Application of biotechnology in bean disease management

Beans are notoriously susceptible to diseases, and several are responsible for significant yield and economic losses in Africa. An estimated total of 2,288,000 tons of beans are lost to diseases annually (Wortmann et al., 1998). To design and assess management practices and to develop host resistance to important pathogens of beans, characterisation and understanding of their diverse structure and spatial distribution is important. Identifying some of the disease causing organisms can be difficult and slow using conventional morphological, cultural and pathogenic characteristics. For example, some species of Pythium and Fusarium are often difficult to identify due to the presence of a wide range of species in the soil. However, new biotechnological tools are fast and accurate in detecting and identifying some of these bean pathogens. CIAT and its partners are currently using some of these tools for the detection and characterisation of bean pathogens, thus improving our understanding of their diversity, distribution and role in Africa.

Characterising pathogens

To understand the diversity of Pythium species, over 206 isolates from bean root rot affected areas in Kenya, Rwanda and Uganda have been characterised by molecular methods (sequencing). A total of 38 Pythium species have been identified which include pathogens, saprophytes and biocontrol agents. Five species (*P. spinosum, P. nodosum, P. pachycaule, P. torulosum* and *P. salpingophorum*) were observed for the first time to be pathogenic to beans. Distribution mapping shows *P. ultimum var. ultimum* to be the most frequently occurring species in the region.

Similar biotechnology techniques are also being used to establish the identity and relationship of Pythium species from bean and other crops. In collaboration with colleagues in CIAT Colombia, a multiplex PCR assay for six common Pythium species that are pathogenic to beans has been developed. This molecular tool is efficient in species identification, as it enables simultaneous detection and identification of several species in one assay. Molecular markers to detect and identify *Fusarium solani f.sp phaseoli* fungus, (which cause disease in beans), have been identified. These are used to detect and distinguish strains that infect beans from those that either infect other host crops or are saprophytes (Figure 1 overleaf).

Characterisation of the angular leaf spot organism *Phaeoisariopsis griseola*, a major foliar pathogen of beans in Africa, is based on a lengthy method using virulence of each isolate on 12 differential cultivars. However, in collaboration with colleagues in CIAT Colombia, a standardised molecular system has been developed to characterise the pathogen more efficiently. Locus-specific microsatellite derived markers that can distinguish between different groups of *P.*
identified. This has facilitated the and Mexico 54, MAR 1) have been some genotypes (G 10474, G 10909 against angular leaf spot disease in environmental factors. For example, not influenced by climatic or other applicable, MAS is fast, accurate and reducing the breeding process. Where genotypes in early generations, hence used to select indirectly for resistant markers linked to heritable traits are assisted selection (MAS). DNA-based in bean improvement is marker-

**Tools for genetic improvement**

Conventional breeding methods have been used to improve popular cultivars by transferring desired genes (by crossing) and then selecting for their presence in the progenies generated. For some simple traits, this is straightforward as it is easy to detect those off-springs with the desired traits. However, for other traits, conventional methods are difficult, lengthy and less efficient, and especially when handling multiple traits. To overcome these inherent difficulties, biotechnology tools have been developed, based on advances in plant molecular genetics and which offer novel possibilities in the genetic improvement of common bean and other crops. These tools enhance the efficiency, speed and effectiveness of developing new crop varieties. They also facilitate more precise and rapid identification of genotypes carrying certain desirable genes.

**Marker-assisted selection**

One of the biotechnology tools applied in bean improvement is marker-assisted selection (MAS). DNA-based markers linked to heritable traits are used to select indirectly for resistant genotypes in early generations, hence reducing the breeding process. Where applicable, MAS is fast, accurate and not influenced by climatic or other environmental factors. For example, markers linked with resistance genes against angular leaf spot disease in some genotypes (G 10474, G 10909 and Mexico 54, MAR 1) have been identified. This has facilitated the addition, efforts are underway at CIAT to tag genes for Pythium and Fusarium root rots of beans.

**Capacity building**

To improve efficiency and effectiveness of bean breeding programmes in applying modern biotechnology tools, it is crucial to build capacity of the researchers and institutions involved. Potential and capacity for utilisation of new biotechnology tools vary in different countries in eastern, central and southern Africa. However, limitations to wide application include underdeveloped infrastructure, lack of trained researchers and technicians, laboratory chemicals and equipment, and limited institutional and financial support.

Whereas, it is desirable that each national research institution acquires the minimum infrastructure to apply biotechnology tools, the reality is that for the moment this is not practical. A networking approach has been successfully used among national bean programmes in Africa where resources and responsibilities are shared, complementing efforts to overcome institutional limitations, while taking advantages of programmes with comparative advantages. A similar strategy is being put in place in the application of biotechnology tools. As capacity improves, the division of roles will gradually evolve with the use of biotechnology tools becoming more decentralised, leaving the regional facilities to carry out more specialised functions. In Uganda, NARO working in collaboration with IPGRI, CIAT and IITA has established a biotechnology centre. This facility offers an opportunity for scientists and students from the region to conduct group training attachments and postgraduate research, and is central to the application and scaling up of MAS.