Overview

Pigeonpea is an important crop in Asia, Africa, and Central and South America, grown on nearly 5 million hectares worldwide. Despite its importance for food security in the world’s poorest regions, it has been under-researched in the past.

Rapid advances in genetic improvement of pigeonpea have been constrained by the lack of genomic resources coupled with low genetic diversity in the primary gene pool.

Pigeonpea was neglected until 2005, when intensive efforts by ICRISAT, the CGIAR Generation Challenge Programme, the US National Science Foundation, the Indian Council for Agricultural Research (ICAR) and several other programs led to the development of significant genomic resources.

In 2011, a global team comprising several organizations from China, Europe and the USA, and led by ICRISAT, sequenced the pigeonpea genome.

The innovation

- **Illumina** – a next generation sequencing technology – was used to generate the draft genome assembly of pigeonpea genotype ICPL 87119 (popularly known as Asha).
- This technology was used to generate a 237.2 Giga base pair of sequence, which, along with Sanger-based Bacterial Artificial Chromosome-end sequences and a genetic map, was assembled into scaffolds representing about 73% (605.78 Mega base pair) of the pigeonpea genome.
- Genome analysis led to the identification of 48,680 pigeonpea genes. A few hundred of these are unique to the crop and relate to drought tolerance, an important trait that can be transferred to other legume crops.
- The research identified 309,052 simple sequence repeats (SSRs), and 23,410 SSR primers were designed.
- Similarly, after aligning the transcript sequences from 12 genotypes, a total of 28,104 novel single nucleotide polymorphisms (SNPs) were identified.
- The completion of the pigeonpea genome has made a significant contribution to the genomic resources available for pigeonpea.

The impact

- The availability of a genome sequence opens up new avenues for pigeonpea improvement.
- The genome sequence will help harness pigeonpea’s genetic diversity by identifying molecular markers and genes for targeted traits, and will allow researchers to develop superior varieties and parental lines of hybrids.
- It will also be useful in identifying germplasm lines or advanced breeding lines with a broader genetic base for future breeding programs.
- Modern genetics and breeding approaches such as genotyping by sequencing, marker-assisted recurrent selection and genomic selection will improve the efficiency of pigeonpea breeding.

Comparison of pigeonpea and soybean chromosomes based on their genome sequences.