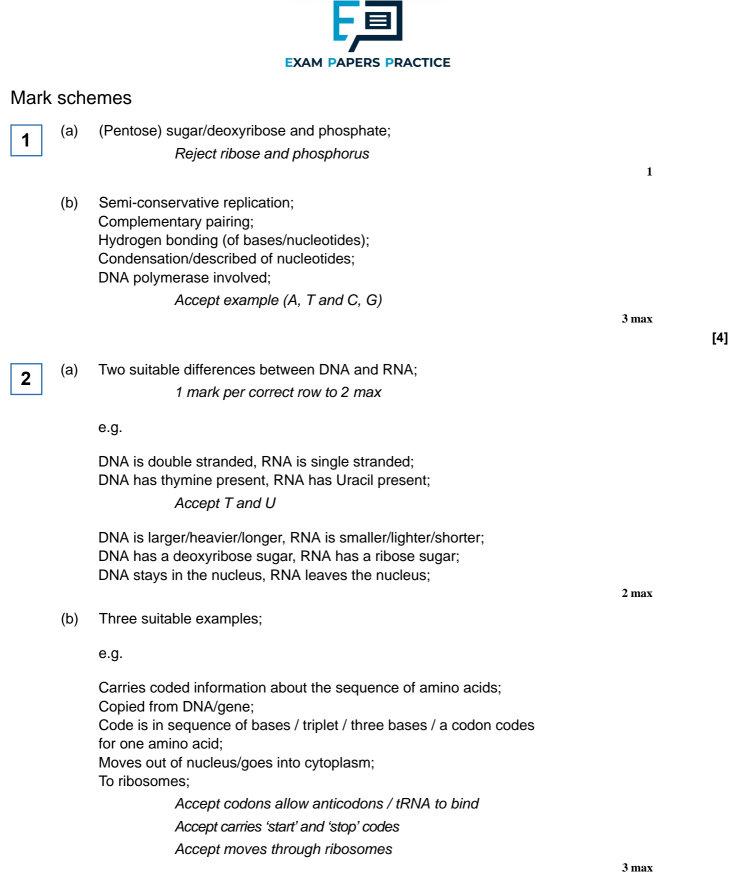


Nucleic acids 2

Level: AQA A Level 7402 Subject: Biology Exam Board: Suitable for all boards Topic: Nucleic acids 2 Type: Mark Scheme

To be used by all students preparing for AQA A Level Biology 7402 foundation or higher tier but also suitable for students of other boards.





(a) Any two of:

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

(b) Base sequence (on DNA/in gene); Determines sequence of amino acids; By determining base sequence on (messenger) RNA; Code is a triplet code/three base code for an amino acid; 2 max (C) Pairs of chromosomes/two chromosomes; With genes for same features / with same genes; At same loci / in same sequence; Accept same alleles 2 max [6] Decreases by 50%; (a) Per generation / per division; Only accessible if linked to first marking point OR ¹⁵N makes up ½ after 1 division; Makes up 1/4 after 2nd division; 2 (b) In DNA replication strands separate; Each acts as template (for formation of new strand); One strand in each new molecule / semi-conservative replication; New strands made using ¹⁴N. 2 max

[4]

2 max



- (a) **D** phosphate;
 - **E** pentose sugar/deoxyribose;
 - F (nitrogenous) base/ organic base/ thymine/adenine/ cytosine/guanine;

In **D** reject phosphorous In **E**

Accept 5-carbon sugar

Reject sugar alone

(b)

| DNA | RNA |
|-----------------|------------------|
| double-stranded | single-stranded |
| deoxyribose | ribose |
| Thymine/T | Uracil/U |
| very large/long | very small/short |

Accept double helix for DNA Accept longer and shorter Need comparison but could be in one box List rule applies.

2 max

3

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6

(a)

| * | * | * | |
|---|---|---|---|
| | | | * |
| | | × | × |

One mark for each correct column Mark ticks only and ignore crosses

- (b) 1. Two marks for box round two hydrogens and one of the oxygens from OH groups on carbons 1 and 4;;
 - 2. One mark from incorrect answer involving any two hydrogens and an oxygen from carbons 1 and 4;

Do not award marks if all atoms concerned are on same carbon atom or are on carbon atoms other than 1 and 4 or where the answer does not have two hydrogen and one oxygen

- (i) Holds chains / cellulose molecules together / forms cross links between (c) 1. chains / cellulose molecules / forms microfibrils, providing strength / rigidity (to cellulose / cell wall);
 - 2. Hydrogen bonds strong in large numbers;x Principles here are first mark for where hydrogen bonds are formed and second for a consequence of this. Accept microfibres

(ii) Compact / occupies small space / tightly packed; Answer indicates depth required. Answers such as "good for storage", "easily stored" or "small" are insufficient.

pentose / 5C sugar = neutral

phosphorus / P = neutral

Phosphate / Phosphoric acid;

1

1

1

1

4

2

2

1

[9]

381 / 384 / 387;

Hydrogen (bonds);

(a)

(b)

(c)

7

(i)

(ii)

(d) (GIn) Met Met Arg Arg Arg Asn;

Deoxyribose;



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

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

Substrate cannot bind / no enzyme-substrate complexes form;

Q Reject = different amino acids are formed

9

(a)

| DNA | V | 2 |
|------|---|---|
| mRNA | × | 1 |
| tRNA | × | 1 |

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

(b) (i) Marking principles

1 mark for complete piece transcribed;

Correct answer UGU CAU GAA UGC UAG

1 mark for complementary bases from sequence transcribed;

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

(ii) Marking principle

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

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

- sugar or phosphate / S-P / nucleotide chain / backbone / original / parent DNA;
 - (ii) X thymine; Y guanine; Z adenine; (Allow T, G and A) Reject: thiamine

[8]

3

1

1



| 10 | (a) | each strand copied / acts as a template; (daughter) DNA one new strand and one original / parent strand; | 2 | |
|----|-----|---|-------|-----|
| | (b) | (i) ¹⁵N / tube B (DNA), more / greater density; (reject heavier) | | |
| | | (ii) DNA with one heavy and one light strand; new / synthesised strand, made with ¹⁴N / light strand; | 1 | |
| | (c) | 32; 28 32 26; | 2 | [7] |
| 11 | | (a) (i) ATA; | 1 | [7] |
| | | (ii) AUA; | 1 | |
| | (b) | tRNA 'clover leaf' shape; (allow reference to loop / folded structure) tRNA standard length; tRNA has an amino acid binding site; tRNA has anticodon available / three exposed bases; tRNA has hydrogen bonds (between base pairs); | 2 max | [4] |
| 12 | (a) | different form of a gene; | 1 | |
| | (b) | hydrogen bonds broken; semi-conservative replication / both strands used (as templates); <u>nucleotides</u> line up complementary / specific base pairing / A and T / C and G; <u>DNA</u> polymerase; | 4 | |
| | (c) | deletion causes frame shift / alters base sequence (from point of mutation); changes many amino acids / sequence of amino acids (from this point); substitution alters one codon / triplet / one amino acid altered / code degenerate / same amino acid coded for; | 4 | |
| | (a) | X, phosphate; | 3 | [8] |
| 13 | (4) | X, prospirate, Y, deoxyribose / pentose / 5-carbon sugar; Z, (nitrogenous) base; (accept named base) | | |
| | (1) | | 3 | |
| | (b) | (specific) hydrogen (bonds); | 1 | |



| (c) | ther | nine 28% so adenine 28% efore 44% cytosine and guanine; efore 22% cytosine; | | |
|-----|-------|---|---|-----|
| | | (idea of equal amounts T and A, C and G – 1 mark, correct answer 2 marks) | 2 | [6] |
| (a) | appi | ropriately placed box; | 1 | [0] |
| (b) | (i) | B; | | |
| | (ii) | А; | 2 | |
| (c) | (i) | determines (sequence of) amino acids / specific protein produced / mRNA formation; | | |
| | (ii) | hydrogen bonds; | 1 | |
| | (:::) | | 1 | |
| | (iii) | stability / protects bases / replication; | 1 | [6] |



15

(a)

(i)

- Joins nucleotides (to form new strand). Accept: joins sugar and phosphate / forms sugar-phosphate backbone Reject: (DNA polymerase) forms base pairs / hydrogen bonds
- (ii) (Prokaryotic DNA)
 - Circular / non-linear (DNA);
 Accept converse for eukaryotic DNA Ignore: references to nucleus, binary fission, strands and plasmids
 - 2. Not (associated) with proteins / histones; Accept does not form chromosomes / chromatin
 - No introns / no non-coding DNA.
 Accept only exons
 Q Neutral: no 'junk' DNA
- (b) (i) 1. Have different genes; Reject: different alleles
 - 2. (Sobases / triplets) are in a different sequence / order; Accept: base sequence that matters, not percentage
 - (So) different amino acid (sequence / coded for) / different protein / different polypeptide / different enzyme.
 Unqualified 'different amino acids' does not gain a mark
 Reject: references to different amino acids formed
 Ignore: references to mutations / exons / non-coding / introns
 - (ii) (Virus DNA)
 1. A does not equal T / G does not equal C;
 Accept: similar for equal
 Accept: virus has more C than G / has more A than T
 - 2. (So) <u>no</u> base pairing;
 - 3. (So) DNA is not double stranded / is single stranded.

2 max

2 max

1

2 max



(a) 1. DNA replicated;

- Reject: DNA replication in the wrong stage
- 2. (Involving) specific / accurate / complementary base-pairing; Accept: semi conservative replication
- 3. (Ref to) two identical / sister chromatids;
- 4. Each chromatid / moves / is separated to (opposite) poles / ends of cell. Reject: meiosis / homologous chromosomes / crossing over Note: sister <u>chromatids</u> move to opposite poles / ends = 2 marks for mp 3 and mp 4 Reject: events in wrong phase / stage
- (b) (i) 1. To allow (more) light through; Accept: transparent
 - 2. A single / few layer(s) of <u>cells</u> to be viewed. Accept: (thin) for better / easier stain penetration
 - (ii) 1. More / faster mitosis / division near tip / at 0.2 mm; Neutral: references to largest mitotic index
 - (Almost) no mitosis / division at / after 1.6 mm from tip;
 Accept: cell division for mitosis
 Penalise once for references to meiosis
 - (So) roots grow by mitosis / adding new cells to the tip.
 Accept: growth occurs at / near / just behind the tip (of the root)
 Accept: converse arguments
- (a) (i) Repeating units / nucleotides / monomer / molecules; Allow more than one, but reject two

1

3

2 max

[8]

4

2

- (ii) 1. C = hydrogen bonds;
 - 2. D = <u>deoxy</u>ribose; Ignore sugar
 - 3. E = phosphate; Ignore phosphorus, Ignore molecule

16



| | | | | ` |
|-----|---|---|---|----|
| - 1 | I | L | I | ۱. |
| ١. | I | I | I | , |
| | | | | |

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

Spelling must be correct to gain MP1 First mark = names correct Second mark = % correct, with <u>adenine as 34%</u>

- (b) (i) 153;
 - Some regions of the gene are non-coding / <u>introns</u> / start / stop code / triplet / there are two DNA strands;

2

1

1

1

2

2

[8]

Allow <u>addition</u> mutation Ignore unqualified reference to mutation Accept reference to introns and exons if given together Ignore 'junk' DNA / multiple repeats

(a) (i) Anaphase

- (ii) 1. Sister / identical chromatids / identical chromosomes; *Reject: Homologous chromosomes separate. Allow any reference to chromatids / chromosomes being identical e.g. same DNA*
 - 2. To (opposite) poles / ends / sides;
- (b) (i) 1. 8.4 / cells with twice DNA content = replicated DNA / late interphase / prophase / metaphase / anaphase;
 Any reference to interphase must suggest towards end of interphase.
 'Chromosomes replicate' is not enough for DNA replicates.
 - 4.2 = DNA not replicated / (early) interphase / telophase / cell just divided / finished mitosis;



(ii) 2.1;

19

| | | | 1 | | |
|-----|---|--|---|--|--|
| (a) | (i) | Spindle formed / chromosome / centromere / chromatids attaches to spindle; | | | |
| | | Chromosomes / chromatids line up / move to middle / equator (of cell); | | | |
| | | Do not award second mark for answers referring to chromosomes 'pairing up'. | | | |
| | | Ignore reference to homologous chromosomes unless context suggests pairing which negates second mark. | | | |
| | | Neutral: Details on nuclear membrane. | | | |
| | | Accept: Diagram for second marking point. | 2 | | |
| | (ii) | Chromosome / centromere splits / chromatids / 'chromosomes' separate / pulled apart; | | | |
| | | To (opposite) sides / poles / centrioles (of cell); | | | |
| | | Reject: Homologous chromosomes separate for first marking point. | | | |
| | | Accept: Diagram for second marking point. | | | |
| | | Chromatids / 'chromosomes' move to poles / sides / centrioles = 2 marks. | | | |
| | | | 2 | | |
| (b) | (i) | Form / replace cells quickly / rapidly / divide / multiply / replicate rapidly; Neutral: Repair cells. | | | |
| | | Answers must convey idea of 'speed'. | 1 | | |
| | (li) | Correct answer = 774 minutes / 12 hours 54mins = 2 marks;; | | | |
| | | Incorrect answer but indicates 3 cell cycles involved = one mark; | 2 | | |
| (c) | Prevents / slows DNA replication / doubling / prevents / slows mitosis; | | | | |
| | | strand not formed / nucleotides (of new strand) not joined ther / sugar-phosphate bonds not formed; | | | |
| | | First marking point must be in context of DNA replication not cell replication. | | | |
| | | Do not negate first marking point if role of DNA polymerase is described incorrectly e.g. Reject: 'joins bases / strands together'. | | | |
| | | Role of DNA polymerase must be correct for last marking point. | 2 | | |

[6]