# Doubled Haploid Technology in Maize (Zea mays): Status and Applications

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#### ABSTRACT

Maize is the third most important staple crop after rice and wheat with enormous diversity and adaptation ability. Hybrid breeding is the most important approach for developing high yielding cultivars in maize. It relies upon the generation of pure inbred lines with desirable traits in quick span to achieve higher genetic gains. The rapidly rising global population and climate change necessitates the development of innovative technologies that can help to safeguard the food security in future. Doubled Haploid (DH) technology is the best approach for rapid development of new inbred lines. DH technology has contributed immensely in the rapid generation of inbred lines and hybrid development. In addition, the use of molecular markers with DH technology resulted into mapping of genomic regions for different traits. The recent development in identification of alternative markers for haploid selection and genome editing approaches will further strengthen the DH technology for commercial maize breeding. This review describes important landmarks of maize DH technology, its applications, and recent advances in utilization of emerging technologies, viz. CRIPSR-cas and genomics approaches for DH technology.

Keywords: Chromosome doubling, Colchicine, Doubled haploid, Hybrid, Reverse breeding

There is need to increase the overall food production by 70% to meet the demands of 9.7 billion populations in 2050 (http://www.fao.org/wsfs/forum2050/wsfs-forum/en/) and that too under diminishing natural resources like arable land and water. The first green revolution was based on adoption of elite varieties with better management practices followed by adoption of hybrids particularly in cross pollinated crops. Hybrids are best candidates to exploit the heterosis at fullest and achieve rapid genetic gains in maize. Effective hybrid seed production relies upon the production of established and prolific inbred lines. Enhancement of genetic gains rely mainly on shortening the breeding cycles. Doubled haploid (DH) technology emerged as the potential technology to boost hybrid seed production through rapid generation of elite inbreds (Prigge et al. 2012). It helps not only in saving the valuable resources and time by rapid attainment of homozygous inbred lines (Prasanna, 2012). DH technology starts with the generation of haploids followed by doubling

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of the chromosomes to attain doubled haploids.

The first spontaneous haploid in maize was reported by Stadler and Randolph in 1929 and Chase was the first to generate maize DH inbreds through spontaneous parthenogenesis in 1946 (Randolph 1932, Chase 1947). Besides this the documentation of 0.1% spontaneous haploid induction rate in maize led to the potential utilization of haploids in hybrid breeding (Chase 1951). However, the breeders faced the constraint of very poor spontaneous haploid induction rate. But interestingly, in 1959, Coe detected a higher induction rate (up to 2.3%) in crosses with inbred line Stock 6 which later served as the progenitor for subsequently developed inducer lines across the globe. Later, Lashermes and Beckert (1988) also derived inducer line WS14 (3–5% HIR) from a cross between lines W23ig and Stock 6. In India, Dr. K.R. Sarkar carried out significant research at IARI on haploid induction and achieved the haploid induction frequency of about 6% (Sarkar et al. 1972). Maize hybrid breeding accelerated in the last decade owing to the remarkable progress in in vivo haploid induction technology (Röber et al. 2005). However, the success of the DH technology for crop improvement depends upon identification of a suitable inducer, possible introgression of genes responsible for haploid induction, standardization of protocol of in vitro methods and chromosome doubling techniques.

Since the origin of DH technology, it has been utilized in diverse crops for derivation of haploids and DH lines. DH mapping population is often used in the construction of genetic maps to identify marker-trait associations, loci/ gene responsible for economically important agronomic traits (Forster and Thomas 2005). Considering the active involvement of various public and private institutions, research labs and commercial seed companies DH technology in maize has turned to become a mature technology platform. Most multinational companies adopted this technique for development of inbred lines and superior cultivars under multi-environmental trials. The International Maize and Wheat Improvement Center (CIMMYT) has been involved in optimization of DH technology for tropical/subtropical areas through Global Maize Program (GMP) launched in 2007 with partnership of University of Hohenheim, Germany. Some other approaches, viz. marker-assisted selections (MAS), transgenic technology, induced mutagenesis also have a potential to accelerate crop improvement in combination with DH technology (Liu et al. 2016). Considering several advantages of DH particularly for 'fast marketing option', it has opened new avenues of benefits for seed industry through reduction of expenses in running breeding operations and accelerating breeding cycles for faster recovery of products.

Fundamental steps in DH Line development (Fig 1)

#### Step 1: Haploid Induction

Haploids can be derived from either male (androgenesis) or female (gynogenesis) gametophytic cells via *in vitro* methods for haploid induction. The haploids derived from male (microspores/immature anthers) and female (ovaries/ovules) gametophytic cells are known as

paternal and maternal haploids, respectively. Although androgenesis has proved a successful technique for DH production in many crop plants, but in maize it achieved little success due to non-responsiveness and dependence on donor genotype, anther stage and pretreatment conditions (Wan and Widholm 1993, Spitko *et al.* 2006). Therefore, *in vivo* haploid induction is a successful method for maize breeding program due to its operational feasibility (Seitz 2005).

In vivo approach of haploid induction includes induction of haploids using parthenogenesis, pollen treatment, inter-specific and intergeneric hybridization, haploidy inducers and directed manipulation of CENH3. The maize DH programme is mainly based on use of indeterminate gametophyte 1 (ig1) mutant and Stock 6-derived lines as inducer lines. The ig1 mutation was first time reported in Wisconsin-23 (W23) that exhibited about 3% haploid induction rates (HIR)

of paternal haploids as compared to 0.1% frequency of maternal haploids (Kermicle et al. 1969). Further molecular studies revealed the localization of ig1 on chromosome 3 which codes for LATERAL ORGAN BOUNDARIES (LOB)-domain protein transcription factors governing lateral organ development (Evans 2007, Husbands et al. 2007). Stock 6-derived haploidy inducer lines have been utilized to a larger extent in maize. The desirable attributes of good maternal haploid inducers in maize are: high haploid induction rate (HIR), plentiful pollen production, easy maintenance, good flowering behavior, proper plant height, disease and pest tolerance and wide adaptation. Initially, a spontaneous HIR of 0.1% was reported in maize (Chase 1947, 1951) which was too low to be utilized for practical breeding in maize. But later on, a much higher induction rate (up to 2.3%) was detected by Coe (1959) in crosses with inbred line Stock 6. It later served as the ancestor of all the successively developed inducer lines in maize (Geiger 2009). Crosses attempted between Stock 6 and W23 resulted in derivation of new inducer lines such as RWS, UH400, MHI and PHI exhibiting high haploid induction rate of around 7-16% (Hu et al. 2016). CIMMYT has also developed second generation tropicalized haploid inducer line (2GTAILs) with 8-15% HIR (Chaikam 2018). The first study on centromere mediated genome elimination was also tested in maize by using the AcGREENtailswap-CENH3 and AcGREEN-CEHN3 transgenes which can complement the phenotype of CENH3 knockout and knockdown lines in maize, respectively (Kelliher et al. 2016). However, HIR in developed inducers was found to be low as compared to commercial inducers available in maize. Quantitative

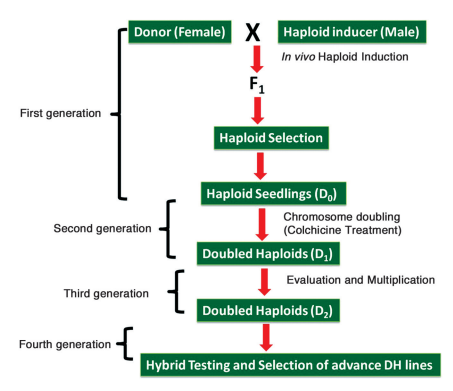


Fig 1 Basic steps for DH line development in maize.

Trait Loci (QTLs) were also identified for HIR in maize among which *qhir1*, *qhir8*, *qhir11* and *qhir12* are major. Recent studies on *qhir11* revealed the role of patatin-like phospholipase known as MATRILINEAL (MTL), PLA1 (PHOSPHOLIPASE A1) and NOT LIKE DAD (NLD) for haploid induction in Stock 6-derived lines (Gilles *et al.* 2017, Kelliher *et al.* 2017, Liu *et al.* 2017). Further investigations suggest that parthenogenesis inducing PsASGR-BABY BOOM-like gene can also be utilized to develop haploid embryos at rate of 25–89% (Conner *et al.* 2017). A list of commercial haploid inducers being utilized for haploid induction in maize breeding programs has been provided in Table 1.

## Step 2: Haploid identification or selection

DH technology gained success in maize breeding programmes due to availability of phenotypic seed coloration marker system that helps in direct identification of haploid seeds after harvesting (differentiate diploid seeds). The basis of induction of haploids has been the presence of dominant anthocyanin marker R1-Navajo (R1-nj) known as 'red crown' or 'navajo' kernel trait encoded by 'red color' gene R1. In the presence of the dominant pigmentation genes A1 or A2 and C2, R1-nj conditions the deep pigmentation of the aleurone layer (endosperm tissue) in the crown (top) region of the kernel. In addition, it causes pigmentation of the scutellum (embryo tissue) (Sarkar and Coe 1966). To be effective, the donor needs to have colourless seeds and the inducer needs to be homozygous for R1-nj and the aforementioned dominant pigmentation genes. A kernel resulting from haploid induction has a red crown (regular triploid endosperm) and a non-pigmented scutellum, whereas a regular F<sub>1</sub> kernel displays pigmentation of both the aleurone (outermost layer of endosperm) and scutellum (Geiger, 2009). In general, genes, that inhibit anthocyanin synthesis, are rather rare in dent maize (unlike flint maize) and, therefore, anthocyanin marker genes in dent maize mostly are well expressed (Rober, 1999). However, this drawback has been negated in the RWS inducer line, which carries a dominant light-independent purple-stem marker. In this case, the inducer is homozygous for the anthocyanin genes B1 (Booster 1) and Pl1(Purple1) which gives lightindependent pigmentation in the coleoptile and root of the F<sub>1</sub> seedlings. Thus, the colourless coleoptiles or root can be identified as haploid type, while the purple coloured ones are diploids at the early development stage (Geiger 2009, Rober et al. 2005). A Stock 6-derived inducer CAUHOI (with 2% HIR and higher kernel oil content of 78 g/kg) was developed to identify haploids based on absence of scutellum coloration of R1-nj scheme and low embryo oil content (Li et al. 2009). The utility of Near-Infrared spectroscopy has also been explored for differentiating haploids and diploid types (Jones et al. 2012). Melchinger et al. (2014) reported an alternative method for identification of haploid seeds based on differences arising in oil content stemming from pollination with high oil inducers. Later, Dong et al. (2018) devised clustered regularly interspaced short

palindromic repeats (CRISPR/cas9) mediated embryo- and endosperm-specific double-fluorescence protein marker for identification of haploids.

#### Step 3: Chromosome doubling

Seeds of haploid type can't be multiplied or selffertilized because they are sterile due to the presence of only one set of chromosomes in their cells. Thus, chromosome doubling is necessary to achieve the fertility in these haploid plants. A low rate of genotype-dependent spontaneous doubling has been reported in maize (Geiger et al. 2006). Therefore, a protocol with mitotic inhibitor colchicine was developed for efficient artificial chromosome doubling (Gayen et al. 1994). Mitotic inhibitors bind to the microtubular protein tubulin and thereby preventing the microtubules from pulling of chromatids towards the poles which in turn results into duplicated haploid genome without cell division. Considering the highly carcinogenic nature of colchicine, other less toxic substrates like herbicides viz. pronamid, trifluralin, oryzalin for artificial chromosome doubling treatments have been developed (Häntzschel and Weber 2010).

# Step 4: Seed multiplication of DH plants

The mitotic inhibitors treated  $D_0$  seedlings produces  $D_1$  seeds under field condition, which represents the completely homozygous line. Five percent of the haploid seeds from source germplasm result into DH lines. This rate of DH conversion depends upon various factors such as genotype background, accuracy of haploid identification system, chromosome doubling procedure and agronomic practices in the green house as well as field conditions.

# Potential Applications of DHs

In recent years, extensive emphasis was laid on shortening of breeding cycles and cost effectiveness. DH technique is an important approach for rapid development of homozygous and homogeneous progenies. It is superior over conventional breeding as it addresses the problem of slow reduction of heterozygosity observed in early segregating generations. DH progenies are homozygous lines and hence serve as good choice for selection and testing under different environmental conditions (Chang and Coe 2009, Liu *et al.* 2016). The various applications of harnessing DH technique have been explained below.

### Genetic mapping studies

QTL mapping can be performed with various populations like F<sub>2</sub>, backcross generations, RIL<sub>S</sub> and DH. However, as compared to other populations DH is most rapidly obtained homozygous population and has edge over F<sub>2</sub> and backcross populations because it provides replicated data over the years and seasons. Therefore, DH is considered as ideal mapping population for construction of genetic maps in various crops (Jeffery and Lübberstedt 2014, Liu *et al.* 2016, Wani *et al.* 2018, Kumar *et al.* 2019, Choudhary *et al.* 2019). Comparative QTL analyses of HIR

Table 1 Detailed list of maize inducer lines for DH line development

| Inducer Name   | Cross/Source  | Origin  | HIR (%)                    | Reference   |
|--|---|---|----------------------------|---|
| Stock 6  | Mexican corn  | Maize Genetics Cooperation<br>Stock Centre, USDA                                | 2.3                        | Coe 1959  |
| KMS (Korichnevy Marker<br>Saratovsky)<br>ZMS(Zarodyshevy Marker<br>Saratovsky)     | Derived from Coe's Stock 6  | Krasnodar Agricultural<br>Research Institute, France,<br>Russia                 | 0.5-3.4                    | Tyrnov and Zavalishina<br>1984  |
| EMK (Embryo Marker<br>Krasnodarsky) or ZMK<br>(Zarodyshevy Marker<br>Krasnodarsky) | Based on Coe's Stock<br>6, PEM (Purple Embryo<br>Marker)  |   | 6 to >10                   | Tyrnov and Zavalishina<br>1984  |
| WS14   | W23ig × Stock 6   |   | 3-5                        | Lasermes and Beckert 1988   |
| KEMS (Krasnador Embryo Marker<br>Synthetic)/ KHI (Krasnodar<br>Haploid Inducer)    | Derived from Coe's Stock<br>6, PEM (Purple Embryo<br>Marker)  |   | 6.0-7.9                    | Shatskaya et al. 1994   |
| MHI (Moldovian Haploid Inducer)  | $KMS \times ZMS$  | Institute of Genetics, Moldova  | 6.5-7.2                    | Eder and Chalyk 2002,<br>Rotarenco <i>et al.</i> 2010                                 |
| RWS  | KEMS $\times$ WS14  | University of Hohenheim   | 8                          | Röber et al. 2005   |
| RWK-76   | WS14 $\times$ KEMS  | (UoH), Stuttgart, Germany   | 9-10                       | Geiger 2009   |
| RWS/RWK-76   | RWS×RWK-76  |   | 9-10                       | Geiger 2009   |
| UH400 (University of Hohenheim 400)  | Derived from KEMS   |   | 8-15                       | Chang and Coe 2009  |
| UH600 (University of Hohenheim 600)  | -   |   | 8.5-12                     | https://plant-breeding.<br>uni-hohenheim.de/<br>en/84531#jfmulticontent_<br>c167370-4 |
| PK6  | Derived from Stock<br>6, WS14, FIGH1 and<br>MS1334 lines  | Diversity et Ecophysiologie<br>des Céréales (GDEC),<br>Clermont-Ferrand, France | 6                          | Eder and Chalyk 2002  |
| HZI1   | Derived from Coe's Stock 6  | Huazhong Agricultural<br>University, Wuhan, China                               | >10                        | Zhang et al. 2008a  |
| CAUHOI   | Stock 6 × Beijing High Oil population   | China Agricultural University   | ~2%                        | Li et al. 2009  |
| JAAS3 (Jilin Academy of<br>Agricultural Sciences 3)                                | Stock<br>6 × M278   | Jilin Academy of Agricultural Sciences, China                                   | 2.5-15.9                   | Cai <i>et al.</i> 2007  |
| PHI 1 (Procera Haploid Inducer 1) PHI2 PHI3 PHI4                                   | MHI × Stock 6   | Procera Agrochemicals<br>Ltd, & Procera Genetics,<br>Fundulea, Romania          | 12.1<br>13<br>14.5<br>12.8 | Rotarenco et al. 2010   |
| TAILs (Tropically adapted inducer lines) like TAIL P1 and TAIL P2                  | (CML494//(RWS×RWK)//<br>(RWS×RWK)<br>(CML494//<br>(RWS×UH400)//<br>(RWS×UH400)  | CIMMYT, Mexico and UoH,<br>Stuttgart, Germany                                   | 8-10                       | Prigge et al. 2011  |
| BHI201   | RWS, RWK-76 and B73   | Iowa State University, USA  | 12-14                      | Frei et al. 2016, Liu et al. 2016   |
| ВНІ306   | RWS, RWK-76, Ames 27451 and PI 340841   |   | 11-14                      | Frei <i>et al.</i> 2016, Liu <i>et al.</i> 2016                                       |
| CAU5   | UH400   | China Agricultural University   | 10                         | Xu et al. 2013  |
| CAU079   | CAUHOI  |   | 9                          | Xu et al. 2013  |
| 2GTAILs (Lines with highest<br>HIR namely 2GTAIL006 &<br>2GTAIL009)                | TAIL7, TAIL8, TAIL9 and (UH400 × RWSCML269) as inducers & CML 269, CML451, CML495, CML395, CKL05017, CK 05022 as non-inducers | CIMMYT, Mexico  | 8-15                       | Chaikam et al. 2018   |

in maize resulted into identification of major QTLs namely *qhir1*, *qhir8*, *qhir11* and *qhir12* which can be pyramided to increase HIR in maize haploid inducers (Prigge *et al.* 2012, Liu *et al.* 2015, Hu *et al.* 2016). DH populations have been utilized for mapping of diverse traits like digestibility, agronomic traits, biotic and abiotic stresses (Table 2). Recently Zhang *et al.* (2019) identified three major QTLs *viz.*, *qSD6-2*, *qSD8-2* and *qSBS1-2* that explained more than 10% of the phenotypic variation for stalk lodging associated traits. With the advent of cheaper sequencing platforms, the mapping studies have shifted to the use of SNP markers as cited from most recent studies (Table 2).

Immortalized  $F_2$  (IF<sub>2</sub>) populations are developed from different crosses made from RILs and/or DH populations and have a potential of permanent use in QTL analysis. IF<sub>2</sub> populations have ideal genetic background and similar to F<sub>2</sub> populations with respect to genetic information, which are used to estimate the additive and dominance effect of QTLs. IF<sub>2</sub> differs from F<sub>2</sub> in respect to replication as unlike IF<sub>2</sub>, F<sub>2</sub> cannot be replicated as the single seed represents a single genotype. In maize breeding program, these populations are useful in dissection of genetic basis of grain yield and its components like kernel quality and kernel architecture (Zhang *et al.* 2014).

Reverse Breeding: The reconstitution or derivation of complementary homozygous parental lines from the elite hybrid (through suppression of meiotic recombination process resulting into varied intact chomosomes from each parent) using DH technology is called as reverse breeding (Dirks et al. 2009). DH technique relying on the haploid production from a heterozygous individual helps to achieve the haploids. Later on, perfectly complementary homozygous DH progenies representing the parental combination of an elite hybrid can be obtained (Dirks et al. 2009, Liu et al. 2016). The complementary homozygous DH lines can be crossed in all possible combinations to regenerate or recreate the similar elite heterozygous genotype. Although it has been not carried out in maize, but its success with Arabidopsis suggests towards its applicability in other crops too (Dirks et al. 2009).

# Marker Assisted Introgression (Gene stacking)

It is a process of incorporating desirable genes into elite lines using molecular markers to enhance the target trait value in particular genotype. Marker assisted backcrossing (MABC) involves molecular marker-based foreground and background selection strategies to introgress the desired genes into an agronomically important elite line. However, it requires a large number of individuals to stabilize or fix the target gene in the recurrent parent genome. Smaller DH population is required for fixation of particular target gene in the last step of MABC. Recently, Chaikam *et al.* (2018) targeted the marker aided introgression (flaking markers-bnlg1811 and umc1917) of qhir1 from TAILs to the elite tropical maize inbreds to develop 2GTAILs. One of the introgressed line, 2GTAIL006 was found agronomically superior as well as exhibited an average HIR of 13.1% over

TAILs (almost 50% superior over the average of TAILs). Thus, MABC in combination with DH technology can help to execute rapid gene pyramiding (Lübberstedt and Frei 2012).

Hybrid breeding through harnessing wild relatives and landraces: Maize landraces are characterized by broad genetic base for biotic and abiotic stresses, wide geographical adaptation, etc. and hence can be used to improve the genetic background of modern elite cultivars (Choudhary et al. 2017). The presence of lethal alleles (masked till homozygous condition) in land races limits their use in crop improvement but DH technology can address this limitation because at haploid level expression of lethal alleles is reduced. Homozygous DH lines contain the fixed allelic variations present in the heterogeneous populations of landraces (Strigens et al. 2013). DH lines created from landraces having high genotypic variance with rapid decay of linkage disequilibrium and absence of population structure can be used for efficient association mapping (Strigens et al. 2013).

#### Plant Varietal Protection

In present scenario, Intellectual Property Rights (IPR) issues are very common in every industry. In this era of growing commercial seed industries, Intellectual Property Protection (IPP) related to Plant Variety Protection (PVP) is gaining importance. The DUS test representing distinctiveness, uniformity and stability has been performed under PVP system of The International Union for the Protection of New Varieties of Plants (UPOV). This test gives the idea about recognizable characters of a particular variety from any other available varieties, which will remain unchanged after its multiplication (UPOV, 2011). The genotypic data of both actual and simulated population created through Single Seed Descent (SSD) and DH process by Smith et al. (2008) was examined during initial and later generations. The inheritance of larger blocks of intact parental chromosomes was observed in DH progenies compared to SSD progenies based on simulation data. The study revealed the possibility of selecting DH progenies which are >90% similar to both initial parental hybrids by third generation.

Genome Editing and DH: A reliable Haploid identification (HID) method or marker is most important component of DH breeding programme (Geiger 2009). Considering the inefficiency of R1-nj marker to express in certain germplasm there was need to look for an alternative HID marker (Chaikam et al. 2015). Lately, a gene called *MATRILINEAL (MTL)/ZmPLA1* has been cloned from maize maternal haploid inducer lines (Kelliher *et al.* 2017, Liu *et al.* 2017). Dong *et al.* (2018) targeted this gene and developed haploid inducer lines using CRISPR/Cas9 system. The inducer lines were then crossed with haploid identification (HID) line (carrying double-fluorescence protein markers) to identify maternal parthenogenesis haploid seeds. The haploid seeds exhibit enhanced green fluorescent protein and DsRED, driven by an embryo-specific promoter (Liu *et* 

Table 2 List of QTLS for different traits using DH populations in maize

|                                  |  | 0   |   |                        |
|----------------------------------|--|---|---|------------------------|
| Cross                            | Major QTLs (Phenotypic Variation Explained in %)   | Traits  | Marker type                                     | References             |
| Nongxi531 × Nongxi110            | Six QTL for oil (4.34–13.13%), six QTL for protein (5.19–6.66%) and five QTL for starch concentration (4.14–7.85%)                               | Grain quality, kernel row number  | SSR   | Zhang et al. 2008b     |
| WBB53 $\times$ KW4773            | One QTL for leaf feeding (25%) and Three QTLs for stalk breakage cumulative 36%)   | Stalk breakage, leaf feeding and plant height against European Corn Borer | SSRs and SNPs                                   | Orsini et al. 2012     |
| IBM2Syn10-DH (B73 $\times$ Mo17) | QTL clusters are located near loci $gln4$ and $gln5$ , which regulate the activity of glutamine synthetase $(5.9-16.5\%)$                        | Agronomic and grain quality traits (genetic response to N deficiency)     | Genotyping-by-sequencing Gonzalez-Portilla 2014 | Gonzalez-Portilla 2014 |
| AGR_9×GSDCRW-1                   | 21 QTLs (Cumulative 38%)   | Node injury against Western Corn<br>Rootworm                              | SNPs  | Hessel 2014            |
| Lim-531 × Lim-789                | Few moderate effect QTLs with a maximum of 13.5% for Glu-Rel   | Biomass compositional and bioconversion characters                        | SNP   | Torres 2015            |
| B73 × Mo17 (IBM) Syn10           | 135 QTLs, 18 known functional genes and 25 candidate genes for flowering time and plant height fine-mapped into a 2.21–4.96 Mb interval.         | For flowering time and plant height traits                                | SNPs  | Liu <i>et al.</i> 2015 |
| $Zheng58 \times Chang7-2$        | 47 QTLs (up to 18.9%)  | Stalk associated traits   | IIIumina Golden-<br>Gate(MaizeSNP3K chip)       | Yujie et al. 2016      |
| Xianyu335 (PH6WC × PH4CV)        | Two major QTLs, $qkll-2$ (17.8%) and $qkl4-l$ (14.2%)  | Kernel length   | SNPs  | Shi <i>et al.</i> 2017 |
| B73 × Mo17 (IBM)                 | Eight QTLs for germination rate, 11 for seedling length, 13 for mesocotyl length, 15 for plumule length, and 18 for coleoptile length (2.5–7.8%) | Germination ability under deep sowing                                     | SNPs  | Liu <i>et al.</i> 2017 |
| $PH6WC \times PH4CV$             | 17 QTLs (Cumulative 35.03%)  | Salinity stress   | SNPs  | Luo et al. 2017        |
| 15TZ-spring × $15XTS$            | Two major QTLs, $qEDI(28.3\%)$ and $qCDI$ (22.6%)  | Ear associated traits like ear diameter                                   | SNPs  | Shi <i>et al.</i> 2018 |
| IBM Syn10-DH (B73 $\times$ Mo17) | Three QTLs namely <i>qSD6-2</i> (10.03%), <i>qSD8-2</i> (13.73%), and <i>qSBSI-2</i> (11.89%)  | Stalk lodging associated traits (stalk diameter, stalk bending strength), | SNPs  | Zhang et al. 2019      |

al. 2014) and an endosperm-specific promoter (Kalla et al. 1994), respectively. The embryo- and endosperm-specific double-fluorescence protein marker can be effectively used in identification of haploid in both mature seeds and young embryo (Dong et al. 2018). This ZmMTL(ZmPLA1) gene is conserved across cereal crops and hence can be used in different crops. CRISPR-Cas9 directed targeting of first exon of the phospholipase in haploid inducer lines resulted into enhancement of haploid induction by 2% (Liu et al. 2017). Similarly, transcription-activator-like effector nuclease (TALEN)-mediated deletions nearby the site of the 4-bp insertion in Stock 6-derived lines enhanced haploid induction up to 12% (Kelliher et al. 2017). Recently, a new approach known as Haploid-Inducer Mediated Genome Editing (IMGE) has been devised for developing genome-edited haploids using CAU5 HI line carrying the CRISPR/Cas9 cassette for ZmLG1 or UB2 as pollinator and B73 as female. Hence the emerging genome/gene-editing technologies will further strengthen DH programmes in maize (Wang et al. 2019, Kumar et al. 2020).

#### Conclusion and future prospects

DH technique is now yielding real and tangible results in both basic and applied biology owing to its various merits such as single step based rapid development of homozygous lines, perfect compliance of developed lines with DUS criteria for variety protection, reduced expenses for maintenance in breeding program, facilitates marker assisted selection, reverse breeding, genome-wide selection and genome editing techniques. Progress and promotion for DH production based maize breeding can be cited from the establishment of centralized maize DH facility at Kiboko (Kenya) by the collaboration of CIMMYT and Kenya Agricultural and Livestock Research Organization (KALRO) and DH facility at Bengaluru, India by collaborating with the Asian institutes. In recent years, there has been shift from conventional methods of inbred development to DH technology derived lines in commercial maize breeding programs. In the US DeKalb640, a double cross (B14 ×  $H2167/H2386 \times H2389$ ) involving 3 DH lines and one conventional inbred was the first accepted commercial hybrid with tolerance to high planting density (Chang et al. 2009). Maize research and breeding can be improved through combining DH technique with hybrid breeding, backcross breeding, genetic mapping, identification of QTLs, transgenic, induced mutagenesis and functional genomics. There is a further scope for identification of suitable inducer lines as well as introgression of genes for haploid induction from temperate to tropical maize as evident from generation of 2GTAILS. Future emphasis should be given on application of centromere mutations to produce haploid plants and identification of alternative markers for haploid selection. Haploid induction can also be combined with mini-chromosome for introducing multiple genes into elite lines. Further, "Super Haploid" inducers can also be developed which are capable of generating haploids with spontaneous haploid genome doubling capability. The integration of DH technology with MAS and GS offers new insights to minimize breeding cycles and maximize genetic gains. The future targets should rely on deeper understanding of the haploid induction mechanism and fine tuning of protocols for CRISP/cas9 for their effective use in enhancing the haploid induction rate in maize.

#### REFERENCES

- Barret P, Brinkmann M and Beckert M. 2008. A major locus expressed in the male gametophyte with incomplete penetrance is responsible for *in situ* gynogenesis in maize. *Theoretical and Applied Genetics* **117**: 581–94.
- Bouchez A and Gallais A. 2000. Efficiency of the use of doubled-haploids in recurrent selection for combining ability. *Crop Science* **40**: 23–29.
- Cai Z, Xu G L, Liu X H, Dong Y L, Dai Y X and Li S H. 2007. The Breeding of JAAS3-Haploid Inducer with High Frequency Parthenogenesis in Maize [J]. *Journal of Maize Science* **15**(1):
- Chaikam V, Nair S K, Martinez L, Lopez LA, Utz H F, Melchinger A E and Boddupalli P M. 2018. Marker-Assisted Breeding of Improved Maternal Haploid Inducers in Maize for the Tropical/Subtropical Regions. *Frontier of Plant Science* 9: 1527.
- Chang M T and Coe E H. 2009. Doubled Haploids. *Biotechnology in Agriculture and Forestry*, Vol 63. A L Kriz, B A Larkins (Eds). *Molecular Genetic Approaches to Maize Improvement*. Springer Verlag, Berlin, Heidelberg, pp. 127–42.
- Chase S S. 1951. Production of homozygous diploids of maize from monoploids. *Agronomy Journal* 44: 263–67.
- Chase S S. 1947. Techniques for isolating monoploid maize plants. *Journal of Botany* **34**: 582.
- Choudhary M, Singh V, Muthusamy V and Wani S H. 2017. Harnessing Crop Wild Relatives for Crop Improvement. *LS: International Journal of Life Sciences* **6**(2): 73–85.
- Choudhary M, Wani S H, Kumar P, Bagaria P K, Rakshit S, Roorkiwal M and Varshney R K. 2019. QTLian breeding for climate resilience in cereals: progress and prospects. *Functional and integrative genomics* **19**: 685–701.
- Coe E H. 1959. A line of maize with high haploid frequency. *American Naturalist* **93**: 381–82.
- Coe E H. 1994. Anthocyanin genetics. *The Maize Handbook*, pp. 279–81. M Freeling, V Walbot (Eds). Springer-Verlag, New York.
- Conner J A, Podio M and Ozias-Akins P. 2017. Haploid embryo production in rice and maize induced by PsASGR-BBML transgenes. *Plant Reproduction* **30**: 41–5.
- Dirks R, Van Dun K, De Snoo C B, Van Den Berg M, Lelivelt C L, Voermans W and Wijnker E. 2009. Reverse breeding: A novel breeding approach based on engineered meiosis. *Plant Biotechnology Journal* 79: 837–45.
- Dong L, Li L, Liu C, Liu C, Geng S, Li X and Xie C. 2018.
   Genome Editing and Double-Fluorescence Proteins Enable Robust Maternal Haploid Induction and Identification in Maize.
   Molecular Plant 11: 1214.
- Eder J and S T. Chalyk. 2002. *In vivo* haploid induction in maize. *Theoretical and Applied Genetics* **104**: 703–08.
- Evans M M. 2007. The *indeterminate gametophyte1* gene of maize encodes a LOB domain protein required for embryo sac and leaf development. *Plant Cell* **19**: 46–62.
- Forster B P and Thomas W T B. 2005. Doubled haploids in genetics and plant breeding. *Plant Breeding Review.* Janick J

- (Ed). 25: 57-88.
- Frei U K, De la Guente G and Lubberstedt T. 2016. Registration of BHI306 maize maternal haploid inducer germplasm. (submitted)
- Gayen P, Madan J K, Kumar R and Sarkar K R. 1994. Chromosome doubling in haploids through colchicine. *Maize Genetics Cooperation Newsletter* **68**: 65.
- Geiger H H. 2009. Doubled haploids. *Maize Handbook, Vol. II: Genetics and Genomics*. J L Bennetzen, S Hake (Eds), Springer Verlag, Heidelberg, New York. pp. 641–59.
- Geiger H H, Braun M D, Gordillo G A, Koch S, Jesse J and Krutzfeldt B A E. 2006. Variation for female fertility among haploid maize lines. *Maize Genetics Cooperation Newsletter* **80**: 28–29.
- Gilles L M, Khaled A, Lafaire J B, Chaignon S, Gendrot G, Laplaige J, Berges H, Beydon G, Bayle V, Barret P, Comadran J, Martinant J P, Rogowsky P M and Widiez T. 2017. A Loss of pollen-specific phospholipase NOT LIKE DAD triggers gynogenesis in maize. *EMBO Journal* **36**: 707–17.
- Gonzalez-Portilla P J. 2014. Genetic analysis of the IBM2Syn10-DH maize population for response to low and high nitrogen input. Graduate Theses and Dissertations, Iowa State University.
- Häntzschel K R and Weber G. 2010. Blockage of mitosis in maize root tips using colchicines-alternatives. *Protoplasma* **241**: 99–104.
- Hessel D A. 2014. Deciphering the genetic architecture of native resistance and tolerance to western corn rootworm larval feeding". *Graduate Theses and Dissertations*, Iowa State University.
- Hu H, Schrag T A, Peis R, Unterseer S, Schipprack W and Chen S. 2016. The genetic basis of haploid induction in maize identified with a novel genome-wide association method. *Genetics* 202: 1267–76.
- Husbands A, Bell E M, Shuai B, Smith H M and Springer P S. 2007. LATERAL ORGAN BOUNDARIES defines a new family of DNA binding transcription factors and can interact with specific bHLH proteins. *Nucleic Acids Research* 35: 6663–71.
- Jones RW, Reinot T, Frei U K, Tseng Y, Lübberstedt T and McClell and J F. 2012. Selection of haploid maize kernels from hybrid kernels for plant breeding using near-infrared spectroscopy and SIMCA analysis. *Applied Spectroscopy* **66**: 447–50.
- Kalla R, Shimamoto K, Potter R, Nielsen P S, Linnestad C and Olsen O A. 1994. The promoter of the barley aleurone-specific gene encoding a putative 7 kDa lipid transfer protein confers aleurone cell-specific expression in transgenic rice. *Plant Journal* 6: 849–60.
- Kelliher T, Starr D, Richbourg L, Chintamanani S, Delzer B, Nuccio M L, Green J, Chen Z, McCuiston J, Wang W, Liebler T, Bullock P and Martin B. 2017. MATRILINEAL, a spermspecific phospholipase, triggers maize haploid induction. *Nature* 542: 105–09.
- Kelliher T, Starr D, Wang W, McCuiston J, Zhong H, Nuccio M L and Martin B. 2016. Maternal haploids are preferentially induced by CENH3-tailswap transgenic complementation in maize. *Frontier of Plant Science* 7: 414.
- Kermicle J L. 1969. Androgenesis conditioned by a mutation in maize. *Science* **166**: 1422–24.
- Kumar P, Choudhary M, Hossain F, Singh N, Choudhary P, Gupta M, Singh V, Chikappa G, Kumar R, Kumar B, Jat S and Rakshit S. 2019. Nutritional quality improvement in maize (*Zea mays*): Progress and challenges. *Indian Journal of Agricultural Sciences* **89** (6): 8950–911.

- Kumar K, Gambhir G, Dass A, Tripathi A K, Singh A, Jha A K, Yadava P, Choudhary M and Rakshit S. 2020. Genetically modified crops: current status and future prospects. *Planta* **251**: 1–27.
- Lasermes P and Beckert M. 1988. Genetic control of maternal haploidy in maize (*Zea mays* L.) and selection of haploid inducing lines. *Theoretical Applied Genetics* **76**: 404–10.
- Li L, Xu X, Jin W and Chen S. 2009. Morphological and molecular evidences for DNA introgression in haploid induction via a high oil inducer CAUHOI in maize. *Planta* **230**: 367–76.
- Liu C, Li W, Zhong Y, Dong X, Hu H, Tian X, Wang L, Chen B, Chen C, Melchinger A E and Chen S. 2015. Fine mapping of *qhir8* affecting *in vivo* haploid induction in maize. *Theoretical* and Applied Genetics 128: 2507–15
- Liu C, Li X, Meng D, Zhong Y, Chen C, Dong X and Tian X. 2017. A 4-bp insertion at ZmPLA1 encoding a putative phospholipase A generates haploid induction in maize. *Molecular Plant* 10: 520–22.
- Liu H, Zhang L, Wang J, Li C, Zeng X, Xie S and Lee M. 2017. Quantitative trait locus analysis for deep-sowing germination ability in the maize IBM Syn10 DH population. Frontier in Plant Science 8: 813.
- Liu Z, Wang Y, Ren J, Mei M, Frei U K, Trampe B and Lübberstedt T. 2016. Maize Doubled Haploids. *Plant Breeding Reviews*, Vol 40, 1st Edn. Jules JanickWiley-Blackwell (Ed).
- Lübberstedt T and Frei U K. 2012. Application of doubled haploids for target gene fixation in backcross programmes of maize. *Plant Breeding* **131**: 449–42.
- Luo M, Zhao Y, Zhang R, Xing J, Duan M, Li J and Zhang H. 2017. Mapping of a major QTL for salt tolerance of mature field-grown maize plants based on SNP markers. *BMC Plant Biology* 17: 140.
- Melchinger A E, Wolfgang S, Friedrich U H and Vilson M. 2014. *In vivo* haploid induction in maize: identification of haploid seeds by their oil content. *Crop Science* **54**: 1497–1504.
- Orsini E, Krchov L M, Uphaus J and Melchinger A E. 2012. Mapping of QTL for resistance to first and second generation of European corn borer using an integrated SNP and SSR linkage map. *Euphytica* **183**: 197–206.
- Prasanna B M, Chaikam V and Mahuku G (Eds). 2012. *Doubled Haploid Technology in Maize Breeding: Theory and Practice*. CIMMYT, Mexico, D.F.
- Prigge V and Melchinger A E. 2012. Production of haploids and doubled haploids in maize. *Plant cell culture protocols*, 3<sup>rd</sup> edn, pp 161–72. V M Loyola-Vargas and N Ochoa-Alejo (Eds), Humana Press, Totowa, NJ.
- Prigge V, Xu X, Li L, Babu R, Chen S, Atlin G N and Melchinger A E. 2012. New insights into the genetics of *in vivo* induction of maternal haploids, the backbone of doubled haploid technology in maize. *Genetics* **190**: 781–93.
- Randolph L F. 1932. Some effects of high temperature on polyploidy and other variations in maize. *Genetics* **18**: 222–29.
- Röber F K, Gordillo G A and Geiger H H. 2005. *In vivo* haploid induction in maize-performance of new inducers and significance of doubled haploid lines in hybrid breeding. *Maydica* **50**: 275–83.
- Rotarenco V A, Dicu G, State D and FuiaS. 2010. New inducers of maternal haploids in maize. *Maize Genetics Cooperation Newsletter* **84**: 1–7.
- Sarkar K R and Coe J E H. 1966. A genetic analysis of the origin of maternal haploids in maize. *Genetics* **54**: 453–64.
- Sarkar K R, Panke S and Sachan J K S. 1972. Development of

- maternal-haploidy-inducer lines in maize (*Zea mays* L.). *Indian Journal of Agricultural Science* **42**: 781–86.
- Seitz G. 2005. The use of doubled haploids in corn breeding. *In: Proceedings of 41st Annual Illinois Corn Breeders' School* 2005. Urbana-Champaign, Illinois, pp. 1–7.
- Shatskaya O A, Zabirova E R, Shcherbak V S and Chumak M V. 1994. Mass induction of maternal haploids. *Maize Genetics Cooperation Newsletter* **68**: 51.
- Shi Z, Zhang R, Xing J, Duan M, Wang Y, Su A and Song W. 2018. QTL mapping of three ear traits using a doubled haploid population of maize. *Plant Breeding* **137**: 706–13.
- Shi Z, Song W, Xing J, Duan M, Wang F, Tian H and Zhang R. 2017. Molecular mapping of quantitative trait loci for three kernel-related traits in maize using a double haploid population. *Molecular Breeding* **37**: 108.
- Shuzhen Z, Zhizeng L and Ding L. 2008. Analysis of quantitative trait loci for grain quality of maize doubled haploid population. *Journal of Agricultural University Hebei* 31(3): 1–5.
- Smith J S C, Hussain T, Jones E S, Graham G, Podlich D, Wall S and Williams M. 2008. Use of doubled haploids in maize breeding: Implications for intellectual property protection and genetic diversity in hybrid crops. *Molecular Breeding* 22: 51–59.
- Spitko T, Sagi L, Pinter J, Marton L C and Barnabas B. 2006. Haploid regeneration aptitude maize (*Zea mays* L.) lines of various origin and of their hybrids. *Maydica* 51: 537–42.
- Strigens A, Schipprack W, Reif J C and Melchinger A E. 2013. Unlocking the genetic diversity of maize landraces with doubled haploids opens new avenues for breeding. *PloS one* 8(2): p.e57234.
- Troyer A F. 2004. Persistent and popular germplasm in seventy centuries of corn evolution. *Corn: Origin, History, Technology*

- and Production, pp 133–232. C W Smith, J Betran, E C A Runge (Eds). Wiley, Hoboken, NJ, United States.
- Tyrnov V S and Zavalishina A N. 1984. Inducing high frequency of matroclinal haploids in maize.
- Doklady Akademii Nauk SSSR 276: 735–38.
- UPOV. 2011.http://www.upov.int/about/en/upov\_system.html.
- Wan Y and Widholm J M. 1993. Anther culture of maize. *Plant Breeding Reviews* 11: 199–224.
- Wang B, Zhu L, Zhao B, Zhao Y, Xie Y, Zheng Z and Wang H. 2019. Development of a haploid-inducer mediated genome editing system for accelerating maize breeding. *Molecular plant* 12: 597–602.
- Wani S H, Choudhary M, Kumar P, Akram N A, Surekha C, Ahmad P and Gosal S S. 2018. Marker-assisted breeding for abiotic stress tolerance in crop plants. *Biotechnologies of Crop Improvement*, Vol 3, pp 1–23. Springer, Cham,
- Xu X W, Li L, Dong X, Jin W W, Melchinger A E and Chen S J. 2013. Gametophytic and zygotic selection leads to segregation distortion through *in vivo* induction of a maternal haploid in maize. *Experimental botany* 64: 1083–1096.
- Zhang Y, Liang T, Chen M, Zhang Y, Wang T, Lin H and Pan G. 2019. Genetic dissection of stalk lodging-related traits using an IBM Syn10 DH population in maize across three environments (*Zea mays* L.). *Molecular genetics and genomics* **294**: 1277–88.
- Zhang Z, Liu Z, Hu Y, Li W, Fu Z, Ding D and Tang J. 2014. QTL analysis of kernel-related traits in maize using an immortalized F<sub>2</sub> population. *PLoSOne* **9**(2): e89645
- Zhang Z, Qiu F, Liu Y, Ma K, Li Z and Xu S. 2008. Chromosome elimination and *in vivo* haploid production induced by Stock 6-derived inducer line in maize (*Zea mays* L.). *Plant cell reports* 27: 1851–60.