**Q1.**Scientists investigated the control of blood glucose concentration in mice. They kept a group of normal mice without food for 48 hours. After 48 hours, the blood glucose concentrations of the mice were the same as at the start of the experiment.

(a)     Explain how the normal mice prevented their blood glucose concentration falling when they had **not** eaten for 48 hours.

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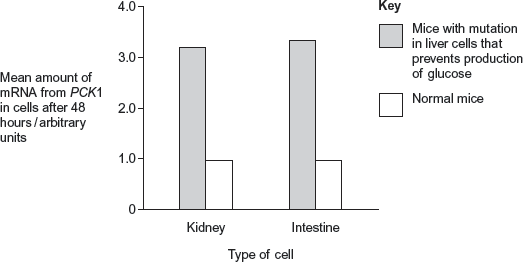
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**(3)**

The scientists then investigated mice with a mutation that prevents their liver cells making glucose. They kept a group of these mice without food for 48 hours. After 48 hours, the mean blood glucose concentrations of the mutant mice and the normal mice were the same.

The scientists investigated how blood glucose concentration is controlled in these mutant mice. An enzyme required for synthesis of glucose is coded for by a gene called *PCK*1. The scientists measured the mean amount of mRNA produced from this gene in cells from the kidneys and intestines of normal mice and mutant mice. They did this with mice that had previously been without food for 48 hours.

The scientists’ results are shown in the graph.



(b)     Use information from the graph to suggest how blood glucose concentration is controlled in the mutant mice, compared with the normal mice.

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**(3)**

(c)     The scientists performed statistical tests on the data shown in the graph, to see whether the differences in the amount of mRNA in cells from normal and mutant mice were significant. Both the probability values they obtained were p<0.01.

Explain what this means about the differences in the amounts of mRNA produced.

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**(2)**

**(Total 8 marks)**

**Q2.**Some mice have diabetes. The diabetes causes the blood glucose concentration to become very high after a meal. Scientists investigated the use of an inhibitor of amylase to treat diabetes.

The scientists took 30 mice with diabetes and divided them into two groups, **A** and **B**.

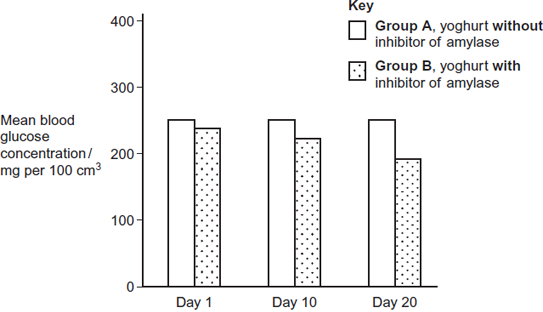
•        **Group A** was given yoghurt **without** the inhibitor of amylase each day.

•        **Group B** was given yoghurt **with** the inhibitor of amylase each day.

Apart from the yoghurt, all of the mice were given the same food each day.

The scientists measured the blood glucose concentration of each mouse, 1 hour after it had eaten. This was done on days 1, 10 and 20 after the investigation started.

The following figure shows the scientists’ results.

  
        Days after the investigation started

(a)     **Group A** acted as a control in this investigation.

Explain the purpose of this group.

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**(2)**

(b)     Apart from the yoghurt, it was important that all of the mice were given the same food each day.

Give **two** reasons why it was important that all of the mice were given the same food each day.

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

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

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**(2)**

(c)     The scientists’ hypothesis was that adding the inhibitor of amylase to the food would lead to a lower blood glucose concentration.

Use your knowledge of digestion to suggest how the addition of the inhibitor could lead to a lower blood glucose concentration.

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**(2)**

(d)     Give **one** reason why these results may **not** support the use of the inhibitor of amylase to treat diabetes in mice.

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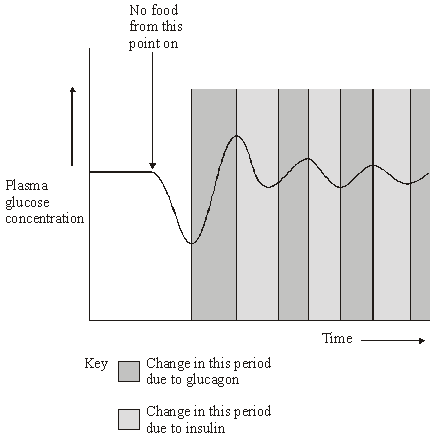
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**(2)**

**(Total 8 marks)**

**Q3.**          Homeostatic mechanisms maintain a constant environment in the body.

(a)     The graph shows changes in plasma glucose concentration that occurred in a person who went without food for some time.



Use evidence from the graph to explain the role of negative feedback in the control of plasma glucose concentration.

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**(5)**

(b)     How does maintaining a constant body temperature allow metabolic reactions in cells to proceed with maximum efficiency?

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**(5)**

**(Total 10 marks)**

**Q4.**          (a)     Explain how insulin lowers the concentration of blood glucose.

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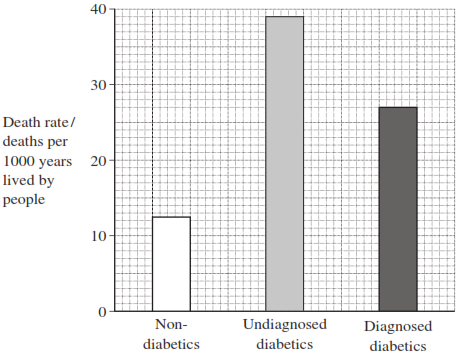
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**(3)**

(b)     Doctors studied a large group of people. They recorded the death rates for non-diabetic people, undiagnosed diabetics and diagnosed diabetics.

They gave the death rates as deaths per 1000 years lived by people.

The graph shows these death rates.



(i)      Calculate the ratio of the death rate of diagnosed diabetics to undiagnosed diabetics.

Ratio .........................................

**(2)**

(ii)     People with undiagnosed diabetes were not receiving treatments, such as insulin injections. Suggest **one** reason for the difference in death rates for undiagnosed and diagnosed diabetics.

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**(2)**

**(Total 7 marks)**

**Q5.**          Exenatide is a drug used for treating Type 2 diabetics. Scientists investigated the effects of exenatide on insulin production.

The scientists used three groups of volunteers who were treated in the following ways.

**Group 1**: healthy, non-diabetics who were injected with exenatide in salt solution   
**Group 2**: Type 2 diabetics who were injected with exenatide in salt solution   
**Group 3**: Type 2 diabetics who were injected with salt solution.

Three hours after these injections, the scientists injected the same amount of glucose into the blood of each volunteer.  
The scientists measured the rate of insulin production by each person before and after injecting the glucose.

(a)     (i)      **Group 1** and **Group 3** were control groups in this investigation.

Explain why each group was used.

**Group 1** ...............................................................................................

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**Group 3** ...............................................................................................

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**(2)**

(ii)     The scientists measured the rates of insulin production per unit body mass.

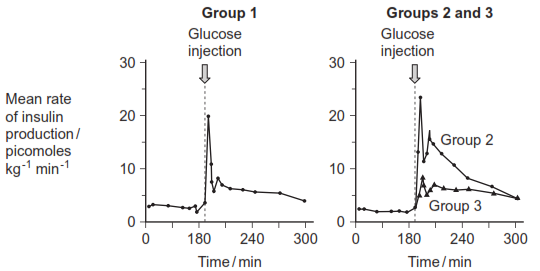
Explain why.

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**(1)**

(b)     The graphs show the mean rates of insulin production for each group.



Suggest how exenatide could help people with Type 2 diabetes.

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**(3)**

**(Total 6 marks)**

**Q6.**          (a)     Describe how insulin reduces the concentration of glucose in the blood.

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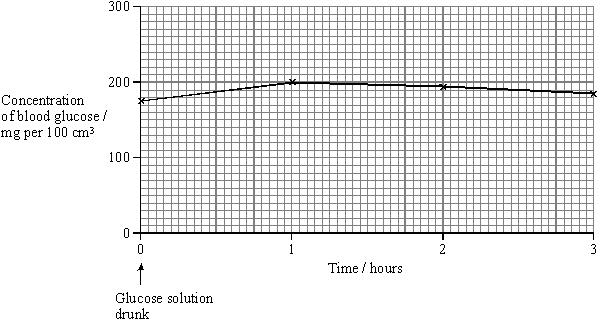
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**(3)**

Some people produce no insulin. As a result they have a condition called diabetes. In an investigation, a man with diabetes drank a glucose solution. The concentration of glucose in his blood was measured at regular intervals. The results are shown in the graph.



(b)     Suggest **two** reasons why the concentration of glucose decreased after 1 hour even though this man’s blood contained no insulin.

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

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

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**(2)**

(c)     The investigation was repeated on a man who did not have diabetes. The concentration of glucose in his blood before drinking the glucose solution was 80 mg per 100 cm3. Sketch a curve on the graph to show the results you would expect.

**(1)**

(d)     The diabetic man adopted a daily routine to stabilise his blood glucose concentration within narrow limits. He ate three meals a day: breakfast, a midday meal and an evening meal. He injected insulin once before breakfast and once before the evening meal.

The injection he used before breakfast was a mixture of two types of insulin. The mixture contained slow-acting insulin and fast-acting insulin.

(i)      Explain the advantage of injecting both types of insulin before breakfast.

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**(2)**

(ii)     One day, the man did not eat a midday meal. Suggest **one** reason why his blood glucose concentration did not fall dangerously low even though he had injected himself with the mixture of insulin before breakfast.

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**(1)**

**(Total 9 marks)**

**Q7.**          Diabetes is a disorder affecting the ability to control blood glucose concentration. One type of diabetes can be due to an abnormality of the insulin receptors in the cell surface membranes of cells in the liver and muscles. A high blood glucose concentration and the presence of glucose in the urine are signs of this type of diabetes.

(a)     (i)      Suggest **one** way in which the insulin receptors might be abnormal.

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**(1)**

(ii)     Explain how the presence of abnormal insulin receptors results in a high blood glucose concentration.

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**(2)**

(iii)     Explain how the kidneys normally prevent glucose appearing in the urine of a non-diabetic person.

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**(3)**

(b)     Twin studies have been used to determine the relative effects of genetic and environmental factors on the development of this type of diabetes. The table shows the concordance (where both twins have the condition) in genetically identical and genetically non-identical twins.

|  |  |
| --- | --- |
| **Concordance in genetically identical twins / %** | **Concordance in genetically non-identical twins /%** |
| 85 | 35 |

(i)      What do the data show about the relative effects of environmental and genetic factors on the development of diabetes?

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**(1)**

(ii)     Suggest **two** factors which should be taken into account when collecting the data in order to draw valid conclusions.

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

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**(2)**

**(Total 9 marks)**

**Q8.**(a)     What is homeostasis?

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**(1)**

(b)     Describe the role of the hormone glucagon in the control of blood sugar concentration.

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**(4)**

(c)     The kidney removes various substances from the blood plasma. The clearance value for a substance is the volume of blood cleared of that substance by the kidney in one minute. This clearance value can be calculated using the equation.



|  |  |  |
| --- | --- | --- |
|  | where the concentration of a substance in the blood is  the concentration of a substance in the urine is  the volume of urine produced is | P g cm-3 U g cm-3 V cm3 per minute |

(i)      Use the equation to work out the clearance value of glucose.

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**(1)**

(ii)     Explain how the activity of the kidney results in this clearance value for glucose.

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**(3)**

**(Total 9 marks)**

**Q9.** The diagram shows some of the events which maintain blood glucose concentration in a mammal.



(a)     Name

(i)      hormone **A**; .........................................................................................

(ii)     organ **B**. ..............................................................................................

**(2)**

(b)     Explain why the events shown in the diagram can be described as an example of negative feedback.

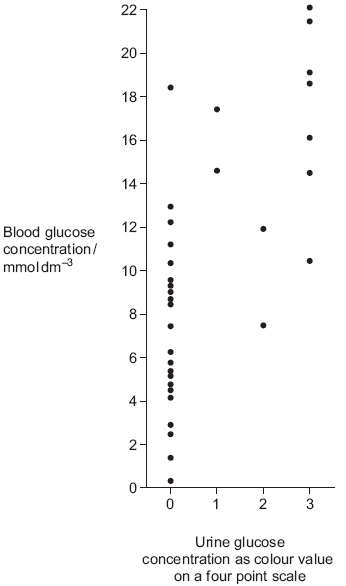
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**(1)**

**(Total 3 marks)**

**Q10.**          (a)     Technicians in a hospital laboratory tested urine and blood samples from a girl with diabetes at intervals over a one-year period. Each time the technicians tested her urine, they also measured her blood glucose concentration. Their results are shown in the graph.



(i)      The girl who took part in this investigation was being successfully treated with insulin. The graph shows that on some occasions, the concentration of glucose in her blood was very high. Suggest why.

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**(2)**

(ii)     Use the graph to evaluate the use of the urine test as a measure of blood glucose concentration.

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**(3)**

(b)     Diabetic people who do not control their blood glucose concentration may become unconscious and go into a coma. A doctor may inject a diabetic person who is in a coma with glucagon. Explain how the glucagon would affect the person’s blood glucose concentration.

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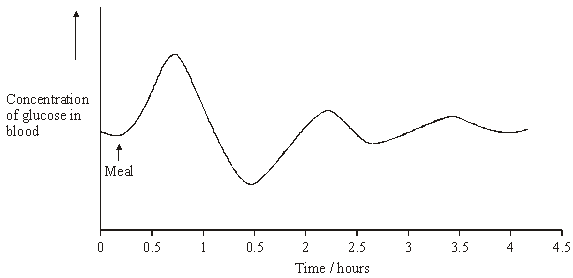
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**(2)**

**(Total 7 marks)**

**Q11.**          (a)     The graph shows changes in the concentration of glucose in a person’s blood following a meal.



Changes in the concentration of glucose are controlled by the hormones glucagon and insulin. Write the letters **X** and **Y** on the graph to show

**X** a time when glucagon secretion would be high;

**Y** a time when insulin secretion would be high.

**(1)**

**S**       (b)     Many diabetics require regular injections of insulin. Describe how bacteria can be genetically modified to produce human insulin.

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**(4)**

**(Total 5 marks)**

**Q12.**          (a)     Sucrose, maltose and lactose are disaccharides.

(i)      Sucrase is an enzyme. It hydrolyses sucrose during digestion. Name the products of this reaction.

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**(2)**

(ii)     Sucrase does **not** hydrolyse lactose. Use your knowledge of the way in which enzymes work to explain why.

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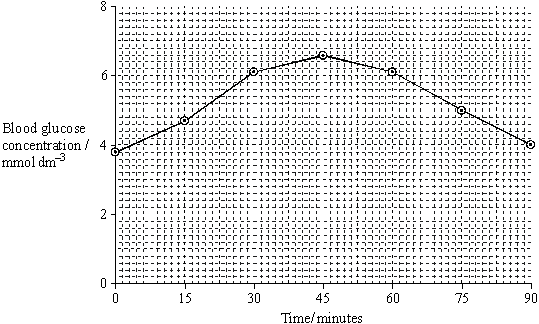
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**(2)**

(b)     A woman was given a solution of sucrose to drink. Her blood glucose concentration was measured over the next 90 minutes. The results are shown on the graph.



(i)      Describe how the woman’s blood glucose concentration changed in the period shown in the graph.

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**(2)**

(ii)     Explain the results shown on the graph.

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**(2)**

**(Total 8 marks)**

**Q13.**          A glucose biosensor is an instrument used to measure glucose concentration. It contains an enzyme called glucose oxidase.

(a)     A glucose biosensor detects only glucose. Use your knowledge of the way in which enzymes work to explain why.

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**(3)**

(b)     It is better to use a biosensor than the Benedict’s test to measure the concentration of glucose in a sample of blood. Suggest **two** reasons why.

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

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**(2)**

(c)     (i)      Diabetes mellitus is a disease that can lead to an increase in blood glucose concentration. Some diabetics need insulin injections. Insulin is a protein so it cannot be taken orally. Suggest why insulin cannot be taken orally.

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**(1)**

(ii)     A drug company produced a new type of insulin. Scientists from the company carried out a trial in which they gave this new type of insulin to rats. They reported that the results of this trial on rats were positive. A newspaper stated that diabetics would benefit from this new drug. Suggest **two** reasons why this statement should be viewed with caution.

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

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**(2)**

**(Total 8 marks)**

**Q14.**          (a)     Adrenaline binds to receptors in the plasma membranes of liver cells. Explain how this causes the blood glucose concentration to increase.

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**(2)**

(b)     Scientists made an artificial gene which codes for insulin. They put the gene into a virus which was then injected into rats with type I diabetes. The virus was harmless to the rats but carried the gene into the cells of the rats.

The treated rats produced insulin for up to 8 months and showed no side-effects. The scientists measured the blood glucose concentrations of the rats at regular intervals. While the rats were producing the insulin, their blood glucose concentrations were normal.

(i)      The rats were not fed for at least 6 hours before their blood glucose concentration was measured. Explain why.

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**(1)**

(ii)     The rats used in the investigation had type I diabetes. This form of gene therapy may be less effective in treating rats that have type II diabetes. Explain why.

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**(1)**

(iii)    Research workers have suggested that treating diabetes in humans by this method of gene therapy would be better than injecting insulin. Evaluate this suggestion.

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**(4)**

**(Total 8 marks)**

**Q15.**(a)     When insulin binds to receptors on liver cells, it leads to the formation of glycogen from glucose. This lowers the concentration of glucose in liver cells.

Explain how the formation of glycogen in liver cells leads to a lowering of blood glucose concentration.

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**(2)**

People with type II diabetes have cells with low sensitivity to insulin. About 80% of people with type II diabetes are overweight or obese. Some people who are obese have gastric bypass surgery (GBS) to help them to lose weight.

Doctors investigated whether GBS affected sensitivity to insulin. They measured patients’ sensitivity to insulin before and after GBS. About half of the patients had type II diabetes. The other half did not but were considered at high risk of developing the condition.

The table below shows the doctors’ results. The higher the number, the greater the sensitivity to insulin.

|  |  |  |  |
| --- | --- | --- | --- |
|  | **Patients** | **Mean sensitivity to insulin / arbitrary units (± SD)** | |
|  | **Before gastric bypass surgery** | **1 month after gastric bypass surgery** |
|  | Did not have diabetes | 0.55 (± 0.32) | 1.30 (± 0.88) |
|  | Had type II diabetes | 0.40 (± 0.24) | 1.10 (± 0.87) |

(b)     The doctors concluded that many of the patients who did not have type II diabetes were at high risk of developing the condition.

Use the data in the table to suggest why they reached this conclusion.

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**(2)**

(c)     The doctors also concluded that GBS cured many patients’ diabetes but that some were not helped very much.

Do these data support this conclusion? Give reasons for your answer.

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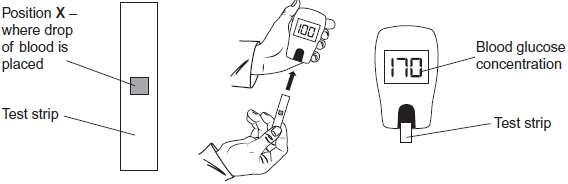
**(3)**

**(Total 7 marks)**

**Q16.Resource A**

A glucometer is a device used to measure blood glucose concentration. A person uses a test strip that goes into the glucometer. They put a drop of blood onto the test strip. There are substances on the test strip that produce a colour change with glucose. The higher the concentration of glucose, the deeper the colour produced. The glucometer measures the depth of colour produced and converts this into a glucose concentration. A new test strip is used for each blood test.

**Figure 1 – glucometer and test strip**

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The following equations show how the substances on the test strip produce a colour change.

Glucose + oxygen    gluconic acid + hydrogen peroxide

Hydrogen peroxide + dye with colour A    dye with colour B + water

Non-diabetics have no glucose in their urine. Diabetics have glucose in their urine if their blood glucose concentration rises above about 170 mg 100 cm–3.

Before the glucometer was available, diabetics used test strips to measure the concentration of glucose in their urine as a means of measuring their blood glucose concentration. When testing urine, the colour of the test strip is compared against a colour chart which gives a glucose concentration range for the colour produced.

**Resource B**

There are two types of diabetes: type 1 and type 2.

•        People with type 1 diabetes do not produce enough insulin.

•        People with type 2 diabetes do produce insulin but have cells which do not respond to insulin.

Doctors use a glucose tolerance test to help diagnose people with diabetes. They start each test after a person has not eaten overnight. They measure a person’s blood glucose concentration.

The person then drinks a solution containing 75 g of glucose. The doctors measure the person’s blood glucose concentration 2 hours later. During the test, the person remains at rest.

**Figure 1** shows three diagnoses that can be made from the results of the test.

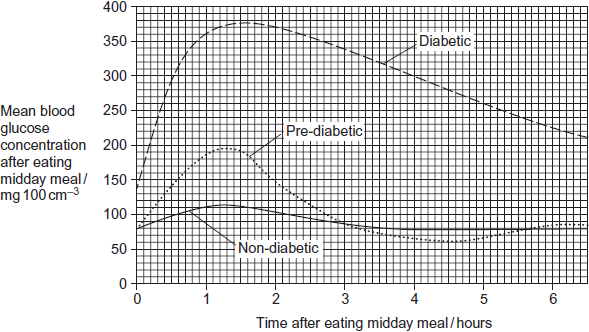
**Figure 2 – glucose tolerance test results and diagnoses**

|  |  |  |  |
| --- | --- | --- | --- |
|  | **Blood glucose concentration after 2 hours / mg 100 cm–3** | **Diagnosis** | **Comments** |
|  | ≤ 110 | Non-diabetic | Low risk for future diabetes |
|  | Between 140 and 200 | Pre-diabetic | High risk for future diabetes. Some doctors recommend that the upper value should be lowered to 180 mg 100 cm–3 |
|  | ≥ 200 | Diabetic | Confirm by doing a second test |

A researcher monitored the mean blood glucose concentration of a non-diabetic, a pre-diabetic and a diabetic after they had each eaten a midday meal.

His results are shown in **Figure 3**.

**Figure 3**



A laboratory worker suspected she had type 2 diabetes but did not have a glucometer.  
Instead she added a drop of her blood to a test strip and used a colour chart to estimate her blood glucose concentration as 140 mg 100 cm–3.

Is it valid to conclude that she did have type 2 diabetes?

Use this information, and **Resource A** and **Resource B**, to explain your answer.

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**(Total 3 marks)**

**Q17.**There are two types of diabetes: type 1 and type 2.

•        People with type 1 diabetes do not produce enough insulin.

•        People with type 2 diabetes do produce insulin but have cells which do not respond to insulin.

Doctors use a glucose tolerance test to help diagnose people with diabetes. They start each test after a person has not eaten overnight. They measure a person’s blood glucose concentration.

The person then drinks a solution containing 75 g of glucose. The doctors measure the person’s blood glucose concentration 2 hours later. During the test, the person remains at rest.

**Figure 1** shows three diagnoses that can be made from the results of the test.

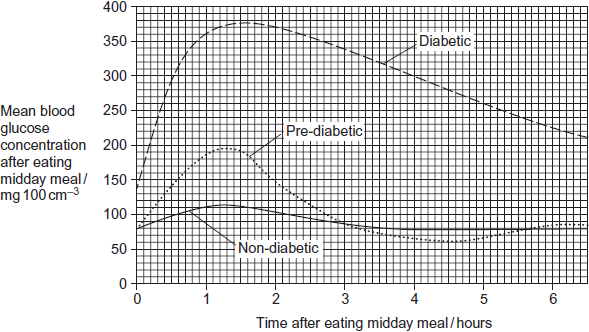
**Figure 1 – glucose tolerance test results and diagnoses**

|  |  |  |  |
| --- | --- | --- | --- |
|  | **Blood glucose concentration after 2 hours / mg 100 cm–3** | **Diagnosis** | **Comments** |
|  | ≤ 110 | Non-diabetic | Low risk for future diabetes |
|  | Between 140 and 200 | Pre-diabetic | High risk for future diabetes. Some doctors recommend that the upper value should be lowered to 180 mg 100 cm–3 |
|  | ≥ 200 | Diabetic | Confirm by doing a second test |

A researcher monitored the mean blood glucose concentration of a non-diabetic, a pre-diabetic and a diabetic after they had each eaten a midday meal.

His results are shown in **Figure 2**.

**Figure 2**



(a)     People with type 1 diabetes are described as being insulin-dependent.  
Suggest why they are described as insulin-dependent.

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**(1)**

(b)     Some people with type 2 diabetes have cells which do **not** respond to insulin.  
Explain how this leads to a reduced ability to regulate blood glucose concentration.

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**(3)**

(c)     During a glucose tolerance test the person remains at rest.  
Why is it important that the person remains at rest?

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**(2)**

(d)     Use **Figure 2** to calculate how many times the maximum mean blood glucose concentration of the pre-diabetic is greater than the maximum of the non-diabetic person. Show your working.

Answer = ..................................

**(2)**

(e)     Give **three** differences between the method used by the researcher to obtain the results in **Figure 2** and the method doctors use to carry out a glucose tolerance test.

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

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

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

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

3 .....................................................................................................................

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

**(3)**

(f)     Some doctors have recommended that the upper value used in the glucose tolerance test should be lowered to 180 mg 100 cm–3.  
Using information from **Figure 1** and **Figure 2**, suggest why.

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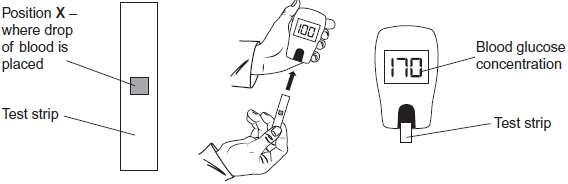
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**(3)**

**(Total 14 marks)**

**Q18.**A glucometer is a device used to measure blood glucose concentration. A person uses a test strip that goes into the glucometer. They put a drop of blood onto the test strip. There are substances on the test strip that produce a colour change with glucose. The higher the concentration of glucose, the deeper the colour produced. The glucometer measures the depth of colour produced and converts this into a glucose concentration. A new test strip is used for each blood test.

**Figure – glucometer and test strip**

****

The following equations show how the substances on the test strip produce a colour change.

Glucose + oxygen    gluconic acid + hydrogen peroxide

Hydrogen peroxide + dye with colour A    dye with colour B + water

Non-diabetics have no glucose in their urine. Diabetics have glucose in their urine if their blood glucose concentration rises above about 170 mg 100 cm–3.

Before the glucometer was available, diabetics used test strips to measure the concentration of glucose in their urine as a means of measuring their blood glucose concentration. When testing urine, the colour of the test strip is compared against a colour chart which gives a glucose concentration range for the colour produced.

(a)     Identify all the substances located at position **X** on the test strip before a drop of blood is added.

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

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

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

**(2)**

(b)     Before the glucometer was available, diabetics used test strips to measure the concentration of glucose in their urine as a means of measuring their blood glucose concentration.

Give **two** reasons why this method of testing urine would **not** give an accurate measurement of blood glucose concentration.

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

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

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

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

**(2)**

**(Total 4 marks)**

**M1.**(a)     1.      Release of glucagon;

2.      Leads to formation of glucose in liver (cells);

*Reject: glucagon breaks down glycogen, or any other biological molecule*

3.      From non-carbohydrates / amino acids / fatty acids.

*Accept: gluconeogenesis / references to glycogen as source of glucose*

**3**

(b)     1.      Mutant mice (mRNA suggests) make a lot of (the) enzyme;

*Accept: PCK1 made (for enzyme made)*

2.      Mutant mice use kidney / intestine (cells) to make glucose;

*Accept: use other organ (than liver)*

3.      Normal mice do this much less / normal mice use liver cells.

**3**

(c)     1.      Differences significant;

*Reject: references to results being significant once*

2.      Probability of difference being due to chance less than 0.01 / 1% / 1 in 100 / probability of difference not being due to chance more than 0.99 / 99% / 99 in 100.

*Ignore: references to 0.05 / 5% / 5 in 100*

**2**

**[8]**

**M2.**(a)     1.      To show the effect of the inhibitor / drug;

2.      To show the effect of yoghurt (on its own does not affect blood  
glucose);

**2**

(b)     1.      Food is a factor affecting blood glucose / different foods contain   
different amounts of starch / glucose / sugar / carbohydrate;

*Accept converse*

2.      To keep starch / fibre intake the same / similar;

*Accept something in food which affects the inhibitor*

**2**

(c)     1.      Fewer E-S complexes formed;

2.      (With inhibitor) less / no starch digested to maltose ;

*Require knowledge that maltose comes from starch*

3.      (So) less / no glucose from maltose;

*Require knowledge that glucose comes from maltose*

*Accept no glucose*

4.      (So) less absorption of glucose (from gut);

**2 max**

(d)     **Suitable reason; with explanation;**

Paired responses – do not mix and match

*Ignore references to correlation does not prove causation,   
it could be due to other factors*

Examples,

1.      Need larger sample / only 30 mice / only 15 mice in each group;

*Accept small sample size*

2.      Might not be representative / anomalies might have a bigger or smaller effect;

*Accept mean not reliable*

***OR***

3.      Investigation only lasted 20 days;

*Experiment was not long enough*

4.      Can’t see what longer term effects are;

***OR***

5.      Fall in blood glucose is small / numbers from graph;

6.      Mice with inhibitor still have a large rise in blood glucose / so don’t know if differences significant;

*Accept differences are due to chance*

***OR***

7.      No stats / SDs / SEs;

8.      So don’t know if differences significant;

***OR***

9.       Blood glucose could continue to fall;

10.     which could be harmful;

***OR***

11.     No group without yoghurt;

12.     So cannot compare to other groups;

**2 max**

**[8]**

**M3.**          **Quality of Communication**

The answers to all sections of this question require the use of continuous prose.  
Quality of language should be considered in crediting points in the scheme. In order to gain credit, answers should be expressed logically and unambiguously, using scientific terminology where appropriate.

(a)     1.      Deviation of a value from norm initiates corrective mechanisms;

2.      fluctuations in plasma glucose concentration detected by hypothalmus / islet cells in pancreas;

3.      initial decrease, no food given (in plasma glucose) stimulates (increased) secretion of glucagon;

4.      increases (in plasma glucose) stimulate (increased) secretion of insulin from β cells as secretors;

5.      correct ref. to interconversion of glycogen / glucose / increased / decreased uptake of glucose by cells (as appropriate) / correct ref to change in membrane permeability;

**5**

(b)     1.      Body temp. / 37 °C  is optimum temp for enzymes;

2.      excess heat denatures enzymes / alters tertiary structure / alters shape of active site / enzyme so substrate cannot bind / eq;

3.      reactions cease / slowed;

4.      too little reduces kinetic energy of molecules / moleculesmove more slowly;

5.      fewer collisions / fewer ES complexes formed’

**5**

**[10]**

**M4.**          (a)     Binds to receptor on target/liver/muscle cell;

*Reject reference to insulin as an enzyme*

Causes more transport/carrier proteins to become active/  
move to (plasma) membrane;

Glucose (diffuses) into cells (and lowers blood glucose);

(Enzymes in cells) convert glucose to glycogen;

Stimulates fatty acids/lipid/fat formation (from glucose);

Raises rate of respiration (in cells), using more glucose;

**3 max**

(b)     (i)      27 : 39;  
1 : 1.44;

*2 marks for 1 : 1.44*

*Accept 0.69 : 1*

*Accept 9 : 13*

**2**

(ii)     One suitable reason; with explanation;

e.g. undiagnosedDiabetic coma/brain cells not enough respiration;Due to low blood glucose/acidosis/dehydration;

Heart attacks/coronary heart disease;Due to faster atheroma formation/damage to arteries;

Kidney failure;Due to damage to blood vessels;

**2 max**

**[7]**

**M5.**          (a)     (i)      Group 1:  To see ‘normal’ response / non-diabetic response /  
as comparison with diabetic response;

Group 3:  To ensure any difference was due to exenatide /  
not due to salt / as comparison to show effect of exenatide on  
diabetes / to ensure effect was not psychosomatic /  
to see placebo effect;

**2**

(ii)     Different mass of person → different amount insulin secreted /  
larger person secretes more insulin / (valid) basis for comparisons  
between people;

*Ignore refs to accuracy*

**1**

(b)     Any **three** from:Increases sensitivity of pancreas cells to glucose;Increases insulin secretion (by pancreas) / similar insulin production  
as healthy / non-diabetic / Group 1;So more stimulation of cells / of liver / of muscles;Causes more glucose uptake (from blood) / blood glucose level  
lowered / kept at normal level / can control blood glucose conc.;Person can consume more carbohydrate / glucose / doesn’t need  
special diet / will not develop symptoms of diabetes;

**3 max**

**[6]**

**M6.**          (a)     insulin binds to specific receptors (on membranes);  
insulin activates carrier proteins / opens channels / causes more  
channels to form;  
insulin increases the permeability of liver / muscle cells / tissues to glucose;  
insulin action results in glucose conversion to glycogen / glycogenesis;

**3 max**

(b)     glucose is used in cell respiration / as energy source / in metabolism;

*(must qualify how glucose is used)*

glucose enters cells / converted to glycogen in cells;  
glucose is excreted / in urine;

*(do not credit no reabsorption of glucose in kidneys)*

**2 max**

(c)     line from 80 mg, increasing but keeping below line for diabetic,  
dropping to 80 mg;

*(line must stablise at, or fluctuate around 80 mg)*

**1**

(d)     (i)      fast acting insulin reduces blood glucose from breakfast;  
slow acting insulin reduces blood glucose from other meals  
before the evening meal / eliminates the need to inject at lunch;

*(must be a reference to the meals)  
(one mark if neither of the above but a clear reference is made to glucose conversion to glycogen);*

**2**

(ii)     glucagon is still active;   
glycogen converted to glucose / glycogenolysis;   
insulin injected at breakfast causes cells to take up glucose  
too slowly for levels to become dangerously low;  
person is not active so little glucose used in respiration;

*(do not credit statements about consuming large breakfasts)*

**1 max**

**[9]**

**M7.**          (a)     (i)      different shape / different tertiary structure /   
different sequence of amino acids;

**1**

(ii)     insulin unable to attach to receptors;  
reduced / no uptake of glucose into cells / no carrier proteins /   
channels for glucose transport;

**2**

(iii)     glucose reabsorbed / absorbed into blood;  
from proximal tubule;  
by active transport / involving membrane carriers;

**3**

(b)     (i)      larger genetic component;

*(must be comparative)*

**1**

(ii)     number of cases studied;  
matched samples;  
age of twins;  
named environmental factor;;

*(allow 2 marks for 2 different factors if no overlap in effect)*

family history of diabetes;  
method of diagnosis;  
same sex in non-identical twins;

**2 max**

**[9]**

**M8.**(a)     Maintaining a constant internal environment;

**1**

(b)     Binds to (specific) receptor;

On muscle / liver cell;

Activation of enzymes (in liver);

Hydrolysis of glycogen;

(Facilitated) diffusion of glucose out of (liver cells) cells;

Increases blood glucose levels;

**4**

(c)     (i)      0 / zero;

**1**

(ii)     1.      Filtration, out of blood (plasma) / into renal capsule;

2.      (Hydrostatic) pressure ;

3.      PCT;

4.      All reabsorbed;

5.      Active transport;

**3 max**

**[9]**

**M9.**          (a)     (i)      glucagon;

*Insist on spelling*

**1**

(ii)     liver;

**1**

(b)     A change to the normal level initiates a response which  
reduces the effect / reverses / acts against the change;

**1**

**[3]**

**M10.**          (a)     (i)      Eaten;

Containing carbohydrate / sugar;

Glucose absorbed from intestine / into blood;

Long time after insulin injection / needs more insulin / has not taken insulin;

Does not convert glucose to glycogen / glucose not taken up from blood;

**2 max**

(ii)     Shows positive correlation / directly proportional;

A range of results for a particular value / values (for different colours) overlap;

Urine test only an arbitrary scale / not directly related to concentration / colour is subjective / few colour values;

*Accept description*

**3**

(b)     Glycogen to glucose / glycogenolysis by activating enzymes;

*If name incorrect this disqualifies.*

Gluconeogenesis;

*Allow explanation in terms of glucose from a non-carbohydrate / named non-carbohydrate source.*

**2**

**[7]**

**M11.**          (a)     On graph:  **X** where glucose level is below norm  
AND           **Y** where glucose level is above norm;

**1**

(b)     EITHER  
1. Use m-RNA + reverse transcriptase to produce gene / (c)-DNA;  
2. Restriction enzyme to cut open plasmid;  
3. Add sticky ends (to insulin gene and to plasmid);  
OR        Allow:  
1. Cut out insulin gene / cut open plasmid with restriction enzyme;  
2. Use same restriction enzyme on second DNA;  
3. Reference to (complementary) sticky ends;  
4. Use ligase to join 2 DNA molecules;  
5. Modified plasmid taken up by bacteria;

**max 4**

**[5]**

**M12.**          (a)     (i)      Glucose;

Fructose;

*Any order.*

**2**

(ii)     Lactose has a different shape / structure;

Does not fit / bind to active site of enzyme / sucrase;

*Only allow a second mark if reference is made to the active site.  
Max 1 mark if active site is described as being on the substrate.*

**OR**

Active site of enzyme / sucrase has a specific shape / structure; Does not fit / bind to lactose;

*Do not accept same shape.*

**2**

(b)     (i)      Rose and fell;

Peak at 45 (minutes) / concentration of 6.6 (mmol dm–3);

**2**

(ii)     Glucose (produced by digestion) is absorbed / enters blood;

Decrease as used up / stored;

**2**

**[8]**

**M13.**          (a)     Enzyme / active site has a (specific) tertiary structure;

Only glucose has correct shape / is complementary / will bind / fit to active site;

(Forming) enzyme-substrate complex;

***Q*** *Allow second mark if candidate refers to correct shape or complementary in terms of the enzyme. Do not allow ‘same’ shape*

***Q*** *Do not allow third mark if active site is described as being on substrate.*

**3**

(b)     (Only detects glucose whereas) Benedict’s detects (all) reducing sugars / named examples;

Provides a reading / is quantitative / Benedict’s only provides a colour / doesn’t measure concentration / is qualitative / semiquantitative;

Is more sensitive / detects low concentration;

Red colour / colour of blood masks result;

Can monitor blood glucose concentration continuously;

***Q*** *Do not credit quicker / more accurate unless qualified.*

***Q*** *Allow Benedict’s detects monosaccharides for first mark point.*

**2 max**

(c)     (i)      Broken down by enzymes / digested / denatured (by pH) too large to be absorbed;

**1**

(ii)     Study not carried out on humans / only carried out on rats;  
Long-term / side effects not known;  
Scientists have vested interest;  
Study should be repeated / further studies / sample size not known;

**2 max**

**[8]**

**M14.**          (a)     1.      Adenylate cyclase activated / cAMP produced / second messenger produced;

2.      Activates enzyme(s) (in cell so) glycogenolysis / gluconeogenesis occurs / glycogenesis inhibited;

*2. Neutral: ‘glucose produced’ as given in the question stem*

*Accept: correct descriptions of these terms*

**2**

(b)     (i)     1.      Glucose / sugar in food would affect the results;

*1. Accept references to starch / carbohydrate*

*Or*

2.      Food / eating would affect blood glucose (level);

*Or*

3.      (Allows time for) blood glucose (level) to return to normal;

*3. Neutral: allows time for insulin to act*

**1 max**

(ii)     Type 2 diabetes is a failure to respond to insulin / still produces insulin / is not insulin-dependent;

**1**

(iii)    (For) – 3 max

*A maximum of three marks can be awarded for each side of the argument*

1.      Avoids injections / pain of injections;

2.      Long(er) lasting / permanent / (new) cells will contain / express gene;

*Ignore references to methodology e.g. sample size not known*

3.      Less need to measure blood sugar / avoids the highs and lows in blood sugar;

4.      Less restriction on diet;

(Against) – 3 max

5.      Rats are different to humans;

6.      May have side effects on humans;

*6. Accept: virus may be harmful / disrupt genes / cause cancer*

7.      Long(er) term effects (of treatment) not known / may have caused effects after 8 months;

8.      (Substitute) insulin may be rejected by the body;

**4 max**

**[8]**

**M15.**(a)     (Formation of glycogen)

1.      Glucose concentration in cell / liver falls below that in blood (plasma) which creates / maintains glucose concentration / diffusion gradient;

2.      Glucose enters cell / leaves blood by facilitated diffusion / via carrier(protein) / channel (protein);

*Not just diffusion*

**2**

(b)     1.      Insulin sensitivity similar to / not (significantly) different from those with diabetes;

*No values for non-obese, so comparisons with ‘normal’ not possible*

2.      Overlap of SDs;

*Accept SE*

3.      Their sensitivity (to insulin also) improved by GBS;

**2 max**

(c)     1.      Sensitivity (to insulin) does increase;

*This part of the question concerns spread of data, not overlap of SDs*

2.      But large SD / large variation (after GBS);

*Accept use of figures / use of SD values to make this point.*

*Ignore ref to SE*

3.      (So) some showing no / little change / get worse;

4.      Do not know what sensitivity to insulin is of non-diabetics (who are not obese);

*Accept ‘normal’ as non-diabetic*

**3 max**

**[7]**

**M16.**1.      Diabetics have (blood glucose) concentration greater than 140 mg cm−3 / than her estimate / estimate suggests she is pre-diabetic;

2.      Colour change is subjective / blood on test strip masks colour change;

3.      Concentration given as a range / estimation is not reliable;

4.      May not have fasted;

5.      May not have had a drink with 75 g glucose;

6.      Only one test carried out;

*No mark for valid or not valid*

**[3]**

**M17.**(a)     Treatment requires person receiving insulin (in some way);

*Accept descriptions e.g. insulin injection*

*Reward idea that insulin must be received, not that it isn’t being produced*

**1**

(b)     1.      No / fewer / abnormal receptors on (cell) membrane;

2.      (So) fewer (glucose) transport proteins;

3.      (So) less glucose can enter (cells);

4.      (So) less glucose converted to glycogen;

*Accept no / fewer enzymes (for this conversion) are activated*

5.      (So, without treatment) blood glucose concentration not lowered when high / above normal;

*Accept converse*

**3 max**

(c)     1.      Movement uses muscles;

2.      Movement increases (rate of) respiration;

3.      Respiration uses glucose / respiration reduces blood glucose concentration;

**2 max**

(d)     1.      Identification of 195 ± 2 and 113 ± 2;

2.      Answer within range of 1.67 to 1.77 (times greater);

*Ignore numbers after two decimal places*

*Correct answer = 2 marks*

**2**

(e)     1.      Meal / uncontrolled intake v 75 g glucose / controlled intake;

*Must have both sides of the story for each point. Marking guidance shows researcher’s method first*

*Idea of could eat anything in meal as against just glucose*

2.      (Concentration) measured over 6 hours / 6+ hours / longer v measured at 2 hours;

3.      (After intake) regular monitoring / several measurements v only measured once / at 2 hours only;

4.      No fasting v fasting before test;

*Credit other descriptions of fasting e.g. went without food as opposed to didn’t have to*

5.      Not (necessarily) at rest v remained at rest;

6.      Tested during afternoon v tested in morning;

*Accept idea of tested at different times of the day*

**3 max**

(f)     1.      Pre-diabetics are at risk of developing diabetes / some pre-diabetics reach a concentration of 180 (mg 100 cm−3) after a meal;

2.      Some pre-diabetics will now be classed as diabetic;

3.      Detection leads to treatment (sooner);

4.      Diabetes damages the body / is life-threatening;

*Accept examples of damage e.g. blindness, heart disease*

**3 max**

**[14]**

**M18.**(a)     1.      Glucose oxidase and peroxidase;

*Both enzymes required*

2.      Dye (with colour A);

*Reject ‘dye with colour B’. Ignore named dyes*

**2**

(b)     1.      Concentration is given as a range (for each colour) / measurement is not precise;

2.      Only measures glucose concentration above normal / above 170 (mg 100 cm−3) (in blood);

3.      170 (mg 100 cm−3) is an average figure / concentration for loss to urine varies (between people);

4.      Difficult to match colour against chart / colour match is subjective;

**2 max**

**[4]**

**E1.**(a)     About a third of students correctly explained how glucagon would be involved in regulation of blood glucose concentrations in the mice and obtained 3 marks. Some students failed to mention what glucagon does, or where it acts. There were some students who got very confused between glucagon and glycogen and others who wrote about glucagon either acting on glycogen to break it down, or had glucagon broken down into glucose. This question showed weaknesses in use of language and terminology by many students.

(b)     This was marked by a very large number of students who wrote about large amounts of mRNA but did not link this to large amounts of the enzyme PCK1. These answers were simply describing the results shown in the graph, not giving a suggestion about how the process is controlled. The examiners required students to show understanding that large amounts of mRNA would (probably) mean large amounts of the enzyme. It should be noted that quite a large number of students wrote about mRNA breaking down glycogen (as an enzyme), or being broken down into glucose. Many students did write about the role of the enzyme in kidney and intestinal cells in producing glucose. However, many of these did not compare this with what happens in the normal mice referred to in the question. Some weak answers simply involved attempts to use rote-learned material about control of blood glucose concentration with no reference to the information in the question.

(c)     This again had students writing about the results / data being significant, rather than the difference in the amounts of mRNA, as given in the stem. Quite a few ignored the ‘less than’ symbol, or read it as ‘more than’. Others ignored the 0.01 and wrote entirely about less than 5%, which did not gain credit.

**E2.**The examiners noted that many students appeared not to have read the main stem of the question carefully.

(a)     Quite a large number wrote about amylase in yoghurt, rather than an inhibitor of amylase. About two thirds identified that **Group A** allowed the effect of the inhibitor to be seen. Very few noted that it also allowed the effect of yoghurt on its own to be seen. Quite a large number simply stated that the group ‘allowed for comparison’.

(b)     60% identified that different foods might affect glucose intake. Very few went on to identify that they could also contain different amounts of starch, the substrate of amylase. Quite a few simply wrote about ‘removing a variable’.

(c)     A third of students obtained both marks. These students displayed a clear understanding of the digestion of starch, including the enzymes involved, the products produced and the impact of the inhibitor. About 40% failed to score, with the commonest error being a statement that starch is ‘broken down’ directly into glucose.

(d)     The examiners had identified six different reasons why the results might not support the use of the inhibitor, based on the context of this question. Many students resorted to rote How Science Works answers about correlation not causing causation, or bias. Nearly half of students failed to score.

**E3.**          **BYA6**

          (a)     Many candidates understood the control of plasma glucose concentration in great detail and were able to reproduce this. However, the question did not simply ask ‘how is plasma glucose concentration maintained?’, but required an explanation of the role of negative feedback in this control. Better candidates realised the distinction and began their answers by explaining the concept of negative feedback and then related this concept to the events shown in the graph. Naturally, they included the method of action of the hormones, but their responses showed a better balance between supplying detail and explaining principles than those who simply answered, once again, the question they assumed had been asked on the basis of one or two key words. Candidates failed to gain marks through confusion between glucagon and glycogen, despite the former being clearly named on the graph. There was also some confusion between what detected the changes in plasma glucose concentration and what subsequently secreted the hormones. Only better candidates were aware of the effect of insulin in stimulating the absorption of glucose from the plasma.

(b)     In this question nearly all candidates knew that enzymes have an optimum temperature, but too many failed to relate this specifically to core body temperature. They were also frequently careless in their description of the effect of temperature on enzymes. The phrase ‘enzymes are denatured by extremes of temperature’ was all too common. Clearly an extreme temperature of –10°C does not denature enzymes. Candidates also need to be more precise in their descriptions of how and why reactions take place. Reactions, in this context, take place because *molecules* of the enzyme and substrate have sufficient kinetic energy to collide frequently and with sufficient energy. Finally, some candidates re-used the term ‘maximum efficiency’ used in the question, without explaining what this meant in terms of rate of reaction.

**BYA7**

(a)     Most candidates were able to state that the release of insulin was triggered by increased plasma glucose concentrations and glucagon was triggered by a decrease. The more able then correctly identified the sites of secretion and their effects on the interconversions between glucose and glycogen. It was disappointing to see few candidates using the graph as instructed by the question and many candidates scored one less mark than perhaps they would have if they had used the graph more. As is often the case, frequent instances of imaginative intermediate spelling of glycogen and glucagon were observed and these were usually too ambiguous to be credited by the examiners.

(b)     This question produced some very comprehensive answers. Candidates had a good knowledge of thermoregulation. Examiners did not credit the term ‘cold centre’ and this term tended to be used only by the weaker candidates. The detail of vasoconstriction was explained very clearly by approximately 50% of candidates. The other 50% had muddled thought on this topic, with constriction of capillaries and veins and movement of capillary networks towards the centre of the body appearing in their answers.

Candidates who did not realise that the question was about enzymes struggled to score marks. The majority of candidates gained some credit. One common error was to refer to the effects of temperature on enzyme activity without direction. Better answers gave the effects of low temperatures quite separately from the effects of high temperatures.

**E4.**          Some 40 percent of candidates scored no marks in part (a). All that was required was the idea of insulin binding to receptors on membranes of target cells, the activation or recruitment of glucose-transporting proteins, more glucose entering cells and this resulting in more conversion of glucose to glycogen. Incorrect responses often began with the hypothalamus detecting blood glucose concentrations and telling the pancreas to excrete insulin. It was also common for candidates to state that insulin converts glucose into glycogen.

In (b)(i), very few candidates were able to calculate a ratio correctly. In (b)(ii), two thirds of candidates were unable to suggest a correct reason why undiagnosed diabetics had a higher death rate than diagnosed diabetics. This appeared to be a case where weaker candidates did not see that they were, effectively, being asked about harmful effects of diabetes; the use of factual recall with understanding.

**E5.**          (a)     In part (i), poor expression quite often made it difficult to interpret candidates’ responses, in particular with control **Group 1** which showed the healthy or non-diabetic response. There was often more success in explaining that control **Group 3** showed that **Group 2**’s response was due to the drug exenatide, or was not just due to the salt solution, or that **Group 3** showed any placebo effect.

In part (ii), two-thirds of candidates appreciated that the amount of insulin produced might vary in people of different sizes and, thus, it was necessary to express the results per unit body mass, in order to allow comparisons to be made.

(b)     The vast majority of candidates correctly interpreted the data in terms of the drug exenatide increasing insulin secretion in type 2 diabetics. Many went on to explain how this increase in insulin could help in the control of blood glucose levels. A few indicated that this was due to increased stimulation of cells to take up glucose. Similarly, some indicated that this would mean the diabetic could have a much less restricted diet. Very few gave a full account and hardly any suggested that the drug might possibly increase the sensitivity of pancreas cells to glucose.

**E6.**          (a)     Generally, this question was answered well, although a lack of precision in some answers cost candidates marks, for example, by failing to refer to receptors that are specific to insulin or by not stating the type of cell where the carrier proteins for glucose are located. Common misconceptions were observed when candidates implied that glucose would only enter cells when insulin was present. Even though this is a topic that is generally covered well by candidates at GCSE, a number still believes that insulin catalyses the production of glycogen and others that insulin is responsible for the breakdown of glycogen. The poor spelling of important technical terms also cost some candidates marks; glycogen and glucagon were most often incorrect.

(b)     The role of glucose in respiration was a commonly given as a correct response and many candidates obtained both marks, usually by including a description about the excretion of glucose in urine. However, a surprisingly high proportion of candidates referred to glucose being lost in urea.

(c)     Most candidates answered this question correctly. A common error was to begin the curve from 175 mg rather than from 80 mg as indicated in the question.

(d)     (i)      Generally, this was answered well by candidates across a wide range of ability levels, but the idea that insulin breaks down glucose was a common misconception.

(ii)     References to the role of glucagon or the process of glycogenolysis were common, although many failed to gain the mark by confusing the spellings of these technical terms.

**E7.**          This question also resulted in the full range of marks.

(a)     (i)      Although most candidates correctly referred to shape as being important, many of these explained their answer in terms of active sites and so could not be awarded the mark.

(ii)     There were many full and accurate answers showing a good understanding of the role of insulin. However, many answers showed considerable confusion. Many candidates stated that glucose was unable to bind to the receptors, that the release of insulin was stopped or that insulin converted glucose to glycogen with no reference to glucose uptake into cells or enzyme activation.

(iii)     This was answered well by the majority of candidates. Most realised that the glucose was reabsorbed, but some stated that the molecule was too large to pass through Bowman’s capsule. Similarly, most candidates stated correctly the site and method of reabsorption. Candidates tended to lose marks as a result of incomplete rather than incorrect answers.

(b)     This was answered well by the majority of candidates. The weaker candidates tended to repeat the data in the table rather than explaining it or did not make a comparison. Most candidates gave at least one valid factor, with many also scoring a second mark. Some failed to score because of vague references to the environment being the same, rather than giving a named environmental factor.

**E8.**This topic area seems generally well understood and many candidates were able to apply their knowledge effectively and gained credit. Careless use of terminology and lack of detail marred the answers of weaker candidates.

(a)     The definition was well learnt by the majority of candidates although some missed the idea of keeping conditions constant or gave a named example rather than the definition.

(b)     Generally this question was well answered but lack of precision in some answers cost candidates marks, for example, by failing to refer to receptors and/or identifying their location. Common misconceptions included the involvement of the pituitary and the hormone directly catalysing the conversion of glycogen to glucose thereby acting as an enzyme.

Weaker candidates confused glucagon with glycogen and usually scored only one mark for the idea of raising blood glucose levels. Diffusion of glucose out of cells into the blood was rarely seen.

(c)     A surprisingly high proportion of candidates correctly used the information given in the stem of the question and their own knowledge to work out the clearance value for glucose and go on to obtain full marks. Many of those unsure of the maths did explain the mechanism of reabsorption of glucose in the kidney and gained two or three marks. The most common omission was to the principle of all the glucose being reabsorbed and weaker candidates were also unsure of the precise location in the tubule where this process actually occurs.

**E9.**          (a)     Candidates generally had no difficulty naming A and B.

(b)     Most candidates were able to provide a sufficiently detailed response to be awarded the mark. There were some responses that were too vague to be awarded credit at this level.

**E10.**          (a)     In part (i), the simple statement that the girl may have eaten a meal rich in carbohydrate gained marks for many; others were able to relate high blood glucose concentration to insulin concentration. Given the scatter of points on the graph, it was perhaps unsurprising that relatively few candidates, in answering part (ii), commented on the positive correlation between the results of the two tests. Many, however, referred to the wide range of glucose concentrations corresponding to individual urine measurements. Candidates should be aware that when required to evaluate it is only fair that the information provided genuinely allows candidates to judge the worth of a particular data set or conclusion. As such, they should consider both how the argument is supported and how it fails to be supported.

(b)     Quality of Written Communication proved a challenge in the final part of this question. The more able candidates gained credit with succinct answers referring to glycogenolysis and gluconeogenesis. Others used the terms but were clearly confused by their meanings, or suggested rather predictably that glucagon could be converted into glucose.

**E11.**          (a)     This question was well answered by a majority of the candidature. The labelling of the graph was clear and candidates produced distinct X and Y marks.

(b)     Good responses to this question told the story well using technical language throughout and obtained the four marks easily. Some candidates used the term ‘restriction enzyme’ but did not state what it was used for (i.e removing the gene for insulin or cutting the bacterial plasmid) so failed to obtain either of the first two marking points. Responses given showed that some candidates had not understood what was taking place and gave lengthy responses which were factually incorrect at every stage.

**E12.**          (a)     Most candidates were able to identify glucose as one of the monomers from which a molecule of sucrose was formed, but there was less certainty about the other. Part (ii) was designed to be accessible to grade E candidates and, in view of this, it was disappointing to see so few gaining full credit. There were a number of predictable errors such as in describing the active site as being on the substrate, and in maintaining that active site and substrate were the same shape. Credit was generally lost however because of a lack of precision in the answers. There were many general references to specificity that were simply worded in terms of sucrose and lactose not being ‘specific to each other’ or enzymes being specific to a particular substrate. Good answers amplified the concepts of shape and fit with appropriate reference to complementarity and the active site of the enzyme.

(b)     It is encouraging to note that most candidates were able to describe the data in the graph with appropriate precision and gained full credit for their answers to part (i). There were, however, candidates who failed to distinguish between the terms ‘describe’ and ‘explain’ and offered inappropriate responses both here and in part (ii). In part (ii), better candidates generally identified the role of absorption in raising the glucose concentration and respiration or storage resulting in the fall after 45 minutes. Difficulties arose where candidates referred imprecisely to sugar, and there were many answers where the examiners were left unclear as to whether glucose or sucrose was being discussed. It was also apparent that many candidates considered the graph to be showing some aspect of enzyme activity and responded in terms of the effect of a particular parameter on substrate or product concentration.

**E13.**          (a)     This question proved to be an effective discriminator with only a third of candidates obtaining maximum marks. Most candidates gained at least one mark often for stating that glucose has a complementary shape to the enzyme in the biosensor. Although many candidates did gain a second mark for referring to the active site a significant number did not and often suggested that an active site is present on the substrate.  
There were, however, some excellent explanations of the way in which enzymes work including references to their tertiary structure and the formation of enzyme substrate complexes.

(b)     The majority of candidates gained at least one mark often by stating that the Benedict’s test only provides a colour or does not measure the concentration of blood glucose. Another common correct response was that the Benedict’s test detects not only glucose but all reducing sugars. However, a number of candidates incorrectly stated that the Benedict’s test detects all sugars. A few candidates indicated that the red colour produced during a positive Benedict’s test would be masked by the colour of the blood. Responses which simply referred to the biosensor being more accurate or quicker without any further details were not credited.

(c)     (i)      Most candidates obtained this mark by suggesting that insulin would be digested or broken down by enzymes. A few candidates did suggest that it may be too large to be absorbed or that it could be denatured by acid in the stomach.  
Candidates who failed to gain the mark often suggested that it would be broken down by amylase.

(ii)     This was very well answered. Almost every candidate obtained at least one mark often by indicating that the study had not been carried out on humans.  
Many candidates then linked this to unknown long term or side effects to gain a second marking point. It was also pleasing to note the number of responses which displayed an understanding of ‘How Science works’ including references to the vested interests of scientists, unknown sample size and the need for further or repeat studies.

**E14.**          (a)     This proved to be an excellent discriminator and just less than half of students gained full credit. This was usually for describing the production of cAMP and its effect on glycogenolysis or gluconeogenesis. Very few students were able to name the enzyme within the plasma membrane as adenylate cyclase. Similarly, there was sometimes confusion between the activation of this enzyme and intracellular enzymes. Some students thought that adenylate cyclase directly affects glycogenolysis or gluconeogenesis. This said, the correct use of scientific terms beginning with ‘g’ was generally good and only a minority of weaker students confused ‘glycogen’ and ‘glucagon’. The terms ‘glyconeogeneis’ and ‘glucogenolysis’ also appeared in weaker answers.

(b)     (i)      Most students were aware that glucose in food would affect the results or that eating would affect the blood glucose concentration. Very few students referred to the importance of allowing the blood glucose concentration to return to normal.

(ii)     Students who failed to score typically stated that type 2 diabetes is not a genetic disease.

(iii)    The most common advantages given for gene therapy were the avoidance of injections and the longer-lasting effect. The most common disadvantages given were the possible side effects, which were usually linked to the virus, and that the long- term effects are not known. Surprisingly, relatively few students made reference to the fact that rats are different from humans. Weaker students seemed to rely on ‘stock’ *How Science Works* answers to gain credit for the ‘against’ side of the evaluation. They typically referred to unknown sample sizes and the influence of unnamed ‘other factors’. These were not credited.

**E15.**(a)    80% of students failed to score. Most repeated the stem of the question and wrote that formation of glycogen leads to a lowering of blood glucose. Others drifted into explanations of the mode of action of insulin and the role of the pancreas. The examiners were looking for the idea that formation of glycogen lowers the concentration of glucose in liver cells below that in the blood. As a result, there is a diffusion / concentration gradient for the entry of glucose into liver cells (from the blood).

(b)     Students scored much better on this part. Most noted that the insulin sensitivity was similar in both groups of patients. Some backed this up with observations about overlap of standard deviations. Again, the examiners accepted standard errors here but it should not be assumed that this applies in all cases. About 40% of students got both marks and a similar percentage got one mark.

(c)     It was heartening in this part to see many students using knowledge of standard deviations to answer this question correctly. Nearly 40% obtained all three marks and these students all included statements about the large standard deviations of the means after surgery and how these showed that some patients’ sensitivity to insulin did not increase after surgery. In this context, the examiners ignored references to standard errors, since it was the spread of data about the mean(s) that was important.

**E16.**Many students produced sound responses to the final question using information from the resources appropriately.

**E17.**(a)     Almost all students appreciated what was meant by ‘insulin-dependent’.

(b)     Although the wording used by many students did not entirely match that of the Marking Guidelines, there was often a good understanding of the scientific principles demonstrated to support the assessments made.

(c)     It was appropriate to consider exercise as equivalent to movement. The majority of students appreciated that such was based on respiration and that respiration uses glucose.

(d)     In written papers, students often struggle with calculations. However, in this case, the majority produced an answer that fell within the stipulated range.

(e)     It was pleasing to see that most students gave both sides of the story to give differences between the two methods. It was not appropriate to give credit where this was not the case. Many students were able to make full use of the resource material and obtain full marks for this question.

(f)     This part proved to be more challenging for many students although some excellent accounts were seen. A synthesis of ideas from the resource material was the key in this question as opposed to lifting relevant aspects for comparison with the previous part, a skill that was more demanding and reflected by the mark range seen.

**E18.**(a)     Most students were able to identify the correct substances but it was an error of assessment not to cancel a mark when an additional substance, such as hydrogen peroxide, was also included in the list.

(b)     This question proved to be demanding for many students despite the possible reasons exceeding the maximum mark possible.