Minireview

**Brassica genomics: a complement to, and early beneficiary of, the Arabidopsis sequence**

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**Abstract**

Those studying the genus *Brassica* will be among the early beneficiaries of the now-completed *Arabidopsis* sequence. The remarkable morphological diversity of *Brassica* species and their relatives offers valuable opportunities to advance our knowledge of plant growth and development, and our understanding of rapid phenotypic evolution.

**Beyond Arabidopsis**

Less than 20 years after DNA-level studies [1] highlighted the small DNA content and simple organization of the genome of *Arabidopsis thaliana* - an ephemeral weed-like member of the *Cruciferae* family - an essentially complete sequence of its genome has been announced [2]. This, the first complete genome sequence for a flowering plant, provides the means to investigate in unprecedented detail the common features that plants share with one another and with other life forms, as well as the unique features that adapt particular taxa to diverse life histories (such as the largely sessile nature of plants). Many botanists are especially interested in studying how the interaction of natural and human selection has enabled a small subset of plants to be ‘domesticated’ to provide food, feed, and fiber. Such investigations should be greatly facilitated by the availability of plant genome sequences.

**The economic and morphological importance of Brassica**

Crops of the genus *Brassica* (tribe Brassiceae), which are in the same taxonomic family as *Arabidopsis thaliana*, are widely used in the cuisine of many cultures, due, in part, to the many choices of edible forms in the genus. Economically, *Brassica* is loosely categorized into oilseed, vegetable, and condiment crops. *B. napus, B. rapa* (formerly *campestris*), *B. juncea*, and *B. carinata* provide about 12% of the worldwide edible vegetable oil supplies [3]. *B. oleracea* and *B. rapa*, the so-called ‘cole crops’, comprise many of the vegetables in our daily diet. Several of these vegetables have extreme morphological characteristics. Examples of such morphologies include the enlarged inflorescence of cauliflower (*B. oleracea* subspecies *botrytis*) and broccoli (*B. oleracea* subspecies *italica*); the enlarged stem of kohlrabi (*B. oleracea* subsp. *gongylodes*) and marrowstem kale (*B. oleracea* subspecies *medullosa*); the enlarged root of turnip (*B. rapa* subspecies *raphiota*); the enlarged and twisted leaves of Pak-choi (*B. rapa* subspecies *chinesis*) and Chinese cabbage (*B. rapa* subspecies *pekinesis*); and the enlarged single apical bud of cabbage (*B. oleracea* subspecies *capitata*) or the many axillary buds of Brussels sprout (*B. oleracea* subspecies *geminifera*) [4]. Finally, the seed of *B. nigra* is utilized as a condiment - mustard. *Brassica* species are a valuable source of dietary fiber, vitamin C, and other possible salubrious factors such as anticancer compounds [5]. Estimates of the
economic importance of *Brassica* species are conservative, because several cole crops such as collards are cultivated primarily for local or home use, but are nonetheless a dietary mainstay in low-income communities where other fresh vegetables can be prohibitively expensive.

**The current state of Brassica genomics**

The genomes of diploid *Brassica* species are 3-5 times the size of the *Arabidopsis* genome, ranging from 0.97 pg/2C (468 Mb/1C; where C is haploid DNA per nucleus) for *B. nigra* to 1.37 pg/2C (662 Mb/1C) for *B. oleracea*. The amphidiploid genomes, which have two sets of chromosomes from each parent species, range from 2.29 pg/2C (1,105 Mb/1C) to 2.56 pg/2C (1,235 Mb/1C) [6,7]. Relationships among the three diploid species - *B. rapa* (syn. *rapa*; 2n = 20, genome AA), *B. nigra* (2n = 16, genome BB) and *B. oleracea* (2n = 18, genome CC) - and three amphidiploids - *B. napus* (2n = 38, genome AACC), *B. juncea* (2n = 36, genome AABB) and *B. carinata* (2n = 34, genome BBCC) - are well known [8]. Rapid-cycling strains of *B. oleracea* have been developed that have a life cycle as short as that of *Arabidopsis thaliana*, making them easier to study [9].

Neutral DNA polymorphisms are common among conspecific genotypes of most *Brassica* species, and at least 15 molecular maps have been produced, using at least 900 different publicly available *Brassica* and *Arabidopsis* DNA probes (which are each used with similar efficacy). Virtually all the probes hybridize to two or more loci, even in diploid *Brassica* species; however, only a subset of loci segregates for allelic variation in any one mapping population. The subset of genetically mapped loci can, therefore, be used to infer the locations of many additional sequence tagged sites (STSs) [10].

The comparative organization of the chromosomes of *Brassica* and *Arabidopsis* is well studied. On the basis of 186 corresponding loci, about 19 chromosome structural rearrangements differentiate *B. oleracea* and *A. thaliana* orthologs, suggesting that chromosomal tracts of about 20-25 cM remain largely colinear in the two taxa [10]. Microsynteny studies involving comparative genetic and physical mapping of specific chromosome segments have shown largely conserved gene order in *Arabidopsis* and *Brassica*, but some disruption in gene content by deletions or insertions [11,12]. Furthermore, duplicated partial gene clusters are commonly found in both *Arabidopsis* and *Brassica* species [13-16]. Comparative sequencing has permitted orthology assignment of some duplicated segments in *Arabidopsis* and *Brassica* species [17].

**Expectations of comparative Arabidopsis-Brassica genomics**

Comparative data help to highlight how tools from *Arabidopsis* might be used to identify genes that may directly account for *Brassica* quantitative trait loci (QTLs). An example of this from one of our labs involves the flowering-time genes. In *B. rapa*, a major QTL for flowering time was found in a region homologous to the top of chromosome 5 in *Arabidopsis* where several flowering-time genes are located [18]. After backcrossing, the *B. rapa* locus segregated as a discrete character, and, by comparative fine mapping, it was shown to correspond to the *Arabidopsis* flowering-time gene *fcl* [19]. In *B. oleracea*, 15 QTLs controlling flowering time were aligned to *Arabidopsis* genomic regions containing flowering-time genes [20]. Ten of these QTLs occurred in regions corresponding to several segments of the *Arabidopsis* genome containing genes that affect flowering: the top of chromosome 5, which contains 7 flowering genes (*tfl1, flc, tfl2, co, fy, art1 and emf1*); three regions of chromosome 1 containing *efs, fha, and gi*; and a region of chromosome 3 containing *hy2* and *vrnl*. Similarly abundant sets of *Arabidopsis* candidates have been found for *Brassica* QTLs that affect leaf and overall plant architecture [21]. The *Arabidopsis* genome sequence provides a valuable resource for identifying and further evaluating sets of candidate genes that may account for the genetic control of complex traits in *Brassica*.

**New research opportunities from Brassica genomics**

Comparative genomics has advanced the discovery and understanding of conserved genes, but most sequencing projects have also revealed many rapidly evolving ‘orphan’ genes of unknown function and evolutionary history. Such rapidly evolving genes in *Drosophila*, mammals, and several other species are important for reproductive success, cell-cell recognition, and cellular response to pathogens [22,23]. Because expressed sequence tag (EST) databases for *Brassica* remain small (as of 27 January 2001, a search of the National Center for Biotechnology Information (NCBI) databases [24] for ‘Brassica’ reveals only 5,868 matches), the abundance and possible importance of rapidly evolving genes in accounting for key differences between *Brassica* and *Arabidopsis*, or among different *Brassica* species and genotypes, remain unknown.

Naturally occurring *Brassica* genetic variants may prove useful for new investigations into plant development, particularly through the study of the effects of alleles that lack variation in *Arabidopsis*. Many *Brassica* phenotypes are under much more complex genetic control than similar phenotypes in *Arabidopsis*. For example, the discovery that the joint effects of two mutations in *Arabidopsis*, *CAULIFLOWER* and *APETALA1*, could produce a plant with a curd-like inflorescence [25] is in contrast to the much more complex genetic control of curd architecture reported for *Brassica* [20]. The high level of duplication in the *Brassica* genome [10] may permit mutations in loci that are under tight selective constraints in *Arabidopsis*. Although early hints that there may be large-scale chromatin duplications in
Arabidopsis [26,27] have been borne out by the complete sequence [2,28-30], gene loss or divergence has been substantial; most Brassica genes, however, are represented in two to three or even more copies. Investigations into the expression and function of redundant genes in Brassica species should shed light on the role of genome replication in the phenotypic divergence of the genus.

Brassica species also provide the opportunity to study rapid genome changes associated with polyploidy. Each of the amphidiploid species - B. napus, B. juncea and B. carinata - can be resynthesized by hybridizing the diploid species and then doubling the chromosomes [31]. This results in completely homozygous polyploid lines that should not segregate, but variation in DNA restriction fragments [32] and phenotype [33] has been observed among progeny derived from self-pollination of single resynthesized polyploids. As Brassica and Arabidopsis genes share, on average, 87% sequence identity [16], it may be possible to survey large numbers of Brassica genes for changes in expression using DNA microarrays based on Arabidopsis gene sequences. Comparisons of newly derived polyploids with their exact diploid progenitors should provide insight into the consequences of polyploidy on plant evolution.

Brassica is distinguished from many other plant taxa by its remarkable propensity to evolve new morphological variants rapidly, as is evident from the inter-fertility among common vegetables such as cabbage, cauliflower, broccoli, Brussels sprouts, and kohlrabi. In much the same manner that the dog genome project promises to contribute to the identification of genes conferring morphological differences among mammals [34], Brassica genomics might provide new insights into the size and shape of plants. To this end, the complete sequence of Brassica’s close relative, Arabidopsis thaliana, should prove to be a powerful tool.

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