D1.3 MUTATIONS AND GENE EDITING

Guiding Questions

How do gene mutations occur?

What are the consequences of gene mutation?

Linking Questions

How can natural selection lead to both a reduction in variation and an increase in biological diversity?

How does variation in subunit composition of polymers contribute to function?

D D

Theme: Continuity + Change

Level of Organization: Molecules

Written and drawn by:

PETER MARIER



SL LEARNING OUTCOMES

D1.3.1	Gene mutations as structural changes to genes at the molecular level	Distinguish between substitutions, insertions and deletions.
D1.3.2	Consequences of base substitutions	Students should understand that single-nucleotide polymorphisms (SNPs) are the result of base substitution mutations and that because of the degeneracy of the genetic code they may or may not change a single amino acid in a polypeptide.
D1.3.3	Consequences of insertions and deletions	Include the likelihood of polypeptides ceasing to function, either through frameshift changes or through major insertions or deletions. Specific examples are not required.
D1.3.4	Causes of gene mutation	Students should understand that gene mutation can be caused by mutagens and by errors in DNA replication or repair. Include examples of chemical mutagens and mutagenic forms of radiation.
D1.3.5	Randomness in mutation	Students should understand that mutations can occur anywhere in the base sequences of a genome, although some bases have a higher probability of mutating than others. They should also understand that no natural mechanism is known for making a deliberate change to a particular base with the purpose of changing a trait.
D1.3.6	Consequences of mutation in germ cells and somatic cells	Include inheritance of mutated genes in germ cells and cancer in somatic cells.
D1.3.7	Mutation as a source of genetic variation	Students should appreciate that gene mutation is the original source of all genetic variation. Although most mutations are either harmful or neutral for an individual organism, in a species they are in the long term essential for evolution by natural selection. NOS: Commercial genetic tests can yield information about potential future health and disease risk. One possible impact is that, without expert interpretation, this information could be problematic.

HL LEARNING OUTCOMES

D1.3.8	Gene knockout as a technique for investigating the function of a gene by changing it to make it inoperative	Students are not required to know details of techniques. Students should appreciate that a library of knockout organisms is available for some species used as models in research.
D1.3.9	Use of the CRISPR sequences and the enzyme Cas9 in gene editing	Students are not required to know the role of the CRISPR—Cas system in prokaryotes. However, students should be familiar with an example of the successful use of this technology. NOS: Certain potential uses of CRISPR raise ethical issues that must be addressed before implementation. Students should understand that scientists across the world are subject to different regulatory systems. For this reason, there is an international effort to harmonize regulation of the application of genome editing technologies such as CRISPR.
D1.3.10	Hypotheses to account for conserved or highly conserved sequences in genes	Conserved sequences are identical or similar across a species or a group of species; highly conserved sequences are identical or similar over long periods of evolution. One hypothesis for the mechanism is the functional requirements for the gene products and another hypothesis is slower rates of mutation.

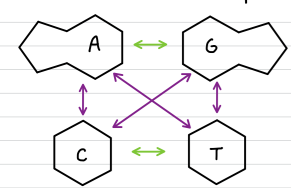
D1.3.1—Gene mutations as structural changes to genes at the molecular level.

D1.3.2—Consequences of base substitutions. D1.3.3—Consequences of insertions and deletions

gene mutation: permanent structural changes to genes at the molecular level (i.e. change in DNA nucleotide base sequence)

Base Substitution mutation: replacement of one base in the coding section of a gene with another

can result in



Transversions point mutation that changes a pyrimidine nucleotide (single ring) to a purine (double ring) or vice - versa

Transitions: point mutation that changes a purine nucleotide (single ring) to another purine or a pyrimidine (double ring) to another pyrimidine.

More common than transversions (as same ring structure kept) and more likely to result in silent mutations and persist as single-nucleotide polymorphisms

single-nucleotide polymorphisms (SNPs): a variation at a single base in a DNA sequence present in more than 1% of the population. SNPs can occur in both coding and non-coding sequences. If it occurs in a gene it may or may not result in a new amino acid and polypeptide due to degeneracy of genetic code

Individual 2

ACCAT ACCAC

INDINITION

ACCAT ACCAC

INDINITION

INDI

Base Substitution mutations can be classified based on the resulting consequence:

- Same-sense (silent) mutation: base substitution which alters the codon for an amino acid into another codon for the same amino acid due to degeneracy of genetic code.

 No effect on the polypeptide produced and the phenotype
- missense mutation: base substitution which alters the codon for an amino acid into another codon for a <u>different</u> amino acid. Changes the polypeptide produced. If the new amino acid causes the protein to fold differently it may alter protein function and the resulting phenotype (ex: Sickle cell anemia)
- nonsense mutation: base substitution which alters the codon for an amino acid into a <u>STOP</u> codon, causing translation to end prematurely. Polypeptide produced is shortened which may alter its function or more likely make it non-functional, altering the phenotype (ex: Cystic fibrosis)
- Insertion mutation: addition of one or more nucleotides within the base sequence. Effect depends on the position and how many nucleotides are added (ex: HTT gene and Huntington's disease)
 - number inserted is not multiple of 3

 Results in a frameshift mutation where the reading frame (grouping of codons) is shifted, altering many/all amino acids downstream of mutation. The earlier in the sequence it occurs the greater the impact. Protein produced is very different from original and is likely non-functional

TACCGGACGTTTCGCACC

AUGGCCUGCAAAGCGUGG

Met - Ala - Cys - Lys - Ala - Trp

TACCTGAGGACGTTTCGCACC

AUGGGAUCCUGCAAAGCGUGG

Met -Gly - Ser - Cys - Lys - Ala - Trp

extra codon (s) added, resulting in a longer

polypeptide. May alter protein function.

• number inserted is multiple of 3

Majority of protein unaltered

TACCGGACGTTTCGCACC

AUGGCCUGCAAAGCGUGG

Met - Ala - Cys - Lys - Ala - Trp

TACCGGGACGTTTCGCACC

AUGGCCCUGCAAAGCGUGG

Met - Ala - Leu - Gln - Ser - Val

TACCGGACGTTTCGCACC

AUGGCCUGCAAAGCGUGG

Met - Ala - Cys - Lys - Ala - Trp

TACC××ACGTTTCGCACC

AUGG UGCAAAGCGUGG

Met - Val - Gln - Ser - Val

TACCGC ACGTTTCGC ACC

Met - Ala - Cys - Lys - Ala - Trp

TACGGGACGTTTCGCACC

AUG CCC UGC AAAGC GUGG

Met - Pro - Cys - Lys - Ala - Trp

TACCGGACTTTTCGCACC

AUGGCC UGA AAAGCG UGG

Met - Ala

original gene

DNA

mRNA

polypeptide

TACCGGACGTTTCGCACC

AUGGCCUGCAAAGCGUGG

Met - Ala - Cys - Lys - Ala - Trp

Deletion mutation: deletion of one or more nucleotides within the base sequence. Effect depends on the position and how many nucleotides are removed (ex: CCR5 gene and HIV resistance)

number deleted is multiple of 3
 codon (s) deleted, resulting in a shorter
 polypeptide. May alter protein function.
 Majority of protein unaltered

TACCGGACGTTTCGCACC

AUGGCCUGCAAAGCGUGG

Met - Ala - Cys - Lys - Ala - Trp

TAC ××× ACGTTTCGCACC

AUG UGCAAAGCGUGG

Met - Cys - Lys - Ala - Trp

in germ cells and somatic cells. D1.3.7—Mutation as a source of genetic variation

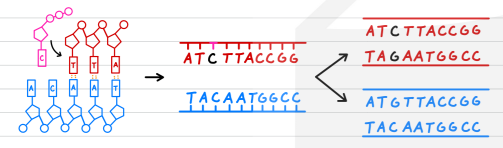
Gene mutations can be caused by a variety of factors:

Scrors in DNA replication and/or DNA repair

DNA polymerases may make errors, leading to mismatched nucleotides, insertions or deletions. Typically errors are identified and corrected via proofreading but on occasion it is not and persists. Thus, the next time to cell replicates, the mutated base sequence will be used as a template and be passed onto next generation of cells.

These spontaneous mutations are rare but normal and expected.

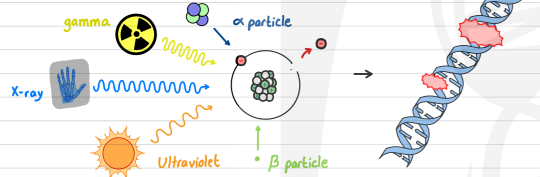
DNA replication DI.I



Mutagen: agent which can induce a change in DNA, increasing mutation frequency above background rate

> Mutagenic ionizing radiation

lonizing radiation are types of high energy (high frequency /short wavelength) radiation: gamma, X-ray, Ultraviolet (UV) as well as a particles and B particles which can displace electrons from atoms, leading to DNA damage such as single-strand or double-strand breaks



> Chemical mutagens

Mutagenic chemicals can induce mutations via a variety of mechanisms such as acting like nitrogenous bases but pairing incorrectly during DNA replication, inserting themselves into DNA causing a distortion in double helix, or by changing pairing properties.



Infectious agents

Infection of organisms by some viruses and some bacteria can result in DNA damage and mutations



HPV infection related to cervical cancer



H. pylori infection related to gastric cancer

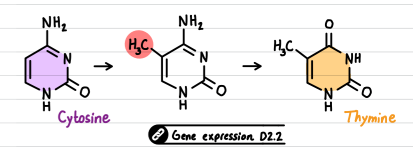
Mutations are <u>random</u>, meaning they are unpredictable.

mutations can occur anywhere in a base sequence of a genome but some areas are more frequently impacted:

• Cytosine - Guanine (CG) rich regions - CpG sites
In order to silence a gene, methylation is done to cause

DNA to become more tightly packed When Cytosine
is methylated it can spontaneously mutate to a

Thymine. CG regions are often methylated and show
higher mutation rates than any other base combination

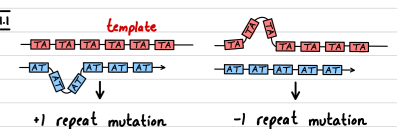


Microsatellites / Short-Tandem Repeats DNA replication DI.I

STRs are repetitive regions in the DNA (ex: ATATAT)

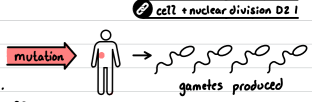
and have higher mutation rates than any other region.

One reason could be due to slippage of DNA polymerase during replication where entire repeats can be gained or lost.



X: No natural mechanism is known for making a <u>deliberate</u> change to a particular base sequence with the purpose of changing a protein or trait. 1.e an organism cannot decide or select sequences to mutate

The consequence of a mutation in humans will depend on which type of cell mutates:



if Somatic cell (body cell) DNA is mutated, only this cell and all resulting daughter cells will carry the mutation. Depending on the gene mutation it may result in cancer (uncontrolled cell division).

As mutation is not present in a gamete, it cannot be inherited by offspring

if germ cell (cells that develop into sperm/ovum) DNA is mutated and it fertilizes an ouum, the mutation will be present in the zygote and all somatic cells of the resulting embryo and adult. Due to meiosis, half of all produced gametes will carry mutation and mutated DNA may be passed onto offspring and inherited



Gene mutations after base sequence, potentially resulting in a new protein. This forms a new allele (gene variant)

members of a species may differ in which alleles they carry (genetic variation) - some being negative, neutral or positive

mutations are the source of genetic variation and a prerequisite for evolution via natural selection, where alleles

associated with advantageous traits are selected for and become more common in population (and vice-versa) -> adaptation

NOS: Commercial genetic tests can provide consumers information regarding disease Susceptibility and health risks

potential issues: X some allele variants may be missing X life insurance companies may use information to raise premiums

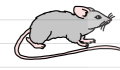
X expert interpretation/genetic counseling required so consumers understand and can make informed decisions

D1.3.8—Gene knockout as a technique for investigating the function of a gene by changing it to make it inoperative. HL D1.3.10—Hypotheses to account for conserved or highly conserved sequences in genes When analyzing the DNA of an organism, determining which sequences are genes (coding) can be done by looking for open reading frames However, how can the function of the gene be determined? START codon STOP codon Gene Knockout: technique that produces a genetically-modified organism with one specific non-functional (inoperative) gene by observing the phenotype and comparing the modified organism to a normal organism, it allows deduction of the function of inoperative gene ex: leptin gene is made LEP Knockout mice eat : hormone leptin inoperative in mice involved in appetite excessively and become obese fhousands of knockout strains in model organisms have been produced, acting as a library for genetic research suppression and regulation of fat stores General process: 1 Observation of phenotype in humans leads to hypothesis - suggesting role of gene 'X'

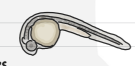
2 test hypothesis experimentally using Knockout for gene 'X' in model organism

3 validate findings in human cells /clinical trials

many different model organisms can be used to experimentally test impact of genes, each offering advantages



mice share 99% of the same genes with humans making them good analogues



zebrafish embryos are transparent, allowing early development to be easily observed



Fruit flies reproduce quickly and in large numbers and have similar genes in few chromosomes - convenient



evolution + speciation A4.1

Yeast are used due to fast reproduction times and case of genetic modification

While genetic variation is present within and across species, some sequences are identical or very similar - showing very little variation

Conserved sequences identical or similar across a species or a group of species

Highly conserved sequences identical or similar over long periods of evolution

ex: cyt c gene codes for cytochrome c; ~ 100 amino acid long protein used in aerobic cellular respiration its sequence and polypeptide product is nearly identical in many organisms

X by comparing gene or polypeptide sequences evolutionary relatedness can be estimated

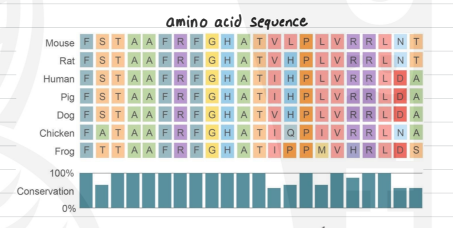
2 main hypotheses for conserved and highly conserved sequences:

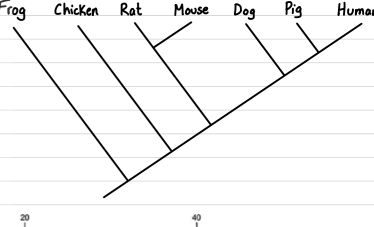
• Functional requirements

gene sequences may code for proteins essential for basic cellular stability, function, or reproduction . change in these sequences could alter the protein's functionality and lead to cell death or impede reproduction, preventing many changes to accumulate over evolutionary history as these are selected against and eventually climinated by natural selection. Akin to survivorship bias as changes aren't seen as they don't persist

Slower mutation rates

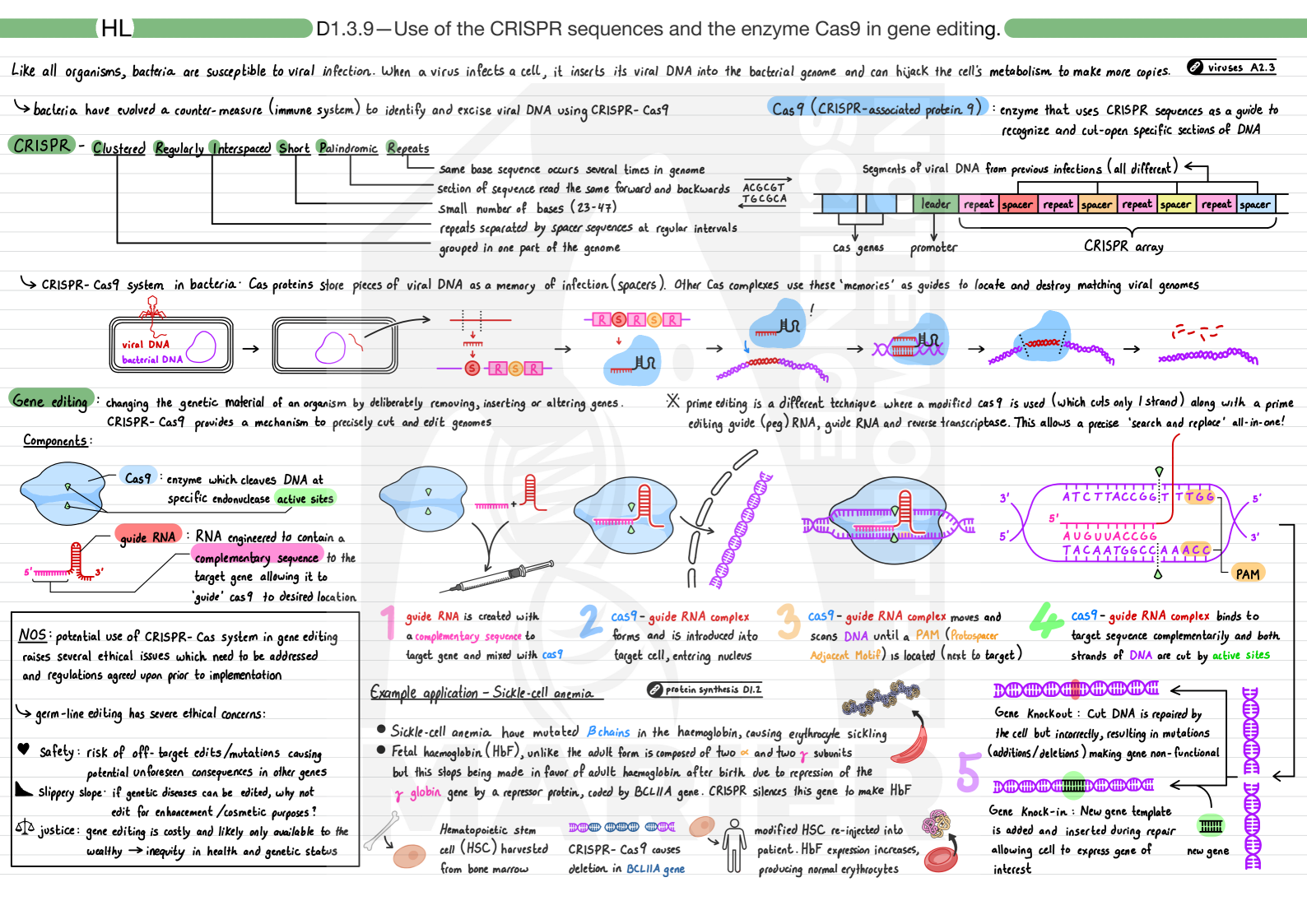
gene sequences are in regions of the genome where mutation rates are low due to higher gene expression. mutation rate is linked to gene expression. highly transcribed genes have lower rates as these regions may have more enhanced proofreading and repair mechanisms





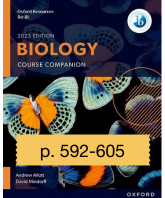
Kulikov et al 2011

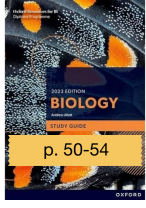
adapted from

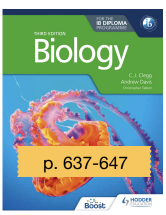


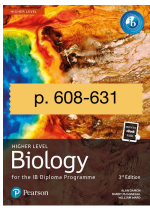


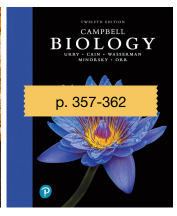
> Textbooks





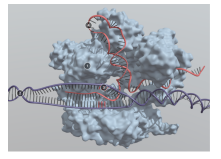








3D models

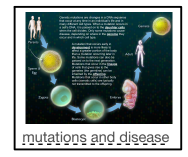


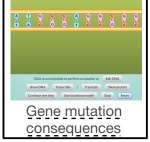
CRISPR/Cas9



Cascade and CRISPR

Simulators / Interactives > Articles







CRISPR in Nature - Innovative Genomics Institute (IGI). (2025, January 20). Innovative Genomics Institute (IGI).

https://innovativegenomics.org/crisprpedia/crispr-in-nature/

Jinek, M., Chylinski, K., Fonfara, I., Hauer, M., Doudna, J. A., & Charpentier, E. (2012). A programmable Dual-RNA-Guided DNA endonuclease in adaptive bacterial immunity. Science, 337(6096), 816-821.

https://doi.org/10.1126/science.1225829

Kulikov, A. V., Shilov, E. S., Mufazalov, I. A., Gogvadze, V., Nedospasov, S. A., & Zhivotovsky, B. (2011). Cytochrome c: the Achilles' heel in apoptosis. Cellular and Molecular Life Sciences, 69(11), 1787-1797. https:// doi.org/10.1007/s00018-011-0895-z

Lino, C. A., Harper, J. C., Carney, J. P., & Timlin, J. A. (2018). Delivering CRISPR: a review of the challenges and approaches. Drug Delivery, 25(1), 1234-1257. https:// doi.org/10.1080/10717544.2018.1474964

What Is CRISPR: The Ultimate Guide To CRISPR Mechanisms. Applications, Methods & More. (n.d.). Synthego. https:// www.synthego.com/learn/crispr