not to these bases forming the restriction sites. However, most candidates were able to equate the frequency of cutting to the small size of the resulting fragments.

**E20.**          (a)     This was poorly answered, despite the fact that similar questions have been set on many previous BYA3 papers. Many candidates simply stated that an antigen was a marker on a cell, without indicating its molecular nature. Few referred to its role in producing an immune response, simply limiting their answer to cell recognition.

(b)     In part (i) many candidates started both lines at the same point on the axes. Some who did this also failed to indicate which was the first injection and which was the second. In part (ii) many candidates thought the second injection was a test to see if the first injection worked. Some thought memory cells were not produced until the second injection. There were also references to antigens changing and malaria having two hosts. In (iii) it was very clear that most candidates did not understand what an antigen is. Most candidates thought that the vaccination was attenuated and made with a weakened strain of malaria which cannot cause disease. There were also frequent references to the antigens having been killed.

(c)     Many candidates suggested looking for burst/damaged red blood cells, or looking for the antigens or antibodies in the blood. Only a minority of candidates correctly realised that parasites would be visible under the microscope.

**E21.**          Part (a) was well known by most candidates. However, in (b)(i), very few gained both marks for the correct answer. Many gave a percentage instead of a number. Where the answer was wrong, it was very difficult for examiners to give credit for correct methodology because most candidates showed little organised working. Many candidates misread the graph, for example reading 74 million instead of 7.4. In (ii), most candidates could see that the percentage being vaccinated was rising each year, while the population was growing more slowly. In (iii), there were many vague answers referring to building up immunity, without any detail of how. Some simply stated that it was a good idea for elderly people to be vaccinated because they are the most at-risk age group. In part (c), there were many good answers, though many candidates referred to this being an attenuated vaccine.

**E22.**          In part (a), there were large numbers of answers which lacked specificity, such as preventing the bacteria escaping. There were the inevitable references to ‘fair tests’ and suggestions that bacteria entering the dish might somehow negate the effect of the antibiotics. In (b)(i), many poor answers were seen. Few commented on slow diffusion. Many simply stated that an antibiotic with a small clear zone could still be effective. In part (ii), most candidates identified **B** and gave a suitable answer. A few candidates chose **A** because it had the biggest clear zone, and some even gave a named antibiotic such as streptomycin or chloramphenicol.

**E23.**          (a)     This was well answered, with many candidates getting two marks and most gaining at least one mark. The failure to gain marks was due largely to the poor use of language with weaker candidates using the term disease rather than antigen.

(b)     Although well answered by many candidates, part (i) caused some difficulties for candidates who gave the well learned "antibodies are specific" response but did not relate this to the context of the question. The data in the stem of (ii) gave problems to candidates who did not precisely identify the antibody to which they were referring. The better responses clearly differentiated between the first antibody and the antibody-enzyme complex. Clarity of written responses also affected part (iii) where the best responses clearly differentiated between the enzyme active site and the antibody-binding site. The poorer responses tended to confuse these and often failed to gain marks as the examiners were unable to determine exactly to which molecule or stage in the process the response was referring.

**E24.**          (a)     Few candidates scored two in the first part and vague references to ‘fighting infections’ were common as was confusion of antigens with antibiotics. In part (ii), some candidates appeared to know the sequence of events but in many cases limited marks were gained because of incorrect biology and discussion of previous infections which ‘the body remembers and is ready to fight off the second time’. B and T cells were confused, the exact role of mitosis was not well known and few explained the specific antibody synthesis by plasma cells. Some answers used bacteria as the infective agent.

(b)     This section scored the majority of the marks for the question in most scripts. Problems arose mainly where candidates failed to apply their knowledge to the situation. As a result, the gene was transferred to a plasmid which was put into maize or left in a test tube. In a few cases, the first part of the question led to suggestions of infecting the maize with various blood cells.

(c)     Several good answers referring to passive immunity were seen here but many failed to gain marks through lack of precision in their answer.

(d)     Examiners were looking for more than a simple response about cost here. Valid ideas on the ethics of this work or the possibility of transferring a disease were awarded marks.

**E25.**          (a)     Generally answered quite well, though, because the stem clearly indicated the yield of antibodies was higher with the secondary response, answers that simply stated ”more antibodies produced’ were not credited. Examiners were looking for the faster rate of production as this was not given in the stem.

(b)     Although this question was attempted by most candidates it showed that some thought antibodies can die.

(c)     The better candidates tended to respond in terms of eliciting an immune response to the antivenom and gained the mark. Some candidates responded in terms of ethical issues and vegetarianism. Examiners found many of these responses were too vague or below the level needed to gain credit.

**E26.**          (a)     This proved an excellent discriminator. Some achieved three marks but there was a tendency to include interpretation rather than method or to suggest impractical steps such as adding the test plate to the solution. Flow charts seem to be unfamiliar to many candidates.

(b)     Whilst a few scored two here, many failed to gain marks by discussing enzymes and active sites, failing to realise that antigens with a specific shape were involved.

**E27.**          (a)     There were many excellent answers to this part of the question that described phagocytes engulfing microorganisms and the subsequent role of lysosomes. Where difficulties arose, they not infrequently stemmed from too much, rather than too little, knowledge and there were some extremely involved and often confused answers that wrote in great depth about antigen presentation, opsonisation, b cells and t cells, none of which were relevant in the context of this question. The treatment of immunology should be confined to the principles set out in section 3.1.6 of the specification. Candidates will only be required to recall information explicitly described in this section.

(b)     From the evidence in the answers to part (i), most candidates appeared to appreciate that a person might be infected with different strains of the influenza virus. Only the better candidates, however, were able to explain this in terms of memory cells. Less convincing answers were seldom expressed in appropriate scientific terminology.  
Antigen, antibody and antibiotic were often used interchangeably in these responses, and there were many references to ‘attacking while the immune system was still weak’.  
In part (ii), candidates frequently ignored the instruction to ‘Use your knowledge of antigens’ and merely identified Strain 1 as the most frequent. There were however some excellent answers.

**E28.**          (a)     This part of the question attracted some very detailed descriptions of everything to do with the immune system, including T-lymphocytes, suppressor cells and the role of macrophages. However, in some of these accounts the main points were missed. Many candidates appeared of the opinion that plasma cells made memory cells. Others considered that lymphocytes were antibodies. In a few cases, plasma cells sometimes unfortunately became plasma or even plasmids. The idea that the B cells clone and these differentiate into plasma cells or memory cells seemed to elude quite a few candidates.

(b)     Most candidates correctly identified the glycoprotein and gave an acceptable reason.

**E29.**          (a)     This generally proved a good opening question, but weaker responses were less specific about the nature of the ‘chemical’, or just referred to a ‘substance’ that stimulates an immune response.

(b)     It was not uncommon to see all marks achieved. Better candidates appreciated that the B-lymphocytes would divide by mitosis, or produce clones, and then produced an accurate description of the production of antibody-secreting plasma cells and memory cells. Weaker responses discussed the role of T-lymphocytes, pathogens, plasmids or assumed that memory cells already existed.

(c)     While many could correctly identify the glycoprotein as the component, the reason provided was often unconvincing. At a simple level, glycoproteins were on the surface of the virus -not the cell - but few identified that difference in shape, from body proteins, would enable recognition of the antigen. An accurate calculation was rarely seen with few able to achieve both marks and some did not attempt it. Lack of clear working meant that it was not possible to award any credit in many cases.

**E30.**          (a)     Most candidates were able to extract the relevant information from the passage and explain that many of those with the illness failed to see a doctor. The concept of a ratio in part (ii) proved difficult for some to understand. Although there were many correct answers there were many that should have been considered, at best, to have been improbable.

(b)     Many candidates were able to comment on the identical nature of the sugars in the bacterial antigens and on the surface of nerve cells. For some, this led to a comprehensive account of antibody binding and the formation of an antibody-antigen complex. Others rather lost their way at this stage and did no more than suggest that this led to antibodies ‘attacking’ the nerve cells.

(c)     There were many lengthy accounts presented in answer to this part of the question.  
Able candidates frequently described the entire process of ventilation and extended their answers onto additional sheets before eventually arriving at a point where they describe the effects of paralysis. It was clear, however, that many candidates had little idea of the precise role of the diaphragm. It was not infrequently described, for example, as ‘pushing the ribs up and out’. Such statements as ‘breathing in causes the diaphragm to flatten’ were common and revealed confusion between cause and effect.

**E31.**          (a)     Although it was evident that most candidates had a good idea of the role of phagocytes, poor use of terminology often resulted in marks not being awarded. It was common to see responses such as phagocytes ‘fighting disease’ or destroying ‘foreign bodies’ or ‘infections’. Nevertheless, approximately a third of candidates obtained both marking points.

(b)     (i)      Most candidates obtained one mark for stating that the lungs would not fully inflate or deflate. However, very few candidates obtained a second mark for suggesting that breathing out would particularly be affected or that the rate of diffusion would be reduced. There was some confusion over the meaning of the term *elasticity* with many references to ‘lungs contracting and relaxing’.

(ii)     This proved to be an effective discriminator. The vast majority of candidates obtained at least one mark often by referring to the presence of scarred or fibrous tissue in the lungs. Most candidates then gained a second mark by linking this to a reduction in the surface area for effective gaseous exchange. Fewer candidates specifically referred to the thickening of the alveolar walls but instead mentioned thickening of lung tissue. However, a significant number of candidates were able to link this thickening to a longer diffusion pathway.

(c)     (i)      Most candidates obtained this mark using the information in the passage to explain that lung cancer develops 20 – 30 years after exposure to asbestosis.  
Candidates failing to gain this mark often provided incomplete responses such ae ‘it takes a long time for cancer to develop’.

(ii)     The vast majority of candidates obtained this mark by referring to smoking.

**E32.**          (a)     Over two thirds of the candidates correctly named the nucleus as the structure which confirms that *G. lamblia* is a eukaryotic organism. Ribosome and flagellum were frequent incorrect responses.

(b)     Most candidates gained this mark for stating that the sucker enabled *G. lamblia* to remain in the intestines rather than being moved out as substances passed through. However, some candidates suggested the sucker enabled nutrients to be absorbed or somehow prevented digestion of the organism.

(c)     (i)      Approximately one in five candidates obtained both marks for this calculation. A similar number of candidates did gain one mark for showing some valid working in attempting the calculation.

(ii)     The vast majority of candidates did not obtain this mark as they suggested that the number of cases would increase in the summer due to an increase in water consumption. Very few candidates suggested that an increase in temperature could lead to an increase in the number of *G. lamblia* in contaminated water.

(d)     (i)      Most candidates stated that monoclonal antibodies are the identical with no reference to their shape or the idea that they are specific to a particular antigen.

(ii)     Many candidates obtained this mark often by referring to the complementary shapes of the antibody and *Giardia* antigen. Fewer candidates mentioned the tertiary structure or the variable region of antibodies. Most incorrect responses suggested the antibody has an ‘active site’ or the ‘same’ shape as the *Giardia* antigen.

(iii)     As expected this proved to be a difficult question. Most candidates had difficulty understanding the flow chart and gave a variety of incorrect responses. Despite the information provided in the stem of the question many candidates suggested that the *Giardia* antigen would remain on the plate and cause a positive result.  
Other misconceptions related to the enzyme reacting with the antigen or with the antibodies. Approximately forty percent of candidates obtained one mark for realising that the second antibody with the attached enzyme would remain on the plate. Far fewer of these candidates went on to explain that the enzyme would react with the substrate to provide a yellow colour.

**E33.**          (a)     Only the most able candidates gained full credit on this question. However, most candidates gained one mark for the idea that cervical cancer could be caused by other factors. Unfortunately, some candidates misinterpreted the graph and considered it to show the percentage of women with cervical cancer, rather than the percentage of women with a specific type of HPV. It was very clear that these candidates did not realise that all women in the investigation had cervical cancer. Consequently, this led to responses that were out of context such as that ‘66% of women with HPV16 have cervical cancer’. Better candidates were able to criticise the data. They usually referred to the absence of a control group or suggested that cervical cancer may increase susceptibility to HPV. Weaker candidates often gave vague answers that were not qualified e.g. ‘it does not prove that HPV causes cervical cancer’. Similarly, they did not usually refer to specific types of HPV.

(b)     (i)      Approximately 40% of candidates gained one mark. This was almost always for stating that an antigen stimulates an immune response. Relatively few candidates made reference to the chemical nature of antigens.

(ii)     Just over a third of candidates scored full marks but 60% scored at least two marks. A number of candidates were aware that vaccination causes the production of memory cells or that memory cells remain. However, many candidates had the idea that memory cells ensure a rapid response to the same virus if encountered again. Unfortunately, these points were often poorly expressed by weaker candidates such as in stating that ‘memory cells remember the antigen’ or that ‘they fight the germ quicker’.

(c)     Most candidates suggested that vaccinating young men would reduce the spread of HPV to females. However, it was usually only better candidates who explained this in terms of vaccinated males destroying the virus or not acting as carriers. Weaker candidates usually expressed this idea poorly e.g. ‘vaccinated males cannot be infected’. A number of creditworthy references to herd immunity were made, although it should be noted that this term is not a requirement of this unit. Some candidates suggested that HPV may cause other cancers in males and this was also credited. The most common misconception involved vaccinated males passing on immunity to their children.

**E34.**          (a)     Most candidates had little difficulty obtaining at least one mark by referring to the production of toxins by pathogens. However, a significant number of candidates did not specifically refer to cells or tissues being damaged but instead described how pathogene enter the body or wrote about damage in general terms.

(b)     It was disappointing to find a significant number of candidates describing a vaccine as ‘a weakened form of a disease’. However, most candidates did refer to the production of antibodies or memory cells following vaccination. Some candidates did suggest that a vaccine contains antibodies and provides passive rather than active immunity.

(c)     (i)      This question caused little difficulty with the vast majority of candidates able to provide at least one valid factor, often age or gender. Other common correct responses related to obtaining healthy volunteers and individuals who had not been infected with TB or had been previously vaccinated.

(ii)     This proved very difficult. Only a small percentage of candidates obtained this mark by suggesting that the two vaccines have similar antigens. Most candidates simply stated that the two vaccines were most effective when used together.

**E35.**          (a)     Over 90% of candidates gained this mark, usually for ‘virus’. Relatively few referred to ‘fungi’. The few candidates who failed to score usually repeated ‘bacteria’ from the stem of the question or gave a specific example of a bacterium or virus.

(b)     Many candidates gained full marks for this question, although a minority misread the stem of the question and gave ways in which a pathogen could gain entry into the body. The most common mark awarded was for the production of toxins. Unfortunately, some candidates failed to gain the second mark through a lack of detail e.g. ‘damages the body’ and ‘infects cells’.

(c)     (i)      This question was poorly answered by most candidates. Only the most able were aware that monoclonal antibodies were produced by the same B cell or B cell clone. There was evidence of widespread poor expression and responses usually fell into one of two discrete camps. Candidates who focused on the ‘mono’ aspect of ‘monoclonal’ frequently referred to these antibodies ‘only binding to one antigen’ or ‘being produced from a single antibody’. Candidates who focused on the ‘clonal’ aspect usually gave responses that were out of context, such as ‘these antibodies are cloned’ and ‘they are genetically identical antibodies’. Disappointingly, very few candidates mentioned B cells.

(ii)     This question proved to be a good discriminator. The most common mark awarded was for ‘tertiary structure’. Weaker candidates usually went straight into an explanation of why monoclonal antibodies are specific in terms of binding, shape or fit. However, a number of these answers lacked detail regarding what these antibodies bind to. Relatively few of these candidates used the term ‘antigen’. In such cases, the terms ‘bacteria’ or ‘pathogen’ were typically used. A number of candidates also confused antibodies with enzymes, with references to ‘active sites’ and ‘antibodies being complementary to the substrate’. There were, however, some highly impressive answers given by the more able candidates. These usually gained full marks and often gave more detail than that shown on the mark scheme.

(d)     Most candidates gained at least one mark for the idea of reducing the spread of disease. The more able qualified this by explaining that rapid identification allowed infected cattle to be isolated and treated. Relatively few wrote about infected dairy products not being sold. Weaker candidates often gave vague arguments that related to animal rights and economics. These usually involved less distress being caused to the animals, not having to transport the animals or money being saved due to the samples not having to be sent to a laboratory. Such responses were not credited. Similarly, a minority of candidates incorrectly thought that the monoclonal antibodies were being used to treat infected cattle, rather than as a diagnostic tool.

**E36.**          (a)     Phagocytosis was well understood by candidates and many gained full marks. Many, however, wasted time by describing antigen presentation which was not required. Almost all candidates knew that the bacteria would be engulfed and many referred to their inclusion in a phagosome or vacuole. Weaker candidates failed to gain marks through imprecision, e.g., not realising that the lysosomes fuse with the phagocytic vesicle and just writing about them releasing enzymes or not specifying what sort of enzymes and then repeating the stem by saying the bacteria were destroyed without any mention of digestion or hydrolysis.

(b)     Many candidates failed to realise that this question was about cell adaptation. There was a lot of confusion between adaptations of the cell and of the intestine wall generally, with many answers focusing on factors such as a good blood supply and maintaining a steep concentration gradient. Such responses were not relevant to this question. Many candidates had a good understanding of glucose co-transport and described this in detail, usually gaining two or three marks, even though their answers were not focused on cell adaptation. Most scored one mark for understanding a large surface area was involved but many attributed this to villi rather than to microvilli. Better candidates gained a further two marks for explaining the cells would have a large number of mitochondria that provide the ATP for active uptake.

**E37.**(a)     Candidates’ knowledge of classification allowed many to make valid statements in their answers to part (a) about cats and cheetahs being from the same family or both being feline. Occasional candidates incorrectly referred to cats and cheetahs belonging to the same species. In part (b), some candidates were able to interpret the grafting of skin from one part of an animal to another as a test to see whether rejection would occur in these circumstances. The word ‘reaction’ was not considered to be synonymous with the specific biological meaning of rejection.

(b)     Candidates could have taken one of two approaches in answering part (a). They could either have concentrated on the speed of rejection or on the closeness of the genetic relationship between relevant animals. Despite this, this part of the question was not answered well and responses tended to lack the necessary precision to gain credit. Most candidates responded to the word reliable in part (b) with a suitable comment about the size of sample, but there were a few responses that were correctly worded in terms of the duration of the observation. Although many of the answers to part (c) were correctly based on the inference that cheetahs must share similar antigens as skin grafts were tolerated between animals, responses to part (d) were often poor. There were many confused accounts that failed to reflect the fundamental idea that proteins such as antigens are coded for by DNA and so any variation in the amino acid sequence of the protein implied a variation in the DNA coding. Candidates rarely answered in these simple terms.

**E38.**          (a)     Two thirds of candidates were aware that the vaccine needed to be given before girls are sexually active or likely to carry HPV. However, this was often poorly expressed by weaker candidates e.g. ‘this is when girls reach puberty or are sexually mature’. Other typical responses attempted to explain why the vaccine was given in general terms, rather than to this specific age group e.g. ‘to prevent girls developing cancer later in life’.

(b)     One third of the candidates had the idea that different types of HPV have different antigens. However, only better candidates wrote about the consequence of this in terms of memory cells and antibodies. Weaker candidates were let down by poor expression. Responses such as ‘the immune system does not recognise the virus’, ‘memory cells do not remember the virus’ and ‘antibodies cannot fight the virus’ were seen. A minority of candidates thought that the vaccine contained antibodies.

(c)     Nearly half of the candidates had the idea that more memory cells or more antibodies would be produced. However, only the very best candidates mentioned both for full credit. Two misconceptions were seen in the responses of weaker candidates. The first was that each injection of the vaccine was based on a different strain of HPV and would therefore provide ‘better immunity’. The second involved safety. They thought that ‘if given all at once, the immune response would be too strong’. Similarly, many unqualified references to the primary and secondary responses were seen.

(d)     Two thirds of the candidates gained at least one mark for stating that cancer takes many years to develop or that it takes time for young girls to become sexually active. Candidates who failed to score often wrote that ‘it takes time to develop immunity’ or ‘it takes many years to obtain data’. Some weaker candidates wrote about immunity being passed on to offspring or that it would take many years to vaccinate 80% of 12- 13 year olds.

(e)     75% of candidates gained full credit for stating that cancer can be caused by other factors or that the vaccine may not work. Weaker candidates often repeated information in the passage and wrote that smear tests are needed to remove abnormal cells before cancer develops. A minority thought that cancer is infectious and that smear tests detect the virus.

(f)      Only better candidates were aware that people who are vaccinated will destroy the virus or not act as carriers. Very few of these candidates appreciated that people who are not vaccinated are therefore more likely to meet people who are vaccinated. Many candidates simply repeated the stem of the question and stated that ‘if vaccinated, people cannot spread HPV to others’. Weaker candidates often referred to the ‘disease’ being destroyed rather than the virus.

**E39.**(a)     Relatively few candidates appeared to be aware that points on a graph should be joined with straight lines if it is felt that the position of intermediate points cannot be predicted reliably. Given that this decision had been made by candidates in drawing their graphs in stage 2, this was somewhat surprising.

(b)     Although many candidates were able to describe how the curve rose to a maximum value at 180 units or a dose of 0.25 g per kg, a significant number missed the point plotted for a zero dose. Other candidates misread the second point as representing a dose of 0.5 g per kg.

(c)     It remains disappointing that so few candidates can calculate percentage increase or decrease. There were many incorrect answers to this question, frequently from otherwise sound candidates.

(d)     Most candidates appeared to appreciate that calculating the dose per unit mass allowed differences in mass to be considered and a comparison to be made. Many responses, however, failed to gain credit because of the vague use of terms such as “bigger mice” and “size” rather than mass.

(e)     It would appear that some candidates had been taught about the immune response in much greater detail than required by the specification. This additional detail tended to confuse rather than help the candidates and reduced their marks for this question. It was relatively uncommon to see three marks awarded for what should have been a straightforward account. Common errors made by less able candidates involved the confusion of antibody and antigen or failing to identify the antigens as being on the surface of the sheep red blood cells.

(f)     Most candidates correctly pointed out that this investigation was carried out on mice and, therefore, the results might not apply to humans but only the better candidates were able to suggest a second valid reason.

**E40.**          (a)     (i)      This caused little difficulty for most candidates with the vast majority gaining at least one mark for suggesting that *C. difficile* is resistant to antibiotics. Although many candidates realised that the other bacterial species would be killed, they failed to gain a second mark by not stating that there would be an increase in the number of *C. difficile*.

(ii)     Most candidates gained this mark by suggesting that the immune system would be less effective. There were several answers linked to older people taking lots of antibiotics. These responses were not credited.

(b)     Although the majority of candidates obtained at least one of the two marks available, there was still some confusion, particularly with weaker candidates, about the precise role of methicillin. Most candidates realised it was a competitive inhibitor but a significant number referred to it possessing an active site. Approximately forty percent of candidates provided a clear accurate explanation of competitive inhibition by methicillin.

(c)     (i)      The majority of candidates had little difficulty explaining that some of these patients were already ill and this illness could be the cause of death.

(ii)     The vast majority of candidates gained this mark by describing the increase in the number of deaths up to 2006 followed by a decrease.

(iii)     Less than a third of candidates could correctly calculate the percentage increase in the number of deaths caused by MRSA in Wales from 1996 to 2006. A small percentage of candidates obtained a single mark for reading figures from the graph but almost sixty percent scored zero.

**E41.**          (a)     Nitrogen-fixing was the commonest wrong answer in this question. The majority of responses were correct.

(b)     This question was answered poorly because students did not think through the processes that were taking place in the reed bed. There were many incorrect responses referring to processes in the reeds that result in the formation of nitrates from ammonia / nitrite. Some then went on to gain one mark for active transport of these nitrates into the plant roots. Better students correctly linked the use of ATP from aerobic respiration in the active transport of nitrates, and wrote clearly and concisely. There was a surprising amount of confusion between diffusion and active transport, with active transport being said to be needed to diffuse nitrogen-containing substances from areas of high to low concentration. The oxygen was also thought to create a concentration gradient to allow the roots to take up the nitrogen-containing substances by diffusion.

(c)     There were some very clear answers to part (i) from students who understood that too fast a flow would not allow time for the nitrification to occur, hence the decrease in concentration of nitrates. There was also not enough time for the saprophytes to decompose the sewage to release ammonium compounds. Some failed to mention the ammonia being converted. Other answers suggested that the soil would become waterlogged, preventing the action of the nitrifying bacteria, or that the reeds would take up more of the nitrates or that numbers of denitrifying bacteria would increase, converting the nitrate to nitrogen gas. A number thought that if the flow was too fast, the reeds would be unable to take up the nitrates, so they would end up in the lake. The fast flow was also thought to reduce the oxygen concentration in the water, thus preventing the action of the nitrifying bacteria. There was also confusion with leaching and eutrophication. There were only very occasional references to the bacteria being washed away by the fast flow. The fast flow was also said to maintain a steep diffusion gradient and increase uptake by the plant roots.

In part (ii), it was clear that many students had learnt this topic thoroughly and included all marking points. Weaker students could not explain the increase in decomposers breaking down the dead plants and using up the oxygen in the water in their respiration. The algae were often described as ‘feeding’ on the nitrates. A common incorrect reason for the death of the fish was a lack of food once the plants in the lake died. A minority of students had no understanding of the process of eutrophication and thought that dehydration and osmosis caused the fish to die or that high nitrate concentrations were toxic to both fish and algae. Increasing concentrations of carbon dioxide were also thought to be responsible for the death of the fish.

**E42.**The starting point for questions requiring longer responses must be careful determination of precisely what is required. This question required students to explain how vaccines protect people against disease but few could resist the temptation to describe in great detail everything they knew about immunology. This often resulted in the allocated space being filled with material that, at the very best, could only be regarded as of marginal relevance. Most students should have been able to access the first three points on the scheme and indicate that antigens on weakened or dead pathogens stimulated the production of memory cells. The fact that credit was not always awarded stemmed from interchangeability of the terms pathogen and disease, and uncertainty over the origin and nature of memory cells. The second part of the mark scheme referred to the generation of a secondary immune response. Those students who finally arrived at this concept, often did no more than offer a few passing thoughts at the very end of the page or on an extra sheet. As always with questions on this topic, the use of language was often far from convincing and there were many references to antibodies "fighting" and memory cells "remembering".

**E43.**          (a)     (i)      Many candidates gave a generic answer, failing to refer to the passage as instructed. These candidates often scored only one mark for explaining that an antigen causes an immune response. Candidates who scored both marks used the information given to explain that, in this example, the antigen was a protein on Chlamydia.

(ii)     In this question, candidates were more confident in using the information from the passage and most gained at least a mark for explaining that the proteins on the Chlamydia cell and the heart were similar. There were candidates who confused antigens with antibodies and even enzymes but many candidates gained a second mark, usually by explaining that antibodies would attack the heart muscle cells. There were a number of excellent answers that showed a clear understanding of the immune response.

(b)     A number of candidates did not go further than the information given in the question, simply stating that the vaccination would prevent Chlamydia infection. This was not credited. Candidates who considered the information in the passage wrote about the possibility of preventing atheroma or, if the human proteins were similar to those in the mouse, the risk of causing heart disease. There were also creditworthy references to the cost of a vaccination campaign being higher than alternative methods of reducing the incidence of Chlamydia. Unfortunately, many limited their answer to just one factor rather than evaluating the suggestion as instructed.

**E44.**(a)     Most students correctly identified the evidence as relating to four polypeptide chains. Incorrect answers usually centred on the presence of variable regions or of hydrogen bonds. There was some evidence of the difficulties that students find in interpreting diagrams with numerous references to two polypeptide chains.

(b)     Most students clearly appreciated that an antigen is able to bind to an antibody to form an antigen-antibody complex. Not all, however, were able to identify the binding site of the antibody as having a complementary shape to the antigen. Many of the less able students confused antibodies with enzymes. Use of the term active site rather than binding site was perhaps understandable, but many went considerably beyond this in writing of substrates and enzyme-substrate complexes. There were also many students who failed to maintain the necessary focus and wrote at length of plasma cells, memory cells and vaccines.

**E45.**A number of misconceptions about the immune system, immune response and vaccination were commonly expressed by students in different parts of this question. The same, or similar, misconceptions have been seen in previous papers and have been commented upon in the reports.

(a)    (i)      Many students did not focus on a comparison of a vaccine consisting of whole cells that had been heat treated with one containing only parts of the bacterial cell. The former might not have been killed, or might contain toxins, whereas the latter could not be alive and would not contain toxin. Many students felt that the introduction of whole cells with their many antigens might overwhelm the immune system. These students did not seem to appreciate that our immune systems are exposed to multiple antigens on a daily basis and are not overwhelmed.

(ii)     The examiners were looking for the idea that a whole-cell vaccine would contain many different antigens and each would lead to the production of an antibody specific to it. Only about a fifth of students obtained both marks here. Most students simply wrote that there would be more antigens and thus more antibodies. This made it impossible for examiners to know whether or not they understood the ideas of different antigens and different, specific antibodies.

(b)     This part produced roughly equal percentages of students obtaining 3, 2, 1 or 0 marks. Many started with the assumption that people with whooping cough would have no antibody against the toxin; because if they had the antibody, they wouldn’t get ill. Indeed, many stated that only people who had been vaccinated against whooping cough could have antibody against the toxin. In essence, they did not seem to understand that an immune response occurs naturally when someone becomes infected with a pathogen.

(c)     Many appeared to find it difficult to accept that a doctor’s diagnosis might be wrong and wrote about vaccines not working and the bacterium mutating. About a quarter of students did spot that the scientists’ work suggests there might not be a real rise in whooping cough cases and that this might be linked to mis-diagnosis. Another quarter identified one of these points.

**E46.**          (a)      It was clear that many students had learned to define key terms and these students gave clear and accurate answers. Others were less specific, most commonly stating that a pathogen was “something” that caused disease. This was insufficient to gain the mark.

(b)      Many students gained all four marks. The main reason for failing to gain marks was for not specifying the idea of fusion of the lysosome with the phagocytic vesicle. Some students, however, failed to read the question properly and wrote in more general terms about B cells and T cells or lymphocytes, ignoring phagocytes.

(c)      Poor terminology let down students who clearly understood the concepts. Too many used active site terminology and were unable to gain the first marking point. The second marking point tended to be gained more often with the majority of students correctly referring to antibody and antigen being complementary, fitting and binding. Those who did not get this mark failed to do so by writing about the antibodies binding to the pathogens rather than to the antigens.

**E47.**(a)    (i)      This was done well by many students. Where the mark was not given it was usually because a student stated that the antigen was foreign but did not go on to add that it would cause an immune response.

(ii)     Difficulties with this question were linked to poor understanding of the ways in which substances pass through membranes. Weaker answers referred to the antigens not being ‘allowed’ through, rather than incompatibility between the shape of the antigen and the shape of trans-membrane protein channels.

(b)     Many students scored full marks. Rather than microfold cells being the route by which vaccines could enter the body, weaker responses included ideas such as the vaccine being given as treatment for a disease, or people being injected with microfold cells.

**E48.**The questions in each part of this question related to the short comprehension passage. Very many students appeared to pay little attention to the contents of the passage and many did not read the questions carefully enough

(a)     This produced some very good answers where students clearly described invasion by pathogens, their reproduction, invasion of host cells and production of toxins. They then linked these to death of host cells and tissues. About a third obtained three marks. Some students concerned themselves with the immune response that they had been told the people with AIDS did not have; they scored very poorly.

(b)      A commonly stated misconception in this question was that vaccines are given after infection, to treat rather than to prevent.

(i)       Nearly forty percent scored one mark, either for the idea of there not being time for antibodies to destroy HIV, or for there not being time for memory cells of the immune system to respond. Few gave complete explanations; for example, following up the antibody point by stating that antibodies do not / cannot enter cells to follow HIV.

(ii)      Students did better in this question and both lines of argument were seen. The commonest correct answers involved different antigens being produced and antibody not being able to bind any more. Students’ powers of expression frequently let them down here.

(c)     This question was notable for the use of rote answers; “there might be side effects”, “it might cost too much”, “it has only been tested on animals”, “it’s not ethical.....”. The question asks about these types of vaccines, the ones discussed in the passage, with their various characteristics. A few very good answers were seen, where students did discuss, for example, the possible perils of an attenuated virus as a vaccine, when HIV is stated to show a lot of variability. In a ‘real world’ context, it was interesting to note that almost no students appreciated that one would become HIV positive if one had been vaccinated. Encouragingly, one of the more common acceptable observations was that vaccination might encourage unsafe practices, which could spread HIV if the vaccine did not work.

**E49.**Parts (a)(ii), (b)(i), (b)(ii) and (c) proved to be good discriminators.

(a)    (i)      Nearly all students scored one mark and this was usually for suggesting that this method prevented the scientists from being biased. Unfortunately, the second suggestion provided by some also related to the scientists, rather than the volunteers; for example, ‘they may have a vested interest’. Students who scored a second mark often referred to reducing the placebo effect or psychological effects.

(ii)     One-third of students scored full marks. The most common mark points awarded were for suggesting that different types of cigarette contain different amounts of nicotine, different amounts may be absorbed, different amounts may be excreted and that the volunteers may have different blood volumes. Unfortunately, a lack of precision let down some students; for example, ‘they smoked different cigarettes’ and ‘they had different body masses’.

(b)    (i)      Just over half of students scored at least two marks. This was usually for mentioning that nicotine will not bind to the protein, so the smoker will not feel rewarded. Surprisingly, few students clearly expressed the idea that the vaccine stimulates the production of antibodies to nicotine, or that these antibodies bind to nicotine. A common misconception seen in weaker responses was that the vaccine *contains* antibodies to nicotine. Similarly, some students thought that *this* vaccine contained a weakened strain of bacteria. Generally, there were three incorrect approaches to this question, which were all due to not reading the introduction carefully enough. The first was that the vaccine *causes* the release of reward chemicals, meaning that a person would not need to smoke. The second was that the vaccine stops smokers from feeling addicted, rather than stopping them from feeling rewarded. The third was that the antibodies to nicotine bind to the protein in the brain, instead of to nicotine. Unfortunately, some students had the second mark disqualified for referring to the protein receptor in the brain as an enzyme. A minority also thought that the ability of the vaccine to stop people smoking could be spread within the population to other smokers by herd immunity.

(ii)     Just under half of students scored at least two marks. The most common mark points awarded were for appreciating that people choose to smoke, the vaccine would be expensive and less money would be needed to treat the effects of smoking. Relatively few referred to it being unethical not to treat smokers, or that money would be better spent in preventing people from smoking. Students who failed to score often gave vague responses; for example, ‘if it is free, more people will stop smoking’ and ‘it will prevent people from dying of cancer’. A minority suggested that the vaccine should not be used at all, due to the Government losing millions of pounds each year in tax on cigarettes. Some of the weakest responses did not answer the question set. These typically contained stock How Science Works phrases such as, ‘it is only one study’ and ‘we do not know the sample size’.

(c)     It was disappointing that only one-fifth of students scored at least three marks. Again, weaker responses often contained stock How Science Works phrases, which did not apply specifically to this investigation. The question clearly asked students to *use the data* to evaluate the statement made by the journalist. This said, many students did note that high antibody responders are more likely to stop smoking. Many also realised that the placebo group and low antibody responders had a similar percentage of volunteers who stopped smoking. The next most accessible mark point was that the volunteers may start smoking again after five or six months. Better responses also noted the peak, or drop, in the concentration of antibodies. However, some failed to mention when this occurred, or quoted an incorrect time from the graph. Relatively few students suggested that only a small proportion of the population may be high antibody responders, or that the large sample size produced more reliable or representative results. Overall, it was evident that many students did not analyse the data in the graph and table in enough detail, particularly in relation to the timing of events.

**E52.**(a)    It was pleasing to see many good answers to this part that focused on how bacteria are destroyed by phagocytes. Some students drifted into general accounts of the immune response and others began by writing at length about how phagocytes find bacteria. About 30% obtained all three marks. It was common for students to be vague or wrong about the role of lysosomes. It was not uncommon to see references to lysosomes fusing with bacteria, rather than with the vacuole containing the bacteria. The examiners were looking for references to hydrolytic or digestive enzymes destroying the bacteria, rather than just enzymes breaking down bacteria.

(b)     80% obtained both marks. Those who failed to score usually included features of eukaryotic cells in their answers.

**E54.**(a)     Over 40% gave the correct response of regulator protein. Students who failed to score often gave a generic response about what should be in a vaccine, ‘antigen’ being insufficient for this mark. Students should ensure that they relate their answer to the information given in the question.

(b)     Students were expected to have seen from the passage that the LDL would bind to a receptor on the surface membrane of the liver cell and then use their knowledge of the structure of the membrane to suggest a suitable route into the cell, for example through a carrier protein or channel protein. Many students failed to use the information in the passage and therefore could only gain one mark. A significant number of students recognised that the LDL would be lipid soluble and could pass through the phospholipid bilayer. This alternative was also credited.

(c)     This question discriminated well with weaker students struggling to interpret the information given in the passage. They frequently confused the antibody-antigen response with enzymes, referring to active sites on the antibody or antigen and the formation of enzyme-substrate complexes. There was also mention of receptor "cells" and "antigens on the cell surface membrane of the regulator protein" by students who clearly had a poor understanding of the molecules. However, the more able students were able to gain two marks, usually for referring to the antibody binding to the regulator protein which prevented it from binding to the receptor.

(d)     It was clear that many students had learnt a generic answer to this type of question. Most students knew that the control group would be treated the same as the experimental group but the generic response of ‘give a placebo’ was insufficient because the answer needs to be related to the information in the question. The mark scheme required students to write about injection of saline without the monoclonal antibody. The second marking point, ‘treated the same’ was sometimes expressed as ‘given the same diet’, or the ‘same amount of exercise’ or ‘have the same amount of LDLs’. It is worth noting that the same treatment will include the medical experience and measurements taken, so identifying a particular control variable, such as diet, in isolation, was insufficient to gain this mark.

**E57.**(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.

**E58.**(a)     Almost 80% of students scored both marks, in a question which tested straightforward recall. Some described pathogens entering cells and reproducing without going on to clarify the damage that would have been caused to the cells. A minority misinterpreted the question and described two ways in which pathogens were transmitted.

(b)     The context of this question proved difficult for many students with fewer than half the students explaining that water would move out of the bacterial cell by osmosis because of the water potential gradient. A large number incorrectly wrote about water being drawn out of the blood and washing away the bacteria and many argued that water would enter the bacteria causing osmotic lysis. Few students went on to explain why the loss of water would kill the bacteria