

Gene Editing - II

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What is CRISPR?

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• CRISPR stands for Clustered Regularly Interspaced Short Palindromic Repeats.

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- CRISPR Cas9 is the most prominent genome editing technique .
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- It allows researchers to permanently modify genes in living cells and organisms.

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- This can be used to correct mutations at precise locations in the human genome to treat genetic causes of diseases. \n
- Correcting the mutation in an embryo ensures that the child is born healthy and the defective gene is not passed on to future generations. \n

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How does it work?

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• The gene editing tool has two components :\n\n

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- 1. a **single-guide RNA (sgRNA)** that contains a sequence that can bind to DNA.
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- 2. the **Cas9 enzyme** which acts as a molecular scissor that can cleave DNA.

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- In order to selectively edit a desired sequence in DNA, the sgRNA is designed to find and bind to the target.
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- The genetic sequence of the sgRNA matches the target sequence of the DNA that has to be edited.

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- Upon finding its target, the Cas9 enzyme swings into an active form that cuts both strands of the target DNA. \n
- One of the two main DNA-repair pathways in the cell then gets activated to repair the double-stranded breaks. \n
- While one of the repair mechanisms result in **changes** to the DNA sequence, the other is more suitable for **introducing specific sequences** to enable tailored repair.

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- In theory, the guide RNA will only bind to the target sequence and no other regions of the genome.
- But the CRISPR-Cas9 system can also recognise and cleave different regions of the genome than the one that was intended to be edited. \n
- These "off-target" changes are very likely to take place when the geneediting tool binds to DNA sequences that are very similar to the target one. \n
- Though many studies have only found few unwanted changes suggesting that the tool is probably safe, researchers are working on safer alternatives. \n

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Why is CRISPR- Cas9 system significant?

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• Normally, if sperm from a father with one mutant copy of the gene is fertilized in vitro with normal eggs, 50% of the embryos would inherit the condition.

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• However, when the gene-editing tool was used, the **probability of inheriting the healthy gene increased** from 50 to 72.4%. There was also no off-target snipping of the DNA. \n

- The edited embryos developed similarly to the control embryos indicating that editing does not block development. \n
- Clinical trials are under way in many countries to use this tool for treating cancer.

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- It was shown in mice that it is possible to shut down HIV-1 replication and even eliminate the virus from infected cells.
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- In agriculture, a new breed of crops that are gene-edited will become commercially available in a few years.
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- Given all these, making gene editing possible in human reproductive cells deserves serious considerations in terms of legal, social and ethical consequences.
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Source: The Hindu

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