

## Birsa 101 CRISPR Gene Therapy

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### Why in News?

Recently, The Ministry of Tribal Affairs, along with CSIR and CSIR-IGIB, held a workshop on Birsa-101, during the Janjatiya Gaurav Utsav 2026.

- **Birsa-101** - India's first indigenous *CRISPR* (Clustered Regularly Interspaced Short Palindromic Repeats) based gene therapy for **Sickle Cell Disease (SCD)**.
- **Nomenclature** - Named after **Bhagwan Birsa Munda**, marking his 150<sup>th</sup> birth anniversary.
- **Developed by** - CSIR-IGIB (Council of Scientific and Industrial Research - Institute of Genomics and Integrative Biology), with support from the **Ministry of Tribal Affairs**.
- **Technological framework** - Transferred to **Serum Institute of India** for manufacturing.
- **Key Features** - It works like a *precise genetic surgery*, capable not only of curing Sickle Cell Disease but also transforming treatment pathways for several hereditary disorders.
- Uses engineered *enFnCas9*, a sharper and safer molecular tool than *SpCas9* used in Casgevy.

***enFnCas9*** - (enhanced *Francisella novicida Cas9*) is a next-generation *CRISPR* gene-editing platform engineered for ultra-high specificity and efficiency.

***SpCas9*** (*Streptococcus pyogenes Cas9*) is the most widely used *CRISPR-Cas9* genome editing enzyme.

- **Genome-wide assays** showed up to 1,000-fold fewer off-target events

compared to conventional CRISPR tools.

*Genome-wide assays scan an organism's entire DNA sequence at once. This allows scientists to identify genetic mutations, measure gene activity, and map molecular interactions across the whole genome.*

- **Mechanism of Action**

- **Direct Root Correction** - Birsa 101 corrects the single genetic misprint causing sickle cell disease.
- **Stem Cell Editing** - The therapy extracts and edits blood-forming stem cells from the patient's bone marrow to fix the defect.
- **Permanent Cure** - Engrafted cells produce healthy blood for patient's lifetime subject to long-term clinical follow-up.
- **Affordability** - It is designed to be more affordable than expensive foreign gene therapies like Casgevy and Lyfgenia.
- **National Mission** - India aims to eliminate sickle cell disease by **2047**, heavily focusing on tribal welfare across critical belts like Jharkhand, Chhattisgarh, Madhya Pradesh, and Odisha.

## Quick Facts

### Sickle Cell Disease

- A hereditary blood disorder caused by a *single typo in the genetic code*.
- Red blood cells twist into a *crescent or sickle shape, clogging blood vessels*.
- Leads to *oxygen starvation of organs*, painful crises, and long-term damage to heart, lungs, and kidneys.

### CRISPR

- It is a revolutionary gene-editing technology, works like molecular scissors (Cas9 protein) guided by RNA to cut defective DNA.
- The cell repairs itself using a corrected DNA sequence.
- Enables scientists to locate and fix errors in DNA with extraordinary precision.

## References

1. [India Education Diary | Birsa 101](#)
2. [ORF | Birsa 101](#)
3. [India Today | Birsa 101](#)



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