

basic education

Department:
Basic Education
REPUBLIC OF SOUTH AFRICA

NATIONAL SENIOR CERTIFICATE

GRADE 12

LIFE SCIENCES P2

NOVEMBER 2023

.co.43

MARKS: 150

TIME: 21/2 hours

This question paper consists of 15 pages.

INSTRUCTIONS AND INFORMATION

Read the following instructions carefully before answering the questions.

- 1. Answer ALL the questions.
- 2. Write ALL the answers in the ANSWER BOOK.
- 3. Start the answers to EACH question at the top of a NEW page.
- 4. Number the answers correctly according to the numbering system used in this question paper.
- 5. Present your answers according to the instructions of each question.
- 6. Do ALL drawings in pencil and label them in blue or black ink.
- 7. Draw diagrams, tables or flow charts only when asked to do so.
- The diagrams in this question paper are NOT necessarily drawn to scale. 8.
- 9. Do NOT use graph paper.
- You must use a non-programmable calculator, protractor and a compass, 10. 1505.CO.FD where necessary.
- 11. Write neatly and legibly.

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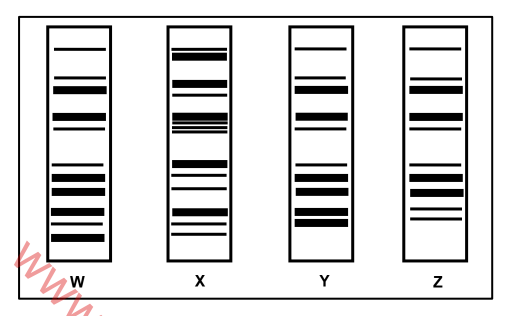
SECTION A

QUESTION 1

- 1.1 Various options are provided as possible answers to the following questions. Choose the answer and write only the letter (A-D) next to the question numbers (1.1.1 to 1.1.9) in the ANSWER BOOK, e.g. 1.1.11 D.
 - 1.1.1 The base pairing in DNA was discovered by ...
 - Α Watson and Wilkins.
 - В Franklin and Wilkins.
 - C Franklin and Crick.
 - D Crick and Watson.
 - 1.1.2 Agene codes for the production of ...
 - Α a chromosome.
 - an allele. В
 - C DNA.
 - a protein.
 - Which ONE of the following is a characteristic of stem cells? 1.1.3
 - They are easily obtained from any organ. Α
 - В They divide by meiosis.
 - С They are haploid.
 - D They can be stimulated to form any type of cell needed.
 - 1.1.4 The chances of having a female child in humans is ...
 - Α 25%
 - В 50%
 - C 75%
 - D 100%
 - 1.1.5 Which ONE of the following is part of the reason why colourblindness is more common in males than in females?
 - The allele for colour-blindness is recessive and located on Α the X-chromosome.
 - В Colour-blind males have two copies of the allele for colourblindness.
 - C The allele for colour-blindness is recessive and located on the Y-chromosome.
 - D Fathers pass the allele of colour-blindness to their sons only.

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The DNA profile of four individuals are given below. 1.1.6

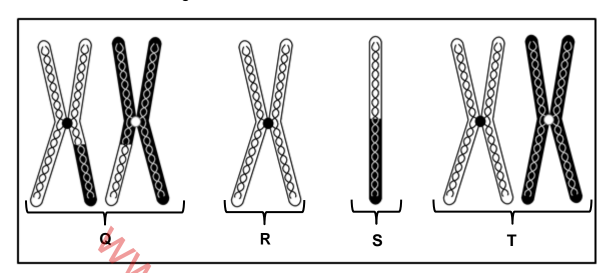


Which individuals are possible members of the same family?

- X and Z only Α
- X, Y and Z only В
- W, Y and Z only С
- W, X and Y only
- When two plants heterozygous for a characteristic are crossed, 1.1.7 the expected ratio is:

Apecieu raii	0 13.	CO
Dominant phenotype		Recessive phenotype
3	:	1
1	:	3
1	:	2
1	:	1
	Dominan	

Copyright reserved Please turn over 1.1.8 The diagram below represents the structure of chromosomes at different stages of meiotic cell division.



Which ONE of the following chromosomes would be found in a cell during late Anaphase II?

Α

Q

- В R
- S С
- Т
- The scientist who discovered Little Foot is ... 1.1.9 -3°.CO.49
 - Α Lee Berger.
 - В Raymond Dart.
 - С Ron Clarke.
 - D Robert Broom.

 (9×2) (18)

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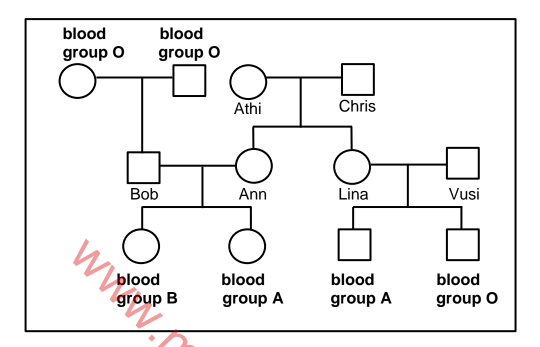
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- 1.2 Give the correct biological term for each of the following descriptions. Write only the term next to the question numbers (1.2.1 to 1.2.10) in the ANSWER BOOK.
 - 1.2.1 The position of a gene on a chromosome
 - 1.2.2 The type of evolution characterised by long periods of little or no change followed by short periods of rapid change
 - 1.2.3 The natural shape of a DNA molecule
 - 1.2.4 The type of bond found between two amino acids
 - 1.2.5 The type of vision shared in primates that allows for depth perception
 - 1.2.6 The type of dominance which results in an intermediate phenotype in the heterozygous condition
 - 1.2.7 The fluid of the nucleus where free nucleotides are found
 - 1.2.8 A tangled mass of chromosomes located within the nucleus
 - The division of the cytoplasm after a nuclear division 1.2.9
 - 1.2.10 The name for the X and Y sex chromosomes in humans (10×1) (10)
- 1.3 Indicate whether each of the descriptions in COLUMN I apply to A ONLY, B ONLY, BOTH A AND B or NONE of the items in COLUMN II. Write A only, B only, both A and B or none next to the question numbers (1.3.1 to 1.3.3) in the ANSWER BOOK.

	COLUMN I	COLUMN II
1.3.1	A genetic disorder caused by a chromosomal mutation	A: Haemophilia B: Colour-blindness
1.3.2	The importance of meiosis	A: Formation of gametes B: Halving of the chromosome number
1.3.3	The organelle where DNA is found in plants	A: Mitochondria B: Chloroplast

(6) (3×2)

Copyright reserved Please turn over 1.4 The diagram below shows the inheritance of blood groups in a family.



- Name the type of diagram shown. 1.4.1 (1)
- 1.4.2 Give the number of alleles that control blood groups. (1)
- 1.4.3 How many generations are represented in the diagram? (1)
- Lina's genotype is I^Ai. 1.4.4

State ALL the possible genotypes of Vusi. (2)

1.4.5 Give the genotype of Bob.

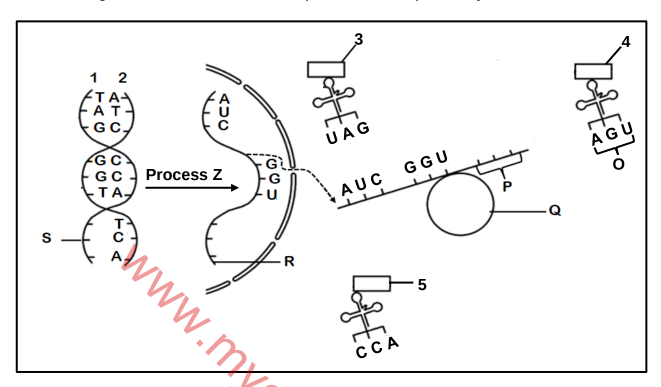
1.4.6 Give the name of the individual which displays co-dominance. (2)

(8)

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(1)

1.5 The diagram below is a schematic representation of protein synthesis.



1.5.1 Identify:

> (a) Process **Z** (1)

> Molecule R (1) (b)

> Organelle Q (c) (1)

1.5.2 Give the collective name of nitrogenous bases O. (1)

1.5.3 Determine the sequence of the nitrogenous bases at area S. (1)

1.5.4 Which strand (1 or 2) was used as a template for the formation of molecule R? (1)

(1) 1.5.5 Which amino acid (3, 4 or 5) will be brought to area P?

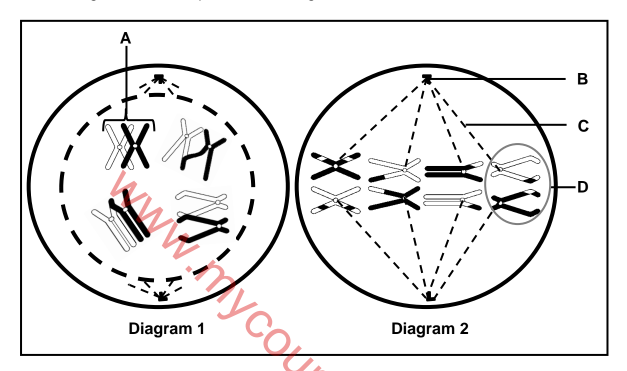
1.5.6 Name the type of sugar that forms part of the structure of molecule R. (1) (8)

> **TOTAL SECTION A:** 50

SECTION B

QUESTION 2

2.1 The diagrams below represent two stages of meiotic cell division.



2.1.1 Name structure:

(a) B	C		(*	1)
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- 2.1.2 Identify the phase represented in Diagram 1. (1)
- 2.1.3 Give THREE reasons for your answer to QUESTION 2.1.2. (3)
- 2.1.4 Describe the process taking place at **A**. (3)
- 2.1.5 (a) Identify the phase represented in Diagram **2**. (1)
 - (b) Describe the difference in the events that take place in the phase mentioned in (a) and the same phase during mitosis. (2)
- 2.1.6 Describe the results at the end of meiosis if the chromosomes at **D** failed to separate. (3) (15)
- 2.2 Describe the process of DNA replication. (6)

2.3 Read the information below.

A gene, VKORC1, codes for a blood-clotting factor in humans. This gene is made up of 163 amino acids.

A mutation occurred that affected amino acid 128 and 139, the sequence CTG changed to CAG and the TAT became TCT. This mutation has been transmitted as an autosomal dominant characteristic through the generations.

The mutation has resulted in resistance to Warfarin drugs in humans. Warfarin is used in the treatment of thrombosis. Thrombosis results in the formation of a blood clot in the artery. Warfarin causes the thinning of blood to break down the blood clot.

- 2.3.1 Give ONE piece of evidence from the information that shows that the mutation for this gene occurred in the DNA molecule. (1)
- 2.3.2 How many nitrogenous bases code for the VKORC1 gene? (2)
- 2.3.3 Describe what is meant by an autosomal dominant allele. (3)
- The table below shows the amino acids and their corresponding 2.3.4 codons.

CODONS	AMINO ACID
GAC	Leu
UCU	Ser
AUA	Try
GUC	Gln
AGA	Arg
ACA	T rp
CAG	Gln
UAU	Phe

Explain:

How the mutation on the VKORC1 gene resulted in resistance to Warfarin in humans

The effect of this mutation on humans with thrombosis (b) (3)(14)

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2.4 Polydactyly is a condition that leads to extra fingers or toes. It is caused by a dominant allele.

A man who is heterozygous for polydactyly has a wife who is not polydactyl.

Using the letters \mathbf{R} and \mathbf{r} , do a genetic cross to show the percentage chance that their children will have polydactyly.

(6)

In summer squash plants, white fruit colour **(B)** is dominant over yellow fruit colour **(b)**, and round fruit **(D)** is dominant over oval fruit **(d)**.

A summer squash plant that is homozygous for white and round fruit is crossed with a plant that is homozygous for yellow and oval fruit.

2.5.1 State the:

- (a) Genotypes of the P₁-parents (2)
- (b) Phenotypes of the F_1 -generation (2)
- 2.5.2 Two plants that are heterozygous for both characteristics were crossed.
 - (a) Give ALL the possible genotypes in the **gametes** that will be formed.

(2)

(b) How many plants in the next generation are likely to have yellow and oval fruit?

(1)

2.5.3 Give the possible genotypes of both parents that must be crossed if a farmer wants summer squash that are white with oval fruit only.

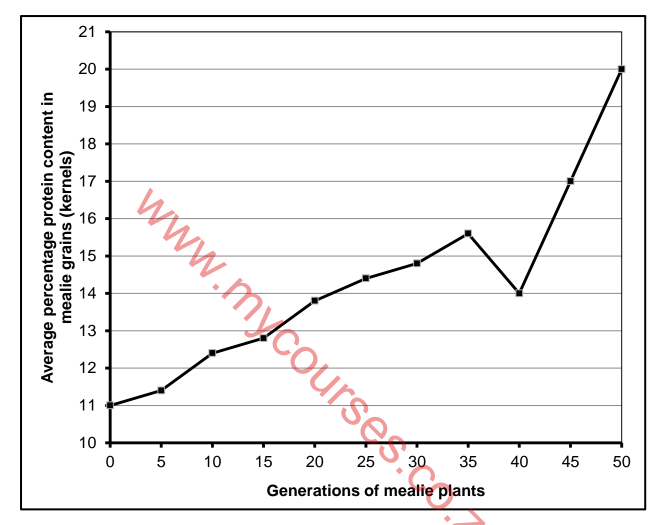
(2) **(9)** [**50**]

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QUESTION 3

3.1 The graph below shows the results of artificial selection for protein content in mealie plants over 50 generations.



3.1.1 Describe how this farmer did artificial selection of the mealie plant. (3)

3.1.2 What was the average percentage of the protein content in the mealie grains (kernels) at the 15th generation? (1)

3.1.3 By how many times did the average percentage of the protein content in the mealie grains (kernels) increase between the 40th and 50th generation? Show ALL working. (2)

3.1.4 Describe ONE way in which the process of artificial selection is different from genetic engineering.

3.2 Describe Darwin's theory of evolution by natural selection. (7)

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(2) **(8)** 3.3 An investigation was done to determine the relationship between the height of the head and bite force in lizards.

The procedure was as follows:

- The scientists collected 120 lizards with similar characteristics that were around the same reproductive age in different habitats.
- Their body characteristics and DNA were analysed to determine if they belonged to the same species.
- 40 lizards belonged to species **A**, 36 to species **B** and 44 to species **C**.
- Each species was kept in its cage with environmental conditions similar to their habitats.
- The height of the head was measured for each lizard and averages calculated for each species.
- Using a Kistler force, the bite force of each lizard in each species was measured five times and the average calculated for each lizard and each species.

The results are shown in the table below.

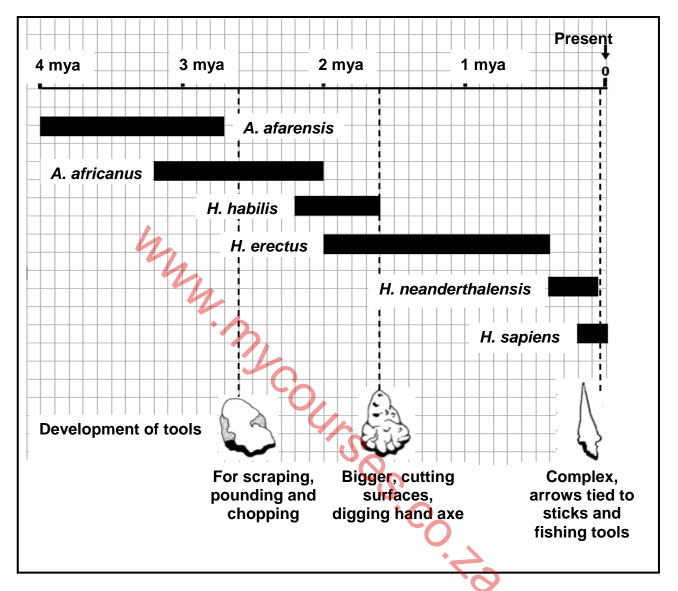
Species	Height of the head (mm)	Bite force (N)
Α	10,3	12,4
В	10,7	14,3
С	13,2	20,4

3.3.1 Identify the:

3.3.1	identity the.	
	(a) Independent variable	(1)
	(b) Dependent variable	(1)
3.3.2	State TWO factors that were kept constant for this investigation.	(2)
3.3.3	Apart from the sample size, state ONE way in which the reliability of the results was ensured for this investigation.	(1)
3.3.4	The height of the head was different in each species of lizard.	
	Name the type of variation displayed by this characteristic.	(1)
3.3.5	Describe the relationship between the height of the head of the lizards and the bite force.	(2)
3.3.6	Which species (A , B or C) would be expected to be feeding mainly on tough fibrous plants?	(1)
3.3.7	Explain your answer to QUESTION 3.3.6.	(2)
3.3.8	Which species (A, B or C) would be most suited to live in narrow areas between the rocks?	(1)

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3.4 The diagram below shows a timeline of different hominid species and the development of tools.



3.4.1 Which species in the diagram above existed/survived for the longest period of time? (1)

3.4.2 Calculate the period (million years) in which the *A. afarensis* and *A. africanus* coexisted. Show ALL working. (2)

3.4.3 Name the species that was also known as the handyman. (1)

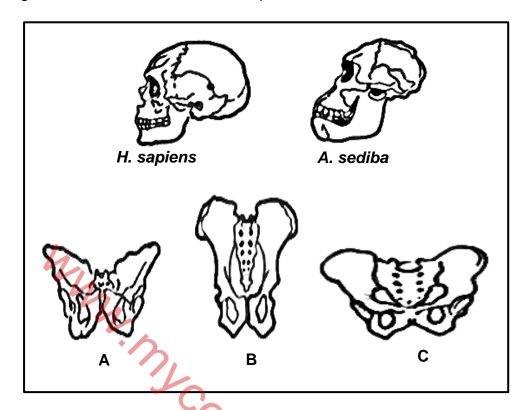
3.4.4 State TWO uses of the tool that was developed 2,6 mya. (2)

3.4.5 Identify TWO species that used the most complex tools. (2)

3.4.6 Explain how the changes in brain size over time relates to the development of tools. (3)

(11)

3.5 The diagrams below show the skulls and pelvises of different hominids.



3.5.1	State the genus name of A. sediba.	(1)
3.5.2	Describe the shape of the spine of <i>H. sapiens</i> .	(1)
3.5.3	A. sediba is thought to be a transitional species.	
	State what is meant by a transitional species.	(2)
3.5.4	Give the LETTER of the pelvis that would be representative of <i>A. sediba</i> .	(1)
3.5.5	Explain your answer to QUESTION 3.5.4.	(2)
3.5.6	Explain the significance of the change in prognathism from <i>A. sediba</i> to <i>H. sapiens</i> .	(5) (12) [50]

TOTAL SECTION B: 100 GRAND TOTAL: 150